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**Measurements of
volatile organic
compounds at a
suburban ground site**

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Measurements of volatile organic compounds at a suburban ground site (T1) in Mexico City during the MILAGRO 2006 campaign: measurement comparison, emission ratios, and source attribution

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Abstract

5 Volatile organic compound (VOC) mixing ratios were measured with two different instruments at the T1 ground site in Mexico City during the Megacity Initiative: Local and Global Research Observations (MILAGRO) campaign in March of 2006. A gas chromatograph with flame ionization detector (GC-FID) quantified 18 light alkanes, alkenes and acetylene while a proton-transfer-reaction ion-trap mass spectrometer (PIT-MS) quantified 12 VOC species including oxygenated VOCs (OVOCs) and aromatics. A GC separation system was used in conjunction with the PIT-MS (GC-PIT-MS) to evaluate PIT-MS measurements and to aid in the identification of unknown VOCs. The VOC measurements are also compared to simultaneous canister samples and to two independent proton-transfer-reaction mass spectrometers (PTR-MS) deployed on a mobile and an airborne platform during MILAGRO. VOC diurnal cycles demonstrate the large influence of vehicle traffic and liquid propane gas (LPG) emissions during the night and photochemical processing during the afternoon. Emission ratios for VOCs and OVOCs are determined from early-morning enhancement ratios and compared to emission ratios calculated from the PMF results. Average emission ratios for non-oxygenated species relative to CO are on average a factor of ~ 2 higher than measured for US cities. Emission ratios for OVOCs are estimated and compared to literature values the northeastern US and to tunnel studies in California. Positive matrix factorization analysis (PMF) is used to provide insight into VOC sources and processing and to estimate OVOC emission ratios. Three PMF factors were distinguished by the analysis including the emissions from vehicles, the use of liquid propane gas and the production of secondary VOCs + long-lived species. The total PIT-MS signal was summed to estimate the fraction of identified vs. unidentified VOC species.

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1 Introduction

Recent estimates of the global emission of volatile organic compounds (VOCs) into the atmosphere range from about 1200 to 1600 TgC yr⁻¹ (Olivier et al., 2005; Goldstein and Galbally, 2007; Reimann and Lewis, 2007; Williams and Koppmann, 2007; Yokelson et al., 2008). Major sources of atmospheric VOCs include vegetative emission, biomass burning, and the use of fossil fuels. On a global scale, VOC emissions are dominated by isoprene, which accounts for about 65% of biogenic emissions and 40% of total VOC emissions (Guenther et al., 1995, 2006; Williams and Koppmann, 2007). Biomass burning, both natural and as the result of human activity, and fossil fuel use are each thought to account for about 10% of global VOC emissions (Reimann and Lewis, 2007). Recent estimates suggest the contribution from biomass burning could be as large as 500 TgC/year (Yokelson et al., 2008). Regardless of global emissions, any of these sources can dominate locally and regionally. Anthropogenic VOC emissions often dominate in areas of high population density and contribute significantly to urban air pollution (Olivier et al., 2005; Reimann and Lewis, 2007). Air pollution is of growing international concern and contributes to approximately 2 million premature deaths per year worldwide (World Health Organization, 2008).

In and around heavily populated urban areas, VOCs released as a result of fossil fuel use are significant contributors to the formation of photochemical smog and ground-level ozone. Ozone, a secondary pollutant that is formed by photochemical reactions involving VOCs and nitrogen oxides (NO_x), has direct adverse effects on human health and can damage agricultural crops (Finlayson-Pitts, 1997; Krupa et al., 2001). Urban VOCs can also contribute to formation of particulate pollution that has significant detrimental effects on human health (Brunekreef and Forsberg, 2005; van Zelm et al., 2008). VOCs and particulate emissions related to human activity and the subsequent formation of secondary organic aerosol (SOA) have the potential to affect climate (Ramanathan et al., 2007). Mechanisms of SOA formation from urban VOCs are not well understood (de Gouw et al., 2005; Volkamer et al., 2006; Hallquist et al., 2009).

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The Mexico City Metropolitan Area (MCMA), a high-altitude sub-tropical megacity with a population of about 18 million people, is an urban center where a dense population and a local geography that restricts transport, contribute to the city's significant air quality problems. Hydrocarbon measurements in 1993, 2002, and 2003 showed highly elevated levels of many anthropogenic VOCs within the city (Blake and Rowland, 1995; Rogers et al., 2006; Velasco et al., 2007). The MILAGRO campaign (Megacity Initiative: Local and Global Research Observations) in March 2006 was designed to address the impact of these emissions on a variety of scales from local to global, and builds on results from smaller multi-investigator campaigns such as IMADA-AVER and MCMA-2003 (Edgerton et al., 1999; Molina et al., 2007, 2010).

During MILAGRO a variety of different instruments and techniques were used to quantify VOCs from both fixed sites and mobile platforms (Heald et al., 2008; Apel et al., 2010). VOC measurements were made by two different ground-based instruments at the sub-urban T1 site (Fast et al., 2007): an in-situ gas chromatograph with flame ionization detection (GC-FID) was used to measure light hydrocarbons and a proton-transfer-reaction ion-trap mass spectrometer (PIT-MS) was used to measure acetonitrile, aromatics and oxygenated VOCs (de Gouw et al., 2009). Here, these measurements are compared with canister sample analyses and with proton-transfer-reaction mass spectrometry (PTR-MS) measurements made from the Aerodyne mobile laboratory and from the US Department of Energy G1 aircraft. Because the Aerodyne mobile laboratory and the G1 aircraft also sampled near other surface sites and because canister samples were collected at many surface sites and from the NASA DC-8 and NCAR C-130 aircraft, these comparisons can be used to evaluate the consistency of VOC data obtained throughout the campaign (Kleinman et al., 2008; Fortner et al., 2009; Karl et al., 2009; Apel et al., 2010). In addition, this study supplements our understanding of the specificity of proton-transfer-reaction mass spectrometry (PTR-MS) measurements in a dense megacity with a complex VOC composition that challenges the analytical capabilities of this technique (de Gouw and Warneke, 2007).

platforms at or above T1. A PTR-MS instrument measured VOCs from the Aerodyne mobile laboratory while it was located at the site on 20–21 March. Another PTR-MS instrument measured VOCs from the DOE G-1 aircraft during multiple over flights of T1. Figure 1 gives an overview of when the different VOC measurements at T1 were made. Sampling frequency, averaging periods and limits of detection are summarized in Table 1. A complete list of species quantified can be found in Table 2.

2.2 Instrumentation

2.2.1 NOAA GC-FID

Five-minute integrated samples of light hydrocarbons were collected cryogenically every 15 min and injected into a gas chromatograph equipped with a 50 m DB-5 alumina column divided in half to allow back flushing, which shortens analysis time. A flame ionization detector was used to quantify 18 VOCs in the C2-C6 range with a detection limit of ~5 pptv. Calibrations were made 1–2 times per day from a gas standard. A detailed description of this instrument is given elsewhere (Goldan et al., 2000). Samples were collected from an air intake approximately 6 m a.g.l. that was shared with the PIT-MS system described below. Both instruments sampled outside air pumped through 1/4" PFA Teflon tubing at a rate of 3 SLM. The precision of the measurements is estimated to be 5% from the reproducibility of calibration measurements from one calibration mixture during the campaign. The accuracy is estimated to be 10% based on the variability between different calibration mixtures.

2.2.2 NOAA PIT-MS

The PIT-MS instrument uses an ion source and drift-tube reactor similar to that of a commercial PTR-MS instrument, but is equipped with an ion trap mass spectrometer instead of a quadrupole mass filter (Warneke et al., 2005a, b). During MILAGRO, the PIT-MS provided simultaneous measurements of product ions with mass to charge

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ratios (m/z) from 33 to 225 amu at a frequency of 5 min⁻¹. Calibrations using a gas standard were performed automatically every 2 h and manually using multiple dilution steps approximately once per week. Twelve different VOCs and OVOCs were quantified based on previous identification in the literature (de Gouw and Warneke, 2007). Final PIT-MS data were reported as 5-min averages for easy comparison with the GC-FID data and to improve signal-to-noise.

The measurement precision is calculated using ion-counting statistics as detailed elsewhere (Hayward et al., 2002; de Gouw et al., 2003; de Gouw and Warneke, 2007). Towards the end of the MILAGRO campaign, degradation of the PIT-MS secondary electron multiplier (SEM) detector resulted in increased noise and higher limits of detection for this period (Table 1). Accuracies for species quantified by PIT-MS are estimated at 20% for all species except acetic acid and naphthalene (50%) due to a lack of a calibration standard for the latter compounds. Benzene was detected by PIT-MS during MILAGRO but suffered from interference from clusters of water with protonated acetic acid and was not reported in final data for the campaign.

In order to distinguish isomers and identify unknown compounds detected by PIT-MS, a GC pre-separation method (GC-PIT-MS) was used on seven occasions predominantly at times when morning maxima in VOC mixing ratios occurred. In GC-PIT-MS mode, the PIT-MS instrument is coupled to a gas chromatograph fitted with a temperature programmed DB-624 column (Warneke et al., 2003). Air samples are collected in a liquid-nitrogen cooled cryogenic trap prior to injection onto the GC column. GC-PIT-MS produces a chromatogram for each of the about 180 product ion masses monitored by PIT-MS. While measuring the column effluent, the PIT-MS was operated with a time resolution of 2 s. Typical VOC peak width was 4 s or less and therefore peaks were under-sampled and cannot be used for quantification.

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2.2.3 UCI canisters

Whole air samples were collected by the University of California, Irvine in 2 L electro-polished, pre-treated, evacuated stainless steel sampling canisters. Each canister was filled for 3 h and was pressurized to about 100 kPa. After sample collection, the canisters were returned to the University of California, Irvine, where they were analyzed for CH₄, CO, hydrocarbons, halocarbons and alkyl nitrates. Detection limits for all species compared here were less than 3 pptv with precisions between 1% and 4%. For more detailed descriptions of the UCI measurements we refer to (Colman et al., 2001). Canister samples were also collected at the T0 ground site and onboard the NASA DC-8 and NCAR C-130 aircraft, and analyzed using the same methodology.

2.2.4 Aerodyne PTR-MS

A commercial PTR-MS (Ionicon Analytik, Austria) instrument was deployed on the Aerodyne mobile laboratory during the campaign (Rogers et al., 2006; Herndon et al., 2008). The mobile laboratory was parked at the T1 site for approximately 40 h from 04:00 UTC on 20 March 2006 until 18:00 UTC on 21 March 2006. Calibrations were made at regular intervals using a gas calibration standard and average calibration factors were applied to the data after the campaign. Two scan modes were used for the mobile laboratory PTR-MS. In the first mode, 24 scans of 9 masses were made each with a one second dwell time. This was followed by 12 cycles of the full mass range (20–160 amu) with a 0.1 s dwell time. The data obtained in both modes were averaged on a 1-min time basis for comparison to PIT-MS measurements during this period.

2.2.5 DOE PTR-MS

The Pacific Northwest National Laboratory deployed a commercial PTR-MS on the DOE G1 aircraft. Eleven VOCs were measured with a frequency of 0.1 s⁻¹. PTR-MS dwell times on the aircraft ranged from 0.5–1.0 s per mass and calibrations were typically done at least twice per flight day. Thirty-eight over-flights of T1 occurred during MILAGRO.

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2.2.6 Carbon monoxide measurements

Carbon monoxide measurements were made at the T1 ground site by the Georgia Institute of Technology using a modified Thermo Electron 48C CO monitor (Parrish et al., 1994). CO measurements were reported on a 1-min time base. The precision and accuracy of these data are estimated to be ± 5 ppbv and $\pm 5\%$, respectively.

CO was also quantified by gas chromatography from the UC Irvine canisters. These measurements agreed with the in-situ measurements averaged over the canister sampling periods to within 5% ($r^2 = 0.90$).

2.3 Positive Matrix Factorization (PMF)

2.3.1 PMF method

The PMF algorithm solves the bilinear, receptor-only, unmixing model, with positively-constrained factor values, and has been widely used for factor analysis and source-apportionment of both particulate matter and VOC measurements (Zhao et al., 2004; Brown et al., 2007; Engel-Cox and Weber, 2007; Reff et al., 2007; Lanz et al., 2008; Ulbrich et al., 2009; Slowik et al., 2010). Some concepts that are relevant to the understanding of this work are briefly described here. For additional details about the method, the reader is referred to the above references.

For PMF analysis, data are assembled into a 2 dimensional $m \times n$ matrix \mathbf{X} such that each of the i rows contains the measured VOC mixing ratios at sampling time t_i and each of the j columns contains the time series of a sampled VOC $_j$. A corresponding matrix is assembled specifying the measurement precision (“uncertainty”) for each point in the data matrix (σ_{ij}). The bilinear unmixing model represents the measured VOC concentrations as the sum of the contributions of p factors, each of which is comprised of a chemical profile (f) and a factor time series (g), such that for each point in the data matrix (\mathbf{X}_{ij}):

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$$\mathbf{x}_{ij} = \sum_p g_{ip} f_{pj} + e_{ij} \quad (1)$$

where e_{ij} is the fit residual for each matrix (Paatero and Tapper, 1994; Paatero, 1997; Ulbrich et al., 2009).

The PMF algorithm finds solutions of the model by minimizing a “quality of fit” parameter Q defined as:

$$Q = \sum_{i=1}^m \sum_{j=1}^n (e_{ij}/\sigma_{ij})^2 \quad (2)$$

The minimum expected value of Q/Q_{exp} is obtained when all data elements have been fit within their uncertainty (i.e., $e_{ij}/\sigma_{ij} \sim 1$), thus Q_{exp} should be approximately $m \times n$ for large datasets. Q values can be normalized to Q_{exp} , such that the expected best fit would have $Q/Q_{\text{exp}} \sim 1$. The number of factors that best represent the dataset is ultimately chosen by the user, commonly based on both (1) quantities such as Q/Q_{exp} that characterize the quality of the reconstruction, and (2) the physical plausibility of the factors. PMF solutions for a given number of factors are not mathematically unique, i.e. linear transformations (“rotations”) of the factor time series and source profiles may result in a acceptable fit to the data with similar but slight larger values of Q (Paatero and Hopke, 2009). A subset of approximate linear transformations can be explored in PMF using the FPEAK parameter.

In this study, we use the PMF2 algorithm in robust mode with default convergence and outlier criteria values. We evaluate the analysis using the recently-developed PMF Evaluation Tool (PET) (Ulbrich et al., 2009).

2.3.2 Data preparation

PIT-MS measurements were averaged over GC-FID sampling periods (Fig. 1, Table 1) for the period from 11–27 March 2006. Periods when either instrument was off line

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were excluded from the PMF analysis. The data matrix for the analysis presented here consisted of 851 simultaneous measurements (rows) from the GC-FID and PIT-MS instruments, arranged so that 18 columns contain the time series of 18 species from the GC-FID, 12 columns contain the time series of 12 VOCs quantified by PIT-MS, and the final 35 columns contain the time series of 35 PIT-MS ion signals not quantified (65 columns total). The 35 PIT-MS ion signals chosen were not identified as specific molecules, but all did show a significant signal during the campaign.

For the GC-FID data we used 5 pptv or 5%, whichever number was larger, to calculate errors for PMF analysis. Errors for the PIT-MS data were calculated using ion counting statistics (de Gouw and Warneke, 2007). The error for 11 PIT-MS masses with signal to noise ratios (SNR) less than 2 was increased by a factor of 4, as is typical for the PMF method (Paatero and Hopke, 2003; Ulbrich et al., 2009). None of the measurements considered in the analysis had a SNR < 1. Error was increased for only one quantified PIT-MS mass (m/z 129, naphthalene). Results of PMF analysis and an estimate of the uncertainties involved in this analysis are discussed in Sect. 4.

3 Data quality evaluation

The quality of VOC data obtained at T1 was evaluated using direct measurement comparisons between the two NOAA measurements and other measurements where possible. The selectivity of the PIT-MS data was evaluated using GC-PIT-MS analyses.

3.1 Measurement comparisons

3.1.1 NOAA GC-FID vs. UCI canisters

The on-line GC-FID measurements were compared to UCI canister measurements for 17 compounds. For the purpose of these comparisons, the on-line GC-FID data were averaged over the 3-h sampling times of the UCI canisters. Examples are shown for

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propane, ethylene and acetylene (Fig. 2a–c). The data in Fig. 2 were fit using 2-sided, orthogonal distance regressions (ODR). The resulting regression slopes (s), and values of the linear correlation coefficient (r^2) are summarized in Table 2 for all 17 overlapping VOC measurements. For all VOCs, the degree of correlation between the in-situ and canister measurements was high ($r^2 > 0.87$) and most measurements agreed within the 10% accuracy of the in-situ measurements. Notably, the canister measurements of acetylene (Fig. 2c) were systematically higher than those made in-situ. The reasons for this difference are unknown; calibration standards were not compared for the purpose of this study. Some of the scatter around the best fit may be explained by differences in sampling times since the GC-FID only sampled for one 5-min-period every 15 min while canister sampling was continuous for 2–3 h.

3.1.2 NOAA PIT-MS vs. UCI canisters

Both the PIT-MS and the UCI canisters reported mixing ratios for aromatic compounds. An example of the comparisons is shown for toluene in Fig. 2d. The PIT-MS data were averaged onto the canister sampling times for the purpose of this plot. The two measurements correlated well and agreed quantitatively within 7%. The other PIT-MS measurements of aromatics correlated well but were systematically higher than the UCI canister measurements (Table 2). The reasons for these differences are unknown; calibration standards were not compared for the purpose of this study. PTR-MS measurements of higher aromatics represent the sums of many different isomers, each with its own calibration uncertainty. As a result, inter-comparisons tend to show larger differences than for single compounds (de Gouw and Warneke, 2007).

3.1.3 NOAA PIT-MS vs. Aerodyne PTR-MS

The Aerodyne mobile laboratory collected data at the T1 site for two days during the campaign (Fig. 1). Six VOCs were measured by both the NOAA PIT-MS and the Aerodyne mobile laboratory PTR-MS. Figure 2e shows the measurement comparison for

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acetaldehyde. Slopes and correlation coefficients for all 6 VOC comparisons are summarized in Table 2. Correlation coefficients (r^2) for all 6 compared compounds were greater than 0.80.

PIT-MS measurements of methanol were systematically 32% lower than those of the Aerodyne PTR-MS. This difference is similar to the difference observed (35%) between the calibration standards used with each of the two instruments. The PIT-MS measurements of reported aromatics were systematically 40–60% higher than the Aerodyne PTR-MS measurements (Table 2) and are outside of the combined measurement uncertainties. Unlike methanol, differences in calibration standards do not appear to explain differences between measurements of aromatics.

3.1.4 NOAA PIT-MS vs. DOE PTR-MS

The DOE G1 aircraft flew over the T1 ground site 38 times during MILAGRO. All overpasses occurred in mid-afternoon when VOC mixing ratios for primary species were at or near daily minima. Aircraft overpass altitudes ranged between 800–2800 m a.g.l. with typical well-mixed daytime boundary layer depths of at least 2000 m. Measurements of boundary layer heights indicate that overpasses of T1 occurred within the boundary layer (Shaw et al., 2007; Fast et al., 2009).

Eight VOC measurements were compared for the 15 overpasses when both instruments were operating. Overpasses were defined as measurements made by the DOE-G1 instrument within a horizontal distance of 5 km from the ground site. DOE-G1 PTR-MS measurements for each VOC during a particular over flight were averaged and compared to PIT-MS ground measurements averaged within a window of ± 10 min from the over flight period. An example of the comparison between surface and airborne measurements is shown in Fig. 2f for acetone. Due to the low number of data points, linear least-squares fits of the data were constrained to an intercept of zero for these comparisons. Slopes and r^2 values for fits are shown for all 8 VOCs in Table 2. The inter-comparison is challenging due to: (1) low VOC mixing ratios in a narrow concentration range, (2) the large vertical separation, and (3) short duration of the over-flights.

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Therefore, correlation coefficients for individual VOCs were low ($r^2 < 0.55$), but most species except the C9-aromatics did not show large systematic differences between the two measurements.

3.2 GC-PIT-MS

The NOAA PIT-MS was operated in GC-PIT-MS mode seven times during MILAGRO to identify VOCs and evaluate PIT-MS measurements at T1. The GC-interface was similar to that used in a previous study (Warneke et al., 2003); however, the present application is unique in the sense that a chromatogram is obtained at each detection mass of the PIT-MS. Most GC-PIT-MS analyses occurred early in the day coincident with peak VOC concentrations. Figures 3 and 4 show selected chromatograms from the use of this method for PIT-MS signal attribution. Individual masses of interest are discussed in the following section.

The chromatographic peaks observed at masses 33 (methanol), 45 (acetaldehyde), 59 (acetone), 93 (toluene), 107 (C8-aromatics), and 121 amu (C9-aromatics) were very similar to the results from previous GC-PTR-MS measurements in urban air (Warneke et al., 2003; de Gouw and Warneke, 2007) and provided no evidence of important interferences for any of these compounds.

Acetonitrile is measured by PIT-MS at 42 amu. Interferences from the reaction of alkanes and alkenes with O_2^+ ions have previously been observed using GC-PTR-MS (de Gouw et al., 2003; de Gouw and Warneke, 2007). Interferences from propane were evident in GC-PIT-MS chromatograms from Mexico City where mixing ratios of propane reached as high as 250 ppbv. From the GC-PIT-MS measurements it is difficult to quantify the sensitivity of the signal at mass 42 amu to propane. However, scatter plots of the signal at 42 amu vs. propane showed no correlation between the two measurements. We conclude that while propane contributed to the signal at 42 amu, most of the variability in the 42 amu signal was due to acetonitrile.

Even masses are typically associated with nitrogen-containing compounds in PTR-MS measurements. The chromatogram at 54 amu contained a small peak that is

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attributed here to acrylonitrile. However, a larger fraction of the signal at 54 amu in on-line measurements comes from $\text{H}_3\text{O}^+(\text{H}_2\text{O})_2$ clusters at 55 amu; water cluster ions show a tail towards lower masses in PIT-MS. Two unidentified even-mass compounds were detected at 46 and 62 amu by GC-PIT-MS but no conclusive identification was established. Low mixing ratios and isotopic interferences at adjacent masses from more abundant compounds (acetaldehyde and acetic acid respectively) prevent even tentative quantification of these even mass compounds.

Formic acid has previously been reported as the dominant contributor to the PTR-MS signal at 47 amu (Veres et al., 2008). Organic acids do not elute from the GC column used in GC-PIT-MS and therefore cannot be positively identified by this technique. However, several other compounds (ethanol, dimethyl ether, and formate esters) are detectable by GC-PIT-MS at 47 amu. These compounds did not appear in the chromatograms during MILAGRO and it is unlikely that they are major contributors to the signals on this mass.

The presence of the gasoline additive, methyl tert-butyl ether (MTBE) at 57 amu has been previously observed in field measurements in the United States and Mexico City and confirmed in laboratory studies (Karl, 2003; Warneke et al., 2003, 2005b; Rogers et al., 2006). During MILAGRO, GC-PIT-MS confirmed that the signal at 57 amu was dominated by a fragment of MTBE (62%) with minor contributions from the butenes (multiple peaks, 23%) and acrolein (15%), which compounds have been previously observed at this m/z in PTR-MS (Fortner and Knighton, 2008). MTBE also interfered (16%) with the measurement of methyl ethyl ketone (MEK) at 73 amu during the campaign. Previous work has shown that only about 0.2% of the signal from MTBE appears at its protonated parent ion (89 amu) and GC-PIT-MS showed no evidence of MTBE on this mass.

Acetic acid has been quantified previously at 61 amu using PTR-MS (de Gouw et al., 2003). The chromatogram at 61 amu does not show acetic acid because it does not elute from the column. Two other species were observed in the chromatogram, tentatively identified as methyl formate and ethyl acetate. The latter compound is also

observed at 89 amu and has been previously reported in Mexico City (Rogers et al., 2006; Fortner et al., 2009). Ion counts observed at T1 in GC-PIT-MS chromatograms are at least an order of magnitude smaller than those that would be expected from PIT-MS online measurements of 1–2 ppbv VOC at 61 amu. We conclude that the online signal at 61 amu is dominated by acetic acid with only minor interference from methyl formate and ethyl acetate.

Isoprene mixing ratios measured from UCI canisters revealed average integrated mixing ratios of about 50 pptv \pm 100% with maximum values at night and minima about noon local time. A small signal from isoprene was present in GC-FID chromatograms but this compound was not quantified by GC-FID because it was frequently below detection limit. GC-PIT-MS chromatograms did show peaks consistent with extremely small mounts of isoprene (69 amu), methyl vinyl ketone + methacrolein (71 amu) and one monoterpene (at 81 and 137 amu), possibly limonene. Isoprene, however, cannot be separated chromatographically (by GC-PIT-MS) from furan, and the chromatogram at 71 amu also showed contribution from several C5 alkenes. Additionally, measurements at 69 and 71 amu were highly correlated with primary species such as aromatics and alkenes, suggesting that non-biogenic sources dominate these mass signals in the MCMA. We conclude that PTR-MS measurements at masses 69 and 71 are not reliable indicators of biogenic emissions in the MCMA.

The chromatogram at 131 amu revealed the presence of two unknown compounds with relatively short retention times. On-line measurements show large intermittent spikes at this mass that often correlate with peaks (in order of decreasing signal magnitude) at 109, 145, 140, 122 amu and weak signals at numerous masses above 157 amu. The origin of this signal is unknown but could be due to a local source of halogenated hydrocarbons. Retention times suggest that these halogenated hydrocarbons, if present, have at least one fluorine substitution. Fragmentation of protonated, high-mass halogenated hydrocarbons in PTR-MS is probable, but poorly characterized and this finding remains speculative.

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Signals from aromatic compounds in GC-PIT-MS chromatograms are shown in Fig. 4. Peaks for small aromatics can be clearly identified for benzene (79 amu), toluene (93 amu), xylene isomers (107 amu), and ethyl benzene. (107 amu). Compound identification consistent with C9 aromatics (121 amu) and some of the C10 isomers (135 amu) can be assigned by retention times with some confidence (Fig. 4b). Some of the peaks in chromatograms at 135 amu and all of those at 149 amu are tentatively identified in Fig. 4c. C11 aromatics probably dominate the signal at 149 amu although the identification of specific peaks remains tentative (Fig. 4).

PIT-MS measurements at 129 amu are reported here as naphthalene: few other hydrocarbons have this molecular mass, and high concentrations of gas-phase naphthalene exceeding 0.3 ppbv ($1.5 \mu\text{g m}^{-3}$) have been previously been reported for Mexico City (Marr et al., 2006). Maximum mixing ratios of naphthalene measured by PIT-MS were as high as 0.8 ppbv, however this compound was not calibrated during the campaign. GC-PIT-MS has a low sensitivity to naphthalene and it is very likely that ambient levels of naphthalene were below the limit of detection for the method. No peaks were observed in the GC-PIT-MS chromatogram at mass 129.

4 Results and discussion

4.1 Diurnal VOC patterns

VOC mixing ratios at T1 had a pronounced diurnal pattern during MILAGRO that is summarized graphically by chemical family in Fig. 5. Large, direct emissions and a shallow boundary layer resulted in a maximum concentration for most compounds just before local sunrise (07:00 local time). After sunrise, mixing ratios for primary emissions decreased rapidly due both to photochemical oxidation and dilution into an expanding daytime boundary layer (Fig. 5a–c). Compounds with shorter lifetimes reached their minimum values more quickly and their normalized profile minima were closer to zero. Similar diurnal patterns were evident in the three-factor PMF solution discussed below.

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Photochemical production of oxygenated species leads to a very different diurnal profile. The balance of the production and the dilution in an expanding boundary layer causes mixing ratios for most OVOCs to decline more slowly after sunrise than those with exclusively primary sources (Fig. 5d). The atmospheric processing of primary emissions and production of secondary products is discussed in detail elsewhere (de Gouw et al., 2009).

The fraction of the PIT-MS signal that could not be attributed to specific compounds is shown in Fig. 6 as a function of the time of day. The unidentified signal averaged 18% of the total for the entire campaign, a result which is consistent with previous estimates of unidentified signal from field measurements with PIT-MS (Warneke et al., 2005b). During the campaign, the majority of PIT-MS signal can be attributed to compounds clearly identified using GC-PIT-MS (Fig. 6).

4.2 Urban emission ratios

4.2.1 Hydrocarbons

The early morning maximum in most VOCs is due to the accumulation of primary emissions in a shallow boundary layer in the absence of significant chemical removal. Therefore, these data are very useful for the determination of urban emission ratios. Most VOCs with primary sources only (hydrocarbons) showed a high degree of correlation with inert combustion tracers such as carbon monoxide (CO) (Fig. 7). Most alkenes and aromatics were also strongly ($r^2 > 0.70$) correlated with carbon monoxide (Table 3). Propane, n-butane and i-butane correlated poorly with CO, as their emissions are likely dominated by leakage of liquid propane gas (LPG) rather than combustion (Blake and Rowland, 1995).

Here we derive emission ratios from the slopes of two-sided (ODR) fits of the VOC data vs. CO. Table 3 summarizes these emission ratios for all measured hydrocarbons. It should be noted that the resulting LPG species emission ratios have substantially larger uncertainties than other VOCs due to much lower correlation with CO. Emission

ratios for important VOCs not measured by GC-FID or PIT-MS (e.g. ethane, C7 and higher alkanes, and benzene) were calculated from UCI canister measurements of these compounds and CO.

4.2.2 Oxygenated VOCs

5 Many OVOCs have both direct emission sources and are formed photochemically. In an urban environment, sources of OVOCs from combustion are likely minor compared with those from industrial, evaporative and particularly photochemical sources. As a result, poor correlation with CO is expected and observed for most OVOCs (Fig. 8). Fits made in the plots of morning OVOC mixing ratios vs. CO tend to parallel the lower
10 edge of the data (closest to the CO axis) suggesting a method for estimating OVOC measurements that might be useful in future studies.

Emission ratios vs. CO were calculated from slopes of ODR fits for measurements made between 04:00 and 07:00 local time in order to minimize the effects of photochemical production. The results are shown by solid, black lines in Fig. 8 and summarized in Table 4.
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4.2.3 Comparison of emission ratios with those from US cities

Emission ratios for hydrocarbons vs. CO from measurements made in the United States (Baker et al., 2004; Warneke et al., 2007) are added to Fig. 7 and to Table 3 for comparison purposes. For most alkenes and aromatics, Mexico City emission ratios to CO are approximately a factor of two larger than corresponding values from the United States. The difference is probably due to differences in automobile emission systems and fleet ages between the two countries. CO to CO₂ ratios of about
20 4.5 ppbv ppmv⁻¹ were observed during the MILAGRO campaign (Vay et al., 2009); a value similar to the ratio observed in plumes originating in China and twice as large as the 1–2 ppbv ppmv⁻¹ observed in industrialized nations like Japan and the United States (de Gouw et al., 2004; Takegawa et al., 2004). It is worth noting, therefore, that VOC emission ratios per kg fuel burned are probably closer to a factor of 6 ± 2 larger
25 in Mexico City than those for US cities.

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Emission ratios for OVOCs from 2004 measurements in New England and those from a 1999 California tunnel study are shown in Fig. 8 and Table 4 (Warneke et al., 2007; Ban-Weiss et al., 2008). OVOC emission ratios for Mexico City more closely resemble values obtained in the tunnel study than those calculated from measurements in the Northeastern US. Differences between OVOC emission ratios from the MCMA and the Northeastern US could be due to sampling methodology, i.e. the Northeastern US data were largely collected outside city boundaries and although an attempt was made to account for the effects of secondary production using a photochemical lifetime method, the resulting emission ratios may still overestimate the direct emissions. Biogenic emissions may also account for some of the differences observed between Mexico City and New England, particularly for methanol, which has significant biogenic sources (Jacob, 2005; Millet et al., 2008).

4.3 PMF results

4.3.1 The PMF solution

PMF solutions with 1 to 7 factors were examined using the method described in Sect. 2.3. A three-factor solution was chosen based on the plausibility of the results, with factors identified as traffic, LPG leakage, and secondary + long-lived species, described individually in more detail below. This solution has $Q/Q_{\text{exp}} = 4.07$ at FPEAK 0, which suggests that the errors may be somewhat underestimated or that there is substantial variability in the dataset that cannot be modeled well with a few fixed chemical profiles. Q/Q_{exp} increases by less than 10% over the range in FPEAK from -5 to +5. Over the narrower FPEAK range from -3 to +3, Q/Q_{exp} increases by 2% compared to the FPEAK 0 solution. Measurements from each instrument contributed $44 \pm 0.1\%$ (GC-FID) and $56 \pm 0.1\%$ (PIT-MS) to the parameter (Q) over the range from FPEAK -3 to +3. About 20% of the value of Q could be attributed to unidentified PIT-MS signal.

Source profiles for the 3-factor solution are shown in Fig. 9. The first factor, referred to as “traffic” from now on, is comprised of hydrocarbons including highly reactive

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chemical processing of the air mass is at its daily minimum. Other methods used to estimate uncertainties in the analyses, e.g. bootstrapping and seed variation (Ulbrich et al., 2009), were investigated but gave smaller variation in factor profiles. The three-factor solution was investigated as a function of the weighting errors assumed for the GC-FID and PIT-MS data. None of the observed variation changed the conclusions drawn here in any significant way. A more comprehensive analysis of the effects on PMF analysis of relative weighting of the data from different instruments can be found elsewhere (Slowik et al., 2010).

The averaged diurnal variation in the factor time series is shown in Fig. 11. Briefly, the traffic factor consists of aromatics, alkenes, alkanes and acetylene, and is at its maximum in the early morning. The LPG factor consists predominantly of propane, n-butane and i-butane and also has a maximum in the morning. The secondary + long-lived species factor is at its maximum about 2 h later than the traffic factor and falls off much more slowly after sunrise. The averaged diurnal variations in factor time series were overlaid with the VOC data in Fig. 5. The traffic factor closely follows the diurnal variations in the more reactive alkenes and aromatics. The LPG factor is very similar to the measured diurnal variations in propane and other alkanes. The secondary + long-lived species factor does not directly match the diurnal variation of any single oxygenated species, but approximates the ensemble average diurnal variation of the oxygenated species shown.

The PMF reconstructions of the time series of selected VOCs are shown for acetylene, n-butane and acetone at FPEAK 0 in Fig. 12. The variability in acetylene measurement is well described by the PMF time-series reconstruction ($r^2 = 0.89$) and the majority of this compound is attributed to traffic sources. The variability in n-butane is also reasonably well described by the PMF time-series reconstruction ($r^2 = 0.76$). In this case, most of the signal is attributed to LPG leakage. The measurement of acetone and its PMF time-series reconstruction (Fig. 12c) correlate well ($r^2 = 0.93$) and most of the signal is attributed to the secondary + long-lived species factor with a smaller contribution from the traffic factor. Large residuals between measured and reconstructed

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concentrations were frequently correlated with simultaneous spikes in several VOCs without corresponding changes in CO mixing ratios. Examples of such spikes can be seen in Fig. 12 for acetylene and n-butane and they suggest the presence of local emissions, the variability of which is difficult to capture with PMF if the plume composition is sufficiently different from the dominant sources of variation.

Time series reconstructions for other species were compared to the measurements (results not shown here). Linear correlation coefficients were high for C7-C10 aromatics, alkenes and acetylene ($r^2 > 0.85$) and slightly lower for methanol, acetone, acetaldehyde and MEK ($0.82 < r^2 < 0.90$). PMF reconstructions for alkanes and the C11-aromatics correlated reasonably well with measurements ($0.60 < r^2 < 0.88$). Much lower correlations were found for acetonitrile ($r^2 = 0.42$), acetic acid ($r^2 = 0.48$), and naphthalene ($r^2 = 0.55$). Correlations for these species did not improve significantly even when PMF solutions with greater than three factors were considered ($p > 3$). Poor fits for these compounds in the PMF analysis suggest that they have important sources that are not correlated with the three factors identified here. For acetonitrile, the explanation may be the presence of biomass burning sources (de Gouw et al., 2009) that were not identified as a separate factor in the PMF analysis.

4.4 PMF traffic factor

The profile of the traffic factor is shown in Fig. 9 (top panels). The profile is dominated by alkenes and aromatics. C5 and higher alkanes are also present. The only OVOC with significant partitioning to the traffic factor was methanol. Methanol mixing ratios were substantial during MILAGRO and have likely contributions from tailpipe and other vehicular sources previously observed by vehicle emission studies in the MCMA (Rogers et al., 2006) or by industrial sources.

The PIT-MS product-ion signal at 57 amu was predominantly apportioned to the traffic factor, which is consistent with the major contribution of the gasoline additive MTBE to 57 amu (~80%), as determined by GC-PIT-MS (Fig. 3). Other product ion signals with significant presence in the traffic factor are at 69, 71, 83 and 85 amu. The signals at

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69 and 71 amu are typically used to quantify isoprene and its photoproducts MVK and methacrolein in PTR-MS. This analysis suggests that the importance of other species with traffic-related sources (such as the presence of C5-alkenes at 71 amu shown in Fig. 3) at these masses in Mexico City can be shown with PMF analysis. The absence of a PMF factor representing biogenic emissions for compounds measured at m/z 69 and 71 by PIT-MS is consistent with the observations about the signal at these masses (e.g. low mixing ratios, correlation with CO and GC-PIT-MS results) discussed elsewhere in this text.

4.4.1 PMF LPG factor

The LPG factor profile is dominated by the three light alkanes (propane, n-butane and i-butane) associated with liquefied petroleum gas use in the MCMA (Fig. 9, middle panel) (Blake and Rowland, 1995; Vega et al., 2000; Velasco et al., 2007). The reconstruction of n-butane in the 3-factor PMF solution is compared with its measurement in Fig. 12 (middle panel). The three major LPG alkanes also contribute to the traffic factor, possibly because LPG is used as a vehicle fuel (Diaz et al., 2000), and also to the secondary + long-lived species factor, due to their lower reactivity with the hydroxyl radical (k_{OH}) as discussed below. However, the contribution of propane to these factors is strongly dependent on FPEAK (Fig. 10) and therefore not well-quantified by this analysis.

Other compounds apportioned to this factor include other light alkanes, some traffic-related VOCs and methanol, but the amount of signal associated with the LPG factor for these other VOCs decreases as a function of FPEAK. The diurnally averaged profile for the LPG factor has a similar shape to that of the traffic factor, but has a larger magnitude and wider morning maximum (Fig. 11) probably due to the longer photochemical lifetime of major LPG species.

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4.4.3 Estimation of VOC emission ratios using PMF results

Linear regression (ODR) of individual PMF factors plotted vs. CO showed that the traffic factor correlated well with CO ($r^2 = 0.86$) in the predawn period but showed no correlation after noon. The LPG and secondary + long-lived species factors showed poor correlation with CO in the morning but stronger correlations in the afternoon ($r^2 = 0.81, 0.77$ respectively). To estimate ($r^2 < 0.42$) urban emission ratios for VOCs and OVOCs in the MCMA from the PMF results, the sum of the contributions of the traffic and LPG factors were plotted against CO measurements for the 30 compounds quantified and reported during the campaign plus formic acid (m/z 47). The slopes of the resulting plots are presented in Table 3 and 4 for VOCs and OVOCs respectively.

The relative uncertainties (1σ) in the slopes of the linear regression used to calculate PMF-derived emission ratios were small (1–3%) for all compounds. Uncertainty in PMF-derived emission ratios was also assessed using changes in apportionment as a function of the PMF FPEAK parameter. The maximum relative difference in apportionment from FPEAK 0 over the range from FPEAK –3 to 3 for the sum of the traffic and LPG source profiles was larger (4%–100%) than the regression uncertainty for all compounds. The sum of the relative uncertainties for each compound were used to calculate the absolute uncertainties in PMF-derived emission ratios shown in Tables 3 and 4.

Emission ratios calculated from PMF results are compared to values calculated directly from VOC measurements in Fig. 13. Individual points represent VOC species and are colored according to the PMF factor that explained the majority of the variation of that species. Emission ratios estimated for compounds mainly associated with traffic and LPG emissions were systematically lower by 20% than those derived directly from measurements, while the values of PMF-derived emission ratios for secondary + long-lived compounds were larger by an average of 80%. As described in Sect. 4.2.2 above, emission ratios for OVOCs were calculated only from the measured data from 04:00 to 07:00 local time, when the direct emissions should dominate the variability of these

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species. Emission ratios for OVOCs estimated from the traffic + LPG factors from the PMF analysis attempt to include only the directly emitted fraction of the OVOCs. Thus, both methods used here for calculating OVOC emission ratios attempt to correct measurements for secondary production. Uncertainty in the emission ratios calculated from PMF results is dominated by the uncertainties in the PMF results, particularly strongly for OVOCs.

4.4.4 Limitations of the PMF analysis as applied to VOC measurements

Some limitations of the application of PMF to VOC mixing ratios became apparent over the course of this work. First, the bilinear unmixing model makes the assumption that the species identified together in one factor occur in constant relative proportions over the entire measurement period. However, individual VOCs and OVOCs have photochemical lifetimes (k_{OH}) that span several orders of magnitude, and so VOCs emitted from a single source are removed in the atmosphere at very different rates and thus the ratios of their concentrations will change continuously. This helps explain the way individual VOCs are apportioned between the factors. For example, the traffic factor closely matches the diurnal variability of the shortest-lived compounds that have low mixing ratios, while primary hydrocarbons with long lifetimes are partially explained by the secondary + long-lived species factor in proportion to their photochemical lifetimes (Fig. 14), because they are still present in highly processed air. As another example, strict interpretation of the PMF factors (Fig. 9) as sources would suggest that a small portion of toluene and methanol can be attributed to LPG leakage and that benzene and propane are partially attributed to secondary formation, even when this contradicts our knowledge of the sources of these compounds. Thus, the PMF factors identified for this dataset reflect the convolution of VOC sources and VOC lifetimes, and interpretation of the factor profiles strictly in terms of different sources is inappropriate.

Second, for this dataset, the fraction of the variation for some important LPG and OVOC species (e.g., n-butane and acetone) explained by each factor was strongly dependent on the FPEAK parameter (Fig. 10). Known characteristics of sources and

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lifetimes for these VOCs did not help to significantly narrow the range of FPEAKs of plausible or physically reasonable solutions. For some species, therefore, it is difficult to quantify the fraction emitted by particular sources with much certainty.

5 Conclusions

Two ground-based NOAA instruments were deployed during the MILAGRO campaign that provided quantitative analysis of 30 VOCs. Comparison of measurements from these two instruments with canister samples and two mobile PTR-MS instruments was performed and the measurements agreed within uncertainty estimates for most compounds.

PIT-MS provided on-line quantification of 12 VOCs and was used to monitor an additional 150 + product ion masses. In conjunction with a GC pre-separation method, the ion trap provided speciation not possible with quadrupole PTR-MS. PIT-MS also validated the identified compounds commonly quantified by the proton-transfer reaction method. In most cases only minor interferences were observed. About 85% of the total PIT-MS signal arose from identified compounds. Signals at 69 and 71 amu, commonly associated with biogenic emissions at other locations, appear to be dominated by traffic emissions in the MCMA. Product ion masses 69, 71, 75, 85, 87, 130 amu accounted for a large fraction of the unidentified signal from PIT-MS.

Values for VOC and OVOC emission ratios with respect to CO were reported from Mexico City are presented and compared with previously reported values from US cities. Urban VOC emissions in the MCMA are dominated by traffic emissions and LPG usage. Emission ratios to CO for most hydrocarbons are generally about a factor of two larger than in the United States, while emission ratios to CO₂ are likely a factor of (6 ± 2) larger than US values. Urban OVOC emission ratios in Mexico City, calculated by two different methods appear consistent with vehicular sources of these compounds, although measurement uncertainties are large.

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Positive Matrix Factorization analysis performed on the combined data from the two NOAA instruments factors: traffic, LPG usage and secondary production and long-lived species. These factors described much of the variability in VOC mixing ratios, especially for compounds with vehicular and liquid propane gas sources. OVOCs and compounds with long photochemical lifetimes were identified as a single factor in the analysis. PMF solutions with more than 3 factors resulted in splitting of major factors rather than the identification of minor emission sources. The PMF analysis did not identify distinct factors consistent with industrial, biomass burning or biogenic VOC emissions. Factor profiles (grouping of VOC species) are influenced by both sources and photochemical lifetimes of individual VOCs.

PMF results were also used to estimate emission ratios for VOCs and OVOCs. Estimates agreed within uncertainties with the corresponding emission ratios derived directly from measurements for most hydrocarbons typically associated with vehicle usage. PMF-derived emission ratios were about 20% lower and 80% higher for light alkanes and OVOCs respectively. We conclude that PMF describes major VOCs well, but strongly caution against the over-interpretation of PMF factors as strictly corresponding to sources of VOCs because the factors may reflect combinations of sources and photochemical lifetimes, and because solutions may not be unique with respect to rotation.

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Table 1. Limits of detection for individual VOCs quantified during the MILAGRO campaign by PIT-MS and PTR-MS.

Instrument:	NOAA PIT-MS	Aerodyne PTR-MS	DOE PTR-MS	
period:	5 min	1 s	1 s	
date:	11–27 March	28–31 March ^a		
Compound	Limit of Detection (pptv)			
toluene	150	(800)	110	100
Σ C8 aromatics	100	(800)	200	110
Σ C9 aromatics	100	(800)	160	110
Σ C10 aromatics	100	(800)		
Σ C11 aromatics	100	(800)		
naphthalene	100	(800)		
methanol	1500	(1500)	530	
acetaldehyde	300	(1500)	400	350
acetone	150	(800)	230	250
acetic acid	300	(1200)		350
methyl ethyl ketone	300	(2000)		150
acetonitrile	100	(800)		85

^a PIT-MS LODs reported for two periods 11–27 March and 28–31 March. LOD changed due to hardware problems.

Table 2. Slopes resulting from ODR regression of scatter plots between VOCs measured by 2 or more techniques during MILAGRO. Also shown here are 1σ uncertainties in slopes, correlation (r^2) coefficients, and the number of points compared (N).

	best fit slope	r^2	N
	NOAA GC-FID ^b : UCI Canisters ^a		
propane	0.98 ± 0.04	0.96	128
n-butane	1.02 ± 0.03	0.97	128
i-butane	1.02 ± 0.03	0.97	128
n-pentane	1.04 ± 0.03	0.98	128
i-pentane	0.89 ± 0.02	0.98	128
n-hexane	0.94 ± 0.03	0.97	128
ethylene	0.90 ± 0.03	0.97	129
propylene	1.00 ± 0.03	0.97	128
1-butene + 2-methyl propene	0.99 ± 0.04	0.96	128
cis-2-butene	0.95 ± 0.04	0.95	128
trans-2-butene	0.79 ± 0.03	0.96	128
1-pentene	1.18 ± 0.16	0.87	71
cis-2-pentene	1.00 ± 0.06	0.94	77
trans-2-pentene	1.03 ± 0.04	0.96	98
2-methy 2-butene	0.78 ± 0.07	0.90	60
3-methy 1-butene	0.93 ± 0.12	0.87	82
acetylene	0.74 ± 0.04	0.94	129
	NOAA PIT-MS ^c : UCI Canisters ^a		
toluene	1.07 ± 1.74	0.87	112
Σ C8 aromatics	3.38 ± 0.14	0.87	108
Σ C9 aromatics ^d	2.68 ± 0.44	0.72	101

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Table 2. Continued.

	best fit slope	r^2	N
NOAA PIT-MS ^c : Aerodyne PTR-MS			
methanol	0.68 ± 0.11	0.84	1605
acetaldehyde	1.10 ± 0.21	0.81	1695
acetone	1.00 ± 0.14	0.86	1695
toluene	1.61 ± 0.18	0.89	1685
Σ C8 aromatics	1.39 ± 0.18	0.87	1685
Σ C9 aromatics	1.38 ± 0.22	0.84	1685
NOAA PIT-MS ^c : DOE G1 PTR-MS			
acetonitrile	0.93 ± 0.17	0.06	15
acetaldehyde	0.97 ± 0.13	0.18	15
acetone	0.86 ± 0.07	0.50	15
acetic acid	0.50 ± 0.08	0.10	15
methyl ethyl ketone	0.84 ± 0.09	0.36	15
toluene	1.07 ± 0.14	0.51	15
Σ C8 aromatics	0.88 ± 0.17	0.04	15
Σ C9 aromatics	0.30 ± 0.14	0.23	9

^a UCI-Canisters, data from entire campaign.

^b NOAA GC-FID, data from entire campaign.

^c NOAA PIT-MS, data from 4–7 a.m., local time.

^d Calculated using Σ (traffic + LPG) factors; $r^2 = 0.83$ with CO for PMF traffic factor and $r^2 = 0.59$ for LPG factor.

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Table 3. Urban emission ratios for non-methane hydrocarbons quantified during MILAGRO vs. carbon monoxide are compared to PMF-derived emission ratios and values for US cities from the literature. Correlation coefficients (r^2) and the number of points fit (N) are shown for measurement derived emission ratios. Reported errors for measurement-derived emission ratios are 1σ uncertainties in the slope of the ODR fit. Uncertainties reported for the PMF-derived emission ratios were calculated using the method described in the text. Uncertainties reported do not account for calibration uncertainties.

Compound	MILAGRO Emission Ratio Best Estimate 2006 (pptv [ppbv CO] ⁻¹)	r^2	N	New England (Warneke et al., 2007) 2004 (pptv [ppbv CO] ⁻¹)	28 US cities (Baker et al., 2008) 1999–2005	MILAGRO PMF Emission Ratio ^o
Alkanes						
ethane ^a	21.5 ± 10.8	0.50	199	11.62	2.4	
propane ^b	61.7 ± 15.6	0.53	1242	7.73	3.8	43.5 ± 4.1
n-butane ^b	21.7 ± 5.0	0.57	1242	1.69	1.4	17.3 ± 1.5
i-butane ^b	7.2 ± 1.6	0.57	1242	1.01	0.9	6.0 ± 0.5
n-pentane ^b	2.5 ± 0.2	0.81	1241	1.55	1.2	2.6 ± 0.3
i-pentane ^b	3.3 ± 0.4	0.77	1242	3.99	2.9	3.4 ± 0.4
n-hexane ^b	1.49 ± 0.16	0.70	1240	1.07	0.6	1.0 ± 0.1
cyclopentane ^a	0.153 ± 0.008	0.88	166			
cyclohexane ^a	0.164 ± 0.008	0.77	181	0.29		
methyl cyclopentane ^a	0.47 ± 0.02	0.87	194	0.57		
2,2-dimethyl butane ^a	0.36 ± 0.02	0.89	159	0.12		
2,3-dimethyl butane ^a	1.67 ± 0.13	0.72	160	0.27		
2-methyl pentane ^a	1.33 ± 0.07	0.89	165	1.11		
3-methyl pentane ^a	0.90 ± 0.05	0.91	166	1.28		
n-heptane ^a	0.36 ± 0.06	0.69	199	0.40	0.2	
2,4-dimethyl pentane ^a	0.126 ± 0.006	0.87	187	0.17		
n-octane ^a	0.122 ± 0.006	0.33	152	0.20	0.1	
n-nonane ^a	0.065 ± 0.003	0.32	158			
n-decane ^a	0.042 ± 0.002	0.26	182	1.E-04		

Table 3. Continued.

Compound	MILAGRO Emission Ratio Best Estimate 2006 (pptv [ppbv CO] ⁻¹)	<i>r</i> ²	<i>N</i>	New England (Warneke et al., 2007) 2004 (pptv [ppbv CO] ⁻¹)	28 US cities (Baker et al., 2008) 1999–2005	MILAGRO PMF Emission Ratio ^e
Alkenes						
ethylene ^b	7.0 ± 0.4	0.90	1242	4.56	4.1	7.1 ± 0.9
propylene ^b	3.0 ± 0.2	0.86	1242	1.36	1.0	2.8 ± 0.4
1-butene ^b	0.35 ± 0.02	0.87	1236	0.14	0.2	0.32 ± 0.04
2-methyl propene ^b	0.85 ± 0.04	0.91	1236			0.76 ± 0.11
cis-2-butene ^b	0.18 ± 0.02	0.82	1236	0.06		0.16 ± 0.02
trans-2-butene ^b	0.20 ± 0.02	0.82	1235	0.05		0.16 ± 0.02
1-pentene ^b	0.152 ± 0.011	0.88	1211			0.15 ± 0.02
cis-2-pentene ^b	0.100 ± 0.009	0.86	1206	0.05		0.09 ± 0.02
trans-2-pentene ^b	0.190 ± 0.017	0.85	1209			0.17 ± 0.03
2-methyl 2-butene ^b	0.179 ± 0.019	0.83	1202			0.15 ± 0.02
2-methyl 1-butene ^b	0.140 ± 0.011	0.87	1213			0.14 ± 0.02
3-methyl 1-butene ^b	0.044 ± 0.006	0.80	1195			
1,3-butadiene ^a	0.278 ± 0.014	0.84	147			
Alkynes						
acetylene ^b	6.5 ± 0.3	0.90	1242	3.60	3.4	5.0 ± 0.7
Aromatics						
benzene ^a	1.21 ± 0.06	0.94	183	0.62	0.7	
toluene ^c	4.2 ± 0.4	0.69	2563	2.62	2.7	4.0 ± 0.4
Σ C8 aromatics ^c	4.3 ± 0.6	0.71	2563	1.93		3.8 ± 0.5
Σ C9 aromatics ^c	2.8 ± 0.6	0.79	2563	1.07		2.3 ± 0.4
Σ C10 aromatics ^c	0.76 ± 0.15	0.71	2563			0.60 ± 0.14
Σ C11 aromatics ^c	0.16 ± 0.03	0.41	2563			0.12 ± 0.01
naphthalene ^c	0.12 ± 0.01	0.24	2592			0.11 ± 0.01

^a UCI-Canisters, data from entire campaign.

^b NOAA GC-FID, data from entire campaign.

^c NOAA PIT-MS, data from 4–7 a.m., local time.

^d Calculated using Σ (traffic + LPG) factors; *r*² = 0.86 with CO for PMF traffic factor and *r*² = 0.56 for LPG factor.

^e *N* = 851 for emission ratios derived from PMF analysis.

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Table 4. Urban emission ratio estimates vs. CO for OVOCs and acetonitrile calculated by linear regression and from MILAGRO measurements and from the results of PMF analysis. For comparison, literature values from PTR-MS measurements made in New England and from a US tunnel study are also shown. Methods used to calculate these values are discussed in detail in the text. Also shown are correlation coefficients (r^2) and the number of points fit (N). Reported errors for measurement-derived emission ratios are 1σ uncertainties in the slope of the ODR fit. Uncertainties reported for the PMF-derived emission ratios were calculated using the method described in the text. Uncertainties reported do not account for calibration uncertainties.

	MILAGRO Emission Ratio Best estimate 2006 ^a pptv [ppbv CO] ⁻¹	r^2	N	New England (Warneke et al., 2007) 2004					Caldecot Tunnel (Ban-Weiss et al. 2008)				MILAGRO PMF Emission Ratio ^{b,d}
				1999	2001	2006	Diesel	1999	2001	2006	Diesel		
Oxygenates													
methanol	2.1 ± 0.5	0.11	2563	4.0									6.1 ± 2.1
acetaldehyde	1.0 ± 0.3	0.23	2563	0.7	1.0	0.5	0.7	12.0					2.0 ± 0.9
formic acid ^c	0.22 ± 0.11	0.12	2592										0.4 ± 0.3
acetone	0.51 ± 0.13	0.06	2563	2.9	0.7	0.4							1.0 ± 1.0
acetic acid	0.5 ± 0.2	0.09	2563										0.5 ± 0.5
MEK	0.29 ± 0.07	0.13	2563	0.8	0.1	0.1	0.2	1.4					0.5 ± 0.3
Other													
acetonitrile	0.27 ± 0.07	0.24	2563										0.4 ± 0.1

^a Ratios calculated from PIT-MS data collected between 4–7 a.m. LT.

^b Calculated using Σ (traffic + LPG) factors; $r^2 = 0.86$ with CO for PMF traffic factor and $r^2 = 0.56$ for LPG factor.

^c For purposes of PMF, formic acid was included as one of the 65 unquantified PIT-MS measurements.

^d $N = 851$ for PMF-derived emission ratios.

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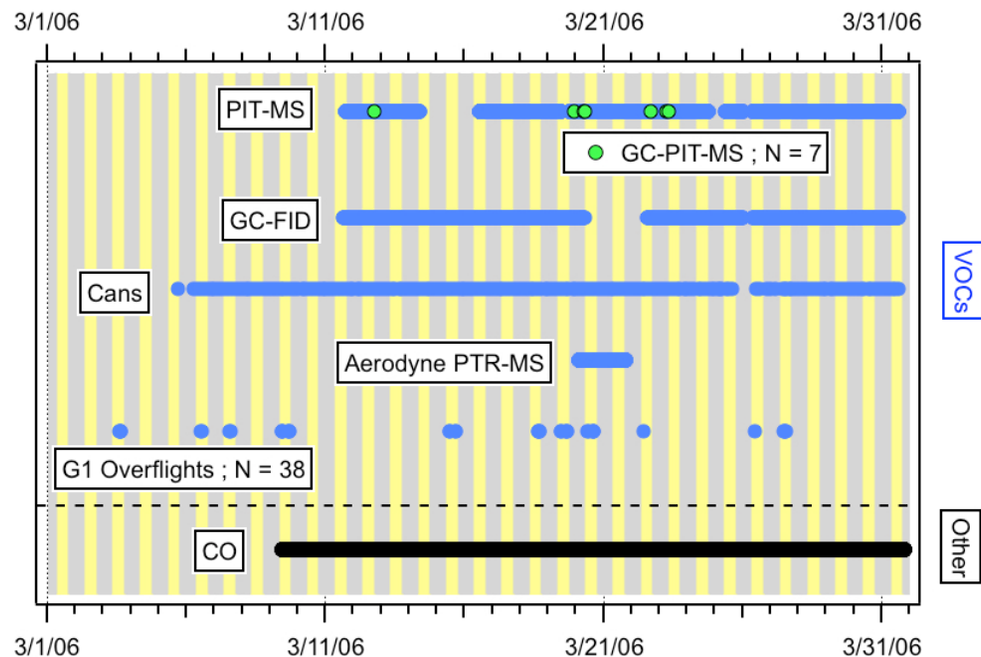


Fig. 1. Sampling periods for the VOC measurement methods and CO at the T1 ground site in Mexico City during the MILAGRO campaign. Approximate daylight hours (7:00 a.m.–19:00 p.m. local time) are shown in yellow.

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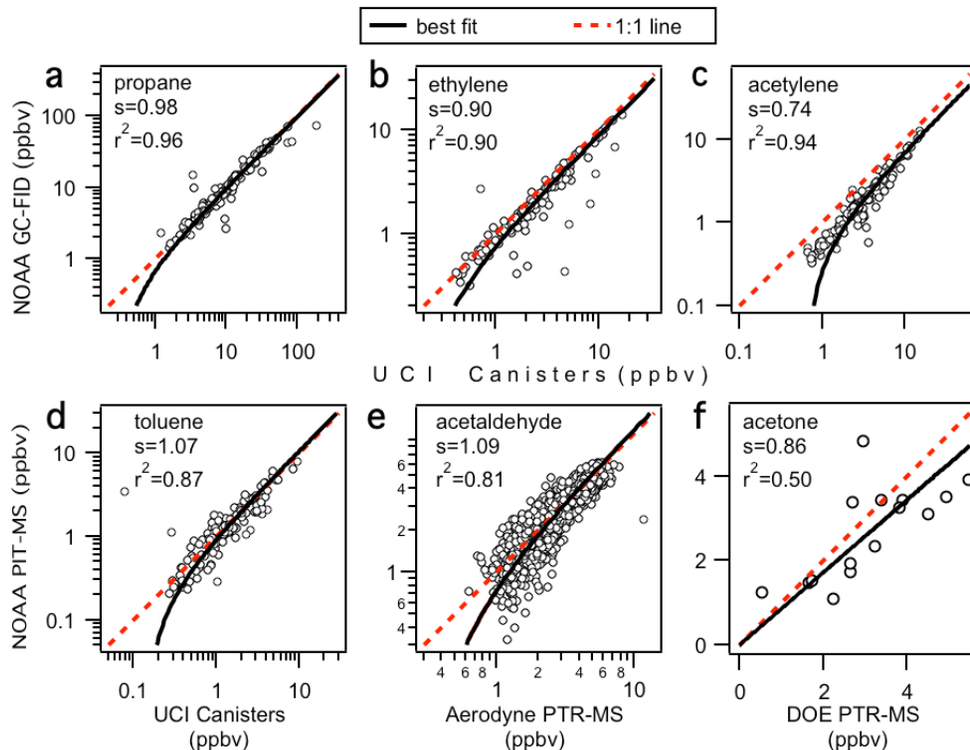


Fig. 2. Selected results of the comparison of VOC measurements at the T1 site during MILA-GRO. Panels **a–c** compare the measurements of the NOAA GC-FID and the UC Irvine canister measurements for the same compound. Panel **d–f** compare measurements made by the NOAA PIT-MS to made from the UC Irvine Canisters and by the PTR-MS instruments on the Aerodyne Mobile Laboratory and the DOE G1 aircraft, respectively. Slopes from linear regressions (s) and correlation coefficients, r^2 , for comparisons are tabulated in Table 2.

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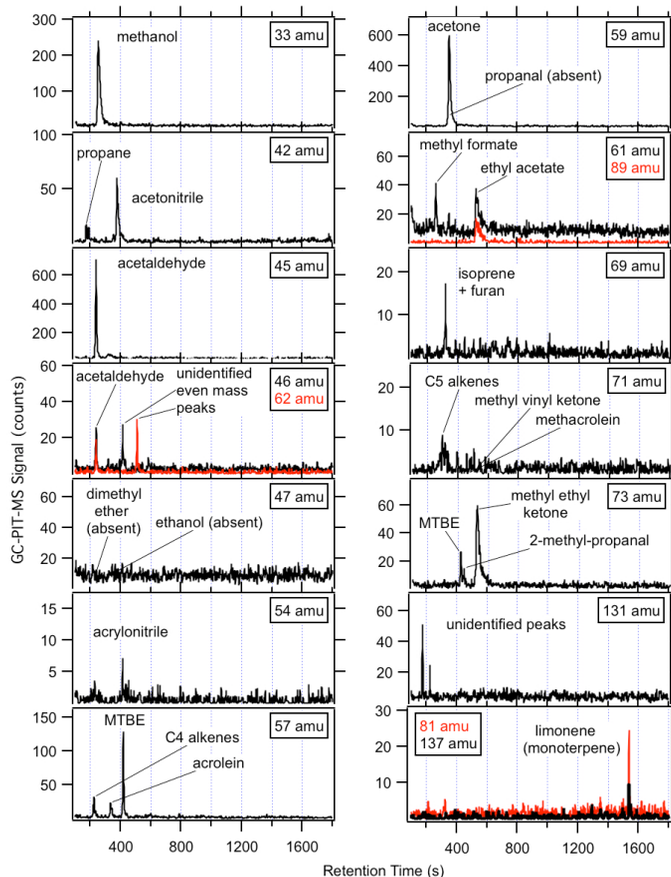


Fig. 3. Select GC-PIT-MS chromatograms obtained during the MILAGRO campaign. Signals reported are sums of six individual chromatograms. The retention time scale is slightly adjusted to be comparable with previously reported values for GC-PTR-MS using the same GC instrument and temperature program (Warneke et al., 2003).

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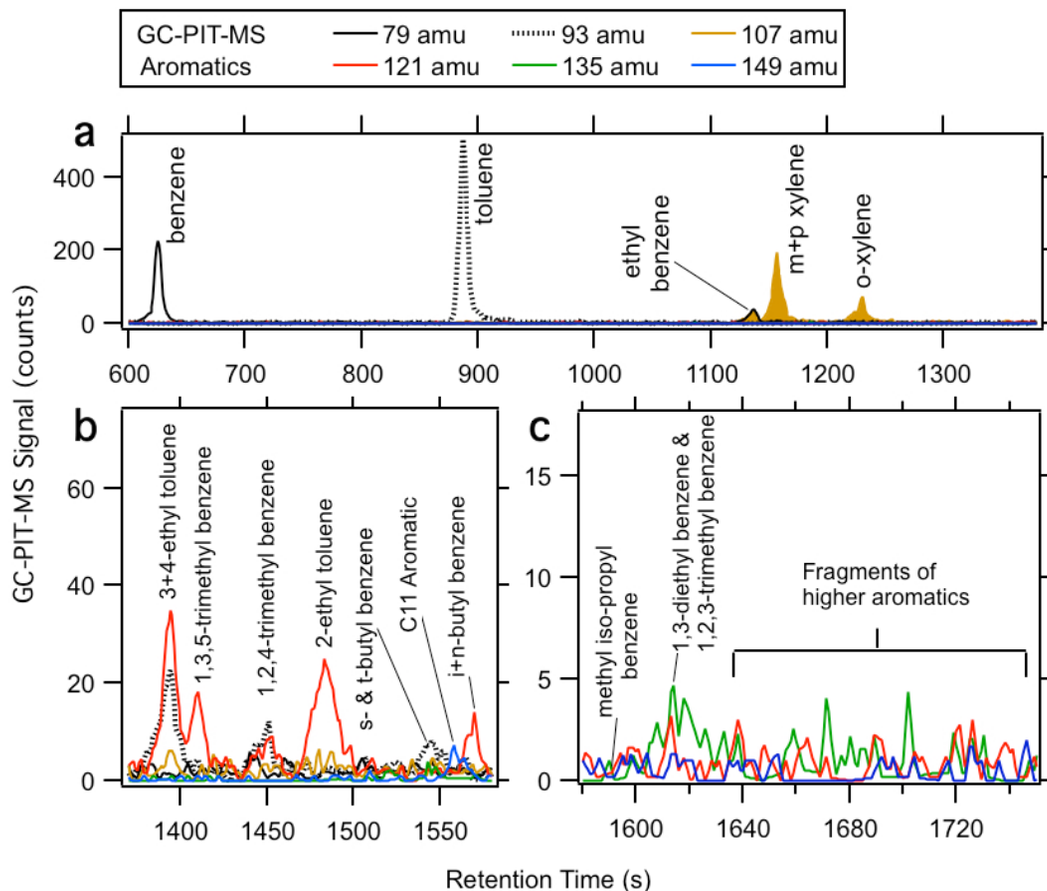


Fig. 4. GC-PIT-MS chromatograms for aromatic measurements from 600–1400 s **(a)**, 1300–1580 s **(b)** and 1580–1750 s **(c)**. Identifications made after 1300 s retention time are more uncertain than those for the C6–C8 aromatics.

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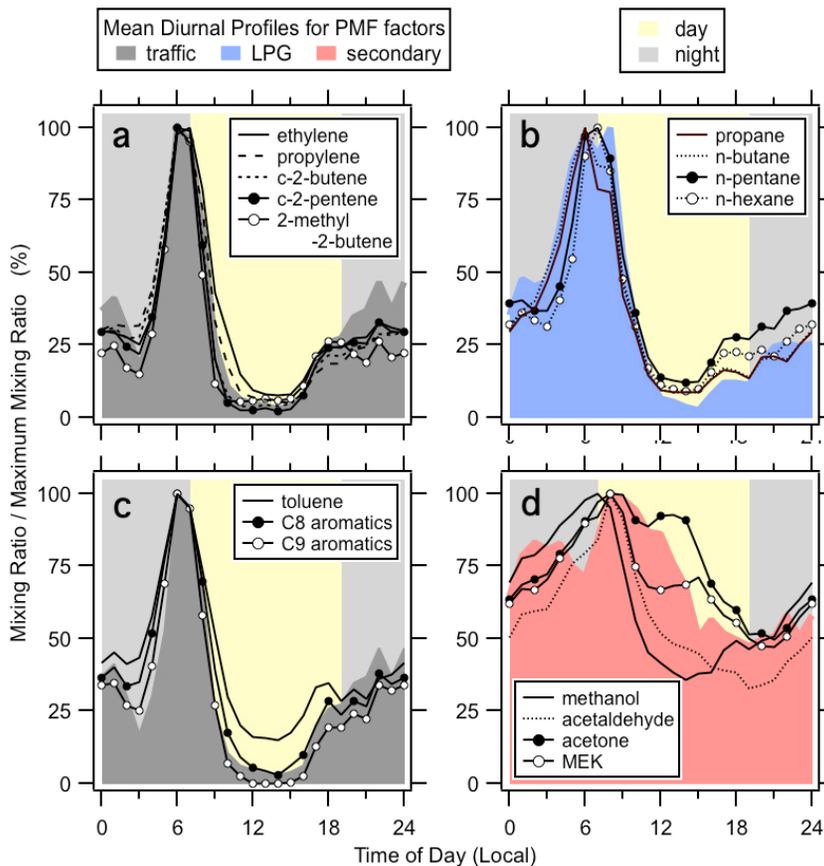


Fig. 5. Measured VOC profiles as a function of local time (UTC – 6h) normalized to the maximum of the mean mixing ratio for individual compounds within chemical families: alkenes (a), alkanes (b), aromatics (c) and OVOCs (d). Normalized mean profiles for PMF factors (FPEAK = 0) are shown as shaded areas for each chemical family.

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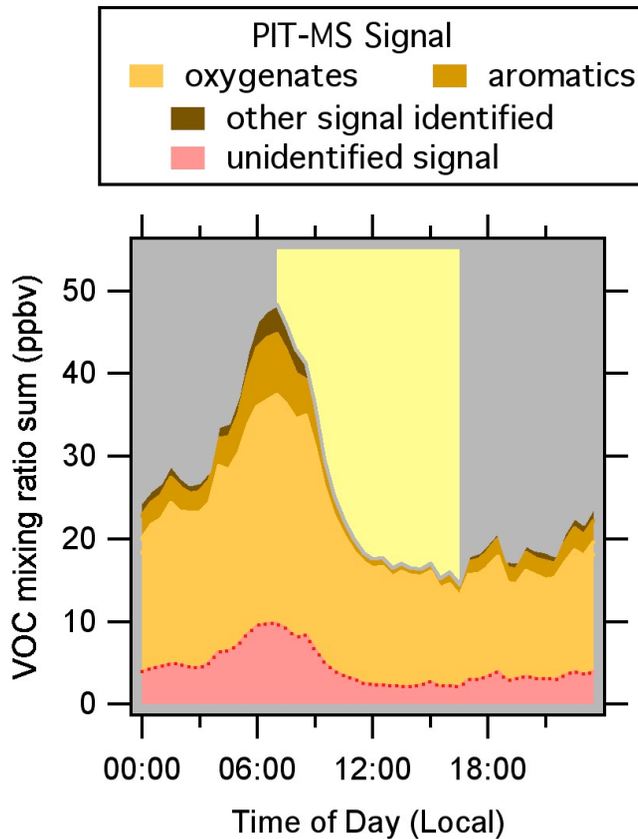


Fig. 6. Average fraction of total PIT-MS signal attributed to oxygenates, aromatics and other identified signal during the MILAGRO campaign. Unidentified signal averages about 18% of the total.

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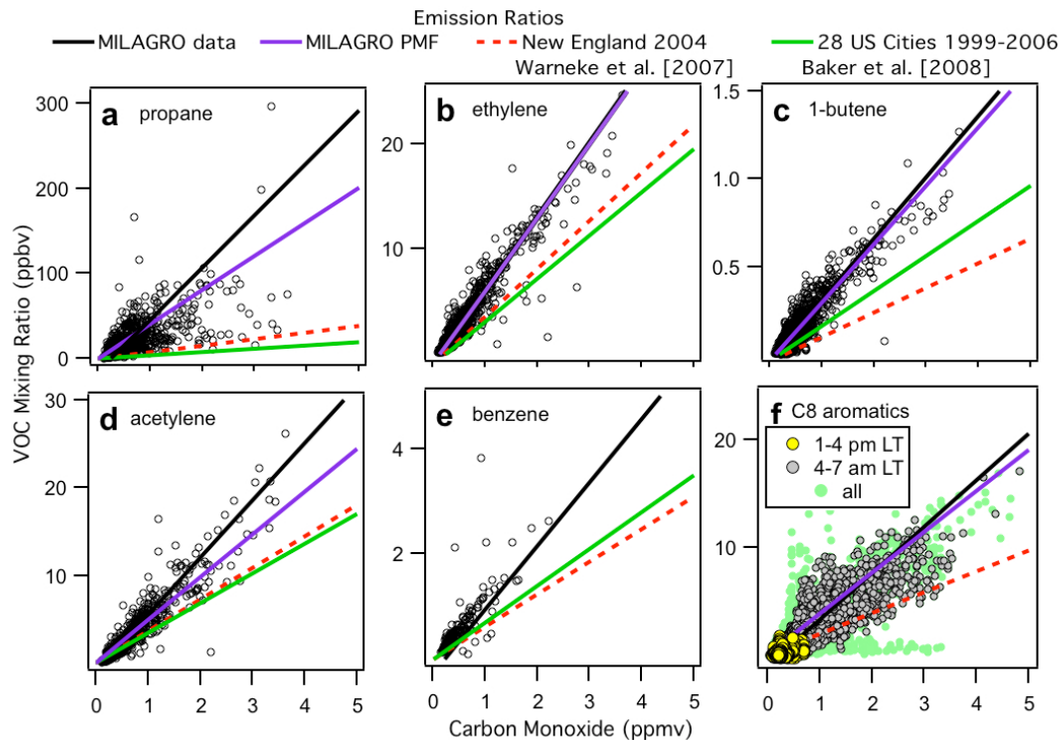


Fig. 7. Selected plots used to calculate urban emission ratios (ERs) with respect to carbon monoxide (CO) for VOC measurements made by GC-FID (**a–d**), canister samples (**e**) and PIT-MS (**f**). Urban ERs calculated from measurements using ODR fits (black) and Positive Matrix Factorization results (purple) during MILAGRO. MCMA ERs are compared to literature values (red, green) obtained for cities in the United States. PIT-MS measurements are shown on a 1-min time base (**f**) to highlight mixing ratio differences prior to local sunrise (grey) vs. those observed during the mid-afternoon (yellow). PMF lines represent ERs obtained by summing the traffic and LPG factors.

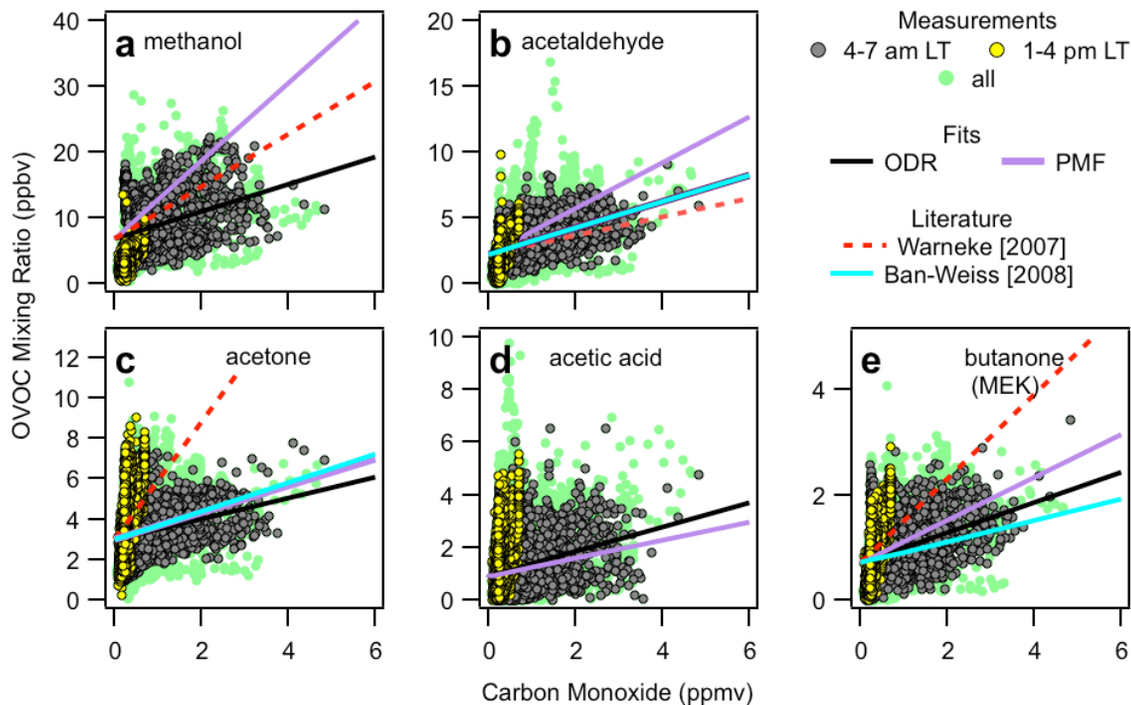


Fig. 8. Plots used to calculate emission ratios with respect to carbon monoxide (CO) for 5 OVOCs quantified by PIT-MS. Emission ratios calculated from measurements using 2-sided regression fits (black) and Positive Matrix Factorization results (purple) during MILAGRO are compared to literature values obtained during field studies in New England (red) and to a California tunnel study (blue). 1-min PIT-MS data show afternoon enhancements in OVOCs typically observed during the campaign. For comparison purposes, lines showing PMF and literature ERs are shown with intercepts obtained from ODR fits of MILAGRO measurements. PMF lines represent ERs obtained by summing the traffic and LPG factors.

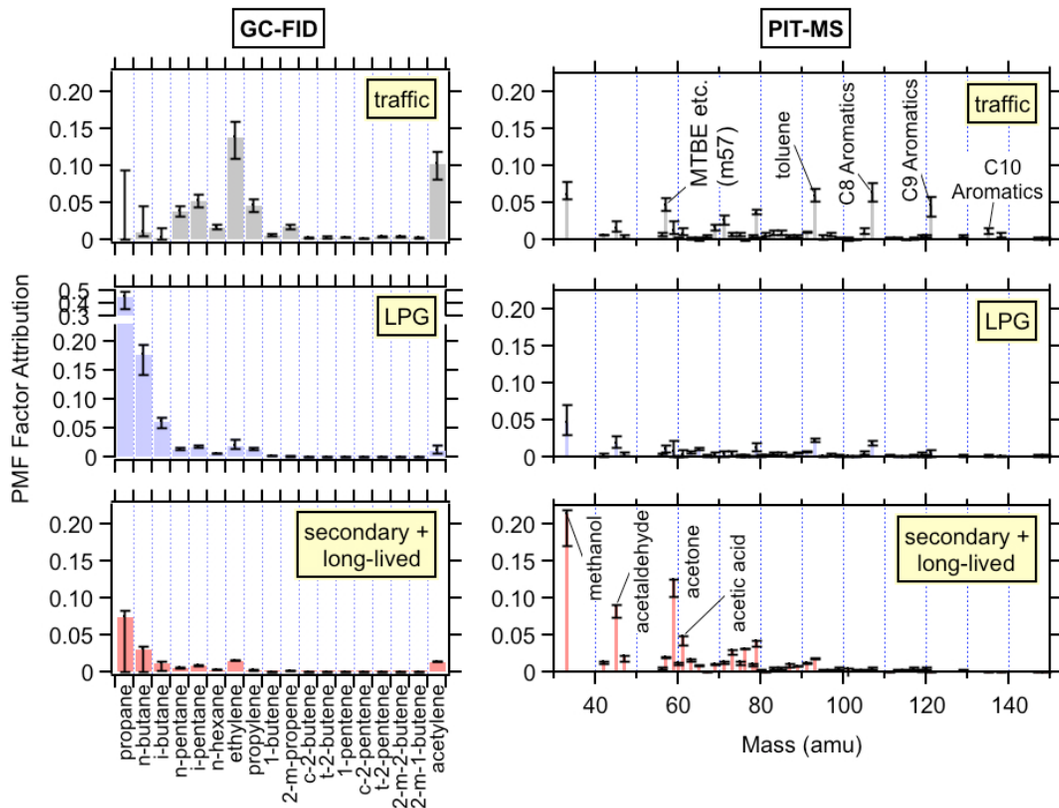


Fig. 9. PMF profiles are shown for compounds measured by the NOAA GC-FID (left column) and NOAA PIT-MS (right column) for the three factors, here labeled traffic, LPG, and secondary + long-lived species. Error bars represent the variation in PMF factors from the FPEAK = 0 solution (solid bars) over the FPEAK range from -3.0 to $+3.0$ as described in the text.

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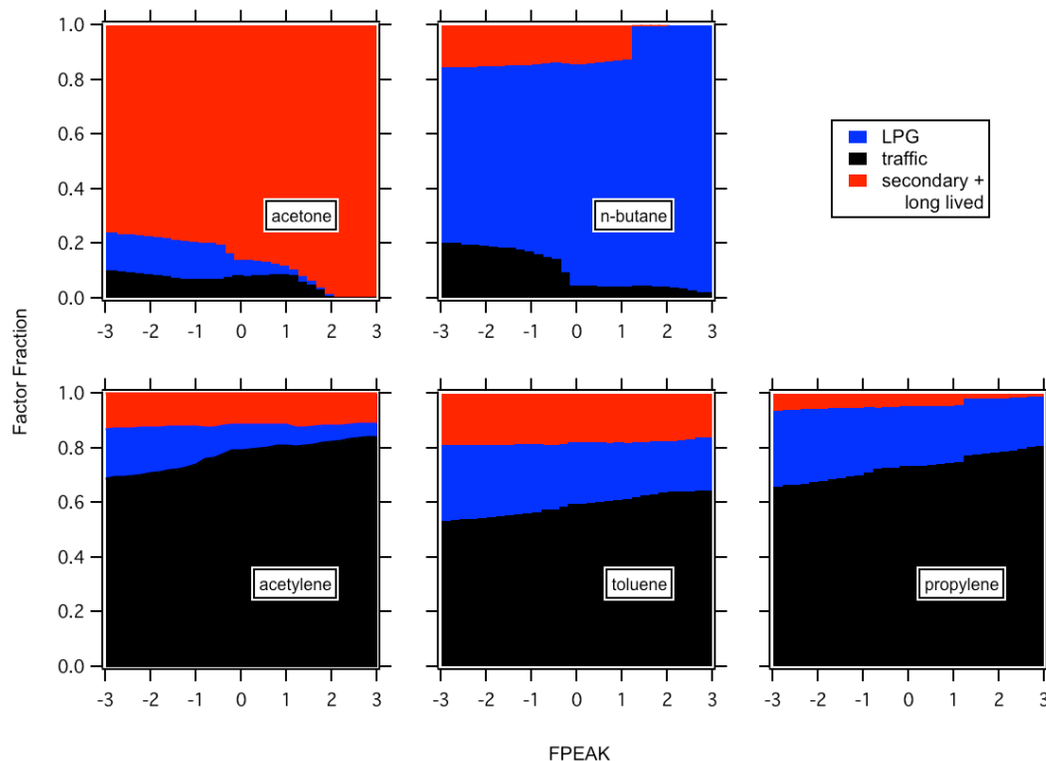


Fig. 10. Variability in apportionment to factors as a function of the PMF rotational parameter FPEAK for acetone, n-butane and three hydrocarbons dominated by traffic emissions (acetylene, toluene and propylene). The change in the PMF quality of fit parameter is less than 2% for the FPEAK values shown.

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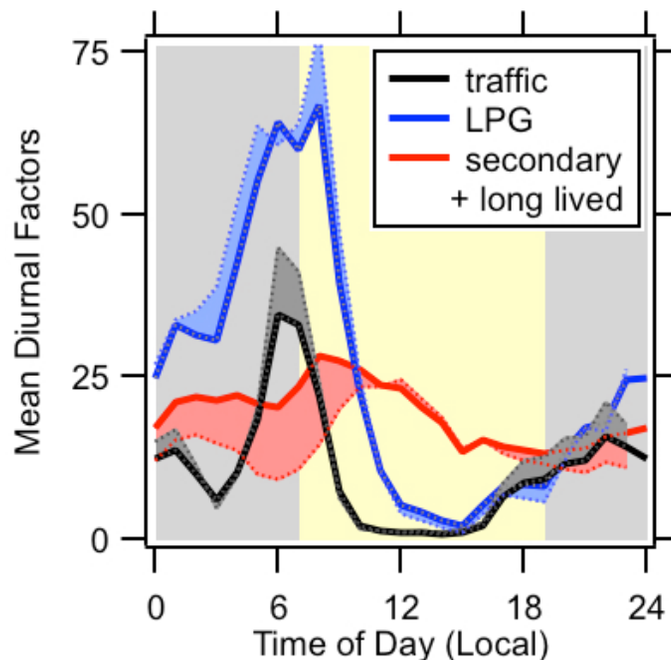


Fig. 11. Mean diurnally averaged profiles for the 3-factor PMF solution obtained from MILAGRO VOC measurements. Solid lines represent the mean diurnal profile for each factor at FPEAK 0 while the shaded regions bordered with dashes show the variation from the FPEAK = 0 solution over the FPEAK range from -3.0 to $+3.0$.

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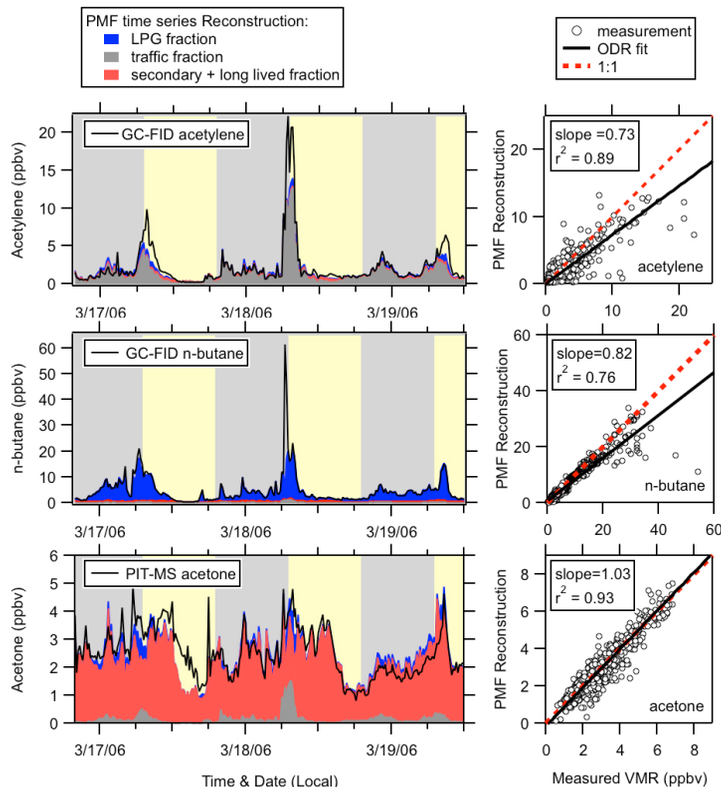


Fig. 12. Selected comparisons between measurements and PMF reconstructions for one period during the MILAGRO campaign. The left column shows time series for representative compounds with dominant apportionment to the traffic factor (acetylene), the LPG factor (n-butane) and the secondary + long-lived factor (acetone). Measurements are shown as solid black traces for comparison. On the right are scatter plots of PMF reconstructions vs. measurements showing single-sided fits (black) and 1:1 lines (red) for the same compounds.

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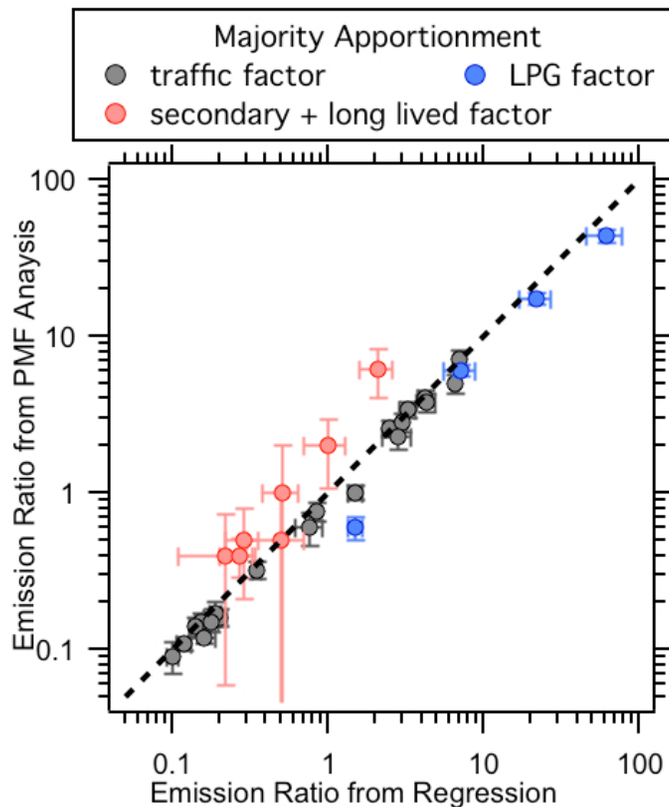


Fig. 13. Comparison of PMF emission ratio estimates obtained by linear regression for all measured compounds. Each point represents one compound while colors represent the dominant PMF apportionment for the respective measurement.

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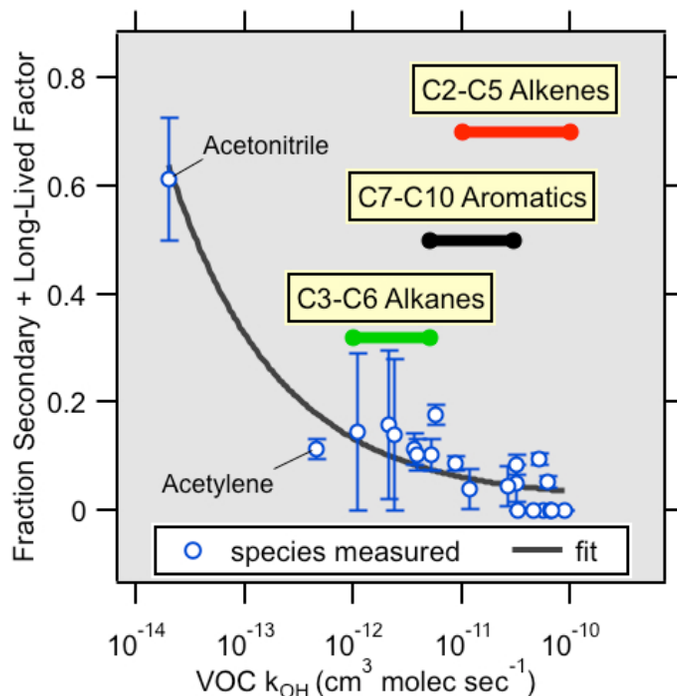


Fig. 14. PMF apportionment of measured compounds to the secondary + long-lived factor plotted vs. photochemical lifetime (k_{OH}) (Atkinson and Arey, 2003). Longer lifetimes are associated with higher secondary + long-lived source apportionment. Colored bars represent the range of lifetimes by compound class for alkanes (green), aromatics (black) and alkenes (red). Error bars represent variability in apportionment due to PMF FPEAK parameter from -3 to $+3$.

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