

**Trace gas and
particle emissions**

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Trace gas and particle emissions from open biomass burning in Mexico

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Abstract

We report airborne measurements of emission factors (EF) for trace gases and $PM_{2.5}$ made in southern Mexico in March of 2006 on 6 crop residue fires, 3 tropical dry forest fires, 8 savanna fires, 1 garbage fire, and 7 mountain pine-oak forest fires. The savanna fire EF were measured early in the local dry season and when compared to EF measured late in the African dry season they were at least 1.7 times larger for NO_x , NH_3 , H_2 , and most non-methane organic compounds. Our measurements suggest that urban deposition and high windspeed may also be associated with significantly elevated NO_x EF. When considering all fires sampled, the percentage of particles containing soot increased from 15 to 60% as the modified combustion efficiency increased from 0.88 to 0.98. We estimate that about 175 Tg of fuel was consumed by open burning of biomass and garbage and as biofuel (mainly wood cooking fires) in Mexico in 2006. Combining the fuel consumption estimates with our EF measurements suggests that the above combustion sources account for a large fraction of the reactive trace gases and more than 90% of the total primary, fine carbonaceous particles emitted by all combustion sources in Mexico.

1 Introduction

Biomass burning (BB) is the largest source of primary, fine carbonaceous particles and the second largest source of trace gases in the global atmosphere (Crutzen and Andreae, 1990; Bond et al., 2004). Industrial and domestic burning of biomass as a fuel (biofuel) occurs globally and year-round; roughly proportional to population, but with much higher use per capita in developing countries (Yevich and Logan, 2003). Open (outdoor) burning is even more widespread geographically, but it occurs mostly during the local dry season. The dry seasons of locations in the Southern Hemisphere (SH) tropics and the mid to high latitudes of the Northern Hemisphere (NH) can start as early as May–June and can last until October–November. The dry seasons in the NH

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tropics fall within the months of February–May; thus, large amounts of open burning emissions are being produced somewhere on Earth in nearly every month. Since most BB occurs in the SH tropics these fires have been studied the most (e.g. Sinha et al., 2003; Yokelson et al., 2008). Less is known about the emissions from the significant amounts of open BB that occur in the NH tropics (i.e. Mexico, Central America, northern South America, the Sahel region of Africa, India, and Indochina (van der Werf et al., 2010)).

As part of the MILAGRO (Megacity Initiative Local and Global Research Observations) project (Fast et al., 2007; Molina et al., 2010), 56 open landscape-scale fires were sampled throughout southern and central Mexico from the air by a USFS Twin Otter and the NCAR C-130 in March of 2006 (Yokelson et al., 2007a, 2009). In addition, 22 fires were sampled from the ground in central Mexico during April–May of 2007 to characterize domestic and industrial biofuel burning (e.g. cooking fires, kilns, etc), crop residue burning, and garbage burning (Christian et al., 2010). All these fires were sampled to meet a variety of objectives that includes: (1) determine the impact of the many different kinds of regional BB on the Mexico City Metropolitan Area (MCMA) and the BB contribution to the MCMA outflow, (2) better characterize the emissions from open and domestic BB in the NH tropics in general, and (3) learn more about the emissions from the significant amount of open biomass burning that occurs globally early in the dry season, but has been relatively undersampled (Hoffa et al., 1999; Korontzi et al., 2003).

Of the 56 fires sampled from the air, the airborne measurements of emission factors (EF) for 20 open fires in the Yucatan that were burning crop residue or tropical dry forest deforestation slash were described by Yokelson et al. (2009). That study also detailed measurements of the chemical evolution of one of the sampled plumes. An additional 8 fires sampled in the airborne work were understory fires in the urban-impacted, mountain pine-oak forests adjacent to the MCMA. These fires and their large impact on the MCMA outflow (accounting for roughly half the $PM_{2.5}$) were described by Yokelson et al. (2007a) and Crouse et al. (2009). The emission factors from the 2007

ground-based fire sampling and a limited assessment of the impact of burning biofuel and garbage in the MCMA, Mexico, and the developing world in general was published by Christian et al. (2010).

The purpose of this paper is to present unpublished results for 25 additional open fires sampled from the Twin Otter during the 2006 airborne campaign and integrate these new data with the previously published results by fire type where applicable. The new airborne data include emission factors for 6 crop residue fires, 3 tropical dry forest fires, 8 savanna fires, 1 dump fire (open burning of garbage), and 7 fires in mountain pine-oak forests that were not impacted by urban emissions. Thus, this paper completes the presentation of the emission factor measurements made during MILAGRO. Coupled with the earlier papers, emission factors are now available for most of the major species emitted by many of the types of burning that are important within Mexico; a country with a high diversity of ecosystems and fire types. While most BB research has focused on savanna and forest fires, in fact, many different fire types are significant in most developing countries (Crutzen and Andreae, 1990; Akagi et al., 2010). In addition, by comparing the EF from these open fires to published EF for fires in similar ecosystems that burned during the late dry season, we find evidence for possible seasonal trends. We briefly describe the location and amounts of the different types of burning and the initial transport of the fire emissions. We conclude with an estimate of the combined impact of open burning and biofuel use in Mexico, which may be relevant to many developing nations.

2 Experimental details

The instruments deployed on the USFS Twin Otter and all of the data-handling procedures were described in detail by Yokelson et al. (2007a). A brief summary is presented next.

The University of Montana airborne Fourier transform infrared spectrometer (AF-TIR) measured samples temporarily detained in the flow-through gas cell to quantify

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water vapor (H_2O), carbon dioxide (CO_2), carbon monoxide (CO), methane (CH_4), nitric oxide (NO), nitrogen dioxide (NO_2), ammonia (NH_3), hydrogen cyanide (HCN), ethene (C_2H_4), ethyne (C_2H_2), formaldehyde (HCHO), methanol (CH_3OH), acetic acid (CH_3COOH), formic acid (HCOOH), and ozone (O_3). The HCOOH emission factors for the fires first presented in this work are based on the new measurement of the HCOOH infrared cross-section now adopted in the HITRAN database (Rothman et al., 2009). The HCOOH emission factors taken from previously published work were rescaled so that all the HCOOH emission factors herein stem from one consistent, updated analysis.

Ram air was grab-sampled into 2-L stainless steel canisters for whole air sampling (WAS) and later analyzed at the University of Miami by gas chromatography (GC) with a flame ionization detector (FID) for CH_4 and the following non-methane hydrocarbons (NMHC): ethane, C_2H_4 , C_2H_2 , propane, propene, isobutane, n-butane, t-2 butene, 1-butene, isobutene, c-2-butene, 1,3-butadiene, cyclopentane, isopentane, and n-pentane. CO was measured in parallel with the CH_4 measurement, but utilized a GC with a Trace Analytical Reduction Gas Detector (RGD). Alternatively, at times, we collected 0.85-L canisters for later analysis at the United States Forest Service (USFS) Fire Sciences Laboratory by GC/FID/RGD for CO_2 , CO , CH_4 , H_2 , and several C_2 – C_3 hydrocarbons. Details of the canister analysis are given by Weinheimer et al. (1998), Flocke et al. (1999), and Hao et al. (1996).

The canister-filling inlet (large diameter, fast flow) also supplied sample air for a Radiance Research Model 903 integrating nephelometer that measured “dry” (inlet $\text{RH} < 20\%$) b_{scat} at 530 nm at 0.5 Hz. The b_{scat} measured in inverse meters at the nephelometer temperature and pressure was converted to b_{scat} at standard temperature and pressure (STP, 273 K, 1 atm) and then multiplied by $208\,800 \pm 11\,900 \mu\text{g sm}^{-2}$ to yield the mass of particles with aerodynamic diameter < 2.5 microns ($\text{PM}_{2.5}$) in $\mu\text{g sm}^{-3}$ of air, based on a gravimetric “calibration” similar to that described in Trent et al. (2000). Our conversion factor is equivalent to a mass scattering efficiency (MSE) of $4.8 \text{ sm}^2 \text{ g}^{-1}$. The MSE obtained in airborne studies of fresh smoke from other fuel types (e.g. Nance

et al., 1993) differ by up to ~20%, which we take as a rough estimate of the uncertainty in our $PM_{2.5}$ values.

An isokinetic particle inlet sampled fine particles with a diameter cut-off of a few microns. Particles of diameter < 1 micron account for nearly all the fine particle ($PM_{2.5}$) mass emitted by biomass fires (Reid et al., 2005). This inlet supplied sample air to two particle impactor samplers (MPS-3, California Measurements, Inc.) that were used to collect aerosol particles having aerodynamic diameters between 0.05 and 0.3 microns onto lacey-carbon transmission electron microscope (TEM) grids. Sampling times in fire plumes were < 1 min. Three TEMs were used for subsequent imaging and compositional analyses of these aerosol particles: a Tecnai F20 (FEI Corp.), a CM 200 (Philips Corp.), and a 2010F (JEOL) (Adachi and Buseck, 2008; 2010). They were operated at an accelerating voltage of 200 kV. Detailed sampling and analysis conditions are described by Adachi and Buseck (2008). The isokinetic particle inlet also supplied a LiCor (Model # 7000) measuring CO_2 and H_2O at 5 Hz and a UHSAS (Ultra High Sensitivity Aerosol Spectrometer, Particle Metrics, Inc.) deployed by the University of Colorado (CU). The UHSAS provided the number of particles in each of 99 user-selectable bins for diameters between 55 and 1000 nm at 1 Hz. All three Twin Otter inlets were located within 30 cm of each other. The nephelometer was not available on the 12 March flight so we used the UHSAS particle counting/size data to indirectly determine particle mass. As in previous studies (Yokelson et al., 2007a, 2009), we assumed all particles to be spheres with diameters determined from a Mie scattering response curve generated with polystyrene latex spheres (PSLs). These results are not corrected for assumed differences in index of refraction, which could undersize the particles by 6% or more in diameter for indices of refraction less than 1.5. The UHSAS only detects high albedo particles (e.g., black carbon is undersampled and, for this study, can be assumed to go largely undetected by the UHSAS). We integrated the UHSAS size distributions to obtain an estimate of the volume of particles (PV_1 , $\mu m^3 cm^{-3}$) in air at 1 Hz and found that the PV_1 (for $PV_1 < \sim 30$) was related to b_{scat}

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measured by the nephelometer as follows:

$$b_{\text{scat}} = PV_1 \times 1.25(\pm 0.25) \times 10^{-5} \quad (1)$$

On March 12, the PV_1 did not exceed $30 \mu\text{m}^3 \text{cm}^{-3}$ in the plume of Fire #3. We used Eq. (1) to convert PV_1 to b_{scat} and then converted b_{scat} to $\text{PM}_{2.5}$ as described above. The resulting $\text{PM}_{2.5}$ value has an estimated uncertainty of about 32%.

2.1 Generalized airborne sampling protocol

The Twin Otter was based in Veracruz with the other MILAGRO research aircraft (<http://mirage-mex.acd.ucar.edu/>). The main goal of the Twin Otter flights was to sample fires throughout south-central Mexico. Background air (i.e. ambient boundary layer (BL) air not in plumes) was characterized when not sampling BB plumes. The continuous instruments operated in real time in background air. The grab sampling instruments acquired numerous spot measurements that were representative of the background air since the continuous instruments showed that the background air was well-mixed on the spatial scale corresponding to the discrete sampling intervals.

To measure the initial emissions from the fires, the aircraft sampled smoke less than several minutes old by penetrating the column of smoke, up to six times per fire, at 150–600 m above the active flame front as terrain allowed. The continuous instruments monitored their species while penetrating the plume. The AFTIR, MPS-3, and WAS were used for spot measurements in the smoke plumes. For every grab sample of the smoke plume we also collected a paired, local, background grab sample just outside the plume. These paired samples allow calculation of the most accurate excess mixing ratios for the smoke (see next section).

2.2 Data processing and synthesis

Grab samples of both the plume and the adjacent background were used to calculate excess mixing ratios (ΔX , the mixing ratio of species “X” in the plume minus the mixing

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ratio of “X” in the background air). ΔX reflects the degree of dilution of the plume and the instrument response time. Thus, a useful, derived quantity is the normalized excess mixing ratio (NEMR) where ΔX is divided by the “simultaneously” measured excess mixing ratio of another species (ΔY); usually a fairly long-lived plume “tracer” such as ΔCO or ΔCO_2 . A measurement of $\Delta X/\Delta Y$ in a plume up to a few minutes old is a molar emission ratio (ER). We computed fire-average molar ER for each individual fire from grab or discrete samples as follows. First, if there is only one sample of a fire then the calculation is trivial and equivalent to the definition of $\Delta X/\Delta Y$ given above. For multiple grab samples of a fire, the fire-average ER was obtained from the slope of the least-squares line (with the intercept forced to zero) in a plot of one set of excess mixing ratios versus another. This method is justified in detail by Yokelson et al. (1999). When the AFTIR and WAS measured the same pair of compounds on the same fire, their data were combined in the plots as shown in Fig. 1a of Yokelson et al. (2009). The CO and CO₂ data from AFTIR and WAS were also used to calculate fire-average modified combustion efficiencies (MCE). MCE is defined as $\Delta \text{CO}_2/(\Delta \text{CO}_2 + \Delta \text{CO})$ and is also equal to $1/(1+(\Delta \text{CO}/\Delta \text{CO}_2))$. We use fire-average MCE in Sect. 3 as an index of the relative amount of fuel consumption by flaming and smoldering combustion, with high MCE indicating a fire with relatively more flaming (Akagi et al., 2010).

A few of the grab-sample based ER were from measurements that were not made on the same instrument. The molar ER to CO₂ for each NMHC measured by U-Miami WAS on the Twin Otter was derived for each fire as follows. The molar ER to CO measured by WAS from a fire was multiplied by the molar $\Delta \text{CO}/\Delta \text{CO}_2$ ER measured on that same fire by AFTIR. CO was measured with high accuracy by AFTIR and WAS. This facilitated coupling data from the two methods.

Emission ratios were obtained from the continuous instruments by comparing the integrals of ΔX and ΔY as the aircraft passed through a nascent smoke plume (e.g. PM_{2.5} from nephelometer and LiCor CO₂). While the instantaneous $\Delta X/\Delta Y$ would be affected by any difference in “real-time” instrument response times, comparing the integrals compensates for this (Karl et al., 2007). When only one pass was made through

the plume of a fire, the ratio of the integrals was taken as the ER. When more than one pass was made through the plume of a single fire, we plotted the integrals versus each other and obtained the ER from the slope; analogous to the grab sample plots. Figure 1 in Yokelson et al. (2009) illustrates the typical analyte levels we encountered and gives examples of ER derivations.

2.2.1 Estimation of fire-average emission factors

For any carbonaceous fuel, a set of ER to CO₂ for the other major carbon emissions (i.e. CO, CH₄, a suite of non-methane organic compounds (NMOC), particle carbon, etc.) can be used to calculate emission factors (EF, g compound emitted per kg dry fuel burned) for all the emissions quantified from the source using the carbon mass-balance method (Yokelson et al., 1999). The carbon mass balance method assumes that all the carbon released to the atmosphere is detected. We approximated total carbon emissions for our EF calculation from AFTIR measurements of CO₂, CO, CH₄, and NMOC and WAS measurements of CO₂, CO, CH₄, and non-methane hydrocarbons (NMHC, a subset of NMOC consisting of compounds containing only C and H and not O, N, etc). The nephelometer PM_{2.5} and measurements made on the C-130 during MILAGRO of the mass fraction of C in BB aerosol (0.48–0.68; Yokelson et al., 2009) were used to estimate the particulate carbon. By ignoring unmeasured gases, we may overestimate the emission factors by 1–2% (Andreae and Merlet, 2001; Akagi et al., 2010). We assumed that all the fires burned in fuels that were 50% C by mass on a dry weight basis (Susott et al., 1996), but the actual fuel carbon percentage could vary by ±10% (2σ) of our nominal value. EF scale linearly with the assumed fuel carbon fraction. Because much of the NO is quickly converted to NO₂ after emission, we also report an EF for “NO_x as NO.” For any species “X” we abbreviate the EF as EFX.

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2.3 Details of flights

For the 28 fires sampled from the air on 6, 9, 12, 17, 18, 22, 23, and 29 of March 2006; the flight paths, fire locations, sampling times, MODIS active fire (“hotspot”) detection rate, flight narrative, and other details were included in previous papers (Yokelson et al., 2007a, 2009). The airborne data first presented in this work were gathered on 4, 8, 11, 20, 25, 27, and 28 of March 2006 during flight missions to the Mexican states of Veracruz, Veracruz and Puebla, Tamaulipas and San Luis Potosi, Chiapas, Oaxaca, Guerrero and Mexico, and Michoacán, respectively. The fire locations for these dates and all the other Mexican airborne fire sampling are shown in Figure 1 along with the six open fires sampled from the ground. Table 1 presents the fire date, identifying number, location, sampling time, hot spot detection information, and a simple fuel description for all the March 2006 airborne fire sampling in Mexico. Also shown in Table 1 is basic information for the 6 open fires sampled from the ground during the spring of 2007 (4 garbage burning and 2 crop residue) with more details available in Christian et al. (2010).

3 Results and discussion

3.1 Emission factors for types of open burning common in Mexico

3.1.1 Crop residue fires

Table 2 shows all the trace gas EF measured for crop residue (CR) fires in Mexico as part of MILAGRO. It includes EF for 6 CR fires measured from the air during March 2006 in the Yucatan published in Yokelson et al. (2009) and EF for 2 CR fires measured from the ground in central Mexico during March 2007 published in Christian et al. (2010). Table 2 also includes previously unpublished EF for 6 CR fires measured from the air during March 2006 throughout south/central Mexico as shown in Table 1.

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We were not able to identify the specific type of crop associated with these fires from the air, but all the CR fires reported in Table 2 were of the type where loosely-packed residue burned mostly by flaming combustion in the field. This is typical of mechanized agriculture and contrasts with much of the crop residue burning in eastern Asia where piles of hand-processed residue burn mostly by smoldering (Christian et al., 2003).

The data in Table 2 can be used to calculate a straight average and standard deviation for the newly-presented CR fires or the Yucatan CR fires separately, with the latter already shown in Yokelson et al. (2009). We found that these average values for the two groups of CR fires were qualitatively similar for all species except acetic acid. Thus, in Table 2 we compute “national average” EF for all the crop residue fires in Mexico. These values apply to a larger geographic region and are based on a larger sample size.

We use two different methods to compute the average values as discussed in detail next. The average fire-integrated MCE is an important parameter to estimate accurately because it reflects the “typical” flaming to smoldering ratio, which has a major influence on nearly all the EF. We make the assumption that our estimate of the average MCE will be more representative if the sample size is larger. Thus, since we measured CO₂ and CO in every fire plume, we take the average MCE for all the CR fires we sampled as our best estimate of the average MCE for all Mexican CR fires. We recognize that our samples are biased temporally (early dry season), geographically (Yucatan and south-central Mexico), and methodologically (12 air versus 2 ground where the latter tends to observe lower MCE). On the other hand we have no valid basis for improving our estimate by excluding certain fires. Thus, we conclude that the average MCE of all our CR fires currently represents our best estimate of the average mix of flaming and smoldering combustion associated with Mexican CR fires.

Because of the strong EF dependence on MCE we need to derive average EF that are consistent with the average MCE based on all the fires. However, for various technical reasons (see Sect. 2), we did not acquire data for every species on every fire. The situation is illustrated in Fig. 2 using 1,3-butadiene, which was measured on only 6

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of the 14 CR fires, as an example. For the six fires with 1,3-butadiene data the average EF was 0.114 g kg^{-1} (open square in Fig. 2) and the average MCE was 0.940. Thus the straight average EF is appropriate for an MCE of 0.940 rather than the average MCE of 0.925 for all CR fires. To derive an EF that is consistent with the average CR fire MCE, we fit a regression line to EF versus MCE using the 6 available measurements and then use the regression equation to calculate an EF at the average MCE for CR fires. This procedure yields an EF of 0.151 g kg^{-1} (filled square Fig. 2), which is consistent with the more extensive MCE information. Thus, for all the species other than CO and CO₂, we plotted the available EFs versus MCE, fit a regression line to those data, and then computed a recommended EF from the fit at the average MCE for all the CR fires (shown in the last column of Table 2). This procedure gives the “straight average” EF when the species was measured on all the fires and an EF that is consistent with the average MCE when a species was measured only on some of the fires. In summary, we recommend using the EF computed at the average MCE although in four cases in this study the single EF measurement made for a species for a fire type has to suffice.

The EF for total PM_{2.5} was measured on many of the CR fires and these EF are also shown in Table 2 along with a recommended average EF_{PM_{2.5}} computed at the average MCE. Particle chemistry details for the CR fires sampled from the ground are shown in Table 4 of Christian et al. (2010) (e.g. EF for metals, elemental carbon (EC), organic carbon (OC), and several anhydrosugars of potential value as smoke markers). In addition, particle chemistry details (major inorganic species, organic aerosol, and black carbon) are shown in Table 3 of Yokelson et al. (2009) for fires #1 and #3 sampled on 23 March and those 2 fires were very likely CR fires.

In this paper we discuss three different methods of estimating the mass of sp²-bonded carbon clusters in the particles, which are responsible for most of the aerosol visible light absorption. Black carbon (BC) emission factors are based on direct optical absorption measurements combined with a mass absorption efficiency. Elemental carbon emission factors are based on thermo-optical, gravimetric measurements that isolate the mass of sp²-hybridized carbon. Soot is identified by its fractal shape and

sp²-carbon bond peak in the electron energy-loss spectra (EELS) in the TEM and is believed to account for most, if not all, black carbon. In this paper we take the EF for EC and BC as equivalent, although this is not always strictly the case. We also use the number fraction of soot particles as a qualitative estimate of the aerosol's tendency to absorb visible light and thereby impact radiative forcing. Detailed discussion of these techniques can be found elsewhere (Reid et al., 2005; Bond and Bergstrom, 2006).

3.1.2 Tropical dry forest (TDF) fires

In Table 3 we show the EF for 6 tropical dry forest deforestation fires sampled from the air in the Yucatan lowlands that were already presented in Yokelson et al. (2009) and the EF for 3 additional tropical dry forest fires that were sampled in more mountainous areas of southern Mexico (see Table 1 or Fig. 1). We have combined all the data and implemented the EF versus MCE regression based approach described above to generate a set of best estimate EF for Mexican tropical dry forest (TDF) fires. In two cases there was only one EF measurement for a species and we show that EF in the “EF at average MCE” column also so that all our best values are together in one column for convenience. The EF published previously for the TDF fires in the Yucatan (Yokelson et al., 2009) and the “all-Mexico” values are very similar, in part because the Yucatan fires account for most of the combined TDF data. The new TDF data now include EF for ethyne and formic acid, which were not available from the Yucatan data. However, these EF are based on only one fire in each case. While most of the national average EF are similar to the Yucatan EF, the national average EF is much higher for NH₃ driven by the large EFNH₃ obtained for the two TDF fires sampled on 11 March 2006 (Table 3). These two TDF fires differed from the seven other TDF fires we sampled in that they were burning understory vegetation in an un-slashed forest. Based on our airborne observations both small deforestation and small understory fires are common in Mexico and nearly all BB emission models (e.g. FINNv1, see Sect. 3.3) are based on moderate resolution remote sensing data or vegetation maps that cannot differentiate

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between small fires of these two types. Thus, including EF from both types in the national average is appropriate. However, a straight average of all our TDF fire samples is equivalent to assuming that ~22% of TDF fires are understory fires and the actual relative frequency of these two fire types at the national scale could differ from what we observed during our one-month airborne search for fires. If the larger EFNH₃ for understory fires was due to higher N content in the understory fuels, we would expect a similar increase in EFNO_x for understory fires (Burling et al., 2010). However, NO_x was below the detection limits for the TDF understory fires we sampled. A lower NO_x/NH₃ ratio is sometimes associated with lower MCE (Goode et al., 2000; McMeeking et al., 2009), but the average MCE for the understory fires (0.926 ± 0.014) was not significantly different from the average MCE for the deforestation fires (0.924 ± 0.016). Finally, we note that Fire #2 sampled on 23 March of 2006 by the NCAR C-130 was probably a TDF deforestation fire and the emissions of several trace gas and particle species from that fire that were not measured on the Twin Otter are reported in Table 3 of Yokelson et al. (2009).

In addition to tropical dry forest, Mexico has remnants of a formerly larger tropical evergreen forest on each coast (Hughes et al., 2000; Jaramillo et al., 2003). We did not locate or sample any fires in the Mexican tropical evergreen forest during our mission. However, Yokelson et al. (2009) noted that the EF they measured for tropical dry forest fires in Mexico are very similar to the EF measured for tropical evergreen forest fires in Brazil (Ferek et al 1998; Yokelson et al., 2008). Thus, we assume that the EF we report here for Mexican tropical dry forest fires are applicable to fires in Mexican tropical evergreen forests.

3.1.3 Mixed crop residue and tropical dry forest fires

During the March 2006 airborne fire sampling in the Yucatan it was often observed that high surface windspeeds contributed to crop residue fires escaping into the adjacent tropical dry forest where they burned mostly understory fuels. The results for the fires associated with this scenario were included in Table 2 of Yokelson et al. (2009).

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In Table 4 we now present these fires as their own category and derive average EF values using the EF versus MCE regression based procedure described above. One might expect that the emissions from these mixed CR/TDF fires would be intermediate between the CR and TDF types. This is observed for a few compounds like methane and ethane, but not for many others. For instance, the mixed fuel fire type has significantly larger EF for NO_x and acetic acid and significantly lower EF for NH_3 than either component fire type. The lower EF for NH_3 is surprising since the largest EFNH_3 of the TDF fires were from the 11 March TDF understory fires, which burned nominally similar understory fuels as in the forest on the margins of the crop residue fires. The increase in EFNO_x by more than a factor of two would also not be predicted by combining contributions from the two fuel types involved, especially since NO_x was below detection limits for the 11 March TDF understory fires. The three fire types had almost identical MCE and there was no seasonal trend in the EF for N-species during our month of sampling. Thus, it is possible that factors other than “fuel type”, MCE, or fuel moisture; such as surface windspeed, may contribute significantly to the mix of emissions produced in open burning. Additional, targeted research would be required to quantitatively estimate any connection between surface windspeed and fire emissions, but if a useful correlation were found it should be feasible to incorporate it in models since the windspeed data is widely available and already commonly used to model sea salt emissions (e.g. Gong et al., 1997).

3.1.4 Savanna fires

We sampled 8 savanna fires from the air during March 2006 in the Mexican states of Chiapas and Guerrero that have not been previously published and the EF are shown in Table 5. We derived study-average EF values at the average MCE as described above. These March 2006 fires occurred during the early part of the dry season and may have burned in fuels that had a higher moisture and nitrogen content than is common during the late dry season. Thus, we compare our Mexican early dry season savanna fire EF to those obtained relatively later in the dry season on African savanna fires using

similar instrumentation and/or sampling methods (Yokelson et al., 2003; Sinha et al., 2003). Table 5 shows that our “early dry” central EF values are 1.7 (or more) times higher for both NO_x and nearly all the observed smoldering species, namely H_2 , NH_3 , CH_3OH , CH_3COOH , C_2H_6 , C_3H_6 , and CH_4 . The early dry MCE is lower so we expect a higher $\text{EFPM}_{2.5}$ (Yokelson et al., 2007a). However, our early dry season $\text{EFPM}_{2.5}$ of $7.6 \pm 2.5 \text{ g kg}^{-1}$ is below the African late dry season EFPM_4 of $10.0 \pm 7.5 \text{ g kg}^{-1}$ (Sinha et al., 2003). If we take the average of the African late dry season value with the Brazilian late dry season savanna fire EFPM_4 of $3.86 \pm 0.36 \text{ g kg}^{-1}$ (Akagi et al., 2010) we obtain $\sim 6.93 \text{ g kg}^{-1}$, which is close to our early dry season value of $\sim 7.6 \text{ g kg}^{-1}$.

HCHO was the only NMOC we observed to have a much lower EF for the early dry season savanna fires, but the decrease is based on just two EF in the present study. EFHCN did not change, indicating that HCN might arise from a non-seasonally varying nitrogen containing compound in the biomass. We note that the seasonal comparison for savanna fires contrasts the tropical forest fire seasonal comparison in Yokelson et al. (2009) where early dry season forest fires had EF for trace gases that were similar to late season forest fires, but EF for $\text{PM}_{2.5}$ that were much lower than for late season forest fires. The lower $\text{EFPM}_{2.5}$ for early dry season forest fires could be related to reduced fire intensity (Yokelson et al., 2009; Reid et al., 1998 and references therein).

In a previous study of seasonal effects on savanna fire emissions, Korontzi et al. (2003) also reported higher EF for fires burning early in the dry season (in Africa) for a few smoldering compounds. Those authors found that the early and late dry season EF both followed one trend with MCE. The difference in EF in their case could be explained by a difference in MCE. In contrast, we find that the EF for early season fires lie well above the trend established by late dry season fires for several compounds (CH_4 , CH_3OH , and NO_x) as shown in Fig. 3. This latter finding could be at least partly due to differences between the fuels in African and Mexican savannas. In summary, so far all observations are consistent with a scenario in which early dry season savanna fires have lower MCE and higher EF for NH_3 , NO_x , and the NMOC produced by smoldering. The larger emissions of NO_x and NMOC could potentially impact the formation of

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secondary organic aerosol (Grieshop et al., 2009; Alvarado and Prinn, 2009) and O₃ (Trentmann et al., 2005).

3.1.5 Pine-oak forest fires

In Table 6 we present the EF measured during March 2006 for 7 pine-oak forest fires in remote mountain locations. Most of these fires were understory fires like those in the MCMA area previously described by Yokelson et al. (2007a), but a few in Chiapas were part of land-clearing operations and therefore, the fuel for those fires included relatively more large-diameter wood. We derive EF values at the average MCE for these fires as a group. For comparison, column 12 (Table 6) shows the average EF measured from the air for the 8 mountain pine-oak fires sampled near the MCMA by Yokelson et al. (2007a). Yokelson et al. (2007a) observed significantly larger emissions of NO_x and HCN (but not NH₃) than what was produced during laboratory experiments burning fuels collected in pine forests located in areas of the US not impacted by urban emissions. They speculated that this difference could be due to nitrogen deposition from the MCMA urban plume. Here, we directly compare the emission factors for the MCMA-impacted pine-oak forest fires to our emission factors for rural pine-oak forest fires in Mexico. We find that the NO_x and NH₃ EF are enhanced in the urban-impacted forests by factors of 2 and 1.7, respectively. However, HCN is not. If we calculate average EF for the rural pine-oak fires that do not include the pine-oak fires in Chiapas (20 March, fires # 1, 2, and 3) where logs may have been a larger component of the fuel, the urban/rural comparison is essentially the same except that the NH₃ enhancement for urban-impacted fire emissions increases. Thus, in these three comparisons, the EFNO_x is consistently elevated by about a factor of two for the urban/deposition impacted forests, but the results for other N-containing species are variable and somewhat inconclusive at this point. As noted above, enhanced NO_x emissions could impact downwind ozone and aerosol formation.

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3.1.6 Garbage burning

Open burning of garbage is an important, but poorly quantified source of emissions in developing countries and in some rural areas of developed countries (Christian et al., 2010). During the 2006 airborne campaign we were able to acquire one sample of the emissions from a small dump fire in Cosolapa, a small rural village in Oaxaca state located ~80 km SW of Veracruz. Christian et al. (2010) reported EF for 4 garbage burning (GB) fires that were sampled extensively from the ground in the northern MCMA during April-May of 2007. In Table 7 we show all the MILAGRO garbage burning EF along with previously published EF from Lemieux et al. (2000) and recommended EF from AP-42 (USEPA, 1995). In addition to CO₂ and CO (used to compute MCE), there were three other species measured in both our airborne sample and the ground-based samples from the MCMA. The emission factors for these species (C₂H₄, CH₄, and NH₃) are plotted versus MCE in Fig. 4, which shows that a single, highly-correlated trend couples the airborne and ground-based measurements. However, with only 5 GB fires sampled our average MCE for open burning of garbage is less constrained than for the other fire types. In addition, for almost a quarter of the species measured, we have only one EF, which precludes a regression of EF vs MCE for those species. Thus, in Table 7 we simply present the average of all our Mexican measurements and also the limited EF data for open burning of garbage from other sources. Additional details and discussion of the chemistry of the particulate emissions from garbage burning can be found in Christian et al. (2010) and in Sect. 3.2, but clearly much more work is needed to determine both the amount of garbage burned and the nature of the emissions.

3.2 TEM analyses of particle chemistry

Some key results of the TEM analyses of our Mexican BB samples have already been reported. The particles emitted by mountain pine-oak forest fires near the MCMA were shown to be internally mixed and have high N content and other properties discussed in Yokelson et al. (2007a). Particles in a plume emitted by a crop residue fire in the

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Yucatan had more sulfur and tarballs (spherical organic particles that are assigned a separate category in TEM analyses) and less chloride after that plume had aged for 10–30 min compared to fresh samples of the plume (Yokelson et al., 2009). The compositions, hygroscopicity, and abundance of tarballs were determined by Adachi and Buseck (2011). Here we report additional findings that summarize the body of our 2006 airborne BB TEM samples and also discuss data from two TEM samples containing garbage-burning emissions.

The composition of ~900 individual, non-volatile aerosol particles from 28 BB samples was analyzed using energy-dispersive X-ray spectrometry (EDS). Most particles consisted of organic aerosol (OA), tarballs, soot, sulfate, potassium chloride, or mineral dust, and many were internally mixed. Using particle shape as the primary criterion, the distribution of these major species was determined by counting their occurrence in 300 particles from each of 70 samples (collected from 34 fires). On average, $27 \pm 21\%$ of particles consisted of soot (Fig. 5). As is typical of soot, those particles consist of tens to hundreds of aggregated spherules that contain poorly ordered, curved, graphitic layers. They absorb sunlight and are significant contributors to climate change (Jacobson, 2001; Chung and Seinfeld, 2002; Adachi et al., 2010). The percentage of particles in a sample that were classified as soot varied even among samples collected from the same fire, reflecting the changing smoke composition that can be emitted by a single fire. The percentage of particles in a sample that were classified as soot increased approximately 4-fold from 15 to 60% as the sample MCE increased from 0.88 to 0.98 (Fig. 6a). This trend is consistent with previous findings that flaming combustion produces the soot (or black or elemental carbon) produced by fires (e.g. Yokelson et al., 1997; Christian et al., 2003; Reid et al., 2005). Conversely, the fraction of OA particles decreased from about 75 to 30% over the same MCE range, confirming that smoldering fires produce aerosol with relatively more organic carbon (Fig. 6b).

In addition to the BB samples, two TEM samples that contained smoke from garbage burning were collected. The samples were collected several minutes apart, with the first being pure GB emissions and the second containing the mixed emissions from

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a crop residue fire and the diluted GB plume. TEM-EDS analysis indicates that their compositions differ (Fig. 7). Sample A in Fig. 7 is of GB and mainly consists of particles containing (a) Mg and Ca; (b) Cr, Fe, and Ni; (c) Al; and (d) Si. There are few soot and OA particles. The filter analyses in Table 4 of Christian et al. (2010) also show that garbage burning produced higher levels of Mg and Ca than the other types of burning investigated, but Cr, Fe, and Ni were not elevated and Al and Si were not measured.

Sample B, which was mostly from a CR fire, but also contained some aged GB emissions due to plume mixing, mainly consists of soot, OA, particles containing Mg and Ca, and their mixtures. Thus, Mg and Ca appear to be consistently associated with GB. The glass commonly used for bottles and windows is about 5.7% Ca and also contains much Si and some Al as a common impurity. Magnesium (the lightest metal) is widely used in manufacturing, often as Mg/Al alloys. About half of the particles in Sample B include Cl. This is consistent with the high Cl content of particles from garbage burning and crop residue fires observed by Christian et al. (2010), which they suggested is likely due to the combustion of polyvinyl chloride and agricultural chemicals, respectively. Some particles are spherical (e.g., aggregated particles in the upper left of Fig. 7b), suggesting that they had been molten and cooled rapidly after emission. This process is similar to the production of coal fly ash; however, these particles could come from other combustion sources.

3.3 The amount and distribution of open burning in Mexico derived from space-based active fire detection

The *Fire Inventory* from NCAR version 1 (FINNv1) model calculates the global biomass consumption due to open burning based on active fire detection with daily temporal resolution and 1 km² spatial resolution (Wiedinmyer et al., 2010). In Fig. 8 we show the 2006 total combusted biomass (estimated by FINNv1) for each Mexican state divided by the state area. A continuous arc with high rates of BB stretches across Mexico to the south of the MCMA for ~2000 km; from the Mexican states of Quintana Roo on the Yucatan peninsula in the east to Jalisco on the west coast. The MCMA itself is also

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a small active zone, potentially because the large population leads to a high rate of accidental fires (E. Alvarado, personal communication, 2009). In Table 8 we show the total biomass combusted in open fires in 2006 in Mexico partitioned into a vegetation classification corresponding to our EF in Tables 2–6. The three most important types of open vegetation burning in the FINNV1 estimate are tropical forest (63.6 Tg), savannas (17.0 Tg), and temperate forest (12.8 Tg). We note that the crop residue fire estimate (2.9 Tg) is the category most likely to be underestimated because the small size and short duration of crop residue fires often leads to a low detection efficiency (Hawbaker et al., 2008; Yokelson et al., 2009 and references therein).

For comparison, Table 8 also includes an estimate for the sum of industrial and domestic biofuel use from Table 7 of Christian et al. (2010) and a crude estimate for garbage burning. The biofuel use estimate of 68 Tg yr^{-1} is the largest annual fuel consumption for the categories considered here. It should be noted that the estimate assumes that growth rates in biofuel use measured in past decades have continued through to the present (Yevich and Logan, 2003). Our rough estimate of the total amount of GB in Mexico is made as follows. Assuming that 0.9 kg of garbage is produced per day per capita in Mexico, based on the lower limit for residents in Mexico City (Ojeda-Benitz et al., 2008), we first calculate that 36.9 Tg of garbage is generated annually by Mexico's population of 112 468 855 (<https://www.cia.gov/library/publications/the-world-factbook/geos/mx.html>). The total annual amount of garbage burned in Mexico depends on the percentage assumed to burn: a number which is not known. McCulloch et al. (1999) estimated that 50% of garbage generated globally was burned, but that percentage included garbage burned in modern incineration facilities in addition to open burning. Assuming a range from 10–50%, the total amount of garbage burned in Mexico is $3.7\text{--}18.5 \text{ Tg yr}^{-1}$. Garbage contains a lot of biomass, but also a lot of plastic and other material (Christian et al., 2010 and references therein). Together BB and GB are estimated to account for roughly 175 Tg of fuel consumption in 2006 in Mexico and based on total hotspots, 2006 was a typical year for open burning in Mexico as shown in Supplement

Fig. 1 (MODIS hotspots for 2003–2010 were obtained from the Mexican Comisión Nacional para el Conocimiento y Uso de la Biodiversidad (CONABIO) website at: http://www.conabio.gob.mx/conocimiento/puntos_calor/doctos/puntos_calor.html, hereinafter cited as “CONABIO” or from FINNV1). The impacts of these emissions are discussed further in Sect. 3.5.

3.4 Airborne search for fires and regional transport of open burning emissions

The distribution and amount of open burning is typically retrieved from space-based detection of burned area or hotspots (Wiedinmyer et al., 2006; Reid et al., 2009; Giglio et al., 2010; van der Werf et al., 2010; Wiedinmyer et al., 2010) and an example of that approach (FINNV1) is described in the previous section. Given the inherent limitations of space-based sensors, it is of interest that our effort to locate fires for emissions sampling provides partially independent information on what burns in Mexico. We followed a simple strategy to locate fires for emissions sampling. If an area had MODIS hotspots (CONABIO) on a particular day and the forecast was for dry weather, it was classified as a location with high potential for fires for the next day’s flight. We attempted to sample fires in all areas of south-central Mexico that had exhibited high numbers of hotspots since 2003 (CONABIO). We sampled fires in all such areas except the valley of the Usumacinta River along the border of Chiapas with the Yucatan peninsula. Significant hotspot occurrence in this area did not begin until April 2006.

Following the above strategy we located more than 60 fires during our airborne search and 56 of these fires were large enough that we could acquire high quality emissions data from the aircraft (Table 1). For each of those 56 fires we checked if the fire had registered as a hotspot in the MODIS active fire product (Justice et al., 2002; CONABIO). A fire was considered to have been detected if there was a hotspot within 2 km of its location from either the Terra or Aqua daytime overpasses. Thus, in some cases the hotspot match could be due to a different fire at nearly the same location, but at a different time. The 2 km threshold was selected to account for the MODIS nadir spatial resolution (1 km), the MODIS point response function, and possible uncertainty

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due to evolution in the spatial distribution of the fire. Despite the fact that we used hotspots to guide our search for fires, only 10 of the 56 fires shown in Table 1 were detected as hotspots. Factors influencing the detection rate are discussed elsewhere (Yokelson et al., 2007a, 2009; Wiedinmyer et al., 2010). Comparison of Figs. 1 and 8 shows that there is good correlation between where we successfully located fires and where the FINNv1 model places the highest BB rates, but systematic measurement of the detection rate for various fire types would be valuable.

Taking the fires we located in the airborne search as one estimate of the distribution of Mexican BB, we explored the impacts of this open burning by running HYSPLIT 5-day forward trajectories (Draxler and Rolph, 2010) from all the open fires we sampled in 2006 and 2007. The starting altitudes were chosen to reflect the typically observed low injection altitudes (all within the boundary layer). Since biomass burning usually peaks in April and May in Mexico (CONABIO); for a few March 2006 fires we also ran forward trajectories starting at the same time of day, but during the months of April and May. The dispersion (not shown) was very similar. We limit the discussion here to potential air quality impacts on the Mexico City metropolitan area (MCMA) since it is Mexico's main population center. During a 2003 experiment focused on MCMA air quality, forward trajectories showed that fires in the Yucatan/Chiapas border region affected the MCMA (Molina et al., 2007). Our trajectories indicate that in 2006, pine-oak forest fires in the mountains adjacent to the MCMA and savanna fires along the Pacific Coast (SE of the MCMA) were more likely to impact the MCMA than emissions from Yucatan fires since the latter bypassed the MCMA. The importance of nearby fires and the low impact of Yucatan fires on the MCMA airshed during 2006 were confirmed using trajectories from other models reported elsewhere (Aiken et al., 2010; DeCarlo et al., 2010; de Foy et al., 2010). The difference in transport patterns between 2003 and 2006 could stem from the El Niño/Southern Oscillation (ENSO) phase: 2003 was an El Niño year while 2006 was ENSO-neutral. The trajectories for the Spring of 2006 could be closer to the long-term average (de Foy et al., 2008). In any case, there is an extensive BB zone to the south of the MCMA in both the FINNv1 results and our airborne fire

search results. This suggests that many different regional transport patterns, with a northerly component, could potentially carry BB emissions to the MCMA.

3.5 Summary of chemistry, climate, and air quality impacts of the different types of combustion in Mexico

5 About 70–80% of open BB occurs in the tropics (van der Werf et al., 2010) along with large amounts of biofuel use and garbage burning (GB); both in rural and urban areas. In addition, the tropics contain an increasing number of megacities due to rapid population growth and urbanization. These emission sources are of critical interest because the tropics are also the key to the oxidizing power of the global atmosphere (Crutzen and Andreae, 2000). The publication of the MILAGRO 2006–2007 BB measurements and the long history of detailed studies of MCMA air quality (e.g. Raga et al., 2001; Molina et al., 2007) suggest that Mexico is very likely the tropical country that is best characterized from an emissions standpoint. Thus, it is of interest to gain insight into the atmospheric chemistry of the tropics by comparing the magnitude of the diverse combustion emission sources in Mexico as a model country for this globally important region.

In Table 8, for each type of open burning sampled in this study, we have coupled the FINNv1 national fuel consumption estimates with our field-measured EF to estimate the total national emissions for several key species. We also show the emissions from national biofuel use and the national urban emissions; both taken from Table 7 of Christian et al. (2010). The latter were based mostly on the MILAGRO 2007 measurements and by scaling the 2004 MCMA emissions inventory (<http://www.sma.df.gob.mx/sma/index.php?opcion=26&id=392>) to the total urban population of Mexico (~75 million) as described in detail therein.

25 The estimated ratio of the annual national primary emissions from BB plus GB to the annual national primary emissions from fossil fuel (FF) combustion in urban areas for some key atmospheric species follows: CO (2.0), PM_{2.5} (47), NH₃ (3.4), NMOC (0.83), and NO_x (0.96) (Table 8, last column rounded to two significant figures). In reality, these

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ratios are highly uncertain since all the underlying BB, GB, and FF fuel consumption estimates are uncertain by at least a factor of two. In addition, secondary aerosol formation is not considered. The evolution of the aerosol/CO ratio was measured in one Mexican BB plume and it more than doubled in a little over one hour (Yokelson et al., 2009). Ratios of secondary to primary aerosol close to a factor of ten have been observed for urban aerosol generated primarily by FF combustion in Mexico (e.g. Salcedo et al., 2006). We also note that the estimate of EF for total NMOC produced by BB and GB in Table 8 is likely a factor of 2 to 3 too low since it relies completely on species that could be quantified with FTIR and WAS. BB studies that also incorporated mass spectrometric sampling (e.g. Christian et al., 2003; Yokelson et al., 2008) typically yield EFNMOC closer to 30 g kg^{-1} for most types of BB; or about twice the largest EFNMOC shown for BB in Table 8. Further, the studies that include mass spectrometric techniques still only account for roughly half the mass of NMOC emitted by BB (Karl et al., 2007). In developed countries, urban emissions from burning FF dominate the total combustion emissions (Forster et al., 2007). FF emissions are also significant in developing countries, but from a tropical (and therefore global) atmospheric chemistry standpoint the emissions from both BB and FF are significant; with our observations confirming that BB is especially likely to dominate the aerosol emissions in the tropics. This is consistent with past estimates that BB produces about 60% of global BC and 94% of primary combustion-generated organic aerosol (Bond et al., 2004).

The climate impacts of BB are complex. Recently, increased attention has been focused on reducing one aerosol component, BC, as a means of achieving rapid, partial mitigation of global warming (Jacobson, 2002; Unger et al., 2010). However, airborne TEM and ground-based filter samples of BB aerosol during MILAGRO confirm that BB aerosol is mostly composed of OA. For instance, with the exception of brick kilns, the 2007 filter-based measurements of BB sources in Mexico found that the EC (a surrogate for BC) emissions were $\sim 12 \pm 12\%$ of the co-emitted OA by mass (range 0.6–36%, based on Table 4 of Christian et al., 2010). The dominant OA emissions contribute to short term cooling through a variety of direct and indirect effects (Reid et al., 2005) and

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even the net forcing of the BC is unclear when indirect effects are considered (Koch and Genio, 2010). BC can undergo long-range transport and increase snow/ice melt rates even when co-emitted with organic species (Hadley et al., 2010). However, BC produced by BB is more hygroscopic and shorter lived than BC produced by FF (Peters et al., 2009). While most of the greenhouse gases emitted by savanna fires are quickly taken up by regrowth of new grasses, deforestation fires are a net source of greenhouse gases (Forster et al., 2007). For these and other reasons, the climate impacts of BB in Mexico and globally will require much more research.

From an air quality perspective, the health impacts of FF emissions are likely magnified relative to most open burning emissions since the FF emissions are produced primarily in the major population centers. However, open GB also occurs in or near urban areas. GB is identified as the main global source of dioxins and likely produces numerous other air toxics (Costner, 2006). In addition, open BB emissions can be transported to major urban areas both in Mexico and globally (Moffet et al., 2008; Aiken et al., 2010; Yokelson et al., 2007b) and the health effects of biofuel use, which mainly consists of open cooking fires built within rural dwellings is a major global public health issue (Dherani et al., 2008).

4 Conclusions

During 2006 and 2007 the initial emissions from 78 fires were measured in Mexico from airborne platforms or a ground-based mobile lab. The fires sampled included examples of most types of open burning and several types of domestic and industrial biofuel use. In this paper we have used the emissions data to derive national-average emission factors (EF) for 6 types of open burning: savanna, tropical dry forest, crop residue, pine-oak (temperate forest), mixed crop residue and tropical dry forest, and garbage burning. The EF for early dry season savanna fires were found to be higher than the EF for late dry season savanna fires for NO_x and many smoldering compounds. Comparison of the emissions from pine-oak forest fires in urban and rural

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areas of Mexico showed that NO_x emissions were about a factor of two higher from the urban-impacted forests; perhaps as a result of nitrogen deposition. The emissions from fires that started as crop residue fires, but escaped to also burn some adjacent forest (possibly due to high surface windspeeds) were examined as a group. The fires that burned both fuel types emitted significantly more NO_x than the fires that burned just one of the two component fuel types. Thus, for NO_x and perhaps some other species, deposition, seasonal phenology, windspeed, or other factors, may play as large a role in controlling EF as vegetation type does.

Both TEM and filter-based techniques were used to analyze numerous samples of Mexican BB aerosol. Both approaches show the overall aerosol composition is dominated by organic species. On average, soot accounts for $\sim 27\%$ of BB aerosol particles by number and EC accounts for $\sim 12\%$ of BB aerosol by mass. When considering all the fires sampled from the air, the percentage of individual aerosol particles that were identified as soot by TEM was positively correlated with MCE. The percentage of organic aerosol particles was negatively correlated with MCE. These observations are consistent with findings that flaming combustion is the source of black carbon and that smoldering combustion produces most of the organic aerosol.

The FINNV1 model and an airborne search for fires both suggest that an arc of intense, March-May, open biomass burning stretches across southern Mexico to the south of Mexico City. Mexico City may be most frequently affected by open BB in the pine-oak forest found in the nearby mountains and savanna fires on the west coast. About 175 Tg of biomass, biofuel, and garbage is burned in a typical year in Mexico and the emissions released to the atmosphere rival or exceed those from urban fossil fuel use for many species. This confirms the important role of biomass burning in the atmospheric chemistry of the global tropics.

Supplementary material related to this article is available online at:
[http://www.atmos-chem-phys-discuss.net/11/7321/2011/
acpd-11-7321-2011-supplement.pdf](http://www.atmos-chem-phys-discuss.net/11/7321/2011/acpd-11-7321-2011-supplement.pdf).

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Table 1. Details of the open fires (not including biofuels) sampled in Mexico during MILAGRO.

Date dd/mm/yy	Fire #	Lat decimal degrees	Long decimal degrees	Time LT	MODIS Overpass ¹		Hotspot Y/N	Platform	Fuels observed from aircraft or ground	Ref ²
					Terra LT	Aqua LT				
Fires sampled from the air in March of 2006										
04/03/06	1	18.3209	-96.2067	1454	1125	-	N	Twin Otter	Crop Residue	
04/03/06	2	18.0509	-96.1922	1511	1125	-	N	Twin Otter	Crop Residue	
04/03/06	3	18.4408	-95.7162	1542	1125	-	N	Twin Otter	Crop Residue	
06/03/06	1	19.0763	-99.0537	1327	1115	1420	N	Twin Otter	Pine-Oak Urban	1
06/03/06	2	19.1739	-99.1903	1332	1115	1420	N	Twin Otter	Pine-Oak Urban	1
06/03/06	3	19.1881	-99.3783	1705-1709	1115	1420	N	Twin Otter	Pine-Oak Urban	1
06/03/06	4	19.0711	-99.2283	1714	1115	1420	N	Twin Otter	Pine-Oak Urban	1
08/03/06	1	20.6510	-97.7880	1621-1624	1100	1405	N	Twin Otter	Crop Residue	
08/03/06	2	20.0480	-97.2830	1646	1100	1405	Y	Twin Otter	Tropical Dry Deciduous Forest	
09/03/06	1	19.3269	-99.4775	1320	-	-	N	Twin Otter	Pine-Oak Urban	1
10/03/06	1	19.6431	-98.3578	1716	1050	1355	N	C-130	Pine-Oak Urban	1
11/03/06	1	22.5314	-99.3219	1444-1455	1130	-	Y	Twin Otter	Crop Residue	
11/03/06	2	22.6132	-99.4895	1502-1512	1130	-	N	Twin Otter	Tropical Dry Deciduous Forest	
11/03/06	3	22.7939	-99.4674	1523-1525	1130	-	N	Twin Otter	Tropical Dry Deciduous Forest	
12/03/06	3	19.1834	-90.7269	1405-1409	1035	-	Y	Twin Otter	Crop Residue	2
12/03/06	4	19.6330	-90.6758	1420-1424	1035	-	N	Twin Otter	Crop Residue	2
12/03/06	5	19.5947	-90.6497	1555	1035	-	N	Twin Otter	Crop Residue	2
17/03/06	1	19.0681	-99.0616	1158-1242	1055	1400	N	Twin Otter	Pine-Oak Urban	1
17/03/06	2	19.3862	-98.6066	1306-1318	1055	1400	N	Twin Otter	Pine-Oak Urban	1
18/03/06	1	19.6685	-97.6232	1452-1511	1140	-	N	Twin Otter	Unknown	
18/03/06	2	19.3456	-98.6851	1546-1639	1140	-	N	Twin Otter	Pine-Oak Urban	1
18/03/06	3	19.4906	-98.0558	1709-1715	1140	-	N	Twin Otter	Pinyon Pine and Juniper	
20/03/06	1	16.6613	-92.1847	1248-1300	1125	1250	N	Twin Otter	Pine-Oak Rural	
20/03/06	2	16.6225	-92.3365	1305-1307	1125	1250	N	Twin Otter	Pine-Oak Rural	
20/03/06	3	16.5068	-92.1999	1314-1339	1125	1250	Y	Twin Otter	Pine-Oak Rural	
20/03/06	4	16.3384	-92.6197	1343-1349	1125	1250	N	Twin Otter	Savanna	
20/03/06	5	16.4534	-92.8539	1402-1409	1125	1250	Y	Twin Otter	Savanna	
22/03/06	1	19.5306	-90.1063	1314-1317	1115	1240	N	Twin Otter	Crop Residue & Tropical Dry Forest Mix	2
22/03/06	2	19.7748	-89.8675	1329-1333	1115	1240	Y	Twin Otter	Crop Residue & Tropical Dry Forest Mix	2
22/03/06	3	19.7703	-89.5177	1344-1348	1115	1240	N	Twin Otter	Tropical Dry Deciduous Forest	2
22/03/06	4	19.8158	-89.4578	1353-1356	1115	1240	N	Twin Otter	Tropical Dry Deciduous Forest	2
22/03/06	5	19.8649	-89.4717	1358-1401	1115	1240	N	Twin Otter	Tropical Dry Deciduous Forest	2
22/03/06	6	20.3552	-88.8383	1424-1427	1115	1240	N	Twin Otter	Tropical Dry Deciduous Forest	2

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Table 1. Continued.

Date dd/mm/yy	Fire #	Lat decimal degrees	Long decimal degrees	Time LT	MODIS Overpass ¹		Hotspot Y/N	Platform	Fuels observed from aircraft or ground	Ref ²
					Terra LT	Aqua LT				
Fires sampled from the air in March of 2006 (continued)										
22/03/06	7	20.8525	-88.4018	1447-1450	1115	1240	N	Twin Otter	Tropical Dry Deciduous Forest	2
22/03/06	8	21.2085	-89.0345	1512-1529	1115	1240	Y	Twin Otter	Crop Residue & Tropical Dry Forest Mix	2
23/03/06	1	19.8266	-89.3825	1410	1020	1325	N	C-130	Unknown	2
23/03/06	2	19.7648	-89.5220	1412	1020	1325	Y	C-130	Unknown	2
23/03/06	3	19.8201	-89.3681	1417	1020	1325	N	C-130	Unknown	2
25/03/06	1	17.8018	-97.9430	1304-1317	-	-	N	Twin Otter	Pine-Oak Rural	
25/03/06	2	18.5907	-96.6805	1428-1432	-	-	N	Twin Otter	Garbage	
25/03/06	3	18.6488	-96.6410	1435	-	-	N	Twin Otter	Crop Residue	
27/03/06	1	18.7020	-100.0799	1246-1254	1130	-	N	Twin Otter	Pine-Oak Rural	
27/03/06	2	17.0270	-99.9755	1353-1354	1130	-	N	Twin Otter	Savanna	
27/03/06	3	16.9111	-99.6913	1404-1407	1130	-	N	Twin Otter	Savanna	
27/03/06	4	16.9302	-99.4475	1414-1422	1130	-	N	Twin Otter	Savanna	
27/03/06	5	16.8788	-99.4733	1427-1428	1130	-	N	Twin Otter	Savanna	
27/03/06	6	16.9103	-99.6917	1433-1439	1130	-	N	Twin Otter	Savanna	
27/03/06	7	17.7585	-98.3360	1638-1642	1130	-	N	Twin Otter	Savanna	
28/03/06	1	19.2787	-101.1498	1141-1145	-	1340	Y	Twin Otter	Pine-Oak Rural	
28/03/06	2	19.6021	-102.3766	1415-1423	-	1340	N	Twin Otter	Pine-Oak Rural	
29/03/06	1	19.6235	-90.3815	1337-1353	1120	1245	Y	Twin Otter	Crop Residue & Tropical Dry Forest Mix	2
29/03/06	2	19.7208	-90.4108	1346	1120	1245	N	Twin Otter	Tropical Dry Deciduous Forest	2
29/03/06	3	19.4230	-90.2315	1400	1120	1245	N	Twin Otter	Crop Residue & Tropical Dry Forest Mix	2
29/03/06	4	19.5128	-89.8192	1412-1416	1120	1245	N	Twin Otter	Crop Residue	2
29/03/06	5	20.0371	-89.8395	1435-1448	1120	1245	N	Twin Otter	Crop Residue	2
29/03/06	6	20.0055	-89.7808	1438-1441	1120	1245	N	Twin Otter	Crop Residue	2
Fires sampled from the ground in 2007										
23/04/07	1	20.01	-99.49	1648-1748	-	-	-	Ground	Garbage	3
24/04/07	1	19.81	-99.22	1304-1841	-	-	-	Ground	Garbage	3
25/04/07	1	19.97	-98.92	1355-1502	-	-	-	Ground	Garbage	3
26/04/07	1	19.70	-98.80	1321-1438	-	-	-	Ground	Garbage	3
30/04/07	1	20.60	-101.22	1246-1334	-	-	-	Ground	Crop Residue	3
01/05/07	1	20.60	-101.22	1426-1518	-	-	-	Ground	Crop Residue	3

¹Blank entries indicate the MODIS scene was heavily impacted by clouds or that the sampled fire(s) were located on the extreme edge of the scene or outside of the scene.

²Yokelson et al., (2007a) = 1, Yokelson et al., (2009) = 2, Christian et al., (2010) = 3. No number is assigned to fires for which EF data is first published in this work.

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Table 2. Emission factors (EF, g kg⁻¹) and modified combustion efficiency (MCE) measured for crop residue fires.

Date	04/03/06	04/03/06	04/03/06	08/03/06	11/03/06	25/03/06	12/03/06	12/03/06	12/03/06	29/03/06	29/03/06	29/03/06	30/04/07	01/05/07	Avg	Stdev	EF at average MCE
Fire #	1	2	3	1	1	3	3	4	5	4	5	6	1	1			
MCE	0.970	0.945	0.870	0.911	0.908	0.950	0.928	0.950	0.956	0.892	0.941	0.937	0.910	0.882	0.925	0.030	
Species	EF	EF															
CO ₂	1767	1728	1545	1640	1636	1723	1631	1722	1729	1603	1679	1689	1628	1577	1664	66	1664
CO	34.66	64.45	146.34	101.68	105.38	58.27	80.04	57.63	50.38	123.20	67.25	71.74	102.17	134.70	85.56	33.75	85.56
NO	-	-	-	2.348	-	-	-	-	-	3.206	1.960	1.322	-	-	2.209	0.788	2.063
NO ₂	3.566	-	-	1.022	-	5.355	4.457	7.077	4.550	4.753	1.253	1.275	-	-	3.701	2.110	3.482
NO _x as NO	2.325	-	-	3.014	-	3.492	2.907	4.615	2.968	5.673	2.777	2.154	-	-	3.325	1.133	3.637
CH ₄	0.859	-	16.422	6.458	4.417	1.726	9.648	1.471	2.067	5.131	4.402	2.709	5.169	6.730	5.170	4.200	5.008
C ₂ H ₄	0.623	-	-	1.453	1.235	0.932	1.465	0.526	0.594	0.962	0.761	0.746	1.512	2.485	1.108	0.561	1.155
C ₂ H ₂	0.193	-	-	0.340	0.218	-	0.292	0.144	0.139	-	0.234	0.202	0.175	0.319	0.226	0.070	0.233
C ₂ H ₆	-	-	-	-	0.817	-	1.476	0.153	0.190	-	0.897	0.662	-	-	0.699	0.493	0.910
C ₃ H ₆	-	-	-	0.452	0.653	-	0.942	0.173	0.205	-	0.630	0.634	0.774	-	0.558	0.267	0.496
HCHO	1.979	-	-	0.523	-	-	-	-	-	-	-	-	2.485	2.467	1.864	0.924	1.845
CH ₃ OH	-	-	3.640	4.297	5.506	0.251	4.397	1.892	0.766	3.530	2.034	1.704	3.704	2.448	2.847	1.582	2.665
CH ₃ COOH	0.890	-	4.183	0.600	-	-	4.643	6.476	7.070	-	3.250	2.388	9.151	6.486	4.514	2.798	4.523
HCOOH	0.682	1.933	1.125	0.554	-	-	1.838	0.294	1.780	-	-	-	0.285	0.526	1.002	0.683	1.003
NH ₃	0.142	1.613	4.015	1.658	-	-	0.910	3.902	1.393	-	0.350	0.339	1.539	2.827	1.699	1.354	1.755
HCN	0.721	-	-	0.019	-	0.476	-	-	-	-	0.277	-	-	-	0.373	0.298	0.158
propane	-	-	-	-	0.283	-	0.436	0.040	0.054	-	0.256	0.184	-	-	0.209	0.150	0.282
isobutane	-	-	-	-	0.030	-	0.032	0.003	0.004	-	0.020	0.012	-	-	0.017	0.013	0.025
n-butane	-	-	-	-	0.086	-	0.097	0.009	0.014	-	0.045	0.048	-	-	0.050	0.036	0.072
t-2-butene	-	-	-	-	0.057	-	0.089	0.011	0.012	-	0.057	0.034	-	-	0.043	0.030	0.057
1-butene	-	-	-	-	0.138	-	0.195	0.041	0.044	-	0.127	0.080	-	-	0.104	0.060	0.134
isobutene	-	-	-	-	0.120	-	0.176	0.022	0.031	-	0.117	0.064	-	-	0.088	0.060	0.117
c-2-butene	-	-	-	-	0.043	-	0.067	0.008	0.009	-	0.043	0.025	-	-	0.033	0.023	0.043
cyclopentane	-	-	-	-	2.E-03	-	3.E-03	1.E-04	4.E-04	-	2.E-03	1.E-03	-	-	1.5E-03	1.2E-03	1.9E-03
isopentane	-	-	-	-	0.034	-	0.014	0.001	-	-	0.010	0.010	-	-	0.014	0.012	0.020
n-pentane	-	-	-	-	0.032	-	0.030	0.003	0.005	-	0.020	0.014	-	-	0.017	0.012	0.025
1,3 butadiene	-	-	-	-	0.175	-	0.214	0.033	0.057	-	0.130	0.072	-	-	0.114	0.072	0.151
PM _{2.5}	-	-	-	-	-	3.79	6.52	-	-	5.78	7.06	3.87	10.14	-	6.19	2.36	6.26
H ₂	-	-	-	-	-	-	-	-	-	4.76	1.66	1.68	-	-	2.70	1.78	2.59

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Table 3. Emission factors (EF, g kg^{-1}) and modified combustion efficiency (MCE) for tropical dry forest fires.

Date	22/03/06	22/03/06	22/03/06	22/03/06	22/03/06	29/03/06	08/03/06	11/03/06	11/03/06	Average	Stdev	EF at average MCE
Fire #	3	4	5	6	7	2	2	2	3			
MCE	0.925	0.923	0.945	0.907	0.926	0.938	0.897	0.912	0.941	0.924	0.016	
Species	EF	EF										
CO ₂	1644	1651	1711	1600	1651	1682	1612	1651	1711	1657	39	1657
CO	84.71	88.22	63.27	104.3	84.26	71.35	118.35	101.22	68.50	87.13	18.20	87.13
NO	2.688	3.866	1.275	1.496	0.877	3.909	2.009	-	-	2.303	1.225	2.311
NO ₂	2.049	3.559	1.248	4.598	3.858	6.253	3.359	-	-	3.561	1.637	3.549
NO _x as NO	4.025	6.187	2.089	4.495	3.394	7.987	4.200	-	-	4.625	1.926	4.627
CH ₄	6.055	7.326	2.557	9.213	6.595	3.718	6.490	5.603	3.572	5.681	2.088	5.682
HCHO	6.661	-	0.886	0.879	2.613	-	-	-	-	2.760	2.726	2.770
CH ₃ OH	2.879	4.527	1.354	4.689	3.325	1.021	3.404	5.535	3.395	3.348	1.483	3.348
CH ₃ COOH	2.534	2.412	1.431	4.678	3.416	-	2.497	-	-	2.828	1.104	2.710
HCOOH	-	-	-	-	-	-	1.823	-	-	1.823	-	1.823
NH ₃	1.869	-	0.539	0.309	0.366	-	1.011	8.259	4.987	2.477	3.032	2.482
HCN	0.302	-	0.225	-	0.172	-	-	-	-	0.233	0.065	0.240
ethane	1.187	-	0.49	1.611	1.065	0.992	-	-	-	1.069	0.403	1.187
ethene	0.992	-	0.742	1.452	1.127	0.925	0.583	-	-	0.970	0.304	0.969
propene	0.521	-	0.423	1.215	0.669	3.924	-	-	-	1.350	1.471	1.263
ethyne	-	-	-	-	-	-	1.142	-	-	1.142	-	1.142
PM _{2.5}	5.48	5.49	2.12	6.59	3.73	3.56	-	-	-	4.50	1.64	4.91
H ₂	3.00	-	2.16	4.32	2.27	1.29	-	-	-	2.61	1.13	2.91

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Table 4. Emission factors (EF, g kg⁻¹) and modified combustion efficiency (MCE) for fires burning mixed crop residue and tropical dry forest.

Date	22/03/06	22/03/06	22/03/06	29/03/06	29/03/06	Average	Stdev	EF at average MCE
Fire #	1	2	8	1	3			
MCE	0.899	0.915	0.936	0.927	0.941	0.924	0.017	
Species	EF	EF	EF	EF	EF	EF		
CO ₂	1607	1627	1671	1656	1695	1651	35	1651
CO	114.4	96.76	72.25	82.44	68.13	86.80	18.98	86.80
NO	4.780	1.620	5.386	2.412	6.345	4.109	2.010	4.109
NO ₂	11.510	2.360	12.430	4.818	9.669	8.157	4.374	8.157
NO _x as NO	12.290	3.159	13.490	4.568	12.650	9.231	4.945	9.232
CH ₄	7.349	6.302	4.151	3.517	4.883	5.240	1.570	5.241
HCHO	-	3.136	-	-	-	3.136	-	3.136
CH ₃ OH	3.834	3.168	2.376	1.694	1.336	2.482	1.030	2.482
CH ₃ COOH	-	3.544	-	6.359	-	4.952	1.991	5.558
NH ₃	0.626	0.080	1.675	-	-	0.794	0.811	1.007
HCN	-	0.102	0.449	-	0.250	0.267	0.174	0.204
ethane	-	-	0.668	0.866	-	0.767	0.140	0.941
ethene	-	-	1.065	0.813	0.433	0.770	0.318	0.998
propene	-	-	0.845	1.386	-	1.116	0.383	1.590
ethyne	-	-	-	0.295	-	0.295	-	.295
H ₂	-	-	2.37	1.44	-	1.91	0.66	1.09
PM _{2.5}	8.29	8.83	10.06	6.21	5.69	7.82	1.83	7.82

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Table 5. Emission factors (EF, g kg⁻¹) and modified combustion efficiency (MCE) measured for early dry season savanna fires in Mexico compared to late dry season data from Africa.

Date	20/3/06	20/3/06	27/3/06	27/3/06	27/3/06	27/3/06	27/3/06	27/3/06	Average	Stdev	EF at average MCE	Africa ¹	Mex/Afr
Fire #	4	5	2	3	4	5	6	7					
MCE	0.934	0.928	0.963	0.914	0.940	0.924	0.914	0.923	0.930	0.016		0.938	0.99
Species	EF												
CO ₂	1671	1667	1740	1625	1678	1626	1623	1652	1660	39.02	1660	1703	0.97
CO	75.61	82.17	42.02	97.49	67.70	84.56	96.66	87.99	79.28	18.05	79.28	71.5	1.11
NO	3.672	4.104	-	-	3.648	-	-	5.100	4.131	0.679	4.217	-	-
NO ₂	2.870	1.257	-	9.416	4.974	-	8.426	5.660	5.434	3.134	4.457	-	-
NO _x as NO	5.544	4.924	-	6.141	6.893	-	5.495	7.191	6.031	0.877	6.093	3.37	1.81
CH ₄	3.450	3.665	-	5.532	3.431	4.442	5.506	2.852	4.126	1.061	3.739	2.19	1.71
C ₂ H ₄	0.892	0.760	1.466	0.845	0.312	0.922	0.741	1.366	0.913	0.364	0.913	1.19	0.77
C ₂ H ₂	0.364	0.399	-	0.058	0.248	0.151	0.096	0.038	0.193	0.146	0.243	0.26	0.94
C ₂ H ₆	1.336	0.589	-	2.504	1.649	-	1.082	-	1.432	0.714	1.371	0.21	6.53
C ₃ H ₆	0.446	0.333	2.303	-	1.298	6.346	2.126	3.969	2.403	2.138	2.474	0.23	10.8
HCHO	0.229	-	-	-	-	-	-	1.363	0.796	0.802	0.598	1.06	0.56
CH ₃ OH	1.966	2.035	-	2.673	1.562	0.992	2.341	3.661	2.176	0.849	1.991	1.17	1.70
CH ₃ COOH	-	-	3.440	-	4.217	9.391	-	-	5.683	3.235	7.503	2.42	3.10
HCOOH ²	0.137	0.350	-	-	-	-	-	-	0.244	0.151	0.273	0.295	0.92
NH ₃	0.730	0.495	-	-	-	-	-	0.205	0.477	0.263	0.572	0.28	2.04
HCN	0.690	0.636	0.202	-	0.767	-	-	0.199	0.499	0.276	0.533	0.53	1.01
PM _{2.5}	8.11	6.48	3.96	10.20	7.76	9.39	10.84	4.43	7.65	2.54	7.65	10.0	0.77
H ₂	1.62	2.05	-	2.56	1.22	-	2.78	1.38	1.93	0.64	1.70	0.97	1.76

¹African late dry season savanna fire emission factors for ethane, propene, and PM_{2.5} from Sinha et al., (2003); H₂ from Andreae and Merlet (2001); remainder Yokelson et al., (2003).

²The formic acid emission factor from Yokelson et al., (2003) was scaled to reflect new reference data (see text).

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Table 6. Emission factors (EF, g kg⁻¹) and modified combustion efficiency (MCE) measured in rural pine-oak forests in Mexico compared to data from urban-impacted pine-oak forests in Mexico.

Date	20/3/06	20/3/06	20/3/06	25/3/06	27/3/06	28/3/06	28/3/06	Average	Stdev	EF at average MCE	Urban Pine- Oak ¹	Urban/Rural Pine-Oak
Fire #	1	2	3	1	1	1	2					
MCE	0.943	0.904	0.869	0.907	0.889	0.903	0.942	0.908	0.027		0.927	1.02
Species	EF											
CO ₂	1693	1593	1509	1603	1549	1597	1673	1603	64.351	1603	1662	1.04
CO	64.76	107.52	145.28	104.58	123.25	109.68	65.11	102.9	29.336	102.9	82.9	0.81
NO	1.903	0.711	2.370	1.635	2.410	1.070	1.128	1.604	0.663	1.604	-	-
NO ₂	1.373	4.379	3.620	2.993	3.742	4.364	1.898	3.195	1.176	3.195	-	-
NO _x as NO	2.798	3.566	4.731	3.432	4.850	3.916	2.309	3.658	0.936	3.657	7.44	2.03
CH ₄	3.522	4.553	11.406	4.500	6.526	5.052	4.367	5.704	2.675	5.704	4.96	0.87
C ₂ H ₄	0.952	1.004	1.802	1.194	1.114	0.473	0.974	1.073	0.395	1.074	0.94	0.88
C ₂ H ₂	0.165	0.274	0.314	0.506	0.519	0.487	0.242	0.358	0.144	0.358	0.19	0.53
C ₂ H ₆	1.271	-	3.959	1.681	1.737	0.416	1.191	1.709	1.200	1.727	0.58	0.34
C ₃ H ₆	1.581	4.329	1.990	1.084	3.241	0.791	1.350	2.052	1.281	2.052	0.5	0.24
HCHO	2.507	2.006	5.381	-	-	-	1.113	2.752	1.845	3.032	2.99	0.99
CH ₃ OH	1.405	2.315	4.243	3.218	3.795	3.216	1.506	2.814	1.101	2.814	2.06	0.73
CH ₃ COOH	-	0.649	-	-	0.265	3.661	-	1.525	1.860	2.610	3.96	1.52
HCOOH	0.280	1.226	1.282	0.390	-	0.176	0.065	0.570	0.541	0.615	1.243	2.02
NH ₃	1.038	0.357	1.144	-	-	0.046	0.006	0.518	0.541	0.539	0.91	1.69
HCN	0.635	1.003	0.532	0.922	1.225	1.099	0.713	0.876	0.256	0.876	1.02	1.16
PM _{2.5}	3.91	11.80	9.17	12.93	17.53	12.40	11.56	11.33	4.13	11.33	11.05	0.98
H ₂	1.36	0.49	5.95	1.47	2.06	1.45	1.43	2.03	1.79	2.03	1.51	0.74

¹From Yokelson et al. (2007a). Their formic acid emission factor has been normalized to reflect new reference data (see text).

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Table 7. Emission factors (EF, g kg⁻¹) and modified combustion efficiency (MCE) measured for open burning of garbage in Mexico compared to other work.

Date	25/03/06	23/04/07	24/04/07	25/04/07	26/04/07	Average	Stdev	AP-42	Lemieux et al., (2000)
Fire #	2	1	1	1	1				
MCE	0.974	0.964	0.911	0.958	0.968	0.955	0.025		
Species	EF	EF	EF	EF	EF	EF			
CO ₂	1538	1404	1270	1385	1409	1401	95		
CO	26.09	33.80	79.10	38.70	29.60	41.46	21.56	42	-
NO	-	-	-	-	-	-	-	-	-
NO ₂	6.865	-	-	-	-	6.865	-	-	-
NO _x as NO	4.477	-	-	-	-	4.477	-	3	-
CH ₄	0.766	1.16	10.3	2.18	1.14	3.109	4.054	6.5	-
C ₂ H ₄	0.322	0.82	4.75	2.20	0.99	1.816	1.779	-	-
C ₂ H ₂	-	0.14	0.72	0.53	0.20	0.398	0.275	-	-
C ₃ H ₆	-	0.36	3.34	0.97	0.36	1.258	1.418	-	-
HCHO	-	0.56	0.48	0.68	0.76	0.620	0.124	-	-
CH ₃ OH	-	0.31	2.81	0.40	0.26	0.945	1.245	-	-
CH ₃ COOH	-	0.58	7.40	0.92	0.78	2.420	3.323	-	-
HCOOH	-	0.05	0.14	0.34	0.19	0.181	0.119	-	-
NH ₃	0.768	0.46	2.52	0.39	-	1.035	1.004	-	-
HCN	0.473	-	-	-	-	0.473	-	-	-
HCl	-	1.65	-	9.80	3.02	4.82	4.36	-	2.4
PM _{2.5} ¹	-	-	20.66	5.00	5.78	10.48	8.82	8	11.3
H ₂	0.091	-	-	-	-	0.091	-	-	-
SO ₂	-	-	-	-	-	-	-	0.5	-

¹This work PM_{2.5} estimated from 1.2 times the sum of species measured on Quartz filters (Christian et al., 2010).

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Table 8. Rough comparison of selected primary emissions from combustion in Mexico.

Combustion Type	Tropical Forest	Savanna	Temperate Forest	Crop Residue	Biofuel ¹	Garbage	National BB+GB	National Urban ¹	BB+GB/Urban
Tg Burned	63.6	17	12.8	2.9	68	11.1	175.4		
EF _{CO}	87.10	79.28	102.90	85.56	58.40	41.46	-	-	-
EF _{PM_{2.5}}	4.91	7.65	11.33	6.26	6.73	10.48	-	-	-
EF _{NH₃}	2.48	0.57	0.54	1.76	0.44	1.04	-	-	-
EF _{NMOC} ²	11.94	15.90	15.16	13.92	4.34	8.11	-	-	-
EF _{NO_x}	4.63	6.09	3.66	3.64	2.04	4.48	-	-	-
Tg CO	5.54	1.35	1.32	0.25	3.97	0.46	12.88	6.68	1.93
Tg PM _{2.5}	0.31	0.13	0.15	0.018	0.46	0.12	1.18	0.025	47.18
Tg NH ₃	0.16	0.010	0.007	0.005	0.030	0.012	0.22	0.065	3.40
Tg NMOC ²	0.76	0.27	0.19	0.040	0.30	0.090	1.65	1.98	0.83
Tg NO _x	0.29	0.104	0.047	0.011	0.14	0.050	0.64	0.67	0.96

¹From Table 7 of Christian et al. (2010) and references there-in.²The emission factor for the sum of identified NMOC. Total NMOC for BB and GB is likely 2-3 times larger (see text).

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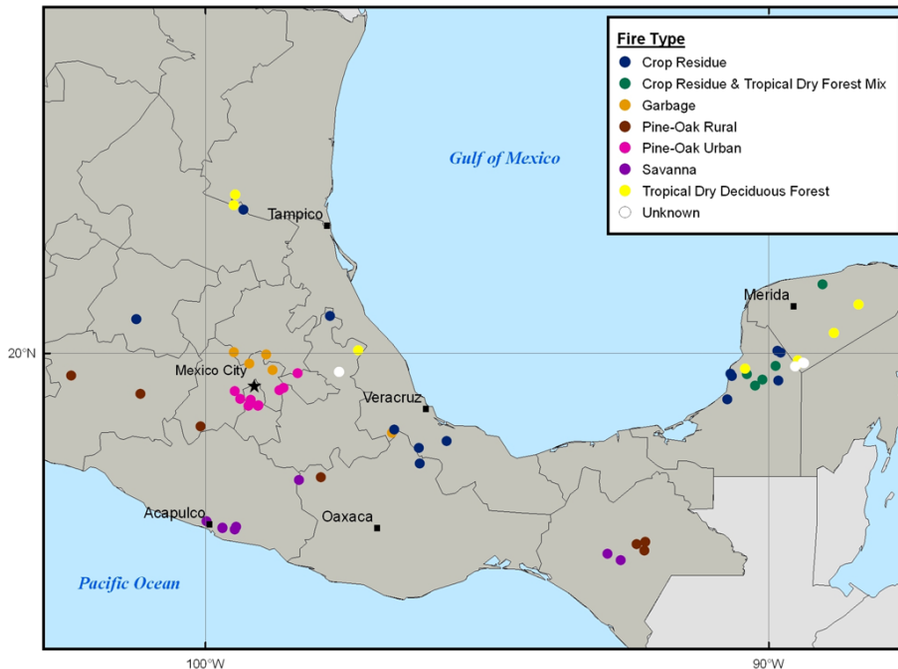


Fig. 1. Map showing location of open biomass burning sampled in Mexico in 2006 and 2007.

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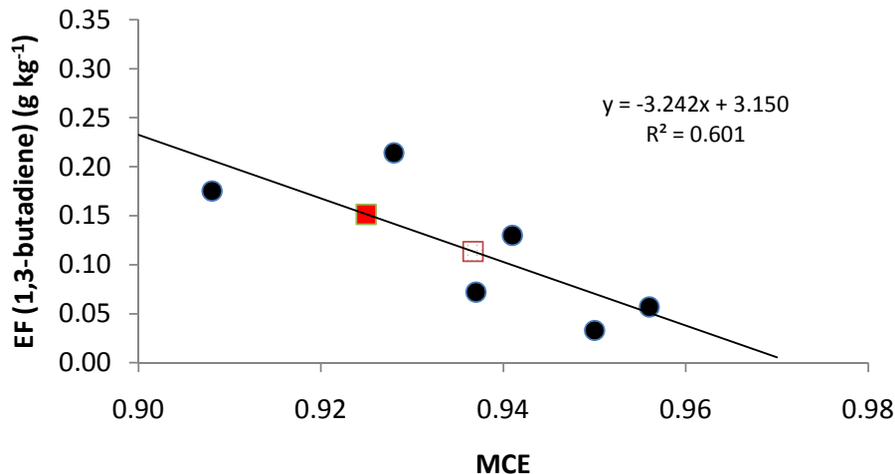


Fig. 2. The solid black circles show all the EF measurements on crop residue fires for 1,3-butadiene and the open red square shows the average of those measurements (0.114 g kg^{-1}). The filled red square is the EF for 1,3-butadiene (0.151 g kg^{-1}) computed from the fit at the average MCE for all the crop residue fires sampled in Mexico. The value from the fit is 32% larger and likely better reflects the average EF (see text).

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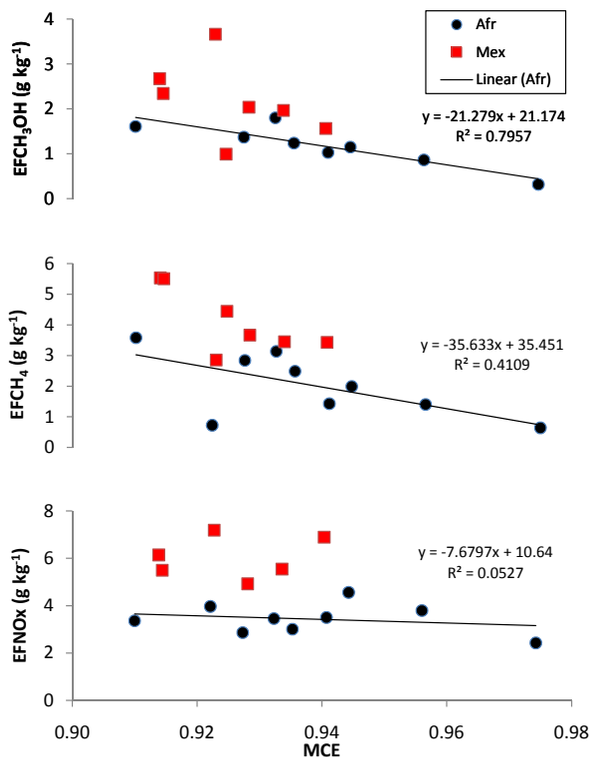


Fig. 3. For several species we show the emission factors measured during the late dry season on savanna fires in Africa (and a fit) for comparison to the emission factors measured on early dry season savanna fires in Mexico. The emission factors for the early dry season fires are higher and lie above the fit for African late season fires for most species as discussed in the text.

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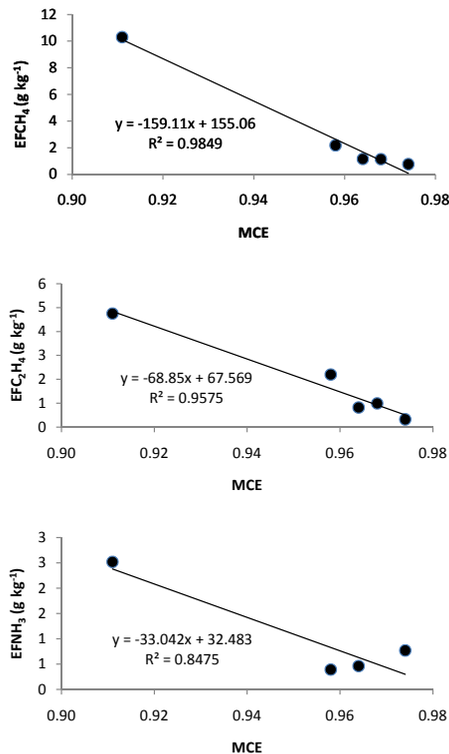


Fig. 4. The 3 species emitted by open garbage burning (in addition to CO and CO₂) that have been measured from both the air (high MCE point) and the ground fit one trend with MCE, but more measurements of this source are needed.

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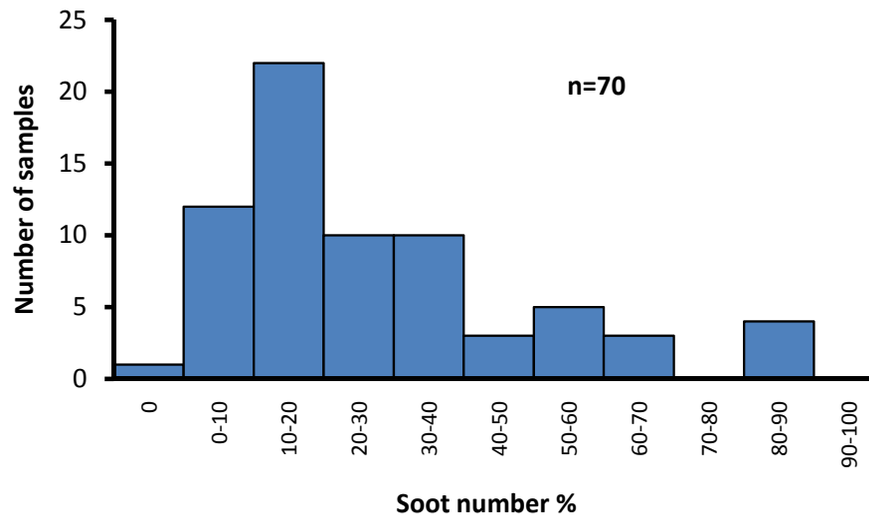


Fig. 5. The number of TEM samples with a percentage of soot particles in the incremental range indicated on the x-axis. On average $27 \pm 21\%$ of the individual BB particles were identified as soot particles (see section 3.2).

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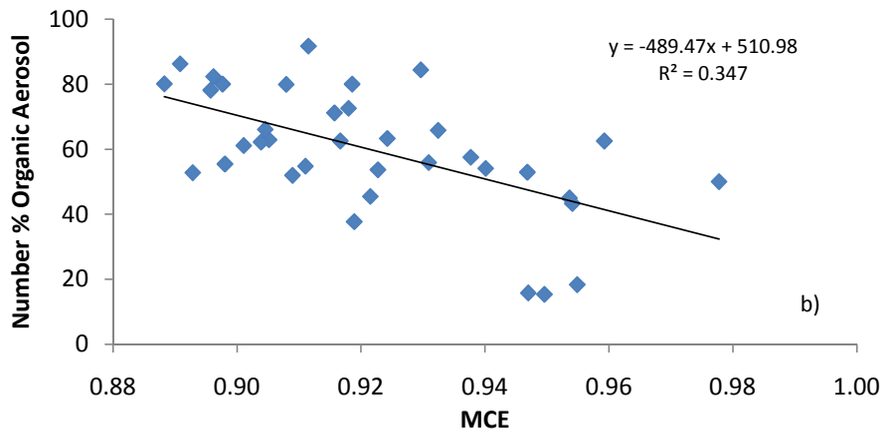
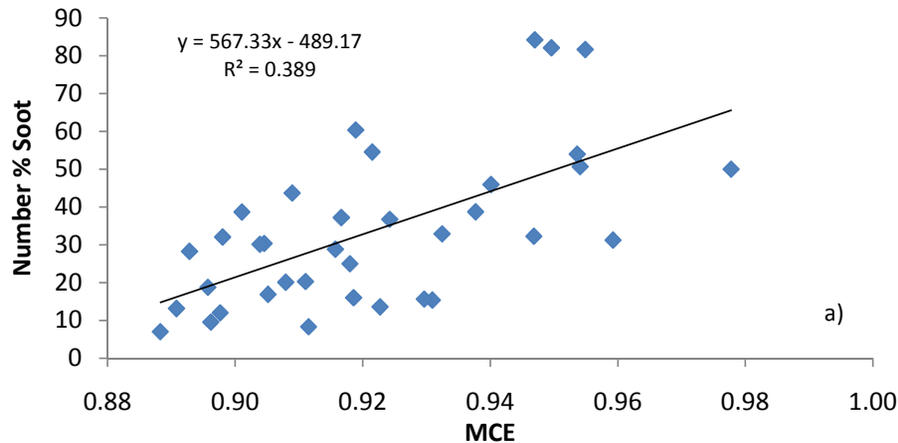


Fig. 6. The percentage of individual BB particles in a smoke sample that are classified as **(a)** soot or **(b)** organic aerosol plotted versus MCE. The tendency for flaming combustion to produce black carbon and smoldering combustion to produce organic aerosol is illustrated.

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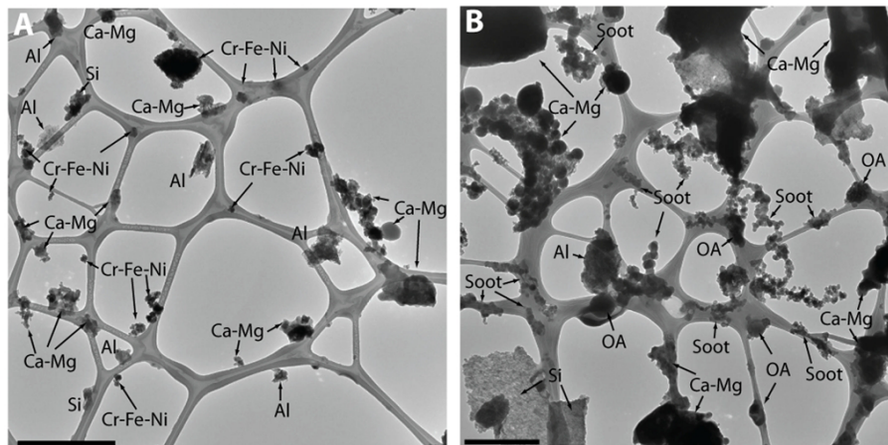


Fig. 7. TEM image of smoke samples from (a) garbage burning and (b) mixture of garbage burning and crop residue burning. The black horizontal bar at the base of each image indicates 1 micron. See Sect. 3.2 for discussion.

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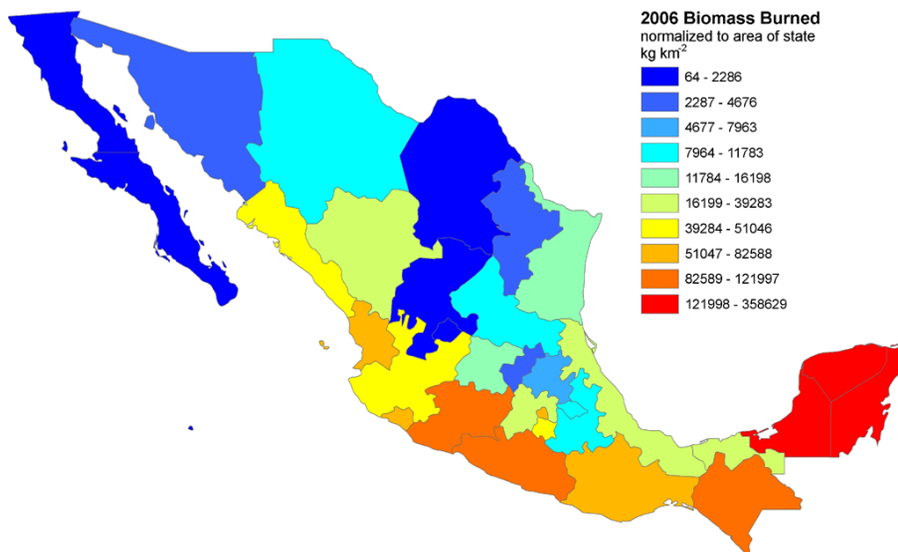
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Fig. 8. Map showing the amount of biomass burned in each Mexican state (as calculated by the FINNv1 model) normalized by the area of the state. The warm colors indicate an arc of high intensity biomass burning south of the Mexico City metropolitan area and stretching about 2000 km from coast to coast.