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**Trace gas and
particle emissions in
central Mexico**

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Trace gas and particle emissions from domestic and industrial biofuel use and garbage burning in central Mexico

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Abstract

In central Mexico during the spring of 2007 we measured the initial emissions of 12 gases and the aerosol speciation for elemental and organic carbon (EC, OC), anhydro-sugars, Cl^- , NO_3^- , and 20 metals from 10 cooking fires, four garbage fires, three brick making kilns, three charcoal making kilns, and two crop residue fires. Biofuel use has been estimated at over 2600 Tg/y. With several simple case studies we show that cooking fires can be a major, or the major, source of several gases and fine particles in developing countries. Insulated cook stoves with chimneys were earlier shown to reduce indoor air pollution and the fuel use per cooking task. We confirm that they also reduce the emissions of VOC pollutants per mass of fuel burned by about half. We did not detect HCN emissions from cooking fires in Mexico or Africa. Thus, if regional source attribution is based on HCN emissions typical for other types of biomass burning (BB), then biofuel use and total BB will be underestimated in much of the developing world. This is also significant because cooking fires are not detected from space. We estimate that ~ 2000 Tg/y of garbage are generated and about half may be burned, making this a commonly overlooked major global source of emissions. We estimate a fine particle emission factor (EF $_{\text{PM}_{2.5}}$) for garbage burning of $\sim 10 \pm 5$ g/kg, which is in reasonable agreement with very limited previous work. We observe large HCl emission factors in the range 2–10 g/kg. Consideration of the Cl content of the global waste stream suggests that garbage burning may generate as much as 6–9 Tg/yr of HCl, which would make it a major source of this compound. HCl generated by garbage burning in dry environments may have a relatively greater atmospheric impact than HCl generated in humid areas. Garbage burning $\text{PM}_{2.5}$ was found to contain levoglucosan and K in concentrations similar to those for biomass burning, so it could be a source of interference in some areas when using these tracers to estimate BB. Galactosan was the anhydro-sugar most closely correlated with BB in this study. Fine particle antimony (Sb) shows initial promise as a garbage burning tracer and suggests that this source could contribute a significant amount of the $\text{PM}_{2.5}$ in the Mexico City metropolitan area. The fuel

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consumption and emissions due to industrial biofuel use are difficult to characterize regionally. This is partly because of the diverse range of fuels used and the thin margins of typical micro-enterprises. Brick making kilns produced low total EFPM_{2.5} (~1.6 g/kg), but very high EC/OC ratios (6.72). Previous literature on brick kilns is scarce but does document some severe local impacts. Coupling data from Mexico, Brazil, and Zambia, we find that charcoal making kilns can exhibit an 8-fold increase in VOC/CO over their approximately one-week lifetime. Acetic acid emission factors for charcoal kilns were much higher in Mexico than elsewhere, probably due to the use of tannin-rich oak fuel. Our dirt charcoal kiln EFPM_{2.5} emission factor was ~1.1 g/kg, which is lower than previous recommendations intended for all types of kilns. We speculate that some PM_{2.5} is scavenged in the walls of dirt kilns.

1 Introduction

In developed countries most of the urban combustion emissions are due to burning fossil fuels. Fossil fuel emissions are also a major fraction of the air pollution in the urban areas of developing countries. However, in the developing world, the urban regions also have embedded within them numerous, small-scale, loosely regulated combustion sources due to domestic and industrial use of biomass fuel (biofuel) and the burning of garbage and crop residues. The detailed chemistry of the emissions from these sources has not been available and the degree to which these emissions affect air chemistry in urban regions of the developing world has been difficult to assess. As an example, we note that Raga et al. (2001) reviewed 40 years of air quality measurements in Mexico City (MC) and concluded that more work was needed on source characterization of non fossil-fuel combustion sources before more effective air pollution mitigation strategies could be implemented. The 2003 MCMA (Mexico City Metropolitan Area) campaign (Molina et al., 2007) and the 2006 MILAGRO (Megacity Impacts Local and Global Research Observations) campaign (Molina et al. 2008) focused on fixed-point monitoring of the complex MCMA mix of pollutants at heavily

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instrumented ground stations and on airborne studies of the outflow from the MCMA region. Explicit source characterization for biomass fires in the MCMA region was part of MILAGRO 2006, but only for landscape-scale open burning (e.g. forest fires in the mountains adjacent to MCMA, Yokelson et al., 2007).

5 Our 2007 ground-based MILAGRO campaign employed an approach that was complementary to most of the earlier work. With a highly mobile suite of instruments, we actively located representative sources of biofuel and garbage burning throughout the MCMA and central Mexico and measured the initial trace gas and particle emissions directly within the visible effluent plumes of these sources. The results should help
10 interpret the data from both the fixed monitoring stations in the MCMA (e.g. T_0 , T_1 , T_2 , etc.) and from aircraft in the outflow (Molina et al., 2008). Our source characterization also has global significance due to the widespread occurrence of these sources throughout the developing world as summarized next.

Recent global estimates of annual biofuel consumption include 2897 Tg/y (Andreae and Merlet, 2001) and 2457 Tg/y (Fernandes et al., 2007), making it the second largest
15 type of global biomass burning after savanna fires. An estimated 80% of the biofuel is consumed for domestic cooking, heating, and lighting mostly in open cooking fires burning wood, agricultural waste, charcoal, or dung within homes (Dherani et al., 2008). The balance of the biofuel is consumed mostly by low-technology, largely unregulated, micro-enterprises such as brick or tile making kilns, restaurants, tanneries, etc. While
20 individual “informal firms” are small, their total number is very large, e.g. ~20 000 brick making kilns in Mexico (Blackman and Bannister, 1998). Thus, this “informal sector” of the economy accounts for over 50% of non-agricultural employment and 25–75% of gross domestic product in both Latin America and Africa (Ranis and Stewart, 1994; Schneider and Enste, 2000).
25

McCulloch et al. (1999) calculated the 1990 garbage production from the 4.5 billion people included in the Reactive Chlorine Emissions Inventory as 1500 Tg. Scaling to the current global population of 6 billion suggests that 2000 Tg is an approximate, present global value. If half of this garbage is burned in open fires or

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incinerators (McCulloch et al., 1999) and it is 50% C, it would add 500 Tg of C to the atmosphere annually. This is about 7% of the C added by all fossil fuel burning (Forster et al., 2007). This crude estimate is fairly consistent with data from the remote Pacific in which $11 \pm 7\%$ of the total identified organic mass in the ambient aerosol was phthalates, ostensibly from garbage burning (Table 6, Simoneit et al., 2004b). It is most economical to burn urban-generated garbage in, or near, the major population centers that produce it. In addition, an estimated 12–40% of households in rural areas of the US burn trash in their backyards (USEPA, 2006). Thus, most garbage burning occurs in close proximity to people, despite estimates that garbage burning is the major global source of some especially hazardous air toxics such as dioxins (Costner, 2005, 2006).

The burning of crop residue in fields is generally considered to be the fourth largest type of global biomass burning with estimates including 540 Tg/y (Andreae and Merlet, 2001) and 475 Tg/y (Bond et al., 2004). Because cities are often located in prime agricultural regions, they may expand into areas where crop residue burning is a major activity and is sometimes the dominant local source of air pollution (Cançado et al., 2006).

In this study we measured the initial emissions of 12 of the most abundant gases, and the aerosol speciation for elemental and organic carbon (EC, OC), anhydrosugars, Cl^- , NO_3^- , and 20 metals from domestic and industrial biofuel use, garbage burning, and crop residue fires. In the following sections the measurements are described in detail and the implications of selected results are discussed.

2 Experimental details

2.1 Source types and site descriptions

We ranged by truck and van from Mexico City ~ 100 km to the north, east, and southeast, and ~ 300 km to the west over the course of about one month in April–May of 2007. The emissions data presented here were obtained from the sources listed

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in Table 1. These include eight indoor open wood cooking fires, two indoor wood cooking fires in Patsari stoves, three charcoal making kilns (from two sites), three brick making kilns, four garbage burns in peri-urban landfills, and two barley stubble field burns.

5 All but one of the open wood cooking fire measurements were conducted in rural and semi-rural homes during actual cooking episodes. The cooking fire in the laboratory of the Interdisciplinary Group on Appropriate Rural Technology (GIRA) was a simulation using an authentic open cook stove and typical fuel wood. For six of the eight homes in which we sampled, the kitchen was housed in a separate building. For the other two,
10 the kitchen was part of the main dwelling with a wall separating it from the sleeping area. Ventilation in all cases was by passive draft through door and window openings, cracks in the walls between boards, and horizontal openings where roof meets wall. Six of the eight kitchens had a dirt floor, seven were constructed of wood and one of brick. A variety of biofuels were available to the homeowners, including wood, corn
15 cobs, corn stalks, and charcoal. The primary fuel in all these homes, and the fuel used in all the fires we measured, was oak or pine collected locally by hand. Cooking fires were built either directly on the ground within a ring of three rocks, or on a mud and mortar, u-shaped, raised open stove. In one instance the “stove” was a dirt-filled metal bucket with rocks on top. A typical food preparation regimen begins with a small,
20 hot, flaming fire to quickly boil a pot of water, which is then loaded with beans and set off to the side to simmer. As the fire begins to die back, the cook begins frying tortillas. Wood is fed gradually to the fire to maintain the right amount of heat and when the cooking ends the fire is generally snuffed out to conserve fuel. A cooking session might last several hours depending on how much food is needed in the next few days.
25 The sample lines of all the instruments were co-located at ~1 m above the fire over the course of the cooking operation. The cook and her youngest children typically remain inside the kitchen for as long as it takes to prepare the food.

The Patsari stove incorporates an insulated fire box that is vented to the outdoors by a metal chimney. It is the product of 15 years of work by GIRA and the Center for

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Ecosystems Research (CIECO) to improve stoves economically (Masera et al., 2005). The stove cuts fuel consumption “per cooking task” roughly in half so its widespread adoption could reduce the total emissions from biofuel use. The chimney provides an approximate 70% reduction in indoor air pollution (Zuk et al., 2007), which is the largest single factor causing mortality in children under five globally (Dherani et al., 2008). It is also of interest that reactions on the chimney surface could modify the emissions (Christian et al., 2007). The chimney does not eliminate all the indoor pollutants because the fire box has an open front that can leak emissions into the room. Also, the chimney emissions may at times be recirculated into the kitchen from outdoors. We sequentially measured first the kitchen air above the stove, and then the chimney emissions from two different Patsari stoves in Pátzcuaro. One was located in a rural kitchen and the other was a newer model located in the GIRA lab’s simulated kitchen.

We sampled three charcoal making kilns in a forested area between MC and Puebla. An excavation ~5 m in diameter is dug by hand and kindling (dry needles, leaves, and twigs) is laid down. Oak logs are stacked in the center and a network of interlaced green oak branches is placed over the top. The excavated dirt is then packed on top to complete the earthen kiln, which has about a dozen vents around the circumference. A kiln of this design yields 200–250 kg of charcoal in about eight days. The supporting oak branches burn away slowly and the kiln must be rebuilt once or more during its lifetime to prevent it from collapsing and smothering the fire. The two kilns at the Hueyitlapichco site were constructed on consecutive days. We sampled them on their second and third day of operation on 19 April, and on their third and fourth day of operation on 20 April. At the San Gaspar site we sampled a single kiln on its fifth day of operation.

Brick making kilns in central Mexico are constructed from bricks. The fire bed and base walls are permanent and often built at the bottom of an excavation, which provides some insulation for the fire bed. There are several large, permanent mortar or concrete “crossbeams” above the fire bed. Green bricks are stacked to a height of several meters

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on the crossbeams (spaced to allow even heat circulation). Brick walls and a roof are then built up around the whole assembly. A fire is lit and fuel is shoveled or thrown in until the desired temperature is reached. Fuel is then added, as needed, to maintain that temperature around the clock for 2–3 days. At varying times each kiln operator uses mortar to seal the walls and most of the roof. Some owners allow the kiln to ventilate freely through the walls and roof for a day before sealing with mortar, claiming this gives a more uniform bake. Others seal the walls and roof before ignition. Kilns number 1 and 2 were burning fuel that was mostly wood waste products that had been hauled onto the site by dump truck. About 90% of this fuel was sawdust by volume. The remainder was divided fairly evenly between wood scraps, plywood, and particle board. A small fraction (less than 1%) was paper and cardboard. Brick kiln number 3 was using only scrap lumber while we made measurements. (We were unable to visit a fourth kiln near Silao that was reportedly burning used motor oil for fuel and a fifth kiln near Salamanca that was burning domestic waste scavenged from a nearby landfill.) The raw material for bricks is soil carved by hand from the ground in the vicinity of the kiln. The soil is mixed with water and manure or other organic waste and stomped barefoot to form a thick paste. The paste is then pressed into a mold and overturned one by one into rows to dry in the sun. Once they are dry enough to handle, the green bricks are stacked (in the shade if possible) and covered to prevent too rapid drying and cracking. Two of the brick kilns were sized to fire 10–12 000 bricks at a time; the third (brick kiln 2) was about three times larger. Kilns of this design are typical for Latin America and Africa, while more efficient designs – and coal fuel – are more common in Asia.

All four garbage burning fires were in the municipal landfills of peri-urban communities north of Mexico City. Only one landfill (Coyotepec, garbage fire 2) was burning when we arrived. This site was evidently excepted from a “no burn” policy that had been in place in northern Mexico state for some months. At the other three sites we ignited relatively small, representative sections of refuse under the direction of local authorities. The landfills held typical household and light industrial refuse. Plastic was by

far the most abundant material present. The following list is an approximate accounting of the composition of the waste stream for these landfills, in roughly diminishing order:

- plastic: bottles, bags, buckets, containers, toys, wrappers, Styrofoam
- paper: newspaper, magazines, cardboard boxes, food containers
- 5 – organic: fruit, vegetables (food waste)
- textile/synthetic fiber: cotton/nylon clothing, scraps
- rubber/leather: neoprene (in one case), sandals, shoes, scraps
- glass: bottles, jars
- vegetation: garden waste, brush, grass
- 10 – metal: soup cans, buckets, oil filters, aluminum foil
- ceramic: cups, dishes, cookware
- other waste materials

It appeared that tires were piled separately and perhaps not burned intentionally at the landfills. Wood was absent from any of the landfills since it is the most common cooking fuel in Mexico. The Tolcayuca landfill (garbage fire 3) was located in a textile manufacturing area and contained a higher proportion of textile waste than the other landfills. Each landfill was attended by people who manually removed items of value, including recyclable plastic bottles and cardboard. The scavenging process was less than perfectly efficient and small portions of the “collectible” waste did get included in the burns. All of our measurements were made from fires burning in the processed refuse from which the bulk of the recyclables had already been removed.

The agricultural waste burns took place in two adjacent, ~2 ha barley fields northwest of Salamanca. The fields had been mechanically harvested so all that remained were

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standing stalks (stubble, ~15 cm) and a mat of broken stalks and chafe, all of it tinder dry. Photographs of many of the field sites described above can be found at http://www.cas.umt.edu/chemistry/faculty/yokelson/galleries/album_Mex/index.html.

2.2 Instrumentation

5 The primary instrument for measuring trace gas emissions was our mobile, rolling cart-based Fourier transform infrared spectrometer (Fig. 2, Christian et al., 2007). It is rugged, easily transported, optionally self-powered, and can be wheeled to remote sampling sites. The optical bench is isolated from the chassis with wire rope shock absorbers (Aeroflex) and holds a MIDAC 2500 spectrometer, White cell (Infrared

10 Analysis, path length 9 m), MCT detector (Graseby), and transfer and focusing optics (Janos Technology). Continuous temperature (Minco) and pressure (MKS) sensors are mounted inside the cell. Other onboard features include a laptop computer, A/D and AC/DC converters, and a 73 amp hour 12 V battery. Sample air is drawn into the cell by an onboard DC pump through several meters of 0.635 cm o.d. corrugated Teflon tubing.

15 Sample air is trapped in the cell temporarily using manual Teflon valves while IR spectra are collected. We used nonlinear least squares, synthetic calibration (Griffith, 2002) to retrieve excess mixing ratios from the spectra for water (H₂O), carbon dioxide (CO₂), carbon monoxide (CO), methanol (CH₃OH), methane (CH₄), ethylene (C₂H₄), propylene (C₃H₆), acetylene (C₂H₂), formaldehyde (HCHO), and hydrogen chloride (HCl).

20 We used spectral subtraction (Yokelson et al., 1997) to retrieve excess mixing ratios for CH₃OH, C₂H₄, C₃H₆, C₂H₂, ammonia (NH₃), formic acid (HCOOH, also denoted FAC), and acetic acid (CH₃COOH, also HAC). At a path length of 9 m the detection limit for most gases was ~50–200 ppb. These gases accounted for all the quantifiable features in the IR spectra. The typical uncertainty for mixing ratios was ±10% (1σ). For CO₂, CO, and CH₄, the uncertainties were 3–5%. More complete descriptions of the system and spectral analyses are given in Christian et al. (2007).

25

After the campaign we checked for analyte losses in the FTIR cell due to adsorption or other reasons (Yokelson et al., 2003). The average NH₃ concentration in the cell

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during one minute of signal averaging (the typical sampling time used in Mexico) was about 71% of its initial level. The average HCl was ~93% of its initial level for the same interval. The ammonia and HCl results reported here have been adjusted upward to account for these cell losses.

5 A basic filter-sampling system with an internal pump (3 L min^{-1}) and an impactor was used to collect fire-integrated $\text{PM}_{2.5}$ on quartz filters. Analyses of the quartz filters were performed in the laboratories of the Research Center for Environmental Changes, Academia Sinica, Taipei, Taiwan. Organic and elemental carbon (OC, EC) were determined with a Sunset Laboratory, Inc. continuous carbon analyzer using
10 thermal-optical transmission (Birch and Cary, 1996; Engling et al., 2006). Anhydrosugars (levoglucosan, mannosan, galactosan) were determined using high-performance anion-exchange chromatography (HPAEC) with pulsed amperometric detection (Engling et al., 2006). Soluble ions were determined with ion chromatography (Hsu et al., 2008b). We analyzed the quartz filters for trace elements using inductively coupled
15 plasma-mass spectrometry (Hsu et al., 2008a; Hsu et al., 2008b).

We did not sample particles with Teflon filters, which are used for gravimetric determination of total $\text{PM}_{2.5}$. However, we did deploy an integrating nephelometer (Radiance Research M903) that measured particle light-scattering at 530 nm and 1 Hz. The nephelometer was calibrated with particle free zero air and CO_2 before and after
20 the campaign. The M903 nephelometer response was attenuated at the highest concentrations we encountered in Mexico. Thus, we applied a correction factor to those high values based on direct comparison in laboratory smoke between the M903 and a TSI 3563 nephelometer, which does have a sufficiently large linear range. The M903 nephelometer output ($b_{\text{scat}}, \text{m}^{-1}$) has been compared directly to gravimetric $\text{PM}_{2.5}$ determinations on cooking fires in both Honduras (Roden et al., 2006) and Mexico (Brauer et al., 1996). For dry, fine particles the conversion factor depends mostly on the EC/OC
25 ratio of the particles. Our average EC/OC ratio (0.284) for cooking fires was very close to that reported by Roden et al. (2006) for their cooking fires (0.267) Thus, we used the average of the two conversion factors from the other cooking fire studies to convert

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light-scattering data from our cooking fires to an estimated total $PM_{2.5}$ as follows:

$$b_{\text{scat}}(530 \text{ nm}, 273 \text{ K}, 1 \text{ atm}) \times 552000 \pm 75000 = PM_{2.5}(\mu\text{g}/\text{m}^3, 273 \text{ K}, 1 \text{ atm}) \quad (1)$$

(The conversion factor is equivalent to a mass scattering efficiency of 1.8). This approach probably gives an uncertainty in our average $PM_{2.5}$ for cooking fires of about 20–30%.

The particles from the other fires were different enough that we did not estimate a total $PM_{2.5}$ for these sources based on light scattering. However, we do report the mass sum of the particle constituents on the quartz filters. In this sum, we multiply the OC by a conservative factor of 1.4 to account for non-carbon organic mass (Aiken et al., 2008). The species measured include most of the major particulate components with the exception of sulfate and ammonium, which accounted for only a few percent of particle mass in other Mexican biomass burning particles (Yokelson et al., 2009). Thus, the sum of detected species is likely not more than 10–30% lower than the total $PM_{2.5}$.

We also deployed a CO_2 instrument (LICOR LI-7000) that was calibrated both before and after the campaign (negligible drift) with NIST-traceable standards spanning the CO_2 range encountered in the field. The CO_2 , nephelometer, and filter sampling systems shared a single inlet (conductive silicon tubing) that was often co-located with the FTIR sample line. In the cases where the FTIR mobility allowed sampling of the emissions at more points than the other instruments, the accurate determination of CO_2 by both the LICOR and the FTIR allowed coupling the two data sets. CO_2 was also used to correlate the particle measurements to the trace gases measured by FTIR as described in detail elsewhere (Yokelson et al., 2009, 2007).

2.3 Calculation of emission ratios and emission factors

An emission ratio (ER) is defined as the initial molar excess mixing ratio (EMR) of one species divided by that of another species, most commonly CO or CO_2 . EMR is simply the molar amount of a species above the background level and is designated with the

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Greek capital delta – e.g. ΔCO , ΔCH_4 , ΔX , etc. Modified combustion efficiency (MCE) is defined as the ratio $\Delta\text{CO}_2/(\Delta\text{CO}_2+\Delta\text{CO})$ and is useful for estimating the relative amounts of flaming and smoldering combustion during a fire, with high MCE corresponding to more flaming (Ward and Radke, 1993). To estimate the fire-average ER for a species “X” we plot ΔX for all the samples of the fire versus the simultaneously measured ΔCO (or ΔCO_2) and fit a least squares line with the intercept forced to zero. The slope is taken as the best estimate of the ER as explained in more detail in Yokelson et al. (1999). Figure 1 is an example of this type of plot showing the $\text{CH}_3\text{OH}/\text{CO}$ ER derived from 10 FTIR samples obtained over the course of a wood cooking fire.

An emission factor for any species “X” (EFX) is the mass of a species X emitted per unit mass of dry fuel burned (g compound per kg dry fuel). EF can be derived from a set of molar ER to CO_2 using the carbon mass balance method, which assumes that all of the burned carbon is volatilized and that all of the major carbon-containing species have been measured. It is also necessary to measure or estimate the carbon content of the fuel. For the fires using biomass fuel we assumed a dry, ash-free carbon content of 50% by mass (Susott et al., 1996). For the garbage fires, which contained only some biomass, we estimated the relative abundance of the materials present from photographs. We then calculated the overall carbon fraction based on those proportions and carbon content estimates for each type of material (IPCC, 2006; USEPA, 2007). Table 2 shows that this procedure resulted in an overall carbon fraction of 40% for the combustible landfill materials. The EF calculations for a charcoal kiln are complex because the fuel carbon fraction increases with time. We used a procedure identical to that described in detail by Bertschi et al. (2003).

EFPM_{2.5} for the cooking fires were calculated by multiplying the fire-integrated PM_{2.5} to CO_2 mass ratio (gPM_{2.5}/gCO₂ as measured by the nephelometer and LICOR) by the EFCO₂ (gCO₂/kg dry fuel as measured by FTIR). A similar method was applied to individual particle species based on net mass loading of fire-integrated filters, volumetric flow, and EFCO₂.

3 Results and discussion

3.1 Cooking fires

Trace gas ER and EF and particle EF based on light scattering for our cooking fires are given in Table 3. The first 10 columns of data are the eight open wood cooking fires plus a column each for the average and standard deviation. The next three columns are the EF and average for the two Patsari stoves as sampled in the kitchen. The last three columns are the analogous data from the outdoor chimney exhaust of the same two Patsari stoves. The EF for individual particle species measured on the quartz filters are given for all the fires in Table 4. Open wood cooking fires are the main global type of biofuel use and we get an idea of the global variability in this source by comparing EF from selected studies for some of the more commonly measured emissions (CO_2 , CO , CH_4 , and PM).

Figure 2 shows EFCH_4 versus MCE (a function of CO and CO_2) for those studies, including this one, where CO , CO_2 , and CH_4 data were all available. (The data in Fig. 2 have been scaled to reflect 50% fuel carbon content by mass.) A range of MCE from about 0.90 to 0.98 (avg 0.946) occurs naturally for individual fires in these studies. This leads to about a factor of 10 variation in EFCH_4 for individual fires, but the study-average values agree reasonably well. Some notes about the studies included in Fig. 2 follow. The Johnson et al. (2008) study was conducted in the same villages in Michoacán where the majority of our cooking fires were sampled. The authors sampled eight open cooking fires and 13 Patsari stoves and reported fire-integrated trace gas emission factors based on gas chromatographic analysis of smoke collected in Tedlar bags over the course of each fire. Zhang et al. (2000) set up a simulated kitchen in China and, using similar sampling methods as Johnson et al., reported fire-integrated emissions from a series of traditional stove types with various common fuels. The Zhang et al. (2000) data in Fig. 2 include only their wood and brush fuel types. Bertschi et al. (2003) reported the average EF for 3 open wood cooking fires in a village in Zambia. Brocard et al. (1996) reported the average EF for 43 open wood cooking fires on

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the Ivory Coast. The Andreae and Merlet (2001) data point is a widely-used global estimate derived from the literature. The Bertschi et al. (2003) EFCH₄ appears higher than the trend and the Brocard et al. (1996) EFCH₄ lower, but these data are consistent with a tendency toward greater variability as the relative amount of smoldering emissions increases in biomass burning fires (Christian et al., 2007; Yokelson et al., 2008).

Particle EF also vary substantially with MCE as seen in Fig. 3, which includes EFPM from three of the same studies that are included in Fig. 2 (Andreae and Merlet, 2001; Johnson et al., 2008; Zhang et al., 2000), as well as two other relevant studies (Roden et al., 2006, 2009). Roden et al. used a combination of nephelometry, absorption photometry, filter collection, and CO/CO₂ instrumentation to measure real-time and fire-integrated EF from 56 fires in various stove types in rural Honduran homes, and 14 laboratory simulations in several stove types. Figure 3 incorporates only their data from 10 traditional, open wood cooking fires in homes. (CO₂ data for calculating MCE for the two Roden et al. (2006, 2009) studies were kindly provided by the authors.) Again there is considerable variability in EF for individual fires, but reasonable agreement between authors on the range and trend with MCE. This body of work on PM suggests a slightly lower average MCE (0.927) than implied in Fig. 2. If we assume a global average MCE in the range ~0.93–0.94, then the trend lines imply global average EF for open wood cooking fires of 4.5±1.4 for CH₄ and 5.7±2.7 for PM. A larger uncertainty in global average EF would result by considering more of the less common fuels (agricultural waste, dung, etc.) and stove types.

For compounds that are major open cooking fire emissions, but difficult to measure by non-spectroscopic methods (CH₃COOH, NH₃, HCHO, CH₃OH, HCOOH), we can compare our current EF from Mexico only to those obtained by open-path FTIR on African open wood cooking fires by Bertschi et al. (2003). The Bertschi et al. (2003) EF were measured at a lower average MCE (0.91) than the average MCE for our fires in Mexico (0.95) and thus, not surprisingly the EF for the smoldering compounds in Bertschi et al. are generally about 2–4 times higher. Averaging the results from these two FTIR-based studies is consistent with the average MCE for cooking fires of ~0.93

derived above.

As mentioned above, the use of improved stoves with chimneys and insulated fire boxes reduces both the total biofuel emissions (due to reduced fuel consumption) and the indoor air pollution. There is also potential for improved stoves to consume the fuel at higher MCE, reducing the EF for smoldering compounds. A further possibility is that the surface of the chimney could scavenge some of the more reactive smoke components before they are emitted to the airshed. To examine these issues we compare the average MCE and EF of the Patsari chimney exhaust to the average MCE and EF for the open fire emissions. The average MCE was lower from our open fires (~0.95) than it was from our Patsari chimney exhaust (0.97). Consistent with the increased Patsari MCE, the EF for CO, CH₄, and the measured NMOC (with the exception of organic acids, C₃H₆, and C₂H₂) were about a factor of two lower from the chimney exhaust. For organic acids, NH₃, and C₃H₆ there was a larger drop (80–95%) in the EF measured from the chimneys that was likely due in large part to losses on the chimney walls. EFC₂H₂ is similar for both sources as it is emitted by both flaming and smoldering (Yokelson et al., 2008) and is not particularly “sticky.” Overall, while only a fraction of the total NMOC emitted could be measured (Yokelson et al., 2008), the sum of the EFNMOC that were measured in this study from the chimney was ~38% of the analogous sum from the open fires. We were unable to measure particle EF from the Patsari chimney. Johnson et al. (2008) also compared EF for open fires to EF for Patsari stoves in their Table 1 (bottom 3 rows). Their data show an increase in MCE from 0.92 (open) to 0.98 (Patsari). They also reported a large reduction in the EF for CO, CH₄, and PM, which was variable depending on the type of Patsari stove sampled. Based on the above, it appears that improved stoves could reduce both fuel consumption (by about half, Masera et al., 2005) and the amount of many pollutants emitted per unit mass of fuel consumed (by at least half).

There is a significant absence in Table 3 of HCN, which is widely used as a biomass burning tracer (Yokelson et al., 2007). HCN is normally well above the detection limits of our FTIR systems for landscape-scale biomass burning (e.g. forest fires, grass fires,

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Yokelson et al., 2007). However, HCN was below our FTIR detection limits for cooking fires in both Africa (Bertschi et al., 2003) and Mexico (current study). A single FTIR sample from a Brazilian stove (Christian et al., 2007) did contain some HCN, but the ER to CO (0.0005) was ~24 times lower than the value for Mexico City area forest fires (0.012, Yokelson et al., 2007). The low HCN/CO ER for cooking fires means that where these fires are common, the biomass burning contribution to total pollution will be underestimated if it is based on an HCN/CO ER appropriate for landscape-scale burning (Yokelson et al., 2007).

Acetonitrile is another useful biomass burning tracer (de Gouw et al., 2001), but cooking fire measurements for this species have not been attempted yet. However, since acetonitrile emissions from other types of biomass burning are usually less than half the HCN emissions (Yokelson et al., 2009), they may also be unusually small from cooking fires. Methyl chloride (CH₃Cl) has also been linked to biomass burning (Lobert et al., 1991), but its emissions are probably much smaller from cooking fires than for other types of biomass burning since wood has much lower chlorine content than other components of vegetation (Table 4, Lobert et al., 1999). Levoglucosan and K (in fine particles) are also used as biomass burning indicators and they were observed in “normal” amounts in the particles from our cooking fires (Table 4) compared to other types of biomass burning. However, as discussed in more detail in Sect. 3.2, levoglucosan and K were also present in similar amounts in the fine particles from garbage burning. Thus, in areas such as central Mexico where garbage burning is common it could contribute a significant fraction of the aerosol levoglucosan or K. The lack of a straightforward chemical tracer for cooking fires is especially significant since these fires will also not be detected from space as hotspots or burned area. In addition, the CO could be underestimated by MOPITT due to the low injection altitude for cooking fire smoke (Emmons et al., 2004) and the short (one-month) lifetime for CO in the tropics. Thus, biomass burning estimates based on HCN or acetonitrile likely underestimate cooking fires (and total biomass burning), while estimates based on levoglucosan or K could be subject to “interference” from garbage burning in parts of the developing

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world. In summary, while survey-based research clearly indicates that biofuel use is the second-largest global type of biomass burning, there is not a simple chemical tracer to confirm this or to independently determine the amount of biofuel use embedded in urban areas of the developing world.

3.2 Garbage burning

Our ER and EF for trace gases emitted by garbage burning are shown for individual fires in the left half of Table 5. Garbage fire 2 had already progressed to mostly smoldering combustion when we arrived. At the other three fires we sampled mostly flaming. Since we don't know the real overall ratio of flaming to smoldering combustion for landfill fires we just calculated the straight average and the standard deviation for all four fires. For the trace gas EF this implies a 3:1 ratio for fuel consumption by flaming/smoldering, which does not seem unreasonable. The EF are computed assuming the waste in these landfills was 40% C by mass. If the %C is higher or lower the real EF would be higher or lower in direct proportion. It is important to note, however, that the ER to CO or CO₂ are independent of any assumptions about the composition of the fuel. The EF for particle species are included in Table 4. Since we only have filter data for three of the four garbage fires, an average of the filter results is equivalent to one-third of the fuel being consumed by smoldering.

We could not find any published, peer-reviewed, direct emissions measurements from open burning in landfills to compare our results to. Data from airborne and ground-based measurements of aerosols over the east Asian Pacific as part of ACE-Asia (Simoneit et al., 2004a, b) revealed significant levels of phthalates and n-alkanes in the aerosols. The presence of these compounds was attributed to refuse burning. A follow up study confirmed these compounds as major organic constituents in both solvent extracts of common plastics and the aerosols generated by burning the same plastics in the laboratory (Simoneit et al., 2005). This indicated their potential usefulness as tracers. However, these are high molecular weight, semi- or non-volatile compounds whose relationship to volatile gaseous emissions is not known.

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The comparison of the garbage burning emissions to biomass burning emissions is interesting. The average ethylene molar ER to CO for garbage burning ($\Delta C_2H_4/\Delta CO$, 0.044) is 3–4 times higher than for our open wood cooking fires (0.013, Table 3) or forest fires near Mexico City (0.011, Yokelson et al., 2007) and is likely a result of burning a high proportion of ethylene-based plastic polymer fuels.

HCl is not commonly detected from biomass burning (Lobert et al., 1999), but the EF_{HCl} in the garbage burning emissions ranged from 1.65 to 9.8 g/kg, a range similar to that for CH₄ in biomass burning emissions. Lemieux et al. (2000) reported a strong dependence on PVC content for HCl emissions from simulations of domestic waste burning in barrels. Their EF_{HCl} was 2.40 g/kg ($n=2$) for waste containing 4.5% PVC by mass, and 0.28 g/kg ($n=2$) for waste with only 0.2% PVC. There was no mention of precautions taken to avoid passivation losses on sample lines, etc. (e.g. Yokelson et al., 2003). In the current study, significant additional chlorine was present in the particles; EF for soluble Cl⁻ alone ranged from ~0.2 to 1.03 g/kg fuel (Table 4). Studies of landfills in the European Union found that the chlorine content of solid waste was about 9 g/kg (Mersiowsky et al., 1999) and that essentially all the chlorine was present as polyvinyl chloride (Costner, 2005), which is 57% Cl by mass. We found that burning “pure” PVC in our laboratory produced HCl/CO in molar ratios ranging from 5:1 to 10:1. Thus, the observed molar ER for HCl/CO in the MCMA landfill fires (0.037–0.19) are consistent with the burning materials we sampled containing ~0.4–4% PVC. Our results also suggest that the majority of the chlorine in burning PVC is emitted as HCl.

Even though the average EC/OC ratio for garbage burning (0.232, $n=3$) is close to that for the cooking fires (0.284, $n=5$), application of the cooking fire conversion factor to the garbage burning light scattering data underestimates the particle mass compared to summing the particle species data. Preliminary work in our lab suggests this could be due to a shift to larger particles in the emissions from burning plastics. We can roughly estimate the EF_{PM_{2.5}} for garbage burning from the particle species data. The sum of the measured particle components averaged 8.74 ± 7.35 g/kg, which, after allowing for unmeasured species, suggests that the EF_{PM_{2.5}} is about 10 ± 5 g/kg.

The average EF_{PM_{2.5}} reported by Lemieux et al. (2000) for burning recycled and non-recycled waste in barrels was 11.3±7.5. The USEPA recommended EFPM for open burning of municipal waste is 8 g/kg (AP-42, USEPA, 1995) based on two laboratory studies from the 1960s (Feldstein et al., 1963; Gerstle and Kemnitz, 1967). We note that the AP-42 recommendations for CO (42 g/kg) and CH₄ (6.5 g/kg) are reasonably close to our values of 45.3±22.8 and 3.7±4.4, respectively. AP-42 also recommends values for SO₂ (0.5 g/kg) and NO_x (3 g/kg).

The EF for EC, OC, levoglucosan, and K for garbage burning had a similar range to the EF for these species for the cooking fires. Levoglucosan is produced from the pyrolysis of cellulose and the landfills contain a lower fraction of cellulose than biomass. However, the levoglucosan emissions per unit mass of paper burned can be considerably higher than those from burning some types of biomass (Table 1, Simoneit et al., 1999). In our data, the average levoglucosan EF from garbage burning is 85% of the EF for cooking fires, which would make it difficult to use levoglucosan to distinguish between these two sources. The other sugars analyzed in this work (mannosan and galactosan) showed more potential promise in this respect as their EF were ~90% lower for garbage burning than for cooking fires. Finally, the garbage burning EF for mannosan was only ~12% lower than the single mannosan EF measurement for crop residue burning. This tentatively leaves galactosan as the most promising sugar of those we analyzed to indicate general biomass burning in the presence of garbage burning.

The garbage burning EF were the most different from the biomass burning EF for numerous metals. With correction for local soil composition, some of these metals could ultimately offer a useful method of assessing the garbage burning contribution to overall air quality. For example, the ratio EF_{garbage}/EF_{cook} for selected particle species was: Sb (555.7), Pb (211.7), Sn (181.9), Cl⁻ (63.7), Cd (33.57), As (20.9), Ca (5.1), and Mg (4.6). We note, however, that the soluble chloride in the one sample of crop residue burning smoke was actually higher than the average value for garbage burning. This could reflect the use of chlorine-containing agricultural chemicals (Sect. 3.4). In

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examining the ratio of the average EF for garbage burning to the average EF for crop residue burning the most elevated metals are antimony and tin (Sb 309.4, Sn 33.6). Thus, initially Sb emerges as a promising tracer for garbage burning.

Both Sb and $PM_{2.5}$ were measured in the MCMA ambient air at T_0 and T_1 during MILAGRO (Querol et al., 2008). The mean mass ratio for Sb/ $PM_{2.5}$ for the March 2006 campaign at these sites was 0.000315. Our mean EF for Sb in $PM_{2.5}$ from pure garbage burning smoke was 0.011 ± 0.008 g/kg. Our estimate of the average $EF_{PM_{2.5}}$ for garbage burning is 10 ± 5 g/kg, implying an Sb/ $PM_{2.5}$ mean mass ratio of ~ 0.0011 for this source. Comparison of the mean mass ratios of Sb/ $PM_{2.5}$ for pure garbage burning and ambient air implies that garbage burning could account for about 29% of the $PM_{2.5}$ in the MCMA. However, we note that this estimate has high uncertainty and that the presence of Sb in the MCMA particulate was attributed by Querol et al. (2008) to the abrasion of tires and brake pads. In addition, higher Sb in the $PM_{2.5}$ might result if there were any manufacturing operations using Sb near the measurement sites. However, our initial crude estimate suggests that garbage burning deserves more attention as a potentially significant contributor to the particle burden of the MCMA airshed. A more rigorous source attribution for garbage burning based on fine particle metal content would require a more complex multi-element approach. The main uses of antimony are as a flame retardant for textiles and in lead alloys used in batteries. Antimony trioxide is a catalyst that is often used in the production of polyethylene terephthalate (PET) and that remains in the material. PET is the main material in soft drink bottles, polyester fiber for textiles, Dacron, and Mylar. The smoke particles from the dump with the highest percentage of textiles (Table 4, garbage fire 3) did have the highest mass percentage of Sb. We noted earlier that at least some of the PET materials (soft drink bottles) were being recycled rather than burned.

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3.3 Industrial biofuel use: brick and charcoal making kilns

3.3.1 Brick making kilns

The particle and trace gas emissions data for brick kilns are in Tables 4 and 5, respectively. The brick kilns we sampled burned mostly biomass fuels and the identities of the emitted NMOC were similar to those from biomass burning. The brick kiln EF were much reduced, likely due to the high MCE and to scavenging by the kiln walls and/or the bricks themselves. It is hard to say how well the emissions from these kilns represent brick making kilns in general because informal industries like brick kilns often burn a combination of biofuel, garbage, painted boards, tires, used motor oil, etc. Though our kilns burned mostly biofuel they emitted a much blacker smoke than any other biomass burning we have observed (EC/OC 6.72, $n=2$). All the photographs of brick making kilns we took and could locate elsewhere showed very black smoke emissions. The high EFCI⁻, but low Sb and other metals for brick kiln 1 suggests that crop waste may have been a fuel component during our measurements or during past uses of the kiln. The elevated Pb from both kilns 1 and 2 may be due to burning painted boards from demolished buildings. Painted boards were identified as a controversial fuel used in some Mexican brick kilns in a report to the USEPA by James Anderson of Arizona State University (<http://www.epa.gov/Border2012/>).

The EFPM_{2.5} must be quite low from our brick kilns as the sum of the species on the two kiln filters was 1.24 and 1.96 g/kg, respectively. Some of the particles being produced in the fire-box may be deposited on the bricks and kiln walls. Despite the low particle emission factors for these kilns, brick making kilns are known to cause locally severe air quality impacts in Mexico as documented by Anderson, who reported PM₁₀ in homes and an elementary school near brick kilns well above 1000 $\mu\text{g}/\text{m}^3$. Blackman et al. (2006) reported that the 330 brick making kilns in Ciudad Juarez (population 1.2 million) produced 16% of the PM and 43% of the SO₂ in the urban airshed. A large reduction in the total emissions from brick kilns is possible at the regional-national

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scale by switching to more fuel efficient designs such as the vertical shaft brick kiln (<http://www.vsbkindia.org/faq.htm>).

To our knowledge, there are no other published data on trace gas and particle emissions for brick making kilns that use wood or cellulose-based waste products as the primary fuel. An inventory of China's CO emissions was constructed following the Transport and Chemical Evolution over the Pacific (TRACE-P) campaign of 2001 (Streets et al., 2003). Those data were recently reevaluated to include a much larger contribution from coal-fired brick kilns (Streets et al., 2006). In a modeling study of aerosol over south Asia, a lack of seasonal variability for Kathmandu was credited to the exclusion of brick kiln emissions from the model (Adhikary et al., 2007). Nepalese kilns are also fueled primarily by coal.

The impact of industrial biofuel use will likely remain difficult to assess for some time. The diverse range of micro-enterprise fuels (biomass, motor oil, tires, garbage, propane, coal, crop residues, etc.) makes it difficult to envision a tracer-based method that would quantitatively retrieve the contribution of this sector of the economy. Survey-based methods, which likely work well for household biofuel use, may be less accurate when applied to highly competitive enterprises operating on thin margins. For example, in the report by Anderson cited above, stockpiled tires were a common sight at brick kilns. However, 100% of owners surveyed responded that they never burned tires while 12% responded that other kiln owners did.

3.3.2 Charcoal making kilns

The particle and trace gas emissions data for the charcoal making kilns are in Tables 4 and 6, respectively. As noted by Christian et al. (2007) the chemistry of the kiln emissions changes over the course of the approximately one-week production cycle. Specifically the molar ratio of total (measurable) VOC to CO increases by about a factor of 8 over this time. Thus, it is most meaningful to compare measurements from the same point in the production cycle, which we have done in Fig. 4. The pattern of increasing total $\Delta\text{VOC}/\Delta\text{CO}$ for Mexico is fairly similar to the trend measured on charcoal

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kilns in Brazil (Christian et al., 2007) and Africa (Bertschi et al., 2003).

Table 6 also includes a simple comparison between the average of the charcoal making EF measured on days 3–5 in Mexico and the average EF for days 1–4 from a Zambian charcoal kiln (Bertschi et al., 2003). The Mexico MCE is slightly higher than the Zambian MCE, but the Mexican EF for acetic acid was 360% higher. The higher acetic acid could be due to oak species being the primary fuel type in Mexico. Tannin content is high in oak species and, as polyphenols, tannins contain abundant hydroxyl and acetyl functional groups. As such they seem likely precursors for pyrolytic formation of acetic acid.

Andreae and Merlet (2001) recommended an EFPM_{2.5} for charcoal making of 4.0 g/kg. The sums of the EF for our PM_{2.5} species in the two kiln filters were below that at 0.56 and 1.65 g/kg, respectively (Table 4). While a thick white smoke emanates from the vents in the kiln walls, much of the particulate matter produced inside probably remains embedded in the dirt walls while most of the gases are vented.

3.4 Crop residue burning

Our emissions data for crop residue fires are in Tables 4 and 5 in the columns labeled “stubble burns.” Yokelson et al. (2009) reported airborne EF measurements for six crop residue fires in the Yucatan peninsula. Since airborne platforms tend to sample smoke with a higher flaming/smoldering ratio than ground-based platforms, we expect that the airborne samples will have a higher MCE and lower EF for smoldering compounds and PM_{2.5}. This pattern is observed. The airborne average MCE was 0.934, while the ground-based average MCE was 0.896. The airborne average EF for smoldering compounds (excluding formic acid) and PM_{2.5} were 65±12% of the average EF measured from the ground. The formic acid average EF was higher for the fires sampled from the air, possibly due to fuel differences. The type of crop residue burning could not be identified from the air. In the future, after analyzing additional airborne samples of crop residue fires from our flights in Mexico, we plan to recommend EF weighted by the relative fuel consumption for flaming and smoldering combustion.

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Table 4 includes a very low EC/OC ratio for the one filter sample of stubble burning smoke. K and Na were very high on this filter and are known to catalytically lower the combustion temperature of black carbon during thermal evolution carbon analysis methods (Martins et al., 1998; Novakov and Corrigan, 1995). The high chloride content in the stubble burn filter may be linked to the use of agricultural chemicals. Typical examples of these products include the herbicide 2,4-D dimethylamine salt (26.6% Cl by mass), the fungicide chlorthalonil (1,3-dicyanotetrachlorobenzene, 53% Cl), and the pesticide Lindane (or Kwell, hexachlorocyclohexane, 73% Cl). Two fires were sampled by the NCAR C-130 during MILAGRO in fuels that could not be identified from the air (fires #1 and #3 sampled on 23 March 2006, Yokelson et al., 2009). The particles emitted by these fires were high in both Cl^- and NO_3^- so they were probably crop residue fires. Fire plays a role in redistributing agricultural chemicals from their original point of application and they then build up in fire-free areas such as the Arctic (Becker et al., 2009).

4 Implications

In this section we discuss the impact of the sources sampled in this work at various scales. For cooking fires we start with national scale assessments for two different scenarios: a mostly rural developing country (Zambia) and a mostly urban developing country (Mexico).

4.1 Cooking fires compared to open burning in a mostly rural developing country

Zambia has a total population of 10 million, of which 60% is rural. As much as one-third of the area of the country can burn in open fires in a single year which, combined with the small population, suggests that cooking fires would be much less important than open burning. However, standard estimation methods reveal otherwise. In Fig. 5 we

compare the dry season emissions from open fires and biofuel use based on the following assumptions: 18.6 Tg/y biomass burned in open fires, 6.4 Tg/y biomass burned in wood cooking fires, and 0.75 Tg/y charcoal use (Bertschi et al., 2003). EF for these combustion types, specific to Zambia when available, are from several sources (Andreea and Merlet, 2001; Bertschi et al., 2003; Christian et al., 2003; Sinha et al., 2004; Yokelson et al., 2008). We divide the annual biofuel emissions by two to estimate the dry season cooking fire emissions, and all the annual open burning emissions are assumed to be generated in the dry season. From Fig. 5 it is apparent that the dry season cooking fire emissions equal or exceed the dry season open burning emissions for four of the 17 species measured from both sources. For 11 of the 17 species, the dry season cooking fire emissions are at least 33% of the total dry season pyrogenic emissions. Thus, for those 11 species the annual emissions are estimated to be larger from cooking fires.

4.2 Cooking fires compared to urban emissions in a mostly urban developing country

Mexico's total population of 100 million is 75% urban. Mexico City (population 20 million, ~double the entire country of Zambia) is the second largest Megacity on earth and an acknowledged major source of pollutants. Mexico has experienced strong rural to urban migration and is considerably more developed than Zambia. We roughly estimate annual biofuel use in Mexico using three tables in Yevich and Logan (2003). From their Table 12 Mexico accounts for 13% of biofuel use in Latin America on an energy basis. From their Table 13 the annual biofuel consumption for Latin America is 358 Tg. And from their Table 16 they estimate a 20% increase in biofuel use every 10 years. We applied this increase to the values from the other tables, which were based on 1988 data. In this way we arrive at 68 Tg/y of biofuel consumption for Mexico, which is mainly cooking fires. We are unsure to what extent this estimate may or may not include potentially substantial industrial biofuel use. We obtained speciated annual emissions for the Mexico City Metropolitan Area (MCMA) from the 2004 MCMA emis-

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sions inventory (<http://www.sma.df.gob.mx/sma/index.php?opcion=26&id=392>). We multiply the MCMA emissions by 75/20 to roughly estimate total annual urban emissions for Mexico. Since both urban and cooking emissions are year round we do not separately calculate dry season emissions because the ratio between the sources would not change. Table 7 summarizes this simple comparison and suggests that ~2 times more NMOC are generated from cooking fires than from urban areas. In addition, PM_{2.5} is estimated to be almost 20 times greater from cooking fires than from urban areas on a national basis. Even if we allowed for a higher degree of secondary aerosol formation in fossil fuel emissions, which may not be the case, the cooking fires clearly dominate. If these estimates are right, then a switch to cleaner burning, more efficient stoves (Sect. 3.1) could provide an enormous reduction of emissions on the national scale.

There is also a possibility that the amount of biofuel use in urban areas is underestimated. Marley et al. (2009) reported that 70% of the carbon in the ambient MCMA aerosol was modern and ascribed this to open biomass burning and garbage burning. However, garbage burning consumes a large fraction of plastics made from fossil fuels. If open burning was the dominant particle source in the MCMA and ventilation rates were similar year round, the PM₁₀ levels should peak in March-May when nearly all the open biomass burning occurs. Instead the PM₁₀ data show at best a weak increase in PM₁₀ during these months (Fig. 6) indicating that a different, year round source of modern carbon could be “embedded” in the urban area. Two possibilities are cooking fires and industrial biofuel use.

4.3 Garbage burning impacts on the local-global atmosphere

We start this section by noting that the prevalence of open burning of garbage may be greater than commonly supposed even in developed countries. As noted earlier, it has been estimated that 12–40% of rural households in the US burn garbage in their backyards (USEPA, 2006). In the US, dump and landfill fires are reported at a rate of 8 400 fires per year (TriData.Corp., 2002). UK landfill operators surveyed by

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Bates (2004) estimated that, at any one time, deep seated fires are occurring at about 80% of landfills.

If we assume that 1000 Tg of garbage with a Cl content of 0.9% are burned each year globally (see introduction and Sect. 3.2), this could volatilize as much as 9 Tg/yr of Cl. This suggests a potential global source of 6–9 Tg/yr of HCl. By comparison, the Reactive Chlorine Emissions Inventory (Keene et al., 1999) estimated that HCl from garbage burning was 2 Tg/yr and that sea salt dechlorination was the main global source at 7.6 Tg/yr. Recent HCl profiles in the marine boundary layer (Kim et al., 2008) may indicate that the sea salt dechlorination HCl source was over estimated. Our measurements indicate that the garbage burning HCl source may have been underestimated. In general, Keene et al. (1999) found that additional HCl sources totaling to 42 Tg/yr were needed to balance the HCl budget. With the above in mind, we propose that garbage burning may be a considerably more important tropospheric source of HCl than previously assumed. We also note that many of the other main HCl sources, such as sea salt and volcanoes, can often be associated with a humid environment and rapid removal of HCl (Tabazadeh and Turco, 1993). In dry environments, such as central Mexico where we measured water mixing ratios as low as 890 ppm, a larger fraction of freshly emitted HCl might react with OH to release Cl atoms. The latter would then react with NMOC. In any case, the HCl from garbage burning in dryer areas could have a longer lifetime and higher relative importance than the same amount of HCl emitted in wetter areas.

We examined data obtained by other MILAGRO investigators for possible evidence of garbage burning. A particles-into-liquid-sampler (PILS) deployed by Georgia Tech at the MILAGRO T1 ground station north of Mexico City during March 2006 observed significant levels of chloride (up to $6 \mu\text{g}/\text{m}^3$) for most of the month, with an average of $0.5 \mu\text{g}/\text{m}^3$ compared to $33 \mu\text{g}/\text{m}^3$ total $\text{PM}_{2.5}$ (Greg Huey, personal communication, 2009). This translates to a mass ratio of 0.015. The average mass ratio of Cl^- to the sum of particle species in our nascent smoke from garbage burning (Table 4) was 0.047 ± 0.011 . Thus, the PILS data is consistent with our Sb-based estimate that

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garbage burning could contribute about one-third of the $PM_{2.5}$ in the MC airshed. However, the PILS chloride may also reflect a contribution from agricultural fires and brick making kilns (Table 4) and volcanoes (e.g. Burton et al., 2007). We note that 3 of the 4 landfills we sampled are within ~ 35 km to the west, north, and east of the T1 site (Table 1). We also note that $EFCI^-$ for brick kiln 1 was high and that this kiln is only ~ 20 km west of T1. In addition, brick kiln 1 was one of many brick kilns in the region.

We also looked for evidence of chlorine atom chemistry in the hydrocarbon ratios measured by whole air sampling. A plot of i-butane versus n-butane for 62 canister samples collected from both airborne and ground based sampling locations in and around MCMA gave an average i-butane/n-butane ratio of 0.33 ($r^2=1.00$, Don Blake, Barbara Barlett, personal communication, 2009). This is consistent with minimal chlorine atom oxidation of alkanes in the air sampled (Kim et al., 2008).

We make two other general points about garbage burning. More work is needed to measure other chlorinated emissions from burning refuse, including CH_3Cl , which is also a proposed biomass burning tracer (Lobert et al., 1991). Secondly, PVC (the primary source of HCl in garbage burning emissions) is also the most important predictor of dioxin emissions from the open burning of domestic waste (Neurath, 2004), so removing PVC from the waste before burning should have multiple benefits.

5 Conclusions

This work measured initial emission ratios and emission factors for trace gas and particle species from five prominent, little-studied combustion sources: wood cooking fires, garbage burning, brick and charcoal making kilns, and crop residue burning. For Zambia, a mostly rural developing country, annual emissions from cooking fires likely exceed those from savanna fires for ~ 11 of the 17 most abundant species measured from both sources. For Mexico, a mostly urban developing country, NMOC from cooking fires are estimated to be substantially higher than from urban fossil fuel sources. Furthermore, cooking fires emit far more primary $PM_{2.5}$ than urban fossil fuel sources on a

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national scale in Mexico. Therefore, the use of improved stoves that reduce both total fuel consumption per cooking task and the pollutant emissions per unit fuel consumption could provide large reductions in pollutant emissions throughout the developing world. Cooking fires produce far less HCN per unit mass of fuel than landscape-scale open biomass burning. Thus, the use of HCN ER or EF appropriate for open burning to estimate regional biomass burning could cause a substantial underestimate of total biomass burning in areas where cooking fires are common (most of the developing world). Acetonitrile emissions from cooking fires are also likely to be much lower than for other biomass burning types. In addition, levoglucosan, K, and CH_3Cl are likely not suitable as biomass burning tracers in areas where garbage burning is common. Since cooking fires are also not detected from space a need exists to identify a chemical tracer for this source, which is currently quantified only by user surveys.

Garbage burning is a globally significant source of particles and trace gases and may be a major global source of HCl. The emissions of particle EC, OC, levoglucosan, and K were similar for garbage burning and cooking fires. Of the three anhydrosugars we measured, galactosan was the one most closely associated with biomass burning. Thus, it shows some promise for distinguishing between garbage burning and general biomass burning. Antimony (Sb) is a potential tracer for garbage burning. Comparison of both the Sb and the Cl^- mass concentrations in the ambient $\text{PM}_{2.5}$ sampled at points in the MCMA airshed with the mass concentration of these species in $\text{PM}_{2.5}$ in fresh garbage burning plumes tentatively suggests that landfill fires may produce about one-third of the $\text{PM}_{2.5}$ in some areas of the MCMA.

Wood-fueled brick making kilns emitted a suite of trace gases similar to those from biomass burning, but with much lower emission factors. Smoke from these kilns had a very high EC/OC ratio. Charcoal making kilns in Mexico, Brazil, and Africa exhibited a consistent pattern of increasing VOC/CO emission ratios over their approximately one week lifespan. The Mexican charcoal kilns produced higher acetic acid to CO ratios than an African kiln, possibly as a result of using tannin rich oak fuel. The $\text{PM}_{2.5}$ emission factor for both kiln types was evidently low relative to other biomass burning

types, possibly as a result of particle-scavenging on the kiln charge and walls. The fuels for brick making kilns vary substantially in ways that will likely remain difficult to quantify and limit the accuracy of regional air quality assessments and global emissions inventories. A single filter sample of smoke from a crop residue fire had very high levels of chloride that probably resulted from the use of agricultural chemicals. More research is needed to identify the nature and fate of the combustion products of agricultural chemicals.

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Table 1. Sampling source types and locations.

Type	Location	Date (2007)	Lat	Lon
Open cook	San Pedro Benito Juárez, Atlixco, Puebla	18 Apr	18.95	−98.55
Open cook	San Pedro Benito Juárez, Atlixco, Puebla	19 Apr	18.95	−98.55
Open cook	San Juan Tumbio, Michoacán	8 May	19.50	−101.77
Open cook	San Juan Tumbio, Michoacán	8 May	19.50	−101.77
Open cook	Comachuén, Michoacán	9 May	19.57	−101.90
Open cook	Comachuén, Michoacán	9 May	19.57	−101.90
Open cook	Comachuén, Michoacán	9 May	19.57	−101.90
Open cook	GIRA lab, Tzetzenguaro, Michoacán	10 May	19.53	−101.64
Patsari cook	GIRA lab, Tzetzenguaro, Michoacán	10 May	19.53	−101.64
Patsari cook	Rancho de Álvarez, Michoacán	11-May	19.54	−101.51
Charcoal kiln	San Gaspar de lo Bendito, Atlixco, Puebla	17 Apr	19.00	−98.54
Charcoal kiln ^a	Hueyitlapichco, Atlixco, Puebla	19 Apr	18.97	−98.56
Charcoal kiln ^a	Hueyitlapichco, Atlixco, Puebla	20 Apr	18.97	−98.56
Brick making kiln	Teoloyucan, Edo. México	24 Apr	19.77	−99.19
Brick making kiln	Barrio México 86, Edo. México	27 Apr	19.41	−98.91
Brick making kiln	Silao, Guanajuato	2 May	20.94	−101.42
Landfill	Soyaniquilpan, Edo. México	23 Apr	20.01	−99.49
Landfill	Coyotepec, Edo. México	24 Apr	19.81	−99.22
Landfill	Tolcayuca, Hidalgo	25 Apr	19.97	−98.92
Landfill	San Martín de las Pirámides, Edo. Mex.	26 Apr	19.70	−98.80
Barley stubble	Rancho de Don Ignacio, Guanajuato	30 Apr	20.60	−101.22
Barley stubble	Rancho de Don Ignacio, Guanajuato	1 May	20.60	−101.22

^a Two separate kilns at one location.

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Table 2. Estimate of the carbon content of Mexican peri-urban landfills.

Category	Relative proportion by volume ^a	Estimated mass fraction ^b	Carbon fraction ^c
plastic	0.65	0.30	0.74
paper	0.10	0.15	0.46
organic (food waste)	0.05	0.05	0.38
textile/synthetic fiber	0.05	0.05	0.60
rubber/leather	0.05	0.05	0.76
glass	0.02	0.05	
vegetation	0.01	0.05	0.50
metal	0.01	0.05	
ceramic	0.01	0.05	
other	0.05	0.20	
net	1.00	1.00	40%

^a Visual estimate of relative volumes of the most prominent waste materials from four Mexican landfills.

^b Rough estimate of relative mass for each material type.

^c Combined estimates from IPCC (2006) Table 2.4 and USEPA (2007) Annex 3 Tables A-125 to A-130.

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Table 3. Normalized emission ratios (ER, mol/mol) and emission factors (EF, g/kg dry fuel) for 8 open wood cooking fires and 2 Patsari stoves in central Mexico.

	Open cook ^a								Open cook		Patsari ^b			Patsari chimney ^c		
	fire 1	fire 2	fire 3	fire 4	fire 5	fire 6	fire 7	fire 8	avg	stdev	fire 1	fire 2	avg	fire 1	fire 2	avg
MCE	0.956	0.919	0.962	0.949	0.933	0.967	0.951	0.959	0.949	0.016	0.952	0.963	0.957	0.966	0.973	0.970
ΔCO/ΔCO ₂	0.046	0.088	0.039	0.053	0.072	0.034	0.051	0.043	0.054	0.018	0.050	0.038	0.044	0.035	0.028	0.031
ΔCH ₄ /ΔCO	0.074	0.092	0.133	0.123	0.103	0.121	0.073	0.100	0.102	0.022	0.124	0.151	0.137	0.086	0.061	0.073
ΔMeOH/ΔCO	0.002	0.012	0.015	0.019	0.020	0.010	0.014	0.013	0.013	0.006	0.005	0.016	0.010	0.004	0.016	0.010
ΔNH ₃ /ΔCO	0.016	0.004	0.037	0.012	0.015	0.006	0.008	0.010	0.013	0.010	0.003	0.003	0.003	0.001	0.001	0.001
ΔC ₂ H ₄ /ΔCO	0.009	0.015	0.013	0.005	0.022	0.012	0.015	0.013	0.013	0.005	0.029	0.030	0.030	0.010	0.017	0.013
ΔC ₂ H ₂ /ΔCO	0.003	0.005	0.007	0.006	0.0004	0.011	0.004	0.010	0.006	0.003	0.038	0.052	0.045	0.008	0.009	0.009
ΔC ₃ H ₆ /ΔCO		0.002			0.0001	0.001	0.003	0.003	0.002	0.001				0.001		0.001
ΔHAc/ΔCO	0.017	0.012	0.012	0.014	0.028	0.008	0.006	0.014	0.014	0.007	0.010		0.010		0.005	0.005
ΔHFO/ΔCO		0.002					0.003	0.001	0.002	0.001					0.0001	0.0001
ΔHCHO/ΔCO	0.006	0.012	0.013	0.011	0.006	0.012	0.013	0.010	0.010	0.003	0.003	0.013	0.008	0.004	0.017	0.011
	ER								avg		stdev		EF		avg	
CO ₂	1743	1660	1749	1721	1687	1760	1731	1742	1724	34	1722	1743	1732	1764	1777	1770
CO	51.5	93.5	43.5	58.4	77.7	38.2	56.2	47.9	58.4	18.5	55.2	42.7	48.9	39.2	31.2	35.2
CH ₄	2.18	4.90	3.30	4.12	4.59	2.63	2.35	2.72	3.35	1.06	3.92	3.67	3.80	1.92	1.09	1.50
MeOH	0.10	1.32	0.74	1.29	1.75	0.43	0.90	0.70	0.91	0.53	0.32	0.76	0.54	0.19	0.58	0.38
NH ₃	0.51	0.20	0.97	0.41	0.70	0.15	0.26	0.29	0.44	0.28	0.11		0.11	0.03	0.03	0.03
C ₂ H ₄		0.87	0.65	0.78	0.40	0.82	0.68	0.70	0.70	0.16	1.62	1.30	1.46	0.40	0.52	0.46
C ₂ H ₂	0.12	0.42	0.26	0.31	0.03	0.37	0.21	0.43	0.27	0.14	1.93	2.04	1.98	0.28	0.27	0.28
C ₃ H ₆		0.31			0.01	0.08	0.28	0.21	0.18	0.13				0.03		0.03
HAc	1.86	2.40	1.15	1.72	4.71	0.65	0.67	1.44	1.82	1.31	1.21		1.21		0.34	0.34
HFO		0.34					0.29	0.11	0.25	0.12				0.01		0.01
HCHO	0.31	1.22	0.63	0.67	0.52	0.49	0.79	0.53	0.64	0.27	0.18	0.60	0.39	0.17	0.57	0.37
NMOC	2.39	6.88	3.43	4.77	7.42	2.85	3.82	4.13	4.46	1.82	5.25	4.71	4.98	1.08	2.29	1.68
total PM ^d	4.94	7.87	8.28	5.82					6.73	1.61						

^a 161 background and indoor sample measurements of nascent smoke from open wood cooking fires in 7 kitchens (*fires 1–7*) and the GIRA lab (*fire 8*).

^b 14 background and indoor sample measurements directly above the fire box of the Patsari stove in the GIRA lab (*fire 1*) and 1 kitchen (*fire 2*).

^c 26 outdoor background and sample measurements at the chimney outlet of the same 2 Patsari stoves.

^d PM measurements were continuous at a sampling frequency of 1–2 Hz.

MeOH-methanol; HAc-acetic acid; HFO-formic acid; NMOC-non-methane organic compounds.

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Table 4. Emission factors (EF, g/kg fuel) for individual particle species^a.

	Open cook (fire 2)	Open cook (fire 3)	Open cook (fire 4)	Open cook (fire 5)	Open cook (fire 6)	Garbage (fire 2)	Garbage (fire 3)	Garbage (fire 4)	Brick kiln (fire 1)	Brick kiln (fire 2)	Charcoal kiln (day 3)	Charcoal kiln (day 4)	Stubble burn (fire 1)
<i>TOT (Thermal Optical Transmission)</i>													
OC	3.77	1.39	2.46	1.43	1.19	10.9	2.13	2.78	0.073	0.283	0.382	1.10	5.92
EC	0.355	0.480	0.667	0.205	0.674	0.381	0.924	0.634	0.596	1.50	0.007	0.031	0.055
EC/OC	0.094	0.345	0.271	0.143	0.568	0.035	0.434	0.228	8.15	5.29	0.018	0.028	0.009
<i>HPAEC (High Performance Anion Exchange Chromatography)</i>													
Levoglucosan	0.901	0.124	0.202	0.111	0.110	0.346	0.290	0.102	0.0004	0.002	0.008	0.119	0.712
Mannosan	0.387	0.010	0.013	0.017	0.033	0.026	0.011	0.004		0.0004	0.001	0.007	0.015
Galactosan	0.180	0.004	0.006	0.008	0.008	0.007	0.001				0.001	0.006	0.028
<i>IC (Ion Chromatography)</i>													
K ⁺	0.0212	0.0296	0.0415	0.0234	0.0151	0.0352	0.0163	0.0129	0.0053	0.0052	0.0060	0.0030	0.2799
Ca ²⁺	0.0056	0.0144	0.0013		0.0001	0.0013	0.0011	0.0004	0.0001	0.0014			
Cl ⁻	0.0088	0.0109	0.0066	0.0038	0.0063	1.03	0.17	0.20	0.5085	0.0538	0.0024	0.0706	0.7207
NO ₃ ⁻	0.0078	0.0074	0.0115	0.0034	0.0033				0.0004	0.0017	0.0007		0.0065
<i>ICP (Inductively Coupled Plasma spectroscopy)</i>													
Fe						0.06937	0.01848	0.02067		0.00859			
Na						0.09713		0.01160		0.00797			0.12498
Mg	0.01038	0.02676	0.00778		0.00232								
K	0.03843	0.07388	0.06657		0.02309		0.03202						0.67046
Ca	0.02657	0.09759	0.02257	0.02244	0.00486	0.32613		0.03058			0.00709		
Sr	0.00036	0.00110	0.00024		0.00008	0.00280		0.00031	0.00007	0.00010			
Ti	0.00108	0.00223				0.00452				0.00065			
Mn		0.00063								0.00016			
Co	0.00004	0.00006			0.00002				0.00006		0.00011		
Ni												0.00057	
Cu	0.00040	0.00042		0.00035		0.00213	0.00035	0.00074	0.00465	0.00017	0.00043	0.00096	
Zn	0.00078	0.00081			0.00052		0.00098	0.00172	0.00066	0.00112			
Cd	0.00002				0.00001	0.00027	0.00059	0.00053	0.00002				
Sn	0.00002					0.00199	0.00345	0.00410	0.00003	0.00006			0.00009
Sb	0.00001	0.00007	0.00001	0.00001	0.00000	0.00212	0.01872	0.01154	0.00002	0.00004	0.000005		0.00003
Pb					0.00003	0.00400	0.00780	0.00460	0.00026	0.00023			
V	0.00008	0.00012	0.00002		0.00003	0.00020	0.00001	0.00002	0.00004	0.00003	0.00010	0.00005	0.00014
Cr		0.00156										0.00350	
As	0.00004	0.00007	0.00010	0.00004	0.00002	0.00287	0.00003	0.00029	0.00002	0.00003	0.00005	0.00023	0.00033
Rb	0.00022	0.00027	0.00031	0.00013	0.00007	0.00021	0.00003	0.00004	0.00002	0.00003	0.00002	0.00009	0.00037
Sum ^b	5.75	2.69	4.27	2.27	2.39	17.22	4.17	4.82	1.24	1.96	0.56	1.65	10.14

^a Data set is limited to those fires for which we collected quartz filters.

^b Sum of masses, excluding anhydrosugars, with OC multiplied by 1.4 to account for non-carbon organic mass.

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Table 5. Normalized emission ratios (ER, mol/mol) and emission factors (EF, g/kg dry fuel)^a for 4 garbage fires, 3 brick-making kilns, and 2 barley stubble burns in central Mexico.

	Garbage burning ^b						Brick kilns ^c					Stubble burns ^d		
	fire 1	fire 2	ER fire 3	fire 4	avg	stdev	fire 1	ER fire 2	fire 3	avg	stdev	fire 1	ER fire 2	avg
MCE	0.964	0.911	0.958	0.968	0.950	0.026	0.952	0.974	0.978	0.968	0.014	0.910	0.882	0.896
$\Delta\text{CO}/\Delta\text{CO}_2$	0.038	0.098	0.044	0.033	0.053	0.030	0.050	0.027	0.023	0.033	0.015	0.099	0.134	0.116
$\Delta\text{CH}_4/\Delta\text{CO}$	0.060	0.228	0.099	0.067	0.114	0.078	0.068	0.098	0.077	0.081	0.016	0.089	0.087	0.088
$\Delta\text{MeOH}/\Delta\text{CO}$	0.008	0.031	0.009	0.008	0.014	0.011	0.022	0.013	0.018	0.018	0.016	0.032	0.016	0.024
$\Delta\text{NH}_3/\Delta\text{CO}$	0.023	0.052	0.017		0.031	0.019	0.001	0.0004	0.001	0.001	0.0003	0.025	0.035	0.030
$\Delta\text{C}_2\text{H}_2/\Delta\text{CO}$	0.024	0.060	0.057	0.033	0.044	0.018	0.005	0.011	0.014	0.010	0.005	0.015	0.018	0.017
$\Delta\text{C}_2\text{H}_4/\Delta\text{CO}$	0.004	0.010	0.015	0.007	0.009	0.004	0.0004	0.003	0.007	0.004	0.003	0.002	0.003	0.002
$\Delta\text{C}_2\text{H}_6/\Delta\text{CO}$	0.007	0.028	0.017	0.008	0.015	0.010	0.003	0.004	0.004	0.004	0.003	0.005	0.005	0.005
$\Delta\text{HAc}/\Delta\text{CO}$	0.008	0.044	0.011	0.012	0.019	0.017	0.002		0.002	0.002	0.001	0.042	0.022	0.032
$\Delta\text{HfO}/\Delta\text{CO}$	0.011	0.002	0.011	0.008	0.008	0.004	0.0004	0.0004	0.0005	0.0004	0.0001	0.004	0.005	0.004
$\Delta\text{HCHO}/\Delta\text{CO}$	0.015	0.006	0.016	0.024	0.015	0.008	0.001	0.002	0.001	0.001	0.0001	0.023	0.017	0.020
$\Delta\text{HCl}/\Delta\text{CO}$	0.037		0.194	0.078	0.103	0.081								
	EF		avg	stdev	EF		avg	stdev	EF		avg			
CO ₂	1404	1270	1385	1409	1367	65	1736	1780	1787	1768	28	1628	1577	1602
CO	33.8	79.1	38.7	29.6	45.3	22.8	55.7	30.2	25.7	37.2	16.2	102	135	118
CH ₄	1.16	10.3	2.18	1.14	3.70	4.44	2.16	1.69	1.13	1.66	0.51	5.17	6.73	5.95
MeOH	0.31	2.81	0.40	0.26	0.94	1.25	1.42	0.39	0.90	0.90	0.30	3.70	2.45	3.08
NH ₃	0.46	2.52	0.39		1.12	1.21	0.03	0.01	0.01	0.02	0.01	1.54	2.83	2.18
C ₂ H ₄	0.82	4.75	2.20	0.99	2.19	1.82	0.26	0.32	0.37	0.32	0.05	1.51	2.48	2.00
C ₂ H ₂	0.14	0.72	0.53	0.20	0.40	0.28	0.02	0.09	0.16	0.09	0.07	0.17	0.32	0.25
C ₂ H ₆	0.36	3.34	0.97	0.36	1.26	1.42	0.28	0.15	0.22	0.22	0.07	0.77	0.77	0.77
HAc	0.58	7.40	0.92	0.78	2.42	3.32	0.21		0.21	0.21	0.01	9.15	6.49	7.82
HfO	0.11	0.30	0.71	0.40	0.38	0.25	0.03	0.02	0.02	0.02	0.01	0.60	1.10	0.85
HCHO	0.56	0.48	0.68	0.76	0.62	0.13	0.08	0.05	0.04	0.05	0.02	2.48	2.47	2.48
HCl	1.65		9.8	3.02	4.82	4.36								
NMOC	2.86	19.8	6.39	3.75	8.20	7.88	2.30	0.48	1.13	1.30	0.92	18.40	15.31	16.85

^a See Sect. 2.4 for details specific to EF calculations for garbage burning.

^b 72 spot measurements from garbage burning in 4 landfills.

^c 77 spot measurements from 3 brick making kilns.

^d 23 spot measurements from 2 barley stubble field burns.

MeOH-methanol; HAc-acetic acid; HfO-formic acid; NMOC-non-methane organic compounds.

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Table 6. Comparison of normalized emission ratios (ER, mol/mol) and emission factors (EF, g/kg dry fuel)^a for 3 charcoal kilns in central Mexico with a charcoal kiln in Zambia.

	Current study ^b						Zambia ^c		Current / Zambia
	day 2	day 3	day 4	day 5	avg	stdev	avg	stdev	
	ER								
MCE	0.818	0.800	0.829	0.809	0.814	0.012	0.783	0.042	1.04
$\Delta\text{CO}/\Delta\text{CO}_2$	0.223	0.250	0.207	0.236	0.229	0.018	0.280	0.071	0.82
$\Delta\text{CH}_4/\Delta\text{CO}$	0.151	0.160	0.273	0.336	0.230	0.090	0.242	0.073	0.95
$\Delta\text{MeOH}/\Delta\text{CO}$	0.155	0.210	0.308	0.142	0.204	0.075	0.111	0.070	1.84
$\Delta\text{NH}_3/\Delta\text{CO}$	0.0032	0.0024	0.0032	0.000	0.002	0.001	0.006	0.002	0.38
$\Delta\text{C}_2\text{H}_4/\Delta\text{CO}$	0.007	0.006	0.010	0.015	0.009	0.004	0.013	0.002	0.72
$\Delta\text{C}_2\text{H}_2/\Delta\text{CO}$		0.001	0.001		0.0007				
$\Delta\text{C}_3\text{H}_6/\Delta\text{CO}$	0.006	0.004	0.010	0.008	0.007	0.003	0.010	0.005	0.68
$\Delta\text{HAc}/\Delta\text{CO}$	0.109	0.164	0.341	0.119	0.183	0.108	0.043	0.031	4.26
$\Delta\text{HfO}/\Delta\text{CO}$		0.002	0.003		0.002		0.003	0.001	0.82
	EF				avg	stdev	avg		
CO_2	612	577	558	582	582	23	542		1.07
CO	87.0	91.7	73.3	87.4	84.9	8.0	96.8		0.88
CH_4	7.52	8.37	11.46	16.77	11.0	4.19	13.4		0.82
MeOH	15.4	22.0	25.8	14.2	19.4	5.48	12.3		1.58
NH_3	0.17	0.14	0.14	0.02	0.12	0.07	0.37		0.31
C_2H_4	0.60	0.55	0.74	1.28	0.79	0.33	1.31		0.60
C_2H_2		0.04	0.06		0.05				
C_3H_6	0.73	0.48	1.08	1.10	0.85	0.30	1.50		0.56
HAc	20.3	32.1	53.6	22.3	32.1	15.2	8.92		3.60
HfO		0.34	0.32		0.33	0.02	0.45		0.73
HCHO							1.06		
NMOC	37.1	55.5	81.6	38.9	53.3	20.6	32.8		1.62

^a See Sect. 2.3 for details specific to EF calculations for charcoal kilns.

^b 36 spot measurements from 3 charcoal kilns.

^c 3 ~1 h measurements over the course of 4 days from a charcoal kiln in Zambia (Bertschi et al., 2003).

MeOH-methanol; HAc-acetic acid; HfO-formic acid; NMOC-non-methane organic compounds.

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Table 7. Comparison of cooking fire and urban emissions for Mexico.

	PM _{2.5}	NMOC	CO	NO _x	CH ₄	NH ₃
Biofuel emission factor (g/kg)	6.73 ^a	54 ^b	58.4 ^a	2.04 ^c	3.35 ^a	0.44 ^a
Annual national biofuel emissions (Tg)	0.46	3.67	3.97	0.14	0.23	0.03
Annual MCMA emissions (Tg)	0.0066	0.53	1.79	0.18	0.24	0.018
Ratio: national biofuels/MCMA	69.11	6.90	2.22	0.77	0.97	1.71
National urban emissions (Tg)	0.02	1.98	6.68	0.67	0.88	0.07
National biofuel/National urban	18.53	1.85	0.59	0.21	0.26	0.46

^a This work.

^b Yokelson et al. (2008).

^c Bertschi et al. (2003).

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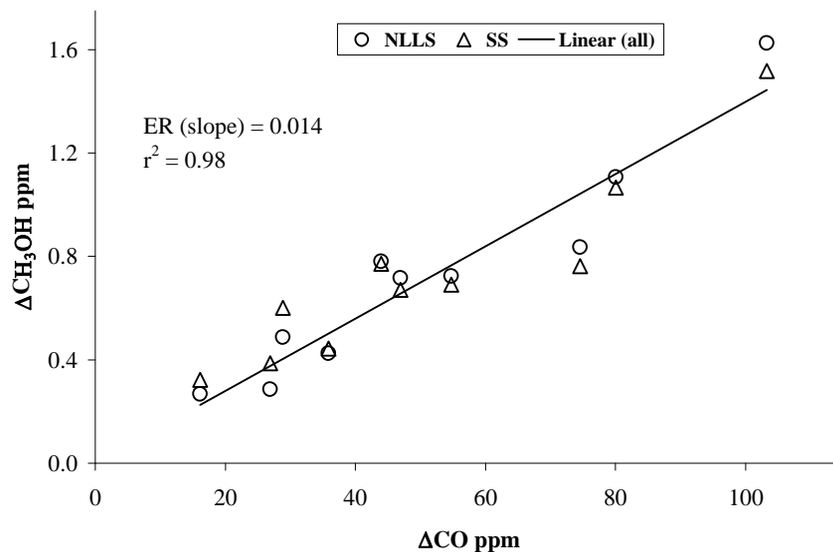


Fig. 1. An example of the determination of the fire-integrated emission ratio for an open wood cooking fire by plotting the excess mixing ratios of methanol versus those of CO. The excess methanol is shown as determined by both nonlinear least squares synthetic calibration (NLLS) and spectral subtraction (SS).

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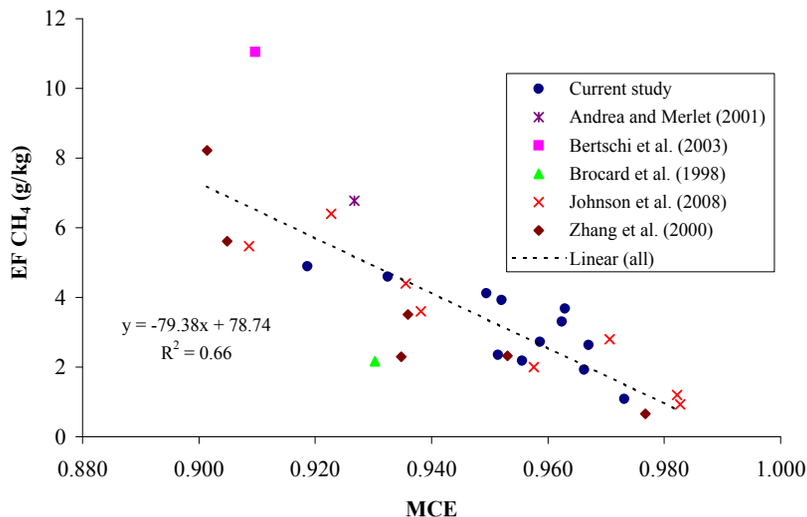


Fig. 2. Variation of the methane emission factor with MCE for open wood cooking fires.

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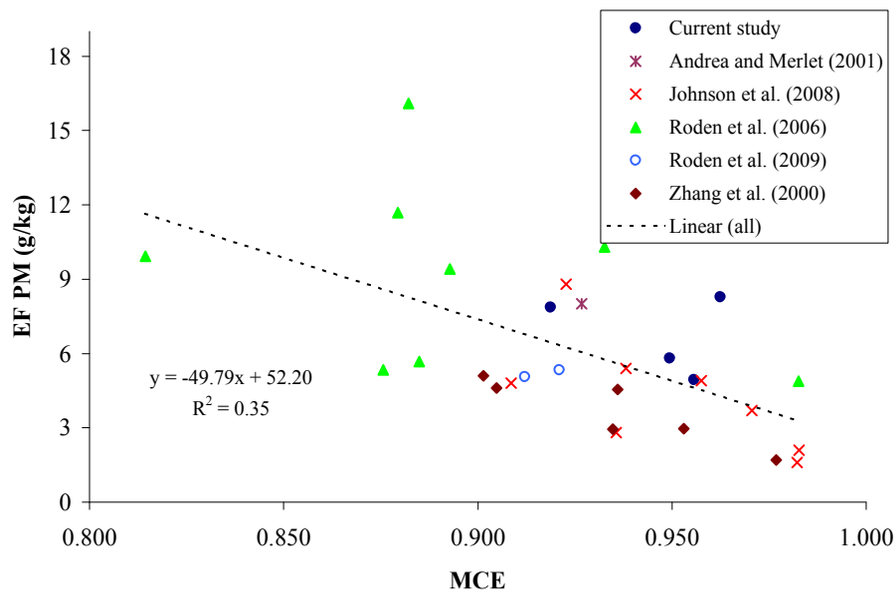


Fig. 3. Variation of the particle emission factor with MCE for open wood cooking fires.

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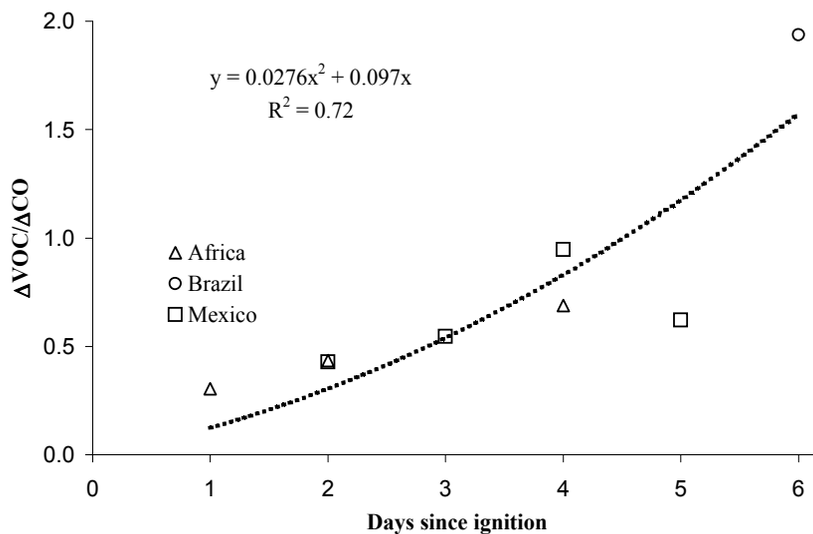


Fig. 4. The sum of $\Delta\text{VOC}/\Delta\text{CO}$ versus approximate kiln age for charcoal making kilns (see Sect. 3.3.2).

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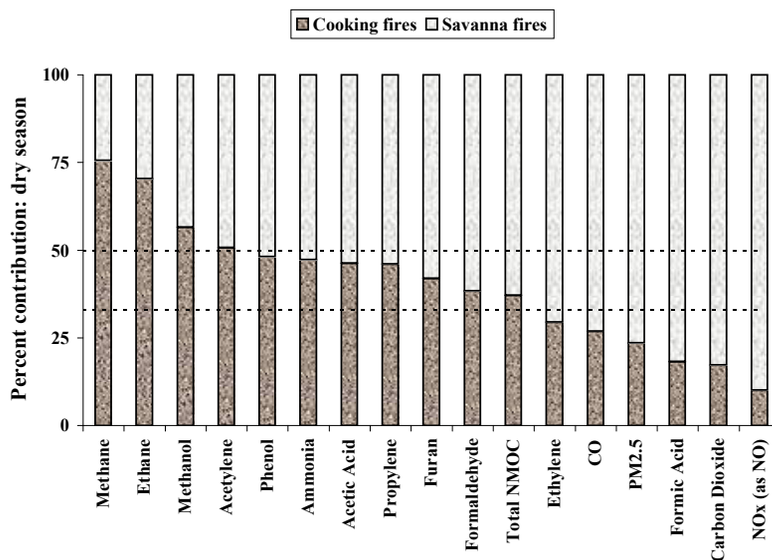


Fig. 5. Percentage of total dry season pyrogenic emissions due to cooking fires or savanna fires in a mostly rural developing country (see Sect. 4.1). If the dry season percentage attributed to cooking fires for a species extends above the 33% line indicated, the total annual emissions of that species will be greater from cooking fires.

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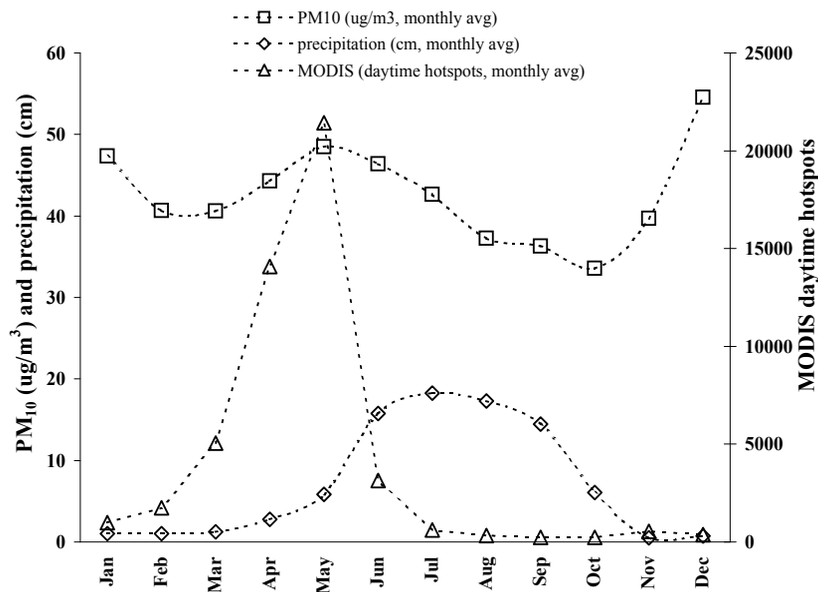


Fig. 6. Time series of monthly average PM₁₀ (Pedregal RAMA station 2003–2008 average, www.sma.df.gov.mx/simat/cambia_base.htm); MODIS daytime hotspots for Mexico (2003–2008 average, www.conabio.gob.mx); and monthly average precipitation for MCMA (see Sect. 4.2).

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