

The Tropical Forest and Fire Emissions Experiment: overview and airborne fire emission factor measurements

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Abstract. The Tropical Forest and Fire Emissions Experiment (TROFFEE) used laboratory measurements followed by airborne and ground based field campaigns during the 2004 Amazon dry season to quantify the emissions from pristine tropical forest and several plantations as well as the emissions, fuel consumption, and fire ecology of tropical deforestation fires. The airborne campaign used an Embraer 110B aircraft outfitted with whole air sampling in canisters, mass-calibrated nephelometry, ozone by UV absorbance, Fourier transform infrared spectroscopy (FTIR), and proton-transfer mass spectrometry (PTR-MS) to measure PM₁₀, O₃, CO₂, CO, NO, NO₂, HONO, HCN, NH₃, OCS, DMS, CH₄, and up to 48 non-methane organic compounds (NMOC). The Brazilian smoke/haze layers extended to 2–3 km altitude, which is much lower than the 5–6 km observed at the same latitude, time of year, and local time in Africa in 2000. Emission factors (EF) were computed for the 19 tropical deforestation fires sampled and they largely compare well to previous work. However, the TROFFEE EF are mostly based on a much larger number of samples than previously available and they also include results for significant emissions not previously reported such as: nitrous acid, acrylonitrile, pyrrole, methylvinylketone, methacrolein, crotonaldehyde, methylethylketone, methylpropanal, “acetol plus methylacetate,” furaldehydes, dimethylsulfide, and C₁–C₄ alkyl nitrates. Thus, we recommend these EF for all tropical deforestation fires. The NMOC emissions were ~80% reactive, oxygenated volatile organic compounds (OVOC).

Our EF for PM₁₀ (17.8±4 g/kg) is ~25% higher than previously reported for tropical forest fires and may reflect a trend towards, and sampling of, larger fires than in earlier studies. A large fraction of the total burning for 2004 likely occurred during a two-week period of very low humidity. The combined output of these fires created a massive “mega-plume” >500 km across that we sampled on 8 September. The mega-plume contained high PM₁₀ and 10–50 ppbv of many reactive species such as O₃, NH₃, NO₂, CH₃OH, and organic acids. This is an intense and globally important chemical processing environment that is still poorly understood. The mega-plume or “white ocean” of smoke covered a large area in Brazil, Bolivia, and Paraguay for about one month. The smoke was transported >2000 km to the southeast while remaining concentrated enough to cause a 3–4-fold increase in aerosol loading in the São Paulo area for several days.

1 Introduction

Biomass burning and biogenic emissions are the two largest sources of volatile organic compounds (VOC) and fine particulate carbon in the global troposphere. Tropical forests produce about one-third of the global biogenic emissions and tropical deforestation fires account for much of the global biomass burning (Andreae and Merlet, 2001; Kreidenweis et al., 1999; Guenther et al., 1995, 2006). Recent estimates of the total amount of biomass burned globally vary from about 5 to 7 Pg C/y (Andreae and Merlet, 2001; Page et al., 2002). The contribution of tropical deforestation fires to total global biomass burning has been estimated as 52% (Crutzen and

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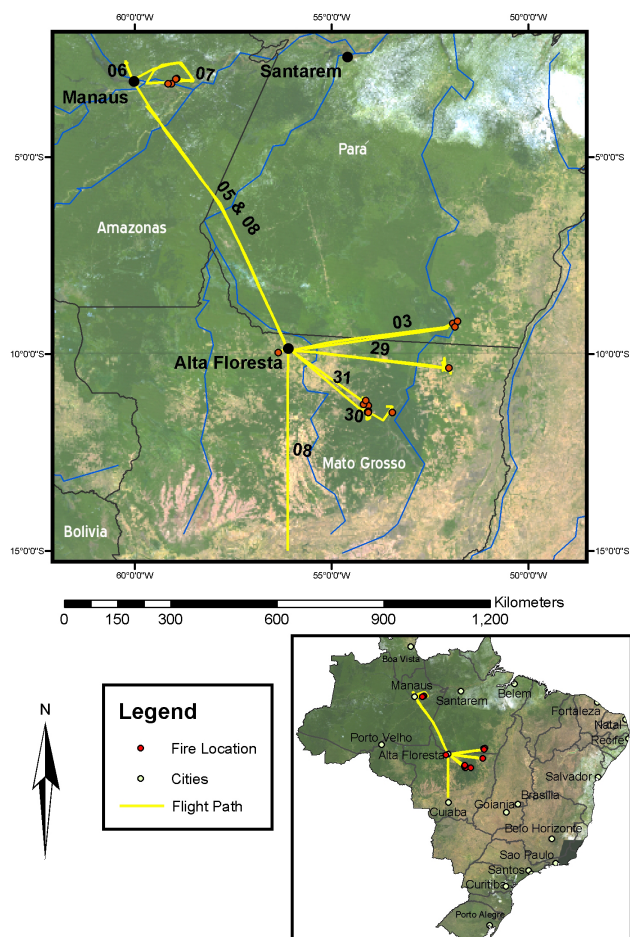


Fig. 1. The TROFFEE flight tracks and the locations of the fires sampled.

Andreae, 1990), 34% (Hao and Liu, 1994), and 15% (Andreae and Merlet, 2001). Thus, factors of 2–3 uncertainty need to be resolved, but these fires consistently emerge as one of the three major types of burning along with savanna fires and domestic biofuel use. A large uncertainty in the estimated area burned is due to uncertainties in remote sensing applications. For example, it is unclear if small fires or understory fires can be quantified from space (Brown et al., 2006), and many fires can be missed from space due to cloud cover, which is common over tropical forested regions. Deforestation fires facilitate land-use change, which alters the biogenic emissions. Thus, to understand regional-global atmospheric chemistry and assess the long-term impact of land-use change, we must thoroughly characterize the smoke emissions from these fires and the different biogenic emissions produced by the primary forest and the various anthropogenic “replacement” ecosystems.

The Tropical Forest and Fire Emissions Experiment (TROFFEE) provided emissions measurements for tropical deforestation fires and tropical vegetation. An overview of

TROFFEE follows. A laboratory experiment was carried out before the field campaigns that intercompared proton-transfer reaction mass spectrometry (PTR-MS), open-path Fourier transform infrared spectroscopy (FTIR), and gas chromatography (GC) coupled to PTR-MS (GC-PTR-MS) on 26 fires burning tropical fuels. The laboratory work helped plan the PTR-MS sampling protocol for the field campaign and instrumentation was available to quantify some particle characteristics not measured in the field. The GC-PTR-MS measured the branching ratios for fire-emitted species that appear on the same mass channel. The laboratory fire and intercomparison results are presented elsewhere (Karl et al., 2007a; Christian et al., 2007a¹).

The TROFFEE field campaigns were in Brazil since it has the most tropical forest and the most deforestation fires. The ground-based field campaigns included measurements of biogenic emissions from pristine forest near Manaus (Fig. 1) (Karl et al., 2007b). The ground campaign also included FTIR emissions measurements on initially-unlifted plumes from 9 biomass fires in the vicinity of Alta Floresta (Fig. 1). These plumes were due to residual smoldering combustion at deforestation sites or pasture maintenance burns or they were from charcoal kilns, cooking fires, burning dung, etc. This element of TROFFEE was motivated by indications from previous field campaigns that initially, unlifted biomass burning plumes might contribute a large portion of the total regional emissions (Kauffman et al., 1998; Reid et al., 1998). The results for unlifted plumes and biofuels are described by Christian et al. (2007b). The ground campaign fires included a planned fire in which Brazilian researchers carried out a “typical” deforestation burn under conditions where the fuel consumption and other aspects of fire ecology could be measured. The emissions from this planned fire were measured by the ground-based FTIR and in the TROFFEE airborne campaign (described next).

The TROFFEE airborne campaign (Fig. 1) consisted of 44.5 flight hours between 27 August and 8 September of 2004 on an Embraer Bandeirante operated by the Brazilian National Institute for Space Research (Instituto Nacional de Pesquisas Espaciais (INPE)). The major instruments deployed on the aircraft included: (1) real-time ozone, condensation particle counter, and mass-calibrated nephelometry (University of São Paulo); (2) PTR-MS (National Center for Atmospheric Research); (3) Whole air sampling in canisters with subsequent GC analysis using flame ionization, mass selective, and electron capture detection (FID, MSD, and ECD; University of California at Irvine); and (4) airborne FTIR (University of Montana). This suite of instruments was well suited for measuring CO₂, CO, PM₁₀, CH₄, NO_x, O₃,

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Table 1. Location and characteristics of fires sampled from the INPE Bandeirante aircraft during TROFFEE 2004 airborne campaign.

Fire name	Date dd/mm	Source location		time period sampled		Fuels observed from aircraft
		Lat dd.ddd	Long dd.ddd	LT	LT	
29 Aug Fire 1	29/08	-10.270	-52.159	13:41:54	14:17:10	slash under partial canopy
29 Aug Fire 2	29/08	-10.357	-52.019	14:30:37	14:43:30	Pasture
30 Aug Fire 1	30/08	-11.315	-54.064	12:56:51	13:00:45	grass and slash piles under partial canopy
30 Aug Fire 2	30/08	-11.459	-54.062	13:04:18	13:13:37	grass and slash piles under partial canopy
30 Aug Fire 3	30/08	-11.479	-54.088	13:20:14	13:20:55	grass and slash piles under partial canopy
30 Aug Fire 4	30/08	-11.491	-54.058	13:29:06	13:36:56	grass and slash piles under partial canopy
SC Fire	30/08	-11.488	-53.458	14:36:25	14:43:59	mixed forest fuels
31 Aug Fire 1	31/08	-11.282	-54.185	13:08:58	13:25:01	mixed forest fuels
31 Aug Fire 2	31/08	-11.183	-54.131	13:30:55	13:44:52	mixed forest fuels
3 Sept Fire 1	03/09	-9.224	-51.918	13:23:32	13:39:36	mixed forest fuels
3 Sept Fire 2	03/09	-9.167	-51.798	13:37:00	13:37:08	mixed forest fuels
3 Sept Fire 3	03/09	-9.311	-51.861	13:52:24	14:02:22	mixed forest fuels
3 Sept Fire 4	03/09	nm		14:13:48	14:14:16	source/fuels not observed from aircraft
3 Sept Fire 5	03/09	nm		13:22:41	13:22:54	source/fuels not observed from aircraft
Planned Fire	05/09	-9.969	-56.345	14:16:42	14:51:08	mixed forest fuels
7 Sept Fire 1	07/09	-3.007	-8.930	11:49:39	11:56:42	mixed forest fuels
7 Sept Fire 2	07/09	-3.011	-58.946	12:01:13	12:01:29	mixed forest fuels
7 Sept Fire 3	07/09	-3.129	-59.056	12:04:50	12:05:58	mixed forest fuels
7 Sept Fire 4	07/09	-3.137	-59.147	12:06:46	12:07:36	mixed forest fuels
Mega-plume	08/09	nm		~11:00	~12:30	source/fuels not observed from aircraft

and >40 non-methane organic compounds (NMOC) including the important biogenic emissions isoprene and methanol.

In phase 1, the aircraft was based in Alta Floresta, Mato Grosso in the southern Amazon (9.917 S, 56.017 W, Fig. 1) from 27 August–5 September where the local dry/burning season was well underway. Regional haze due mostly to diluted biomass-burning smoke of unknown age and the nascent (minutes-old) emissions from 15 fires (mostly deforestation fires) were sampled in the states of Mato Grosso and Pará within about one-hour flight time (~300 km) of Alta Floresta.

In phase 2, the aircraft was based in Manaus, Amazonas (3.039 S, 60.050 W, Fig. 1) from 5–8 September. The local dry season was just beginning there and the air was much cleaner and mostly unaffected by fires; especially in the mornings. The biogenic emissions were sampled from forests, several plantations east of Manaus, and the pristine forest at the ZF-14 tower north of Manaus. The results are discussed and integrated with the ground-based biogenic measurements by Karl et al. (2007b). In addition, four more fires were sampled around noon in the Manaus region. On 8 September from 8–13° S we sampled a smoke plume hundreds of km wide that contained the combined emissions from a huge number of fires. These fires represented a significant fraction of the total Amazon burning for 2004 and they generated a “mega-plume,” which we discuss in detail in Sect. 3.4. All the fires sampled are listed in Table 1. The

TROFFEE flight tracks and individual fires are mapped in Fig. 1. A more detailed map of the 6–7 September flights is given by Karl et al. (2007b).

The fire component of TROFFEE is covered in four initial papers. The lab fire results and the chemistry and impact of unlofted smoke not amenable to airborne sampling are covered in two papers (Christian et al., 2007a¹, b). Karl et al. (2007a) present the instrument intercomparison and the emission ratios of many VOC to acetonitrile, which is thought to be mostly emitted by biomass burning. The main focus of this paper is to provide background on the region and experiment and to detail the airborne measurements of fire emission factors, which are needed as model input and for bottom-up emissions estimates at any scale. Some aspects of the airborne measurements in clean air (relatively unaffected by fires) and haze (dilute/aged smoke) are also given to clarify the regional atmospheric conditions and make our fire-sampling strategy clear.

A major goal of all the TROFFEE fire research was comprehensive sampling of reactive species as close as possible to the source. The rationale for this is given next. Much of the initial interest in fires focused on the climate forcing. In fact, in El-Niño years, the carbon added to the atmosphere by biomass burning may exceed that from fossil fuels (Page et al., 2002). The CO₂ due to tropical deforestation alone may cause an average annual amount of warming that is 20–60% of that caused by the CO₂ from all global

industry (Crutzen and Andreae, 1990) and fires emit more other greenhouse gases (GHG) per CO₂ than fossil fuel use (Christian et al., 2003). Photochemical processing of fire emissions was shown to produce O₃ (Fishman et al., 1991; Andreae et al., 1994), an important GHG (Prather et al., 1994). Particles emitted by fires were found to cause negative forcing both directly (Hobbs et al., 1997) and indirectly by reducing cloud droplet sizes and increasing cloud albedo (Kaufman and Fraser, 1997).

In recent years, the reactivity and the rapid post-emission chemistry of smoke have attracted increasing attention. Early laboratory and field studies of biomass burning had concentrated on measuring the emissions of CO₂, CO, NO_x, and hydrocarbons (Lobert et al., 1991; Blake et al., 1996; Ferek et al., 1998), but later laboratory work showed that 60–80% of the NMOC emissions from fires were actually highly reactive, oxygenated VOC (OVOC) (Yokelson et al., 1996, 1997; Holzinger et al., 1999). The dominance of NMOC emissions by OVOC was then confirmed for all of the major types of biomass burning except tropical forest fires: e.g. savannas, biofuels, agricultural waste, peat, and boreal forest (Goode et al., 2000; Christian et al., 2003; Bertschi et al., 2003a). In addition, field measurements of rapid changes in smoke plume chemistry became available (Goode et al., 2000; Yokelson et al., 2003a; Hobbs et al., 2003). Detailed photochemical smoke models reproduced the observed O₃ formation rate in only some cases and were unable to predict the observed formation of other species such as acetone and acetic acid (Mason et al., 2001; Jost et al., 2003; Tabazadeh et al., 2004; Trentmann et al., 2005). Sensitivity analysis showed that model performance was significantly enhanced by using more complete information on the initial NMOC (mostly OVOC) emissions. About 80% of biomass burning occurs in the tropics, which govern the oxidizing power of the global troposphere (Crutzen and Andreae, 1990). Fires are a major source of CO (the main sink of OH), but the large quantities of OVOC emitted by fires, and the secondary O₃, are HO_x (OH+HO₂) precursors and important oxidants (Finlayson-Pitts and Pitts 1986; Singh et al., 1995). Thus there was a critical need for the first-ever data on OVOC emissions from tropical deforestation fires.

2 Experimental details

2.1 Instrument details

2.1.1 Airborne FTIR (AFTIR) and whole air sampling in canisters

The basic design and operation of the AFTIR system has been described in detail by Yokelson et al. (1999, 2003a, b). A summary description is given here followed by the details of how AFTIR was used to fill canisters. The AFTIR has a dedicated, halocarbon-wax, coated inlet that directs ram air

through a Pyrex, multipass cell. Infrared spectra of the cell contents are acquired continuously (every 0.83 s) throughout each flight and the flow-control valves are normally open, which flushes the cell with outside air every 2–4 s. The fast-acting flow control valves allow the system flow to be temporarily stopped for signal averaging and improved accuracy on “grab samples.” The IR spectra are later analyzed to quantify the compounds responsible for all the major peaks. This accounts for most of the trace gases present in the cell above 5–20 ppbv (Goode et al., 1999).

For TROFFEE, a Teflon valve was added to the AFTIR cell that connected to two options for filling evacuated canisters. For a canister sample of a plume, we used a teflon-diaphragm pump to pressurize the can with gas from the AFTIR cell, which already contained a grab sample of the plume. Pressurizing the cans allows more sensitive and/or a wider variety of analyses and also prevents contamination in the event of a slow leak. Operationally-simpler canister samples of background air were obtained by diverting a portion of the flow through the AFTIR cell into the cans. The .635 cm outside diameter Teflon tubing connecting to the canisters had a pressure higher than the cabin pressure and attached to the can with Ultra-Torr[®] fittings. We flushed the connecting tubing with cell air by loosening the fitting for a few minutes. Once the fitting was retightened the pre-evacuated can was opened and filled to cell pressure within seconds. The filling time of each can was shown by a sharp, (logged) pressure response in the AFTIR cell. The canisters were later analyzed at UCI using GC/FID-MSD-ECD (Colman et al., 2001).

2.1.2 IR spectral analysis

Mixing ratios for H₂O, CO₂, CO, and CH₄ were obtained by multicomponent fits to sections of the IR transmission spectra with a synthetic calibration non-linear least-squares method (MALT 5.2) recently developed by one of the authors (Griffith). To derive excess mixing ratios (ΔX) for the above species in smoke plumes we took the mixing ratio of the species “X” in the smoke plume grab sample minus the mixing ratio of X in the closest grab sample of background air. The use of a nearby background sample for this subtraction is important because it excludes the contribution of the aged smoke that contributes much of the background air in areas heavily impacted by biomass burning.

We used the same background-plume spectra pairs to generate absorbance spectra of the smoke plume samples. Excess mixing ratios are retrieved directly from the absorbance spectra (Hanst and Hanst, 1994). Excess mixing ratios for NO and NO₂ in smoke plumes were obtained from the absorbance spectra using peak integration and a multipoint calibration. Excess mixing ratios for ethylene (C₂H₄), acetylene (C₂H₂), propylene (C₃H₆), methanol (CH₃OH), formic acid (HCOOH), acetic acid (CH₃COOH), ammonia (NH₃), nitrous acid (HONO), hydrogen cyanide (HCN), and ozone

(O₃) were retrieved from the absorbance spectra by spectral subtraction (Yokelson et al., 1997). The spectral subtraction routine used commercial IR reference spectra or multiple reference spectra per species that we recorded in house for NH₃, CH₃OH, CH₃COOH, C₂H₄, and C₃H₆. Excess mixing ratios for C₂H₆ and HCHO were retrieved from the absorbance spectra using MALT 5.2. For most compounds the detection limit was 5–10 ppbv, but for NO_x, HCHO, acetic acid, C₃H₆, C₂H₆, and O₃ it was ~15–20 ppbv.

The spectral analysis routines were challenged by applying them to IR spectra of over 50 flowing standard mixtures. The routines typically returned values within 1% of the nominal, delivered amount. Consideration of the accuracy of the standards, flow meters, and other issues suggests that the absolute accuracy of our mixing ratios is ±1–2% for CO₂, CO, and CH₄ and ±5% (1σ) or the detection limit, whichever is larger, for the other compounds. NH₃ was the only compound noticeably affected by brief storage in the cell, but the NH₃ values have been corrected both for initial passivation of the cell and slow decay during grab-sample storage as described by Yokelson et al. (2003b) and should be accurate to ±10% or the detection limit.

2.1.3 PTR-MS

A detailed description of the PTR-MS instrument is given elsewhere (Lindinger et al., 1998). Briefly, H₃O⁺ ions are used to ionize volatile organic compounds (VOC) via proton-transfer reactions. The value for E/N (E the electric field strength and N the buffer gas density) in the drift tube was kept at about 123 Townsend, which is high enough to avoid strong clustering of H₃O⁺ ions with water and thus a humidity dependent sensitivity. The sensitivity of the PTR-MS instrument during this study was typically on the order of 70 Hz/ppbv (counts per second per ppbv) for acetone and 50 Hz/ppbv for methanol at 2.3 mbar buffer gas pressure with a reaction time of 110 μs and 3–4 MHz H₃O⁺ ions, and thus inferred a signal to noise ratio of 60% at a concentration of 20 pptv and a 2 s integration time. The PTR-MS sampled air through a dedicated, rear-facing, Teflon inlet. About 17 mass channels were monitored during flight with a measurement period for each species of 1–20 s. Higher sampling rates were used in the plumes. More details about the PTR-MS in this campaign are given by Karl et al. (2007a).

2.1.4 Particle, ozone, and auxiliary measurements

A list of the instruments deployed by the University of São Paulo and their measurement frequency follows. (1) DataRAM4 (Thermolectron Corp), which measures the mass of particles with an aerodynamic diameter <10 microns (PM₁₀) and mean particle diameter (microns) at 0.5 Hz. (2) 3-channel nephelometer (RBG) at 0.2857 Hz. (3) 7-channel aethalometer (Magee Scientific) measuring particle absorbance from 950–450 nm every 2 min. (4) Ozone by UV

absorbance (1 min time resolution). (5) GPS (Garmin) measuring UTC time, latitude, longitude, and altitude at 1 Hz. Instruments 1–4 had specialized inlets located on the front belly of the aircraft adjacent to the PTR-MS inlet. The PM₁₀ measurements reported here were measured by the DataRAM4, which is a two-wavelength nephelometer with a built in humidity correction. The instrument has been run side by side with a TEOM (Tapered Element Oscillating Microbalance) under smoky conditions in the Amazon and good agreement was observed.

2.1.5 Flight plans and sampling protocols

While based in Alta Floresta (27 August–5 September) background air (defined here as air not within a visible biomass burning plume) was characterized at various altitudes (up to 3352 m). These were afternoon flights conducted to search for and sample fires and most of the measurements were made below the top of the (hazy) mixed layer. While based in Manaus cleaner background air was sampled during morning flights over a similar altitude range. The Manaus flights included both continuous-spiral and “parking-garage”-type vertical profiles over the instrumented ZF-14 Tower and a constant-altitude “racetrack” pattern that sampled several regionally important ecosystems (undisturbed forest, flooded forest, and various plantations) east of Manaus (Karl et al., 2007b). When sampling background air in either region, the PTR-MS continuously cycled through a suite of mass channels with a resulting measurement frequency for individual species ranging from 10–20 s. Overall, twenty-one canisters were used to “grab” background samples at key locations. The airborne FTIR (AFTIR) was operated either continuously (time resolution of 0.83 to 18 s) or to acquire 133 grab samples of background air.

To measure the initial emissions from fires in both regions, we sampled smoke less than several minutes old by penetrating the column of smoke 200–1000 m above the flame front. The AFTIR system and cans obtained grab samples in the plume (and paired background samples just outside the plume). The other instruments measured their species continuously while passing through the plume. More than a few kilometers downwind from the source, smoke plume samples are “chemically aged” and better for probing post-emission chemistry than estimating initial emissions (Hobbs et al., 2003; de Gouw et al., 2006).

2.2 Data processing and synthesis

Grab samples or profiles of an emission source can provide excess mixing ratios (ΔX, see Sect. 2.1.2). ΔX reflect the instantaneous dilution of the plume and the instrument response time. Thus, a widely used, derived quantity is the normalized excess mixing ratio where ΔX is compared to a simultaneously measured plume tracer such as ΔCO or ΔCO₂. A measurement of ΔX/ΔCO or ΔX/ΔCO₂ made in

a nascent plume (seconds to a few minutes old) is an emission ratio (ER). The ER $\Delta\text{CO}/\Delta\text{CO}_2$ and the modified combustion efficiency (MCE, $\Delta\text{CO}_2/(\Delta\text{CO}_2+\Delta\text{CO})$) are useful to indicate the relative amount of flaming and smoldering combustion for biomass burning. Higher $\Delta\text{CO}/\Delta\text{CO}_2$ or lower MCE indicates more smoldering (Ward and Radke, 1993). For any carbonaceous fuel, a set of ER to CO_2 for the other major carbon emissions (i.e. CO , CH_4 , a suite of NMOC, particulate carbon) can be used to calculate emission factors (EF, g compound emitted/kg dry fuel) for all the gases quantified from the source using the carbon mass-balance method (Yokelson et al., 1996). EFs are combined with fuel consumption measurements to estimate total emissions at various scales. In this project, the primary data needed to calculate EF was provided by AFTIR measurements of CO_2 , CO , CH_4 , and many NMOC. However, the PTR-MS and canister sampling added numerous, important NMOC that were below AFTIR detection limits or not amenable to IR detection. The PM_{10} data allowed inclusion of particle carbon. Next we summarize the methods we used to calculate ER and EF and to couple/synthesize the data from the various instruments on the aircraft.

2.2.1 Estimation of fire-average, initial emission ratios (ER)

The first step in our analysis was to compute molar ER to CO and CO_2 for each species detected in the AFTIR or can grab samples; and molar ER to methanol (justified below) for each species detected by PTR-MS. This is done for each individual fire or each group of co-located, similar fires. If there is only one sample of a fire (as for the canisters) then the calculation is trivial and equivalent to the definition of ΔX given above. For multiple AFTIR grab samples of a fire (or group of fires) then the fire-average, initial ER were 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 (see Figs. 2a and b). This method is justified in detail by Yokelson et al. (1999). We calculated the fire-average MCE for each fire using the fire-average $\Delta\text{CO}/\Delta\text{CO}_2$ and the equation $\text{MCE}=1/((\Delta\text{CO}/\Delta\text{CO}_2)+1)$.

The ER for PTR-MS compounds with respect to methanol were obtained by similar plots except that the integrated excess mixing ratios (ppbv s) for each pass thru the plume were used in lieu of the individual excess mixing ratios (see Fig. 2c). Comparison of integrals provides more accurate ER (Karl et al., 2007a). When two or more compounds appear on the same mass channel, the signal was assigned to each compound using the branching ratios measured by GC-PTR-MS in smoke from tropical fuels burned during the lab experiment. This adds additional uncertainty for these compounds since these branching ratios typically varied by 10–20% from fire to fire during the lab experiments (Karl et al., 2007a).

The ER to CO for the NMOC detected by PTR-MS was derived from a simple two step process. The process is based

on the fact that we have found excellent agreement between FTIR and PTR-MS for methanol, over a wide range of concentrations, in two other studies (Christian et al., 2004; Karl et al., 2007a). An example of the process follows. The ER for acetaldehyde to CO was taken to be the PTR-MS ER “acetaldehyde/methanol” times the AFTIR ER “methanol/ CO .” Multiplying again by the AFTIR CO/CO_2 ratio gave the ratio of the NMOC to CO_2 – as needed for the EF calculation. A slightly different approach was needed to couple the data from the particle instruments. The DataRAM4 measures the STP-equivalent PM_{10} per unit volume ($\mu\text{g}/\text{m}^3$) every two seconds while passing thru a plume. We converted the integrated methanol mixing ratios to an integrated mass (STP) of methanol and ratioed the integrated particle mass to this (see Fig. 2d).

2.2.2 Estimation of fire-average, initial emission factors

We estimated fire-average, initial EF for PM_{10} and each observed trace gas from our fire-average, initial ER using the carbon mass balance method (Ward and Radke, 1993) as described by Yokelson et al. (1999). Briefly, we assume that all the volatilized carbon is detected and that the fuel carbon content is known. For purposes of the carbon mass balance we assume the particles are 60% C by mass (Ferek et al., 1998). By ignoring unmeasured gases we are probably inflating the emission factors by 1–2% (Andreae and Merlet, 2001). We assumed in our EF calculations that all the fires burned in fuels containing 50% carbon by mass. This is in good agreement with previous studies of tropical biomass (Susott et al., 1996), but the actual fuel carbon percentage may vary by $\pm 10\%$ (2σ) of our nominal value. (Emission factors scale linearly with assumed fuel carbon percentage.)

2.3 Overview of Brazilian fires and the fires sampled in the airborne campaign

2.3.1 General fire characteristics relevant to sampling strategies

Conversion of the Amazon primary forest usually starts at the beginning of the dry season (May–July) when the biomass is slashed and dried (Fearnside, 1993). Most of the burns occur late in the dry season (August–October) to achieve high consumption. A typical burn is initiated by starting a line of flame along the outer edge of the slashed area. As the flame front propagates inward, the flame-induced convection column entrains the emissions from both flaming combustion and any nearby smoldering combustion. These emissions can be sampled from an aircraft. In some cases, smoldering can continue after the convection envelope has moved too far away to entrain the emissions or convection from the entire site has ceased. When either of these conditions is met, we term this residual smoldering combustion (RSC). RSC emissions are not initially lofted or amenable to airborne

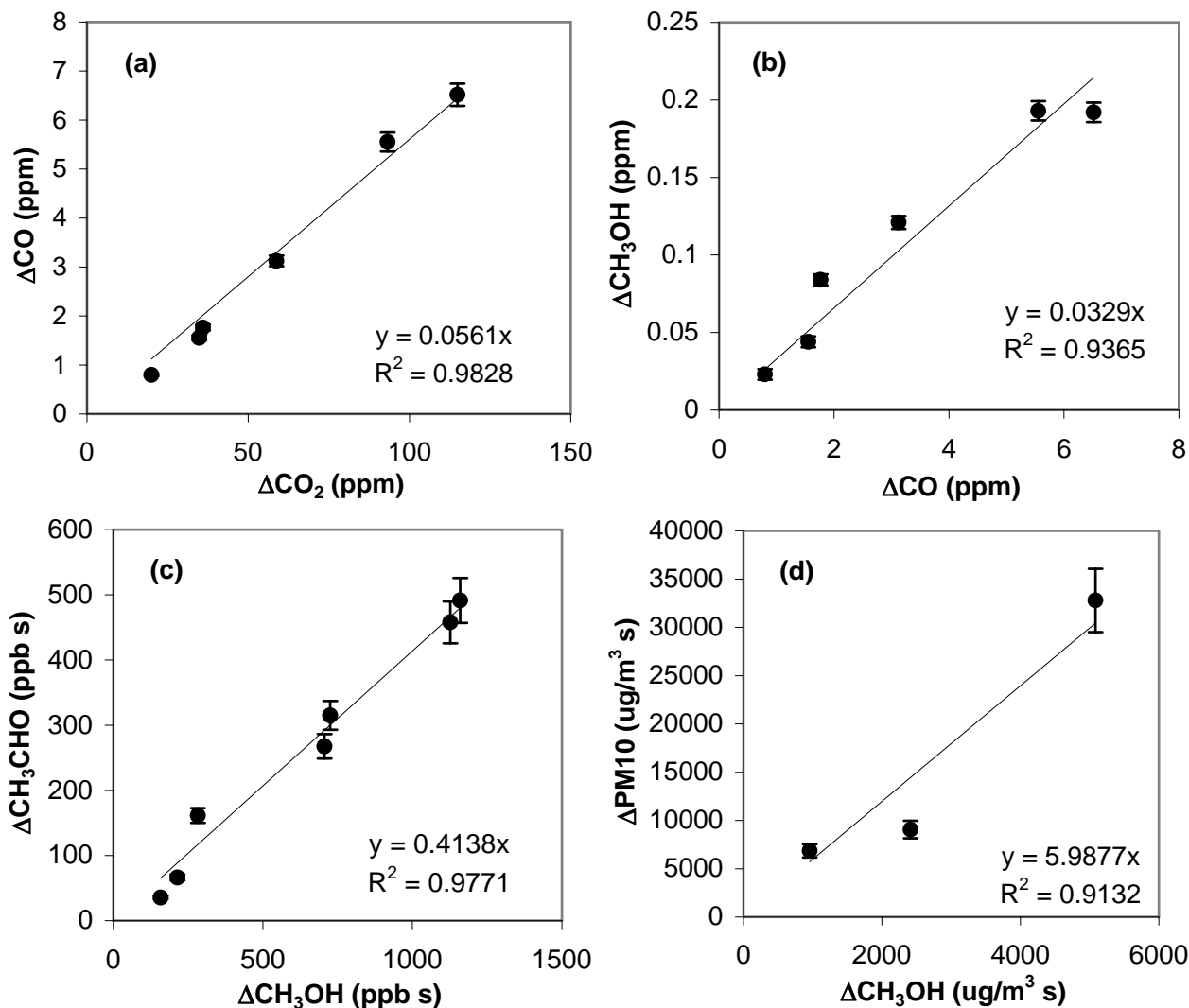


Fig. 2. Examples of the plots used to derive emission ratios (ER) in this work. See Sect. 2.2.1 for details. (a) plot used to derive the ER $\Delta\text{CO}/\Delta\text{CO}_2$ from AFTIR grab samples of the 5 September planned fire. (b) as in a for the ER $\Delta\text{CH}_3\text{OH}/\Delta\text{CO}$. (c) plot for the ER $\Delta\text{CH}_3\text{CHO}/\Delta\text{CH}_3\text{OH}$ from integrated PTR-MS traces during plume penetrations of the 5 September fire. (d) plot used to derive the ER $\Delta\text{PM}_{10}/\Delta\text{CH}_3\text{OH}$ (mass ratio) from integrated PTR-MS and nephelometer traces during plume penetrations of the shifting cultivation fire on 30 August.

sampling. When dry, large-diameter fuels are present RSC may account for a large part of the total biomass consumed (Bertschi et al., 2003b; Kauffman et al., 1998).

2.3.2 Overview of Brazilian biomass burning

This section summarizes Brazilian biomass burning to help assess the representativeness of the fires we actually sampled. Brazil contains $\sim 2 \times 10^6$ km² of savanna (cerrado), mostly in southern Brazil, which is burned every 1–3 years in fires that rapidly consume 5–10 Mg/ha of mostly fine fuels such as grass (Coutinho, 1990; Ward et al., 1992; Kauffman et al., 1994; Andrade et al., 1999). For estimating the emissions

from any global savanna fire, we recommend the tables for savanna fires in Christian et al. (2003) and Andreae and Merlet (2001).

Brazil has $\sim 4 \times 10^6$ km² of evergreen tropical forest mostly in the Amazon basin, which represents $\sim 25\%$ of the world's total "rainforest." Deforestation rates in the Amazon since 1978 ranged from $11\text{--}29 \times 10^3$ km²/y (<http://www.obt.inpe.br/prodes/>). About 85% of the cumulative deforested area for 1978–2005 occurred in the rapidly developing southern and eastern edges of the Amazon basin where the states of Pará and Mato Grosso are located (Fig. 1). Deforestation fires involve large total aboveground biomass (TAGB)

loading averaging ~ 300 Mg/ha of which $\sim 40\%$ is consumed by the fires for a total fuel consumption of ~ 120 Mg/ha. (Ward et al., 1992; Fearnside et al., 1993; Carvalho et al., 1998, 2001; Guild et al., 1998).

Pastures established in previously forested areas of the Amazon are maintained by burning every 2–3 years (Guild et al., 1998). The TAGB can be quite large partly because residual wood debris (RWD) persists for many years. Reported TAGB ranges from 119 Mg/ha (87% RWD) to 53 Mg/ha (47% RWD) in pastures 4–20 years old (Barbosa and Fearnside, 1996; Guild et al., 1998; Kauffman et al., 1998). Large-diameter RWD accounted for $\sim 45\%$ of the fuel consumption in the above studies. Until recently, nearly all deforested areas in the Brazilian Amazon were eventually converted to pasture and the total emissions from Brazilian pasture fires are thought to be comparable to the total emissions from Brazilian deforestation fires (Fearnside, 1990; Barbosa et al., 1996; Kauffman et al., 1998). Globally, deforestation fires associated with shifting cultivation and plantation establishment dominate and pasture fires are relatively less common.

Brazil has recently seen explosive growth in large-scale, mechanized agriculture, especially in Mato Grosso (Cardille and Foley, 2003). Both pastures and forest are converted to croplands for (mostly) soy. In either case, all large-diameter fuels must be removed by the burns. Morton et al. (2006) found that Mato Grosso accounted for 40% of the new deforestation in Amazonia from 2000–2004. Within Mato Grosso from 2001–2004, pasture was still the main use following deforestation, but that fraction was decreasing and direct transition to large areas of cropland accounted for 23–28% of deforestation. Thus, we conclude that the expansion of mechanized agriculture could imply an increase in both the area of individual fires and the fuel consumed per unit area.

A few other less dominant fire-types occur in Brazil. Secondary forests are used in similar fashion to primary forests (Fearnside, 1990, 2000). Lower intensity fires occur naturally in seasonally dry forests, such as the Caatinga in eastern Brazil and these forests are also subject to land-use change (Kauffman et al., 1993). Selective logging promotes fire susceptibility and is increasing in the Amazon (Grainger, 1987; Kauffman and Uhl, 1990; Cochrane et al. 1999; Laurance, 2000).

As discussed in detail by Christian et al. (2007b), RSC could produce a large part of the Amazonian fire emissions and this motivated our simultaneous airborne and ground based campaigns. However, RSC likely occurs mostly on pasture maintenance fires rather than the deforestation fires, which were our main target.

2.3.3 Description of the fires sampled in the airborne campaign

Nearly all the fires we observed in Mato Grosso and southern Pará were related to the expansion of existing, large farms or ranches (Table 1). All but 3 of

these fires were located on the edge of forested areas that were adjacent to large tracts of cleared, often cultivated, land. Casual examination of MODIS visible images of this region reveals that nearly all hotspots are located at the edge of dark-green (forested) areas, adjacent to light-green (cleared) areas (http://rapidfire.sci.gsfc.nasa.gov/subsets/?AERONET_Alta_Floresta/2004252). However, the second fire sampled on 29 August was in a grass meadow and no large fuels were visible from the air. This was probably a maintenance fire for an older pasture. The other exception was two small fires observed on 31 August adjacent to the Xingu River in the center of an indigenous reserve and far from any visible clearings or roads. These fires were likely due to shifting cultivation and the one we sampled is labeled the “SC” fire in Tables 1 and 2. Complete burning of logging slash to prepare for mechanized agriculture can be promoted by bulldozing the fuel into long piles (“windrows”) that were observed from the aircraft on at least one group of fires (30 August Fires 1–4). In all areas, the fires frequently occurred in clusters.

TROFFEE supported a planned, deforestation fire on a farm near Alta Floresta under the supervision of João Carvalho (University of Estadual Paulista) and Ernesto Alvarado (University of Washington). Measurements included fuel consumption, charcoal production, propagation of smoldering combustion, forest flammability adjacent to clearcuts, on-site meteorology, fire effects on groundwater chemistry, and recovery and regeneration of burned areas. The emissions from this fire were sampled by ground-based FTIR (Christian et al., 2007b) and the TROFFEE aircraft (5 September data in Tables 1 and 2). In summary, pasture fires were undersampled relative to their importance in Brazil, but we achieved our objective of comprehensive chemical sampling of the emissions from deforestation fires, which are far more significant globally.

3 Results and discussion

3.1 Characteristics of clean background air

We briefly summarize some of the data obtained in early dry season, clean air near Manaus (see also Karl et al., 2007b). These data are of intrinsic interest and by comparison to data from the more active burning region further south (Sect. 3.2), they highlight the degree to which fires can perturb background air over a large geographic area. Figure 3a shows all the AFTIR CO grab samples from 6 and 7 September, obtained in the Manaus region, which was not visibly impacted by a biomass burning haze before noon. The CO average was 134 ± 13 ppbv. This is a relatively narrow range. Chou et al. (2002) measured numerous CO vertical profiles in nearly the same location in April–May 1987. Their figures indicate that their CO values averaged about 100 ppb. The larger values we observed could be due to a gradual

Table 2. Initial emission factors for the fires sampled at their source during the 2004 TROFFEE airborne campaign. Effective emission factors for the mega-plume, which was sampled downwind from source.

Compound formula or name	29 Aug	29 Aug	30 Aug	30 Aug	31 Aug	31 Aug	3 Sep	5 Sep	7 Sep	Study average EF g/kg	Standard deviation EF g/kg	8 Sep	MPEEF- average ^c # stdev's
	Fire 1	Fire 2	Fires 1–4	SC Fire	Fire 1	Fire 2	Fires 1–5	Planned Fire EF g/kg	Fires 1–4			Mega-Plume Effective EF g/kg	
AFTIR species													
CO ₂	1638	1591	1567	1579	1603	1636	1579	1679	1662	1615	40	1651	0.91
CO	95.72	112.08	133.45	124.82	110.70	93.13	110.52	59.91	72.36	101.41	23.78	87.54	−0.58
MCE	0.916	0.900	0.882	0.890	0.902	0.918	0.901	0.947	0.936	0.910	0.021	0.923	0.61
NO	0.283	nm ^b	0.281	0.514	0.208	0.438	0.746	2.681	nm	0.74	0.877	2.297	1.78
NO ₂	1.979	0.930	1.157	0.509	0.738	2.216	1.393	3.441	4.120	1.83	1.245	1.899	0.05
NO _x (as NO)	1.574	0.606	1.035	0.846	0.690	1.883	1.654	4.926	2.687	1.77	1.359	3.535	1.30
HONO	0.345	0.167	nm	nm	nm	nm	nm	nm	nm	0.26	0.126	nm	nm
CH ₄	4.213	6.916	5.751	7.544	5.323	5.486	7.220	3.353	5.324	5.68	1.380	7.636	1.42
C ₂ H ₄	0.747	1.238	0.958	1.215	0.997	0.809	1.520	0.642	0.454	0.95	0.332	0.378	−1.73
C ₂ H ₂	0.094	nm	0.083	0.101	0.140	0.084	0.172	0.923	0.620	0.28	0.317	0.085	−0.61
C ₂ H ₆	0.548	1.137	0.917	1.157	0.893	0.532	1.478	nm	nm	0.95	0.341	nm	nm
C ₃ H ₆	0.452	0.728	0.424	0.606	0.462	0.317	0.509	0.091	nm	0.45	0.190	nm	nm
HCHO	1.277	1.912	1.674	1.783	1.445	1.517	2.201	1.741	1.409	1.66	0.286	1.004	−2.30
CH ₃ OH	2.077	2.874	2.724	3.371	2.294	2.331	3.002	2.252	2.165	2.57	0.445	2.550	−0.04
CH ₃ COOH	3.134	4.172	3.635	3.590	2.643	3.232	3.190	3.579	3.704	3.43	0.436	9.242	13.33
HCOOH	0.398	0.519	0.377	0.223	0.246	0.508	0.323	0.978	1.715	0.59	0.479	3.266	5.59
NH ₃	1.127	1.364	1.093	1.769	0.653	0.658	1.476	1.236	0.308	1.08	0.460	1.509	0.94
HCN	0.665	0.537	0.699	0.582	0.486	0.409	0.426	2.098	0.184	0.68	0.555	0.169	−0.91
PTR-MS species and PM ₁₀													
acetonitrile	0.574	0.276	0.270	0.381	0.291	0.347	0.485	0.359	0.336	0.37	0.101	nm	nm
acetaldehyde	1.255	1.202	1.167	1.240	0.751	1.041	3.322	1.282	1.202	1.38	0.745	nm	nm
acrylonitrile	0.051	nm	0.038	nm	0.020	0.048	nm	nm	nm	0.04	0.014	nm	nm
acrolein	nm	nm	nm	nm	0.306	0.477	nm	0.808	0.732	0.58	0.232	nm	nm
acetone ^a	0.429	0.525	0.645	0.673	0.235	0.506	0.803	0.694	0.590	0.57	0.167	nm	nm
propanal ^a	0.067	0.082	0.101	0.105	0.037	0.079	0.126	0.109	0.092	0.09	0.026	nm	nm
isoprene ^a	0.236	0.366	0.402	0.396	0.271	0.296	0.625	0.378	0.386	0.37	0.112	nm	nm
furan ^a	0.207	0.320	0.352	0.347	0.237	0.259	0.547	0.331	0.338	0.33	0.098	nm	nm
methylvinyl ketone ^a	0.166	0.499	0.340	nm	0.399	0.318	0.215	0.411	0.436	0.35	0.113	nm	nm
methacrolein ^a	0.066	0.198	0.135	nm	0.158	0.126	0.085	0.163	0.173	0.14	0.045	nm	nm
crotonaldehyde ^a	0.100	0.302	0.205	nm	0.241	0.192	0.130	0.248	0.263	0.21	0.068	nm	nm
methylethyl ketone ^a	0.229	0.469	nm	nm	nm	nm	0.654	nm	nm	0.45	0.213	nm	nm
methyl propanal ^a	0.081	0.165	nm	nm	nm	nm	0.230	nm	nm	0.16	0.075	nm	nm
acetol and methylacetate	nm	nm	0.649	nm	0.840	0.607	0.895	0.700	0.627	0.72	0.120	nm	nm
benzene ^a	0.189	0.381	0.168	nm	0.538	0.176	0.234	0.261	0.172	0.26	0.131	nm	nm
C ₆ carbonyls	0.098	0.307	0.105	nm	Nm	nm	0.241	0.363	0.159	0.21	0.109	nm	nm
3-methylfuran ^a	0.252	0.707	0.434	nm	0.843	0.389	0.668	0.511	0.413	0.53	0.196	nm	nm
2-methylfuran ^a	0.036	0.101	0.062	nm	0.120	0.056	0.095	0.073	0.059	0.08	0.028	nm	nm
hexanal ^a	0.006	0.017	0.010	nm	0.020	0.009	0.016	0.012	0.010	0.01	0.005	nm	nm
2,3-butanedione ^a	0.317	0.790	0.509	nm	0.855	0.490	0.995	0.634	0.659	0.66	0.219	nm	nm
2-pentanone ^a	0.032	0.085	0.052	nm	0.094	0.051	0.106	0.066	0.069	0.07	0.024	nm	nm
3-pentanone ^a	0.014	0.038	0.023	nm	0.042	0.023	0.047	0.029	0.031	0.03	0.011	nm	nm
toluene	0.102	0.109	0.126	nm	0.227	0.096	0.399	0.135	0.368	0.20	0.123	nm	nm
phenol ^a	nm	nm	nm	nm	nm	nm	nm	0.406	0.282	0.34	0.088	nm	nm
other substituted furans	nm	nm	nm	nm	nm	nm	nm	1.095	1.071	1.08	0.016	nm	nm
furaldehydes	nm	nm	nm	nm	nm	nm	nm	0.255	0.256	0.26	0.001	nm	nm
xylenes ^a	0.086	0.092	0.076	nm	0.132	0.060	0.322	0.137	0.115	0.13	0.083	nm	nm
ethylbenzene ^a	0.053	0.084	0.047	nm	0.118	0.044	0.126	0.078	0.052	0.08	0.033	nm	nm
PM ₁₀	17.61	14.43	17.94	20.18	19.81	17.27	26.41	12.53	14.28	17.83	4.121	nm	nm
UCI-Canister species													
OCS	nm	nm	nm	nm	nm	nm	nm	0.0247	nm	0.0247	nm	nm	nm
DMS	nm	nm	nm	nm	nm	nm	nm	0.0022	nm	0.0022	nm	nm	nm
CFC 12	nm	nm	nm	nm	nm	nm	nm	0.0028	nm	0.0028	nm	nm	nm
MeONO ₂	nm	nm	nm	nm	nm	nm	nm	0.0163	nm	0.0163	nm	nm	nm
EtONO ₂	nm	nm	nm	nm	nm	nm	nm	0.0057	nm	0.0057	nm	nm	nm
i-PrONO ₂	nm	nm	nm	nm	nm	nm	nm	0.0010	nm	0.0010	nm	nm	nm
n-PrONO ₂	nm	nm	nm	nm	nm	nm	nm	0.0003	nm	0.0003	nm	nm	nm
2-BuONO ₂	nm	nm	nm	nm	nm	nm	nm	0.0006	nm	0.0006	nm	nm	nm
C ₂ H ₆	nm	nm	nm	nm	nm	nm	nm	0.5600	nm	0.5600	nm	nm	nm
1-Butene	nm	nm	nm	nm	nm	nm	nm	0.0200	nm	0.0200	nm	nm	nm
trans-2-Butene	nm	nm	nm	nm	nm	nm	nm	0.0161	nm	0.0161	nm	nm	nm
cis-2-Butene	nm	nm	nm	nm	nm	nm	nm	0.0202	nm	0.0202	nm	nm	nm

^aA branching ratio has been applied to the signal from a single mass channel as measured by Karl et al. (2007a).^bnm indicates “not measured.”^cThe mega-plume effective emission factor minus the study average emission factor given as the number of standard deviations in the study-average emission factor.

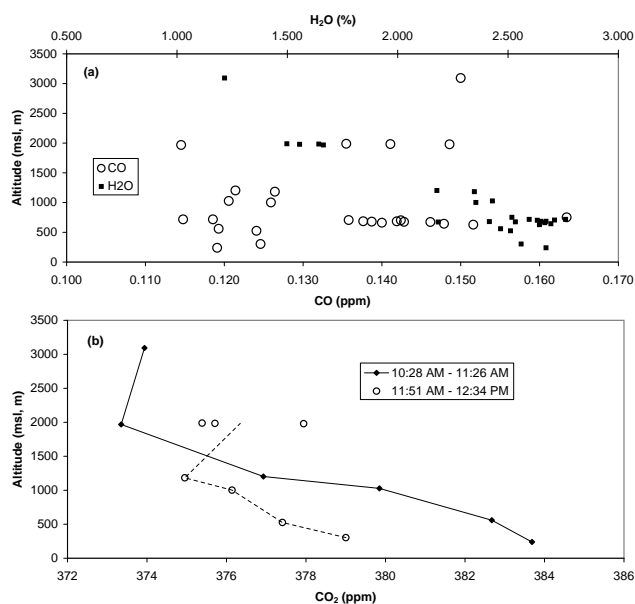


Fig. 3. Clean background air, early in the local dry season, near Manaus on 6 and 7 September, 2004. **(a)** CO and water from AFTIR grab samples of ambient air. Air parcels with high CO above the boundary layer were likely affected by transport from a biomass burning region to the southeast. **(b)** AFTIR vertical profiles for CO₂ above the ZF-14 Tower on 6 September showing progressive depletion by photosynthesis of the CO₂ that builds up overnight from respiration.

increase in pollution in the area and/or the fact that our measurements occurred part way into the beginning of the dry season so that there were probably small enhancements from biomass burning. (A few fires were sampled around noon on 7 September.) Figure 3a also shows the water mixing ratios. The higher altitude CO samples are from above the mixed layer and they show some of the higher mixing ratios. This is consistent with HYSPLIT back-trajectories (Draxler and Rolph, 2003) indicating that the air at this altitude was transported from a region to the southeast with much active burning as suggested by numerous NOAA-12 hotspots. In contrast, HYSPLIT back-trajectories show that the mixed layer air came from the northeast, which was a region mostly free of hotspots.

Figure 3b shows two CO₂ vertical profiles above the ZF-14 Tower northeast of Manaus. One is from late morning and the other is from midday. The profiles are consistent with the CO₂ profiles observed by Chou et al. (2002) in the same region. The morning profiles show CO₂ enhancements at lower altitudes due to nighttime respiration exceeding photosynthesis and, as the day progresses, the enhancements decrease as the forest “draws down” CO₂. Chou et al. (2002) actually observed a CO₂ deficit at lower altitudes by afternoon, but we did not measure afternoon vertical profiles. Our higher altitude CO₂ shows small increases in the later pro-

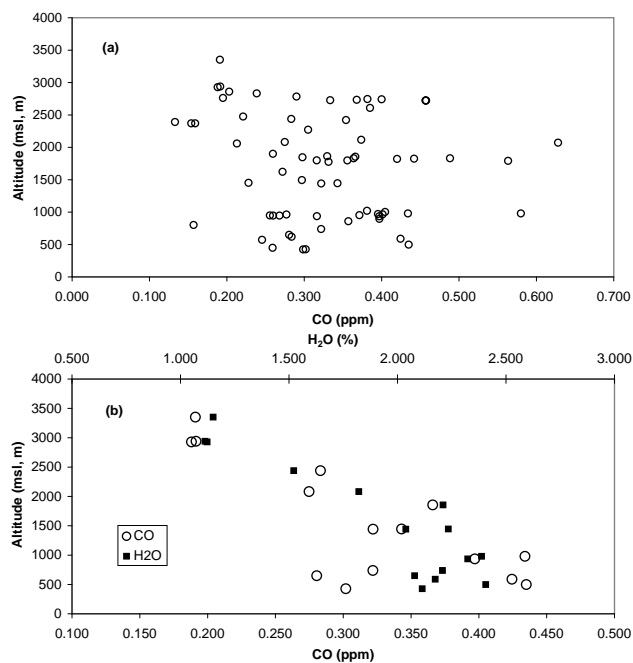


Fig. 4. Regional haze due to biomass fires late in the dry season near Alta Floresta. **(a)** CO from AFTIR grab samples of ambient air vs. altitude for 29 August–5 September. **(b)** as in a for CO and H₂O for 30 and 31 August only, illustrating efficient, initial trapping of the fire-caused haze in the mixed (boundary) layer. (The water vertical profiles can be obtained in other units from the authors.)

file, which could also be consistent with some transport of biomass burning emissions in the upper layer. The main difference between Chou et al. (2002) and our current measurements is the obvious effect of increasing global CO₂. Their 1987 CO₂ values average around 350 ppm, while our 2004 average for the same region is around 380 ppm. Above the ZF-14 tower, our PM₁₀ ranged from $\sim 40 \mu\text{g m}^{-3}$ near the surface to $\sim 30 \mu\text{g m}^{-3}$ near the top of the mixed layer. Our O₃ ranged from 1–10 ppbv near the surface and increased to 20–30 ppbv near the top of the profiles. Our O₃ profile is similar to that of Chou et al. (2002).

3.2 Characteristics of aged regional smoke haze

In contrast to the region near Manaus, the region near Alta Floresta was well into the local dry season and heavily impacted by numerous fires that caused a regional haze of aged smoke sequestered in the mixed layer. (The fire emission factors in Table 2 are derived only from smoke < a few minutes old that was sampled in concentrated, visually-obvious plumes and not from smoke of unknown age that constitutes the regional haze layer.) Figure 4a shows all the CO values from AFTIR grab samples that were not in smoke plumes in this region. The range is from 100–600 ppb with an average and standard deviation of 328 ± 102 ppb. Thus

the background, mixed-layer air in this large fire-impacted region had about 2.5 times as much CO as was found near Manaus. This degree of impact is similar to the impact on dry season CO observations at the same latitude in Africa (Fig. 1b of Yokelson et al., 2003a).

Most of the lower CO values were observed above the mixed layer as can be seen more easily in Fig. 4b. Figure 4b shows the CO and water AFTIR grab sample data in background air for 30 and 31 August. On these days we spent relatively more time above the boundary layer so the vertical patterns are more apparent. The water and CO mixing ratios dropped off with altitude in remarkable correlation. This is consistent with our visual observation that the plumes from active fires rarely penetrated the top of the mixed layer; a limitation that was also observed during the southern African biomass burning season (Yokelson et al., 2003a). Interestingly, the smoky mixed layers in Brazil in 2004 extended to only 2–3 km altitude; much lower than the 5–6 km altitude observed at the same latitude, time-of-year, and local time-of-day in Africa during 2000 (Yokelson et al., 2003a (Fig. 1); Schmid et al., 2003 (Fig. 11)).

We can compare our airborne CO observations in the 2004 regional smoke/haze to a long record of previous airborne measurements in Brazil. In 1979 and 1980 Crutzen et al. (1985) measured CO from 100–400 ppb in haze layers over the Amazon (their Figs. 10 and 11). The 1985 study of Andreae et al. (1988) shows Amazon dry season CO ranging from 150–600 ppb (their Fig. 4). Kaufman et al. (1992) also reported haze layer CO ranging from ~150–600 ppb in 1989 (their Fig. 4). Blake et al. (1996) observed haze layer CO values from ~100–400 during TRACE A in 1992. In 1995 biomass burning was well above average in Brazil. The SCAR-B mission was conducted late in the 1995 dry season as biomass burning peaked and Reid et al. (1998) observed much higher levels of CO than we have presented thus far. Average CO values for flights based in several central Brazil locations ranged from 440–760 ppb (their Table 1).

Our average PM₁₀ values for vertical profiles in the regional haze layer ranged from 70–120 $\mu\text{g}/\text{m}^3$ at 300–500 m to 30–60 $\mu\text{g}/\text{m}^3$ near the top (~3000 m); similar to observations in previous years (Pereira et al., 1996; Reid et al., 1998). Ozone values were about 30 ppbv throughout these haze layers similar to the observations in the CITE-3, Brush-fire, and ABLE-2A studies referred to above. During SCAR-B, however, O₃ ranged from 60–100 ppb, consistent with the more polluted boundary layer present in the late 1995 dry season.

In summary, 2004, through 7 September, had a typical amount of biomass burning haze based on the comparison of our CO, PM₁₀, and O₃ measurements to other measurements from the last 30 years. However, as discussed in Sect. 3.4, our measurements on 8 September probed widespread, unusually high levels of pollutants.

It is also of interest to compare the airborne CO data with the CO data obtained during the same time period

by the ground-based FTIR system (Christian et al., 2007b). The ground-based samples obtained well away from visible smoke plumes return much higher CO values. The average for 25 afternoon samples taken within ~100 km of Alta Floresta from 26 August–8 September was 1.35 ± 1.15 ppm with a range from 0.330 to 4.76 ppm. Gatti et al. (personal communication) monitored CO levels at a pasture site in Rondonia in September and October of 1999 and observed a range of CO from 0.6 to 1.3 ppm. Thus while airborne sampling retrieved the composition of the majority of the mixed layer, more polluted air was found at ground level than would be inferred from airborne measurements. At this time we don't know the thickness of the ground-level layer. Above the mixed layer, the CO tends to drop off sharply to a mixing ratio characteristic of the free troposphere. The African and the Brazilian CO vertical profiles are not shaped like the a-priori CO vertical profile used for MOPITT CO retrievals (Emmons et al., 2004). We speculate that consideration of the actual profile shapes might enhance CO retrievals from space-based instruments.

3.3 Initial emissions from tropical deforestation fires

Since a variety of large changes can occur in smoke chemistry in the minutes to days after emission, segregation of results by sample age and history (to the degree possible) enhances interpretation of the results and comparison with models and other measurements. Thus, only excess mixing ratios measured <~1 km from the fire were used to compute our initial emission ratios and emission factors. Forty-two plume penetrations of this type were made. In contrast to the background-air grab samples discussed above, the excess CO mixing ratios (above background) in the AFTIR, plume grab samples were in the range 1–31 ppmv for ~90% of the samples. Thus, excellent signal to noise was observed on all instruments for each fire for numerous species.

The fire-average, initial emission factors for each compound and fire, along with the fire average MCE, are listed in Table 2. Because NO is rapidly converted to NO₂ (largely due to reaction with O₃ in the entrained background air), we also report a single EF for “NO_x as NO”. We computed this EF from the NO_x/CO₂ molar ER obtained as described in Sect. 2.2.1, but it can also be estimated from Table 2 data using: $\text{EFNO} + (30/46) \times \text{EFNO}_2$. If desired, the molar ER for each fire can be derived from the EF in Table 2 after accounting for any difference in molecular mass.

The timing and extent, and perhaps representativeness, of Brazilian biomass burning in 2004 can be compared to other years using metrics other than the regional CO, PM, and O₃ values discussed in Sect. 3.2. Dating back to at least 1993 a near-continuous, regional record of aerosol optical thickness (Holben et al., 1996; Echalar et al., 1998; <http://aeronet.gsfc.nasa.gov/newaeronet1.html>) and deforestation rates exists. Unfortunately, the Alta Floresta sun photometer was not operational during the peak of the 2004 burning season

(B. Holben, personal communication). The INPE deforestation data, however, shows 2004 (27 429 km²) as the second highest year after 1995 (29 059 km²) – the year of the SCAR-B campaign. Thus, both TROFFEE and SCAR-B were conducted in years when the deforested area was well above the long-term average of $\sim 20\,000$ km². The number of NOAA-12 hotspots (<http://www.cptec.inpe.br/queimadas/>) for 2004 (236 821) is above the 2000–2005 average of 192 569 and just above 2002 (232 921), which was the second-biggest year since 2000. Interestingly, 2002 was the year for another smoke-sampling campaign termed SMOCC (Andreae et al., 2004). While the annual totals for the NOAA-12 hotspots are readily available, they likely underestimate the true number of fires, especially under extreme burning conditions as discussed in Sect. 3.4. In summary, most of our fire sampling was probably conducted under “average” conditions as shown in Sect. 3.2, however, the 2004 annual burning was above average because of intense burning beginning ~ 7 –8 September, which was towards the end of our campaign.

3.3.1 Natural variation in emission factors

In Fig. 5 we plot the fire-average emission factors versus MCE (data from Table 2) for selected compounds. This gives some idea of the natural variation in emission factors that results from deforestation fires burning under a range of vegetative/environmental conditions and with different mixtures of flaming and smoldering combustion. Figure 5a shows NO_x emissions which increase as MCE (and thus flaming combustion) increases. Figures 5b–d show the pattern typical of most of the VOC we measured – the EF for these “smoldering compounds” increased with decreasing MCE. Figure 5e shows that EFPM₁₀ also increases with decreasing MCE. The range in EF (with MCE) for these species is about a factor of two, which is a smaller range than we observed for African savanna fires (Yokelson et al., 2003a). Figure 5f shows that EFCH₃CN did not have a strong dependence on MCE. This is similar to the pattern observed for HCN from savanna fires by Yokelson et al. (2003a). However, like EFHCN, the EFCH₃CN did vary by $\sim \pm 50\%$, possibly due to varying fuel N content. The use of acetonitrile as a biomass burning indicator/tracer is discussed later in this paper and by Karl et al. (2007a).

3.3.2 Comparison with other work

It is most meaningful to compare our study-average, initial emission-factor measurements in nascent smoke from Brazilian deforestation fires with measurements made in August–September of 1990 using a tower-based platform by Ward et al. (1992) during BASE-B; and in August–September of 1995 from an aircraft by Ferek et al. (1998) as part of SCAR-B. We also compare to a widely-used compilation of EF for tropical forests by Andreae and Merlet (2001).

The EFCO₂, EFCO, and, especially, MCE all reflect the overall mix of flaming and smoldering combustion in a fire and thus these parameters can give some idea of the similarity of the combustion characteristics of the fires we sampled to fires sampled previously. This serves as one probe of how representative our fires were of regional fires in general. Ward et al. and Ferek et al. report individual values for flaming and smoldering combustion and it is not always clear if they have a recommended study-average for primary forest fuels. However, our study-average MCE for deforestation fires (Table 2) indicates that they burn with roughly equal amounts of flaming and smoldering (Yokelson et al., 1996). Thus, when necessary, we compare to the average of the flaming and smoldering values given in the other work in the following discussion.

For CO₂ the EF are 1614 \pm 56 (Ward et al., 1992), 1599 (Ferek et al., 1998), and 1580 \pm 90 (Andreae and Merlet). All these values are reasonably close to each other and our study average of 1615 \pm 40. Similarly for CO the previous values are 110 \pm 28, 105, and 104 \pm 20 in excellent agreement with each other and our value of 101 \pm 24. The MCE are 0.903 \pm 0.03, 0.906, 0.906, and our value of 0.910 \pm 0.021. Thus our values are well within the range of previous measurements, but seem to reflect slightly more flaming combustion on average than previous work.

The research fire on 5 September, which was designed to simulate regional fires apparently had a significantly higher MCE than our regional average. However, the higher MCE partly reflected that we did sample the beginning of the fire, but could not finish sampling the full fire (smoldering contributes less at the beginning of a fire) because of aircraft fuel considerations. Our airborne samples showed that MCE initially decreased with time and then stabilized. It is also interesting to note that the fires sampled later in TROFFEE tended to have higher MCE, which could be due to the protracted dry period after unusual rains in mid August. Finally, the plume from the intense burning event sampled on 8 September (see Sect. 3.4) also had higher than study-average MCE. Thus late-season, “higher-MCE” plumes may account for a fair percentage of the total regional biomass burned. On the other hand, prolonged dry spells will desiccate large diameter logs, which tend to burn with a low MCE (~ 0.788 , Christian et al., 2007b) producing initially unlofted smoke. So the real nature of the “total regional smoke” is governed by complex – sometimes competing – trends, which need further analysis.

Rather than an exhaustive species by species comparison with other work for the numerous other trace gases measured, we have tried to summarize the comparison in Fig. 6 and provide some useful guidance. Then a few comments are made about select individual species. Many compounds appear in both our work (Table 2) and the recommendations of Andreae and Merlet (AM). In general, our values are based on a larger number of measurements and should probably be preferred to those in AM who acknowledge basing many

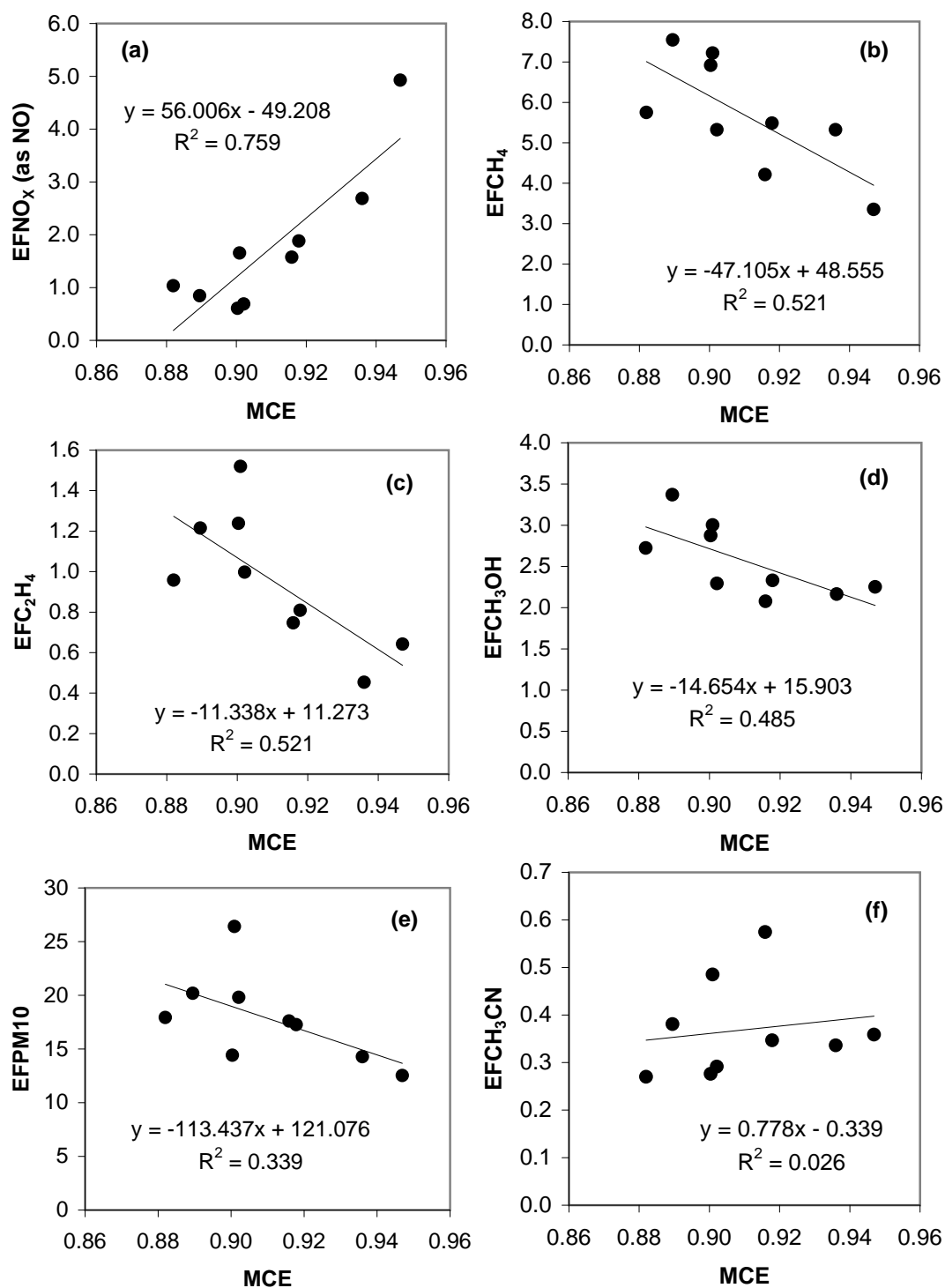


Fig. 5. Fire-average emission factors (EF) plotted versus fire-average modified combustion efficiency (MCE) for the indicated species (data from Table 2). (See discussion in Sect. 3.3.1).

of their values on 1–2 less direct measurements and/or “best guesses” due to a lack of detailed information available at the time. On the other hand, a number of compounds appear

in the AM recommendations that we did not measure during TROFFEE. Most of these are minor plume constituents, but some are of major importance (e.g. SO_2). We recommend

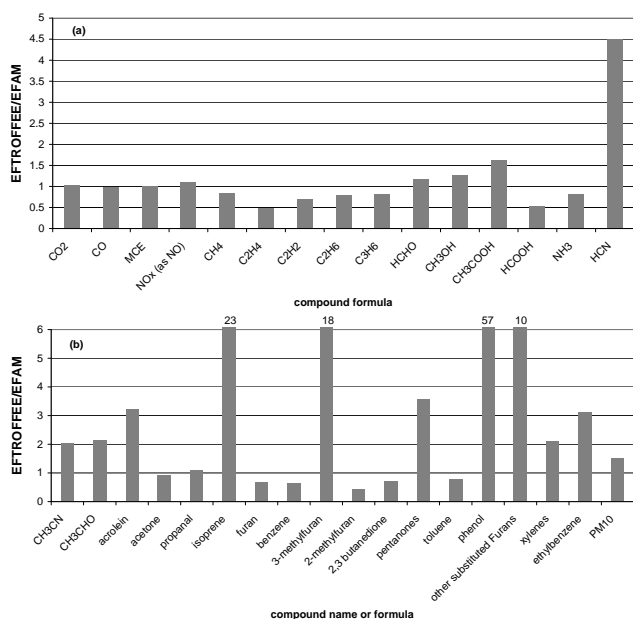


Fig. 6. Comparison of the TROFFEE airborne study emission factors with the recommendations of Andrea and Merlet (2001) (AM) for species in both studies. (AM PM₁₀ is taken as $1.3 \times$ AM PM_{2.5}.) With 8 exceptions, the older (AM) recommendations are within a factor of ~ 2 of the newer TROFFEE EF, which are usually based on more measurements. This suggests that the AM recommendations for species not measured in TROFFEE (e.g. SO₂) are reasonable. **(a)** species measured by AFTIR. **(b)** species measured by PTR-MS and the nephelometer. (When the ratio exceeds the scale shown the value of the ratio is given above the bar.)

using the AM values for compounds we did not measure since there is reasonable agreement between our work and theirs on most of the compounds we both address (see Fig. 6). Finally our work includes data on a number of “new,” significant plume constituents for which information was not previously available. Included in this category are HONO, acrylonitrile, pyrrole, methylvinylketone, methacrolein, crotonaldehyde, methylethylketone, methylpropanal, “acetol plus methylacetate,” furaldehydes, dimethylsulfide, and C₁–C₄ alkyl nitrates (Table 2).

In early fire research it was usually assumed that most of the NMOC were NMHC as was actually the case for industrial combustion of fossil fuels. As mentioned in the introduction, a key discovery of previous FTIR and PTR-MS work was that OVOC accounted for the large majority of NMOC emitted by the fires sampled. A goal of this project was to verify this for tropical deforestation fires. The TROFFEE data show that the molar ratio OVOC/NMHC is about 4:1 – or that OVOC account for $\sim 80\%$ of the NMOC. With the completion of TROFFEE, there are now reasonably comprehensive field measurements of the NMOC emitted by all the major types of biomass burning. The new information

provided on the “universal dominance” of OVOC is significant because of the huge size of the biomass burning source and the reactive nature of OVOC (Mason et al., 2001; Trentmann et al., 2005).

A few comments are made about individual species we measured. An IR signal due to HONO was observed on the lab fires and 2 field fires, but the measurements are semi-quantitative due to a low SNR. However, the presence of any HONO signal is significant since even a small amount of HONO in the initial emissions is a source of OH that speeds up the initial plume chemistry (Trentmann et al., 2005). Our field, study-average HONO EF (0.26 ± 0.13 g/kg) overlaps the other relevant HONO EF we know of (Keene et al., 2006): 0.24 g/kg shrubs, 0.19 ± 0.08 g/kg branches, and 0.14 ± 0.05 g/kg grass.

As mentioned above, the EF for acetonitrile was not strongly correlated with MCE in our field study. Thus, our study-average EF of 0.37 ± 0.10 g/kg seems to be a good estimate for all tropical deforestation fires regardless of MCE. However, our EF for acetonitrile from deforestation fires does differ significantly from recommended EFCH₃CN for other types of burning (e.g. 0.13 g/kg for savanna fires and 4.91 g/kg for burning Indonesian peat (Christian et al., 2003)). In addition, acetonitrile emissions have not been measured for cooking fires, which may be the second largest type of biomass burning. Still, these results suggest that (with attention to the type of fire) PTR-MS acetonitrile measurements could contribute to source apportionment or estimates of the amount of biomass burned using inverse modeling.

The particle emission factors we measured during TROFFEE (PM₁₀, 17.8 ± 4.1 g/kg) are significantly larger than in previous work or recommendations. Ferek et al. (1998) reported a range of EFPM₄ from 2–21 g/kg and a study average of about 11 g/kg for Brazilian deforestation fires. The tower-based measurements of Ward et al. (1992) returned values for EFPM_{2.5} ranging from 6.8 to 10.4 g/kg with an average of about 9 g/kg for forest fuels. Ferek et al. speculated that their higher average and high end values were due to incomplete particle formation being probed from the tower platform. This hypothesis was supported by simultaneous tower and airborne PM measurements on the same Brazilian fires (Babbitt et al., 1996). In that experiment, the airborne EFPM_{2.5} averaged about 11 g/kg while the EFPM_{2.5} measured on the same fires from towers averaged about 4 g/kg. In any case our study average value for PM₁₀, which includes a wider range of particle sizes than the work referenced above, is significantly higher at 17.8 ± 4.1 g/kg. For most types of biomass burning the PM₁₀ values might be expected to be about 30% higher than the PM_{2.5} or PM₄ values (AM, Ottmar, 2001). Applying this factor to the study average of Ferek et al. gives a projected PM₁₀ of about 14 g/kg – still lower than our TROFFEE value. A major reason for the rest of this discrepancy could be related to fire size and intensity. Ferek et al. noted that their largest, most intense fire in

Brazil had a much higher EFPM₄ or PM₄/CO ratio than the other fires they sampled in SCAR-B. They cited their measurements on even larger more intense fires in North America, which had even higher EFPM₄, and proposed that EFPM increase with fire size and combustion intensity. For example they cited EFPM_{3,5} from 15–25 g/kg (implying an average PM₁₀ of ~26 g/kg) for large, intense North American fires (Radke et al., 1991; Hobbs, 1997). In our TROFFEE data, the lowest EFPM₁₀ (12–14 g/kg) are from our smallest fires (5 and 7 September). Our largest EFPM₁₀ (26.4 g/kg) was obtained on 3 September. This plume was the largest and most intense we encountered. Thus we speculate that our larger study-average EFPM values for Brazil could be due to sampling larger, more-intense fires (on average) than in previous studies in Brazil. If correct, this raises two interesting questions: (1) what fire sizes contribute what fraction of the regional biomass burning and (2) is there a trend in fire size related to trends in land-use (Sect. 2.3.2).

A species by species comparison of the emissions for the three main types of burning (savannas, cooking, and deforestation) is beyond the scope of this paper. Here we just point out a few main characteristics of the 3 main types of burning. Cooking fire emissions occur year round and are not initially lofted. The emissions immediately impact human health (Bertschi et al., 2003a). The average MCE is about 0.91 and HCN was not observed from cooking fires. In contrast, savanna fires occur only in the dry season, burn with higher MCE (~0.94), and most of the emissions (including ample HCN) are lofted. Tropical deforestation fires also burn in the dry season (with MCE ~0.91) and generally feature higher smoldering compound emissions per unit mass of fuel, higher fuel loadings, and more emissions per unit area than savanna fires. Specifically, the emission factors for methane, acetic acid, acetaldehyde, acetone, formaldehyde, ethane, methanol, ammonia, and acetonitrile were all ~2–3 times higher for tropical deforestation fires than for savanna fires (Christian et al., 2003). Interestingly, the EF for carbon dioxide, ethylene, HCN, formic acid, acetylene, and propylene were about the same for both fire types. This could imply some value for some of these compounds to serve as rough, global non-biofuel biomass burning tracers, but caution is needed as there are other sources of most of these species (Li et al., 2000; Shim et al., 2007). When compared to savanna fires, the EF for CO is significantly higher for tropical deforestation fires and the EF “NO_x as NO” is significantly smaller for these fires. The NMOC are dominated by OVOC for all types of biomass burning. The smoke from outdoor fires that impacts human health and global climate/chemistry is aged. There are two dry seasons in the tropics: ~February–May in the Northern Hemisphere and ~June–October in the Southern Hemisphere. Only November–January are mostly unaffected by outdoor biomass burning emissions.

3.3.3 Regional-global bottom-up emissions estimates

About 2 million ha of tropical rain forest are burned in an average year in Brazil and ~120 Mg/ha of fuel is consumed in these fires (Sect. 2.3.2). Thus, ~2.4×10¹¹ kg of biomass are burned annually in primary deforestation fires. The last value can be multiplied by any EF in Table 2 for a bottom-up estimate of annual emissions from Brazilian tropical deforestation fires. For instance, 388 Tg and 4 Tg are crude estimates of the average annual CO₂ and PM₁₀ emissions from Brazilian deforestation fires. The amount of biomass burned and the total emissions for each species approximately double if pasture fires are included, although 20–50% even larger emissions than predicted by this type of estimate are warranted for several VOC to account for RSC in Brazilian pasture fires (Christian et al., 2007b). The ~240 Tg of primary forest biomass burned each year is about 20% of the total (1330 Tg) consumed by tropical deforestation fires given by AM. This implies that other countries (Indonesia, Congo, Ivory Coast, etc.) have higher, national, deforestation rates. Assuming that the emissions from Brazilian deforestation fires are similar to those from deforestation fires elsewhere in the tropics, we can use our EF with the AM estimate of fuel consumption (above) to estimate global emissions from deforestation fires. For instance, this implies that about 2148 Tg of CO₂ and 24 Tg of PM₁₀ are emitted by deforestation fires globally on an annual basis. Of course it should be remembered that the emissions from any contributing region are emitted in much less than one year and that the vast majority of these species are too reactive to be well-mixed globally.

3.4 Mega-plume

Section 3.2 showed regional CO falling within the typical range observed in haze layers in previous years until 7 September. In contrast, on 8 September, during transit (at ~2.0 km altitude) from Manaus to Cuiabá we encountered CO values as high as 1172 ppb from about 8.3° S (~11:00 a.m. LT) to 13° S (~12:30 p.m. LT) – a distance >500 km (Fig. 7a). Visibility was often too low to see the ground. Thus, no fires were observed from the aircraft during this time. On 8 September there was also a sharp maximum in the daily NOAA-12 hotspot total for Brazil (<http://www.cptec.inpe.br/queimadas/>) and a very large area with high TOMS Earth Probe aerosol index (AI) appeared (Figs. 7b and c). These observations suggest the presence of either “extreme haze” or a massive plume formed from the combined output of numerous fires. Because many relatively short-lived fire emissions were still present (vide infra), we prefer the latter explanation and have termed this phenomenon a “mega-plume.” The mega-plume or “white ocean” of smoke covered a large area in Brazil, Bolivia, and Paraguay for about one month – draining southeastward. Selected, narrated MODIS visible images of the smoke from this “event” are archived for 6 September to 8 October

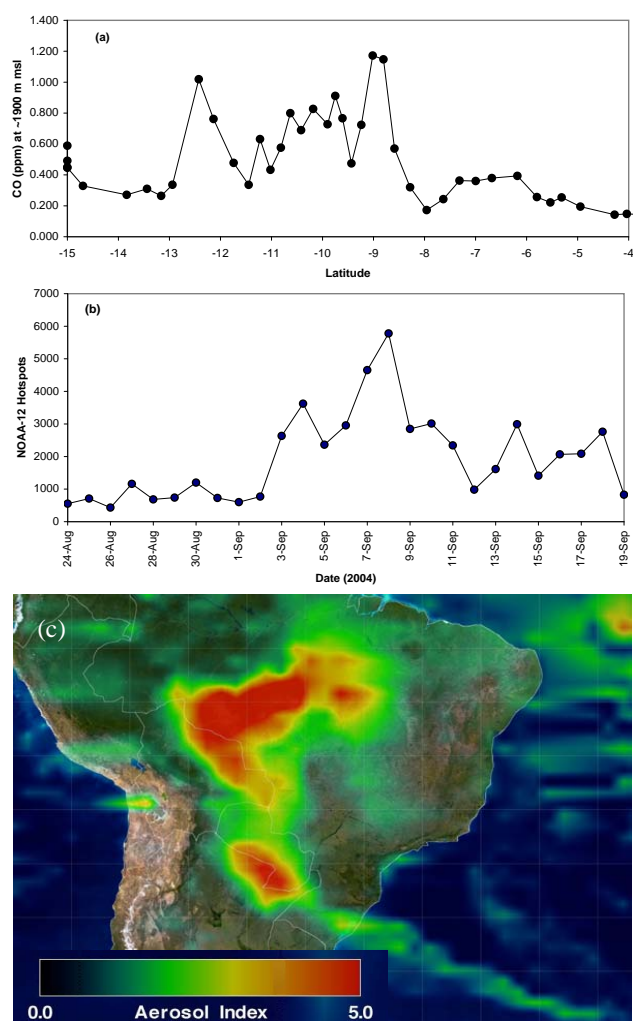


Fig. 7. The mega-plume. (a) CO vs. latitude from the 8 September, 2004 flight showing high CO extending from 8 to 13 south. (b) Daily NOAA-12 afternoon hotspots for Brazil from 24 August to 19 September, 2004. The maximum value occurs on 8 September. (c) TOMS aerosol index on 8 September, 2004.

of 2004 (http://earthobservatory.nasa.gov/NaturalHazards/natural_hazards_v2.php3?img_id=12424) and the TOMS Earth Probe AI was elevated until late September (http://toms.gsfc.nasa.gov/aerosols/aerosols_v8.html). The NOAA-12 hotspots are elevated from ~3–19 September and the hotspots during this period account for roughly one-quarter of the total NOAA-12 hotspots for the year. This suggests that a large fraction of the total biomass burned in 2004 may have produced smoke that was processed in this mega-plume/smoke-ocean event. Brown et al. (2006) describe a similar smoke event in the western Amazon occurring in the fall of 2005. Thus mega-plumes (or smoke oceans) may be fairly common in Brazil and other areas of the tropics when biomass burning is peaking. Whereas smoke plumes usually

age in relative isolation from each other while diluting with regional haze, in the mega-plume scenario, direct mixing of fresh smoke plumes likely dominates. The two chemical processing environments may lead to different outcomes and the latter scenario may be the relevant processing environment for a large part of the total regional emissions.

Only the AFTIR and GPS acquired data on this flight as the other instrumentation had been deployed at the ZF-14 Tower (Karl et al., 2007b). The AFTIR spectra show that the mega-plume contained 10–50 ppbv of numerous reactive species such as NH_3 , NO_2 , CH_3OH , and organic acids and high PM_{10} ($100\text{--}200\ \mu\text{g}/\text{m}^3$) can be inferred from the PM_{10}/CO ERs measured earlier. The mega-plume was not perfectly mixed, however, all the samples showed depletion (relative to CO) of reactive species such as HCHO , C_2H_4 and C_2H_2 . Many samples also showed incipient production of O_3 and both formic and acetic acids. To illustrate the mega-plume chemistry and aging effects, we computed “effective emission factors” (EEF) for the mega-plume that are shown in Table 2. The last sample of clean air (at $\sim 8^\circ\text{S}$, Fig. 7a) was used as the background for the mega-plume samples. The EEF were computed in the same way as EF (Sect. 2.2.2). However, the EEF reflect the temporary composition of the mega-plume at the unknown age that it was sampled. The EEF are not included in the computation of the study-average initial emission factors. Also shown in Table 2, for each species, is the mega-plume EEF minus the study average EF, divided by the standard deviation in the study average EF. As seen in Table 2 the average mega-plume sample was enriched by ~ 6 and 13 standard deviations for formic and acetic acid, respectively. This corresponds to normalized excess mixing ratios relative to CO of about 5% and 1.6% for these species respectively, which can be compared to the study-average initial values of about 1.6 and 0.3%, respectively. $\Delta\text{O}_3/\Delta\text{CO}$ in the mega-plume was most often positive and ranged from -1 to $+5\%$. Potentially similar post-emission smoke chemistry can be probed by comparing the PTR-MS data for initial emissions and the biomass-burning-induced regional haze that we sampled from 27 August to 5 September (Karl et al., 2007a). The haze was less-concentrated and of even more ambiguous age. Nevertheless, it probably had some chemical processes in common with the mega-plume. In the earlier regional haze, Karl et al. (2007a) observed a $\sim 50\%$ increase in the acetone/acetonitrile ratio, but no evidence of secondary formation of methanol or acetaldehyde or other species.

Secondary production of ozone and organic acids in an isolated smoke plume from a forest fire was previously observed in Alaska by Goode et al. (2000). Yokelson et al. (2003a) observed $\Delta\text{O}_3/\Delta\text{CO}$ and $\Delta\text{CH}_3\text{COOH}/\Delta\text{CO}$ rise to $\sim 9\%$ in less than one hour in plumes from African grass fires, but no formic acid was produced. Jost et al. (2003) observed secondary production of acetone in a savanna fire plume. Trentmann et al. (2005) successfully modeled many aspects of the chemical evolution of individual smoke plumes in Africa,

but could not account for the observed secondary production of acetone or acetic acid. The ozone production could only be modeled by increasing the initial VOC (to account for unmeasured VOC) or by invoking a few possible, but unconfirmed, heterogeneous effects. Our observation of fast formation of acetic acid and O_3 in the mega-plume and increased acetone in the regional haze suggests that important, but unknown, chemistry occurs in isolated smoke plumes, mega-plumes, and regional haze.

It is difficult to identify what was burning to create the mega-plume, but some insight is gained from the attempt. The largest group of NOAA-12 hotspots (several thousand in number) for a late-afternoon, 8 September overpass was located near 50 W or ~ 22 h to the east according to HYSPLIT back trajectories (Draxler and Rolph, 2003). The sampled air would have passed this location around 02:00 p.m. LT on 7 September. The afternoon overpass for 7 September shows reduced (compared to the 8th) but very substantial hotspot activity in that same area. Another, closer, large group of NOAA-12 hotspots detected on 8 September was located along 55W only ~ 4 h to the east. This is consistent with the observation of slight aging, but would suggest that the fires were active by $\sim 08:00$ – $09:00$ a.m. The common assumption is, however, that most fires are ignited in the afternoon. Other possibilities are that the relevant hotspots were undetected or that the HYSPLIT back trajectories are highly uncertain in this remote region. To check the former hypothesis we examined the MODIS visible archives available (but hotspot numbers not tabulated) at: http://rapidfire.sci.gsfc.nasa.gov/subsets/?AERONET_Alta_Floresta/2004252. Perhaps surprisingly, even the $\sim 10:30$ a.m. LT Aqua overpass shows hundreds of hotspots along the 55W line and >100 closer hotspots (not seen on NOAA-12 images) that overlap our flight track (approximately along 56 W). This suggests several important things: (1) numerous fires were ignited in the morning on 8 September, (2) much of the smoke we sampled was likely ~ 0 – 4 h old, and (3) the NOAA-12 hotspots sometimes miss significant areas of active burning (at least under extreme conditions). In any case, the mega-plume samples likely probed smoke that was mostly less than 1 day old.

Christian et al. (2007b) observed a study-average MCE for RSC of ~ 0.788 in the ground-based campaign. The mega-plume had a higher MCE of 0.923 suggesting that the bulk of the emissions we sampled were not produced by RSC. The mega-plume MCE is also above our study average MCE of 0.91. This suggests that the smoke we happened to sample from this major burning episode originated from relatively more flaming combustion than in the fires we sampled earlier (except for the planned fire). The humidity during the 8 September flight was also by far the lowest we encountered. By 11:00 a.m. LT the water vapor had dropped below 1% as compared to 1.5–2.5% for afternoon lows on earlier flights. The low humidity and the higher MCE could explain the peak in the hotspots by indicating a strong preference for

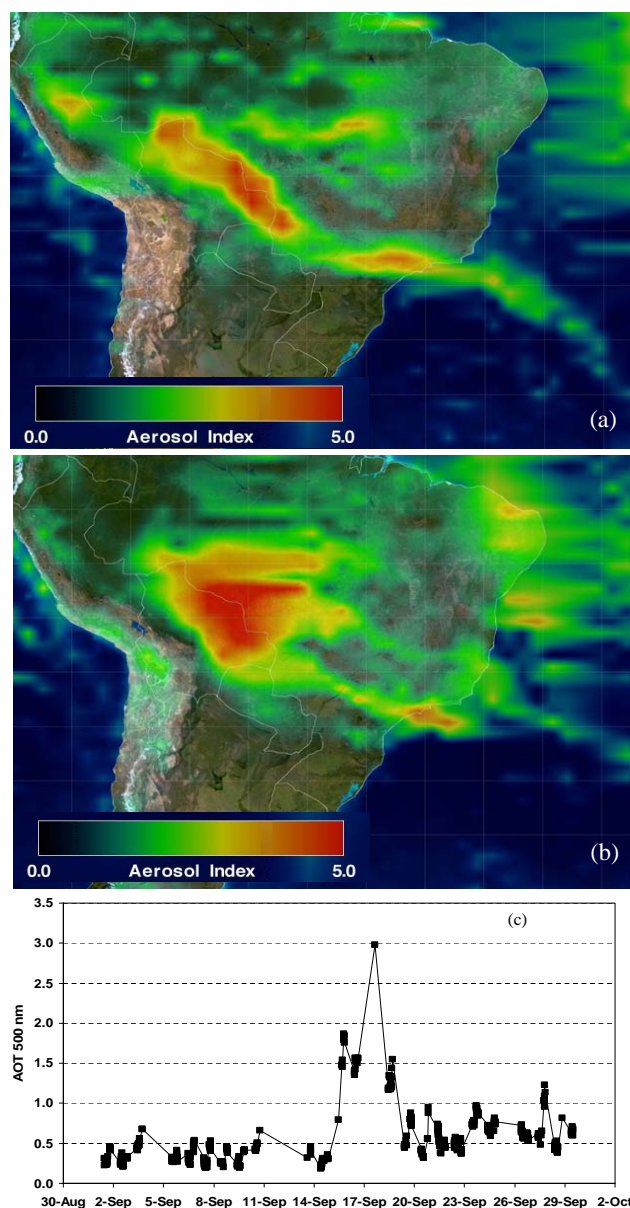


Fig. 8. (a) TOMS aerosol index (AI) on 16 September, 2004 showing concentrated biomass burning emissions (AI ~ 3.5) approaching São Paulo. (b) TOMS AI on 17 September, 2004 showing the most concentrated emissions (AI ~ 3.5) just past São Paulo. (c) São Paulo AERONET aerosol optical thickness for September 2004. The peak (near AOT 3) is in good absolute and temporal agreement with the TOMS AI data.

Brazilian farmers to burn under conditions that promote both flaming and high fuel consumption. In a non-technical summary; landholders wait until conditions are ideal and then do much of their annual burning in the next few days (including burning in the mornings) creating a mega-plume or “white ocean” of smoke that passes over southern Brazil and into the south Atlantic.

There is evidence that the mega-plume had a brief, but major impact on the air quality of São Paulo, which is ~1500 km south of the main burning region. During most of September, the mega-plume exited South America well south of São Paulo (e.g. Fig. 7c). However, the TOMS aerosol index (AI) images for 16 and 17 September show the mega-plume (with AI ~3.5) exiting South America over São Paulo (Figs. 8a and b). Simultaneously, the São Paulo AERONET station recorded a factor of 3–4 increase in aerosol optical thickness above the normal, high value (~1) for this large urban area (Fig. 8c). This incident dramatically illustrates the amount of material transported from Amazonian fires and how long it remains in a concentrated (altered) processing regime.

4 Conclusions

The TROFFEE 2004 airborne campaign in the Brazilian Amazon successfully obtained the first comprehensive emissions data for tropical deforestation fires. It was found that reactive OVOC accounted for about 80% of the NMOC emissions. We recommend emission factors for most of the major species produced by deforestation fires globally. Dry season haze layers in Brazil, due largely to biomass burning, extend to only about one-half the altitudes observed for dry-season smoke layers in Africa. This may have implications for global remote sensing of CO. Very large geographic areas were covered by the type of reactive smoke mixture that has proved hard to handle in photochemical models. Thus, the detailed effects this smoke had when it impacted areas >1000 km to the south would be difficult to predict. Larger fires may emit more particles per unit mass of fuel burned and larger fires may be becoming more common in the Amazon basin (Sect. 2.3.2). Fire emission factors for acetonitrile differ substantially by ecosystem, but could be used with attention to fire type for source apportionment and with inverse modeling to estimate the amount of biomass burned.

The TROFFEE airborne campaign also completed an initial survey of the major fire theatres of the world using new technology and a consistent sampling strategy. The initial emissions from temperate and boreal forest fires and the chemical evolution of two plumes were sampled in 1997 and recommended EF were developed (Yokelson et al., 1999; Goode et al., 2000). In 2000, the chemical evolution of 4 dry plumes and one cloud-processed plume was measured (Yokelson et al., 2003a; Hobbs et al., 2003; Jost et al., 2003) as well as the initial emissions from the two largest types of global biomass burning; savanna fires (Yokelson et al., 2003a) and cooking fires (Bertschi et al., 2003a). In 2001 and 2003, PTR-MS was co-deployed with FTIR and whole air sampling on laboratory fires in savanna, Indonesian, boreal, tropical forest, and temperate fuels and recommended EF for an expanded suite of compounds were produced (Christian et al., 2003; 2004; 2007a¹; Karl et al., 2007a).

We now know the top ~20–50 emissions from each main fire type and have quantified at best ~70% of the NMOC. Of those NMOC, about 70–80% are reactive OVOC and the NMHC are quickly converted to a series of short-lived OVOC intermediates. Rapid, very large changes in smoke composition, that are subgrid for global models, normally occur immediately after emission and there is no evidence for a fixed smoke age at which the chemical composition or rate of change is reproducible. For instance some plumes are characterized by high OH (Goode et al., 2000; Hobbs et al., 2003) and others by low OH (de Gouw et al., 2006). Detailed box models of the initial fast changes have relied on logical assumptions about unmeasured emissions; or proposed possible, but unconfirmed, heterogeneous processes to achieve partial agreement with observations (Tabazadeh et al., 2004; Trentmann et al., 2005).

More biomass burning research is needed including: (1) Airborne plume evolution studies, especially in smoky clouds, with enough instrumentation to constrain models and probe heterogeneous effects, (2) Development and/or deployment of instrumentation to quantify the unknown 30% of NMOC and species like HONO, which evidently strongly impact plume chemistry, (3) Development of well-validated high resolution, smoke chemistry models that can be confidently applied to different regional fire density and smoke transport scenarios and guide the parameterizations needed for global models, (4) Cooking fires are the second largest type of biomass burning, but a fairly large suite of emissions has only been measured on 4 of them: more cooking fires need to be sampled and more species need to be quantified such as acetonitrile and particles, (5) Better understanding of the environmental driving factors for RSC globally, (6) Better knowledge of the fuels that burn in southeast Asia, (7) Validation of space-based fire-related products such as hotspots, burned area, CO, aerosol loading, etc., and (8) Stronger integration of biomass burning measurements into campaigns focused on other issues.

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