



Emissions of trace gases and aerosols during the open combustion of biomass in the laboratory

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[1] We characterized the gas- and speciated aerosol-phase emissions from the open combustion of 33 different plant species during a series of 255 controlled laboratory burns during the Fire Laboratory at Missoula Experiments (FLAME). The plant species we tested were chosen to improve the existing database for U.S. domestic fuels: laboratory-based emission factors have not previously been reported for many commonly burned species that are frequently consumed by fires near populated regions and protected scenic areas. The plants we tested included the chaparral species chamise, manzanita, and ceanothus, and species common to the southeastern United States (common reed, hickory, kudzu, needlegrass rush, rhododendron, cord grass, sawgrass, titi, and wax myrtle). Fire-integrated emission factors for gas-phase CO₂, CO, CH₄, C_{2–4} hydrocarbons, NH₃, SO₂, NO, NO₂, HNO₃, and particle-phase organic carbon (OC), elemental carbon (EC), SO₄²⁻, NO₃⁻, Cl⁻, Na⁺, K⁺, and NH₄⁺ generally varied with both fuel type and with the fire-integrated modified combustion efficiency (MCE), a measure of the relative importance of flaming- and smoldering-phase combustion to the total emissions during the burn. Chaparral fuels tended to emit less particulate OC per unit mass of dry fuel than did other fuel types, whereas southeastern species had some of the largest observed emission factors for total fine particulate matter. Our measurements spanned a larger range of MCE than prior studies, and thus help to improve estimates of the variation of emissions with combustion conditions for individual fuels.

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1. Introduction

[2] Biomass burning emissions are a significant, global source of trace gas and aerosol species in the atmosphere and affect climate, visibility and human health [Crutzen and Andreae, 1990; Naeher et al., 2007; Watson, 2002]. Although biomass burning emissions in the continental United States have been estimated to represent only ~5% of annual average global emissions (computed for 1997–2004) [van der Werf et al., 2006], they play a large role in U.S. urban and regional air

quality, including visibility [McMeeking et al., 2006; Park et al., 2006, 2007; Phuleria et al., 2005; Robinson et al., 2006]. For example, Park et al. [2007] estimated that biomass burning contributed about 50% of the annual mean total particulate carbon (TC) concentrations across the continental United States, with summer wildfires identified as the most important driver of interannual variability in observed TC concentrations [Spracklen et al., 2007]. Further, it is expected that the frequency and magnitudes of wildfires will increase in coming decades in regions affecting the United States [Spracklen et al., 2007], which, along with increased demand for prescribed burning to reduce fuel loads in vulnerable regions [e.g., Haines et al., 2001], will result in increasing impacts from biomass burning.

[3] Model estimates of fire emissions and their impacts require not only burned area and fuel loading inventories, but also fuel-based emission factors (EF) for both gaseous and particulate phase emissions. Emission factors relate the mass of a chemical species emitted to the mass of fuel burned [e.g., Park et al., 2007; Schultz et al., 2008; Wiedinmyer et al., 2006]. EF have been measured in the laboratory and in the field for at least the last 40 years, but they remain a significant source of uncertainty in regional and global estimates of fire

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emissions [Schultz *et al.*, 2008; Wiedinmyer *et al.*, 2006]. Most EF measurements have concentrated on fuels from regions outside of the continental U.S., since these account for the largest fractions of global emissions and thus have the most significant impacts on global tropospheric chemistry. Andreae and Merlet [2001] conducted an extensive literature review and compiled recommended EF for three primary ecosystem types: savanna and grassland, tropical forests, and extratropical forests. These EF have been applied in many modeling studies [e.g., van der Werf *et al.*, 2006]. Although Andreae and Merlet [2001] included North American fuels in their survey, the recommended average values did not necessarily reflect the specific fuel types and combustion conditions most important at U.S. local and regional scales.

[4] Battye and Battye [2002] summarized much of the work reported in the peer-reviewed and gray literature that applied to emissions from U.S. wildland fires, with a focus on field studies, primarily airborne, of emissions from fires in forested regions in the northwestern United States and Alaska, as well as chaparral fires and fires in the southeastern United States [Cofer *et al.*, 1988a, 1988b; Friedli *et al.*, 2001; Hardy *et al.*, 1996; Hays *et al.*, 2002; Muhle *et al.*, 2007; Yokelson *et al.*, 1999]. While field studies have the advantage of measuring emissions from an actual fire, as pointed out by Yokelson *et al.* [2008], they offer only a snapshot in time, space, and combustion phase, and the number of measured species is limited. Controlled laboratory studies can be used to fill in some of these gaps. Some recent laboratory studies of U.S.-relevant fuels have been conducted [Chakrabarty *et al.*, 2006; Chen *et al.*, 2006, 2007; Hays *et al.*, 2002], but we are unaware of published laboratory measurements of emissions from individual chaparral or southeastern U.S. plant species. Earlier laboratory wind-tunnel studies examining several Californian fuels were primarily focused on agricultural waste [Jenkins *et al.*, 1991, 1993, 1996; Turn *et al.*, 1997]. There have been a number of studies focusing on characterization of source profiles, used for source apportionment estimates, for fuels commonly consumed by residential fireplace or wood stove burning, because of their role in urban and suburban air quality degradation [Fine *et al.*, 2001, 2002a, 2002b, 2004; Lipsky and Robinson, 2006; Roden *et al.*, 2006]. Finally, very few studies have presented a comprehensive set of measurements that include both gas-phase and speciated particulate-phase emissions, along with an indicator of combustion conditions.

[5] The Fire Laboratory at Missoula Experiment (FLAME) aimed to fill some of the gaps in available data on emissions from fires in the United States. The study took place at the U.S. Forest Service's Fire Sciences Laboratory (FSL) in 2006 (FLAME 1) and 2007 (FLAME 2). Earlier experiments performed at the FSL examined fire combustion behavior [Freeborn *et al.*, 2008], trace gas emissions [Christian *et al.*, 2004; Goode *et al.*, 1999; Yokelson *et al.*, 1996, 1997] and aerosol emissions [Chakrabarty *et al.*, 2006; Chen *et al.*, 2006, 2007; Engling *et al.*, 2006; Freeborn *et al.*, 2008]. FLAME expanded on this work by including additional fuels and fuel components most relevant to wildland fire and prescribed burning in the United States, and adding/improving measurements of aerosol properties, including emissions of smoke marker species [Sullivan *et al.*, 2008], mercury compound emissions [Obrist *et al.*, 2008], particle size distributions and refractive index (E. J. T. Levin *et al.*,

Biomass burning smoke aerosol properties measured during FLAME 2, manuscript in preparation, 2009), aerosol hygroscopicity, cloud condensation nuclei (CCN) and ice nuclei (IN) activity [DeMott *et al.*, 2009; Petters *et al.*, 2009a, 2009b], and aerosol optical properties [Lewis *et al.*, 2008; L. Mack *et al.*, Measured and retrieved optical properties of biomass smoke from FLAME 2, manuscript in preparation, 2009].

2. Fuel Selection and Treatment

[6] Leaves and woody material from 33 unique plant species (Table 1) were burned individually and in various combinations during FLAME 1 and 2. Fuels that were too moist to burn were dried at 35°–40°C for 48–72 h. The remaining untreated fuels had dried sufficiently during shipping and storage to be used without drying. Fuel moisture (FM, dry weight percent; see auxiliary material Data Set S1) for each fuel as used was determined by holding a sample at 100°C for 48 h and measuring the mass loss.¹ Fuel carbon and nitrogen contents (Table 1) were measured by an independent laboratory. We selected fuels based on their modeled frequency of consumption in wildfires and prescribed fires in the western and southeastern United States and in fire-impacted regions in close proximity to urban areas. We further prioritized selection of species for which little or no peer-reviewed, controlled laboratory emissions data were available.

2.1. Chaparral

[7] Chaparral is a highly diverse ecosystem that is distributed from Baja California to south-central Oregon and covers approximately 6% of the area of California [Keeley and Davis, 2007]. Chaparral-dominated regions coincide with many highly populated areas in California, most notably the Los Angeles and San Diego metropolitan regions, underscoring the need for accurate emission inventories for chaparral fuels. For example, Clinton *et al.* [2006] estimated that ~80% of the fuel consumed by a series of major wildfires in southern California during 2003 were shrubs and duffs. The dominant species within the chaparral ecosystem include chamise (*Adenostoma fasciculatum*) and species in the *Ceanothus* and *Arctostaphylos* genera [Keeley and Davis, 2007]. We tested three fuels representing this ecosystem: chamise, hoaryleaf ceanothus (*Ceanothus crassifolius*), and Eastwood's manzanita (*Arctostaphylos glandulosa*). Samples were collected from the San Jacinto Mountains, about 150 km east of Los Angeles, California (see Table 1). Chaparral fire emissions have been observed from aircraft [Cofer *et al.*, 1988a, 1988b; Hegg *et al.*, 1987], but we are unaware of laboratory measurements that have appeared in the peer-reviewed literature.

2.2. Montane and Subalpine Forests

[8] Montane and subalpine coniferous forests cover major portions of the Sierra Nevada and Cascade mountain ranges [Fites-Kaufman *et al.*, 2007], inland regions of the northwestern United States [Franklin, 1988], and northern Rocky Mountains [Peet, 1988]. These regions encompass many federal Class I areas that are protected against visibility degradation. Species from this ecosystem tested during FLAME

¹Auxiliary materials are available at <ftp://ftp.agu.org/apend/2009jd011836>.

Table 1. Plant Species That Served as Fuels During FLAME

Common Name	Scientific Name	Sampling Location(s)	Carbon Content (%)	Nitrogen Content (%)
Alaskan duff		Tok, Alaska	31	0.5
Black spruce	<i>Picea mariana</i>	Fairbanks, Alaska	55	0.6
Chamise	<i>Adenstoma fasciculatum</i>	San Jacinto Mountain, Calif.	49	1.0
Common reed	<i>Phragmites australis</i>	Cameron Prairie NWR, La.	49	0.5
Douglas fir	<i>Pseudotsuga menziesii</i>	Missoula, Mont.	54	0.5–0.9
Gallberry	<i>Ilex coriacea Ilex glabra</i>	Sandhill Crane NWR, Miss. St. Marks NWR, Fla. Osceola National Forest, Fla.	56	0.8
Grass	<i>various species</i>	Missoula, Mont.	42	3.0
Gray's rabbitbrush	<i>Ericameria nauseosa</i>	Utah	46	1.1
Hickory	<i>Carya nutt</i>	Hillsborough, N. C.	48	2.1
Hoaryleaf ceanothus	<i>Ceanothus crassifolius</i>	San Jacinto, Calif.	48	1.3
Kudzu	<i>Pueraria Montana</i>	Athens, Ga.	47	3.6
Lodgepole pine	<i>Pinus contorta</i>	Missoula, Mont.	42–50	0.3–1.2
Longleaf pine	<i>Pinus palustris</i>	North Carolina, Sandhill Crane NWR, Miss. St. Marks NWR, Fla. Camp Lejeune, N. C.	52	1.1
Manzanita	<i>Arctostaphylos glandulosa</i>	San Jacinto, Calif.	48	0.8
Needlegrass rush	<i>Juncus roemerianus</i>	St. Marks NWR, Fla.	49	1.1
Palmetto	<i>Serenoa repens</i>	St. Marks NWR, Fla. Osceola NF, Fla. Sandhill Crane NWR, Miss.	51	1.0
Peltophorum	<i>Peltophorum inerme</i>	Puerto Rico	48	0.8
Ponderosa pine	<i>Pinus ponderosa</i>	Missoula, Mont.	46–49	0.04–1.3
Puerto Rican fern	<i>Dicranopteris pectinata</i>	Puerto Rico	46	0.4
Rhododendron	<i>Rhododendron minus</i>		51	0.6
Rice straw	<i>Oryza sativa</i>	Douliou City, Taiwan	39–46	0.6–0.9
Sagebrush	<i>Artemisia tridentate</i>	Salt Lake City, Utah	47–51	1.5–2.1
	<i>Hibiscus tiliaceus</i>	Puerto Rico		
Smooth cord grass	<i>Spartina alterniflora</i>	St. Marks NWR, Fla.		
Sugarcane	<i>Saccharum officinarum</i>	Guangdon Province, China	48	1.3
Swamp sawgrass	<i>Cladium mariscus</i>	Big Branch Marsh NWR, La.	48	2.1
Teak	<i>Tectonia grandis</i>	Puerto Rico	44	0.8
Titi	<i>Cyrilla racemiflora</i>	St. Marks NWR, Fla.	54	0.9
Turkey oak	<i>Quercus laevis Walt.</i>	Hillsborough, N. C. Camp Lejeune, N. C.	53	1.3
Utah juniper	<i>Juniperus osteosperma</i>	Utah	49	0.9
Wax myrtle	<i>Myrica cerifera</i>	Sandhill Crane NWR, Fla. St. Marks NWR, Fla.	48–53	1.1–1.4
White spruce	<i>Picea glauca</i>	Fairbanks, Alaska	52	0.8
Wiregrass	<i>Aristida beyrichiana</i>	Sandhill Crane NWR, Miss. St. Marks NWR, Fla. Camp Lejeune, N. C.	48	0.5

included: ponderosa pine (*Pinus ponderosa*), lodgepole pine (*Pinus cortata*), and Douglas fir (*Pseudotsuga menziesii*). We burned needles, woody material, combinations of needles and woody material, as well as litter (dead needles and cones from the forest floor) and duff (partly decayed litter including a portion of the uppermost layers of soil). These species were collected from several rural locations near Missoula, Montana. We also burned a mixture of unidentified grass species collected from a site near the FSL.

2.3. Rangeland

[9] Sagebrush rangeland ecosystems are one of the most widespread in the intermountain west, primarily found in eastern Oregon, southern Idaho, Nevada and Utah [West and Young, 2000]. In addition to big sagebrush (*Artemisia tridentate*), we also burned two other woody species found from this region: Gray's rabbitbrush (*Chrysothamnus nauseosus*) and Utah juniper (*Juniperus osteosperma*). The rabbitbrush and juniper samples were collected near Salt Lake City, Utah. Sagebrush samples were collected from two other areas: an urban setting near the Salt Lake City airport and a rural setting near Missoula, Montana.

2.4. Southeastern Coastal Plain

[10] Forest, rangeland and cropland undergo prescribed burning each year in the southeastern United States [Haines et al., 2001], but wildfires also occur in this region. We burned several species common to the coastal plain region of the southeastern United States, including longleaf pine (*Pinus palustris*), and understory shrubs such as saw palmetto (*Serenoa repens*), gallberry (*Ilex glabra*), and wax myrtle (*Myrica cerifera*). During periods of prolonged drought, fire can spread to dry savannah and wetland ecosystems, so we selected several representative species including titi (*Cyrilla racemiflora*), sawgrass (*Cladium mariscus*), common reed (*Phragmites australis*), wiregrass (*Aristida beyrichiana*) and black needlerush (*Juncus roemerianus*). We also burned kudzu (*Pueraria lobata*), an invasive species that is frequently the target of control efforts, which include prescribed burning.

2.5. Boreal Forests

[11] Boreal forest fires are a major source of carbon to the atmosphere [Kasischke et al., 1995], and their emissions

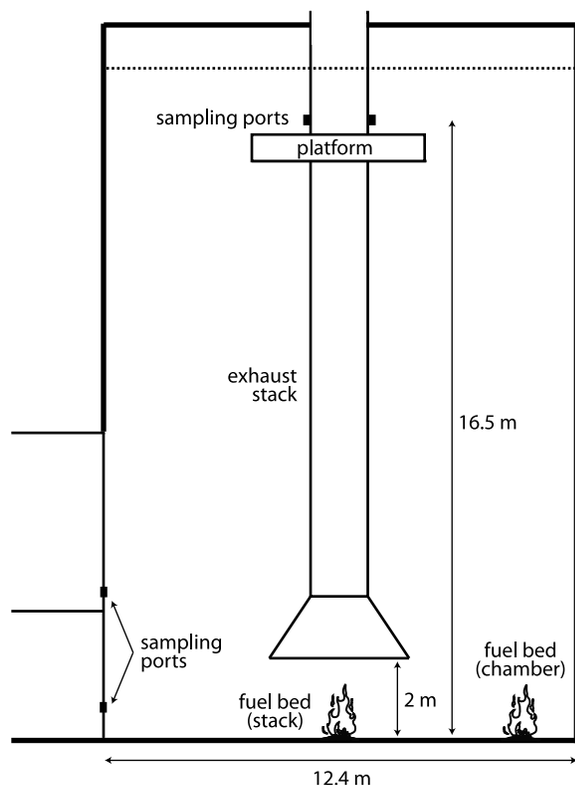


Figure 1. Schematic of the U.S. Forest Service Fire Sciences Laboratory combustion facility, located in Missoula, Montana. Image is to scale. The locations of the fuel bed and of the sampling ports during stack and chamber burns are indicated.

have major impacts on the atmosphere on local and global scales [e.g., French *et al.*, 2002; O'Neill *et al.*, 2002; Pfister *et al.*, 2008; Stohl *et al.*, 2006; Trentmann *et al.*, 2006]. Emissions from boreal North America alone accounted for $\sim 10\%$ of annual average global emissions from 1997 to 2004 [van der Werf *et al.*, 2006] and have been observed to be transported into the U.S. [e.g., Al-Saadi *et al.*, 2005]. White spruce (*Picea glauca*) and black spruce (*Picea mariana*) are ubiquitous conifer species in boreal forests and are commonly found in spruce-fermoss forests that dominate the southern boreal forest zone, which includes a large portion of Alaska [Elliot-Fisk, 1988]. We burned spruce samples collected within ~ 50 km of Fairbanks, Alaska. Wildfires and prescribed burns affect belowground biomass in addition to shrubs and trees, so we also burned samples taken from forest floor (duff), which consisted of the uppermost layer of soil with live and dead feathermoss (*Pleurozium schreberi*). However, we note that we did not have any samples of the underlying peat below the surface duff, which can contribute substantially to total fire emissions [Kasischke *et al.*, 2005; Yokelson *et al.*, 1997].

2.6. Other Fuels

[12] We included a mixture of plants that are frequently burned in Puerto Rico, as biomass burning emissions from this region, as well as from Mexico and Central America, can be transported to the southeastern United States [Kreidenweis *et al.*, 2001]: teak (*Tectona grandis*), sea hibiscus (*Hibiscus*

tiliaceus), peltophorum (*Peltophorum inerme*), sacky sac bean (*Inga laurina*), and fern (*Decranopteris pectinata*). Two agricultural waste products that are burned after harvest were collected in Asia: rice straw (*oryza sativa*) from Taiwan and sugarcane (*saccharum officinarum*) from the Guangdong province of China. Although outside the scope of our general focus on U.S. inventories, emissions from these agricultural wastes have attracted recent interest [Christian *et al.*, 2003; Yokelson *et al.*, 2008] and have been shown to affect air quality in populated regions [Viana *et al.*, 2008; Yang *et al.*, 2006].

3. Experimental Method

3.1. Facility and Burn Procedure

[13] The experiments were performed at the U.S. Forest Service's combustion testing facility at the Fire Sciences Laboratory in Missoula, Montana, which is depicted in Figure 1 and has been described previously [Bertschi *et al.*, 2003; Christian *et al.*, 2003; Yokelson *et al.*, 1996, 2008]. The main combustion chamber is a square room with internal dimensions $12.4 \times 12.4 \times 19.6$ m high and a total volume of ~ 3000 m³. Outside air was conditioned for temperature and humidity and pumped into the chamber prior to each burn. An exhaust stack located at the center of the room begins 2.1 m above the floor and extends through the chamber ceiling. An inverted funnel at the bottom of the exhaust stack narrows from a 3.6 m diameter opening to the 1.6 m stack diameter. Sampling ports that originate near the center of the flow and pass through the walls of the exhaust stack are located ~ 16.5 m above the floor, and are accessed from a measurement platform near the ceiling.

[14] Two types of experiments were performed during FLAME, which we refer to as "stack" and "chamber" burns (auxiliary material Data Set S1). During stack burns, emissions from the fuel bed, located directly beneath the inverted funnel, were drawn through the exhaust stack. Instruments located on the measurement platform continuously sampled through the platform sample ports. Christian *et al.* [2004] used direct observations of gas profiles to confirm that emissions are well-mixed across the stack. In contrast, the combustion room was sealed during chamber burns by closing the exhaust stack. The fuel bed was placed about halfway between the exhaust stack and the chamber wall and a large circulation fan operated in one corner to facilitate mixing. Continuous-sampling instruments were relocated from the measurement platform to laboratories adjacent to the combustion chamber, and drew samples through wall ports. Chamber burns were designed primarily for optical closure experiments not reported here, as those measurements required lower species concentrations and longer sampling periods (~ 2 h) compared to those possible during stack burns, which typically lasted from 5 to 10 min.

[15] The majority of samples burned during stack experiments were placed on a 46×61 cm horizontal metal tray covered with an inert ceramic heat shield. Fuels were stacked horizontally on the fuel bed to facilitate ignition, except for two large fuel mass burns (~ 2500 g) when fuels were stacked in a cylindrical wire cage. The fuel bed was placed on a Mettler-Toledo PM34 balance to monitor its mass as a function of burn time. The initial fuel mass (m_0) and final residual mass ($m_{residual}$), both listed in auxiliary

material Data Set S1 for each burn, were measured with a higher sensitivity Mettler-Toledo PM34-K balance. Initial fuel masses ranged from 25 to 2500 g depending on the objective of the experiment and desired emission concentrations; most were between 100 and 250 g.

[16] We ignited the fuel bed using several methods. During FLAME 1, dry fuels were ignited using a butane pilot lighter applied briefly to the edge of the fuel bed. Fuels with high fuel moistures required the application of a propane torch or heated metal coils for a significant period of time, in some cases continuously, to maintain the fire. Both ignition methods often resulted in a propagating flame front that moved through the fuel bed and simultaneous flaming and smoldering combustion in different parts of the fuel bed. We modified the fuel bed in the FLAME 2 experiments [Sullivan *et al.*, 2008]. Fuels were placed on a lattice of heating tape that was soaked with ~ 15 g of ethanol, which was vaporized and ignited on heating, uniformly igniting the fuel bed. The dense duff core samples still required application of the propane torch to sustain combustion, but all other fuels were ignited effectively using this method. Auxiliary material Data Set S1 provides the components of the plant or plants that were burned during each burn, the ignition method, and the fuel moisture content. We performed three replicate burns for each fuel type during FLAME 1 stack burns and two replicate stack burns during FLAME 2.

3.2. Real-Time Gas Measurements

[17] Real-time measurements of CO_2 , CO, NO, and NO_2 were made at ~ 2 s resolution using three commercial gas analyzers, sampling through aluminum (C gas analyzers) or Teflon lines (NO_x analyzer). Carbon dioxide and water vapor mixing ratios were measured by a Li-Cor Model 6262 nondispersive infrared gas analyzer. Carbon monoxide mixing ratios were measured using a Thermo Environmental Model 48C variable-range gas filter correlation analyzer. Two sets of mixed standards ($[\text{CO}_2] = 362$ ppm, $[\text{CO}] = 0.5$ ppm and $[\text{CO}_2] = 499$ ppm, $[\text{CO}] = 2.7$ ppm) were passed through the analyzers prior to burn ignition for calibration. The mixing ratios of nitrogen oxides ($\text{NO}_x = \text{NO} + \text{NO}_2$) were measured by a Thermo Environmental Model 42 chemiluminescence analyzer. We observed high (>2000 ppb) NO_x concentrations that saturated the analyzer during several FLAME 2 burns and do not report those NO_x data. In some of those cases the NO measurement was valid and is reported. The estimated accuracy/precision of the measurements were: Li-Cor, 1%/0.1%; Thermo Environmental, 2%/1%.

3.3. Trace Gas Canister Measurements

[18] Canister samples of emissions drawn directly from the stack and chamber were analyzed for CO_2 , CO, CH_4 (methane), C_2H_4 (ethene), C_2H_6 (ethane), C_3H_6 (propene), C_3H_8 (propane), three isomers of C_4H_8 (butene), and C_4H_{10} (n-butane) gases with a Hewlett Packard model 5890 Series II gas chromatograph. Background samples were collected in several canisters throughout the day during the experiments and used to calculate the excess mixing ratios of each measured species (e.g., $\Delta\text{CH}_4 = \text{CH}_{4, \text{measured}} - \text{CH}_{4, \text{background}}$). The CO_2 and CO analyses used a 1 mL sample loop to inject the sample, and a 1/8" diameter \times 6 foot Carbosphere (Alltech) column to separate CO_2 , CO, and air with a helium carrier gas at a

flow rate of 16 mL min^{-1} . After separation in the column the sample entered a nickel catalyst methanizer (375°C), which converted the CO_2 and CO to CH_4 , followed by a flame ionization detector (FID) at 350°C . The oven temperature program was isothermal at 100°C . The $\text{C}_1\text{-C}_4$ analyses were performed using a 0.25 mL sample loop, with a $0.53 \text{ mm} \times 30 \text{ m}$ GS-Q (J&W Scientific) column with a helium carrier gas at 6 mL min^{-1} . The oven temperature program for this analysis was 30°C for 6 min, then increasing by $10^\circ\text{C min}^{-1}$ to a final temperature of 90°C for 8 min.

[19] Chromatogram data were processed by Hewlett Packard ChemStation II software. A set of gas standards bracketing the sample concentrations was analyzed with each set of samples to construct a standard curve for each compound. Based on the integrated peak areas, the sample concentrations were calculated from the standard curves. Duplicate analyses were performed every sixth sample to quantify measurement precision error. National Institute of Standards and Technology (NIST) primary standards of CO_2 , CO, and CH_4 were analyzed as samples to measure overall accuracy. Accuracies/uncertainties in the GC analyses were 1%/1% for CO_2 , CO, and CH_4 , and 10%/10% for C_{2-4} gases.

3.4. Trace Gas Denuder Measurements

[20] We measured ammonia (NH_3), nitric acid (HNO_3), and sulfur dioxide (SO_2) concentrations emitted from fires using annular denuders (URG Corporation, Chapel Hill, NC). The denuders operated in series with a filter sampling system (see section 3.5). The sample flow was nominally 10 L min^{-1} and was pulled through a Teflon-coated inlet; Brauer *et al.* [1989] cite efficiencies of 97.3–98.5% for sampling of NH_3 through similar inlets. The HNO_3 denuder was coated with 10 mL of a 1% sodium carbonate + 1% glycerol in a 1:1 methanol/water solution and the NH_3 denuder was coated with 10 mL of a 1% phosphorous acid in a 9:1 methanol/water solution [Perrino *et al.*, 1990; Perrino and Gherardi, 1999]. Coated denuders were dried with N_2 for ~ 20 min. After sampling, each denuder was extracted using 10 mL of deionized water. Extracts were analyzed using a Dionex DX-500 series ion chromatograph. Details of the analysis procedure are given by Yu *et al.* [2006] and Lee *et al.* [2008].

3.5. Particulate Filter Samplers

[21] Three types of filter samplers collected particulate matter on filters during the burns for compositional analysis: a URG annular denuder/filter sampling system (URG, Chapel Hill, North Carolina), a high-volume sampler (Hi-vol; Thermo Anderson, Smyrna, Georgia), and two IMPROVE (Interagency Monitoring of Protected Visual Environments) samplers [Malm *et al.*, 2004]. The Hi-vol and URG samplers were located on the sampling platform during stack burns. During chamber burns, they were moved to the chamber floor, with the Hi-vol samplers on tables to keep the inlets of both samplers at a uniform height (~ 3 m). The IMPROVE samplers had inlets at a similar height, and only ran during chamber burns because of space restrictions on the stack sampling platform. During stack burns, the filter sampler pumps were turned on 30 s prior to ignition and turned off when the fire was

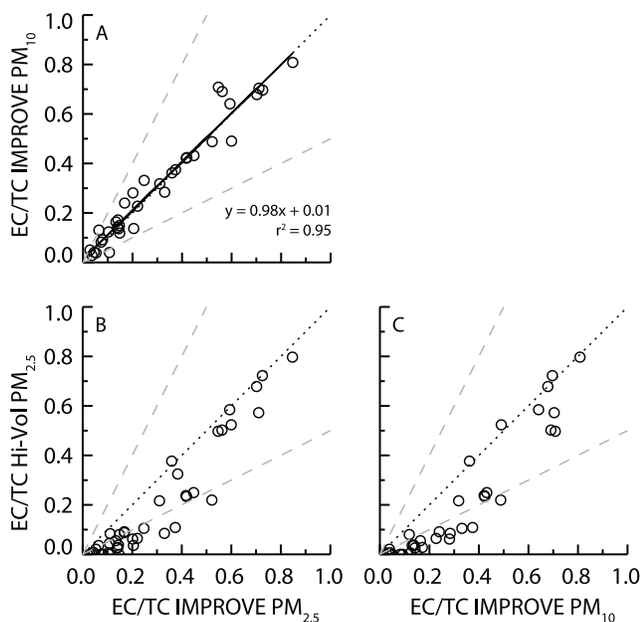


Figure 2. Scatterplots comparing elemental carbon (EC) concentrations, normalized by total aerosol carbon (TC) concentrations, for each thermal optical analysis protocol and/or filter sampler used during FLAME chamber burns. (a) IMPROVE PM₁₀ versus IMPROVE PM_{2.5}. (b) Hi-vol PM_{2.5} versus IMPROVE PM_{2.5}. (c) Hi-vol PM_{2.5} versus IMPROVE PM₁₀. The dashed black line is the 1:1 line, and the two dashed gray lines are the 1:2 and 2:1 lines.

considered extinguished based on visual observations. During chamber burns, the filter sampler pumps were started approximately 4 min after ignition, and individual aerosol samples for each burn were typically collected over 2 h.

[22] The Hi-vol sampler collected samples on quartz filters for thermal optical OC/EC analysis. *Sullivan et al.* [2008] and *Engling et al.* [2006] described the Hi-vol sampler we used during FLAME. The sampler had a nominal flow rate of 1.13 m³ min⁻¹. An assembly of two quartz-fiber filters collected particles divided into two size classes: those with aerodynamic diameters (D_{ae}) > 2.5 μm (coarse mode) and those with D_{ae} < 2.5 μm (fine mode). We only present results from the analysis of the 20.3 × 25.4 cm fine mode filter, equivalent to particulate matter (PM) with D_{ae} < 2.5 μm or PM_{2.5}, because an examination of the IMPROVE filters and volume size distributions [*Sullivan et al.*, 2008; Levin et al., manuscript in preparation, 2009] confirmed that total aerosol mass was dominated by particles in the sub-2.5 μm diameter size range, as expected [e.g., *Ward and Hardy*, 1991]. The quartz filters were wrapped in aluminum foil and baked in an oven over a 36 h period (12 h heating at 550°C + 24 h cool down) prior to sampling to remove any organic contaminants. Punches from the Hi-vol filters were analyzed for the masses of carbon in the OC and EC fractions with a semicontinuous analyzer (Sunset Laboratory, Tigard, Oregon) in “off-line” mode [*Sullivan and Weber*, 2006]. The OC/EC measurements reported here were the average of two 1.4 cm² punches from the same filter to reduce measurement uncer-

tainties associated with sample loading heterogeneity [*Gorin et al.*, 2006].

[23] The URG sampling system consisted of two annular denuders and a filter pack arranged in series, which collected gas and aerosol samples for ion chromatography (IC) analysis [*Lee et al.*, 2004]. The 10 L min⁻¹ sample flow first passed through a Teflon-coated 2.5 μm size cut cyclone to remove large particles, and then through two denuders (section 3.4) and a nylon filter (Gelman Nylasorb, 1.0 μm pore size). A backup cellulose filter coated in citric acid collected any NH₃ lost from the particles collected on the nylon filter. The URG filters were extracted using 6 mL of deionized water. Extracts were analyzed for inorganic species (Cl⁻, SO₄²⁻, NO₃⁻, Na⁺, NH₄⁺, K⁺, Mg²⁺, and Ca²⁺) using two Dionex DX-500 IC systems.

[24] Particles were also collected by two IMPROVE sampling systems during the chamber burns, slightly modified from those used in the IMPROVE network [*Malm et al.*, 2004]. Each system had only A, B and C modules, holding Teflon, Nylasorb, and quartz filters, respectively, and collected particulate matter after PM_{2.5} or PM₁₀ inlets. The C modules held two quartz filters in series to characterize organic aerosol sampling artifacts. During several FLAME 1 burns the IMPROVE modules operated for different time intervals than the other samplers; in those cases smoke species concentrations were corrected using measurements of the room air background concentrations and the total time that room air was sampled. Gravimetric mass was measured from Module A filters following the standard procedure used for samples collected in the IMPROVE network, with relative humidity in the weighing laboratory maintained between 20 and 40%.

3.6. Organic and Elemental Carbon Thermal Optical Analysis Protocols

[25] The OC and EC measurements presented here were obtained using two different protocols. Samples collected by the IMPROVE sampler were analyzed using the IMPROVE_A analysis protocol [*Chow et al.*, 2007], in which the sample was heated to four temperature plateaus (140°, 280°, 480° and 580°C) in pure helium and three temperature plateaus (580°, 740° and 840°C) in 98% helium and 2% oxygen. Analysis of the Hi-vol punches using the Sunset analyzer followed a modification of the NIOSH 5040 protocol [*Bae et al.*, 2004; *Birch and Cary*, 1996]. The sample punch was heated in pure helium to 600°C in 80 s and then to 840°C in 90 s. The sample was cooled for 35 s and oxygen added to the analysis atmosphere (98% He, 2% O₂). Punches were then heated to 550°C in 30 s, 650°C in 45 s, and 850°C in 90 s.

[26] Figure 2 compares EC/TC ratios measured for the IMPROVE PM_{2.5}, IMPROVE PM₁₀, and Hi-vol filter samples collected during FLAME. The good agreement between EC/TC ratios found for the IMPROVE PM₁₀ and PM_{2.5} samples ($r^2 = 0.95$, regression coefficient = 0.98) shows that the EC fraction of TC was similar in both. EC/TC ratios obtained by the same protocol for high EC/TC ratios were strongly correlated, but they disagreed within about a factor of 2 between protocols for samples with low EC/TC ratios, similar to the discrepancies found in biomass burning-impacted samples in previous studies [*Watson et al.*, 2005].

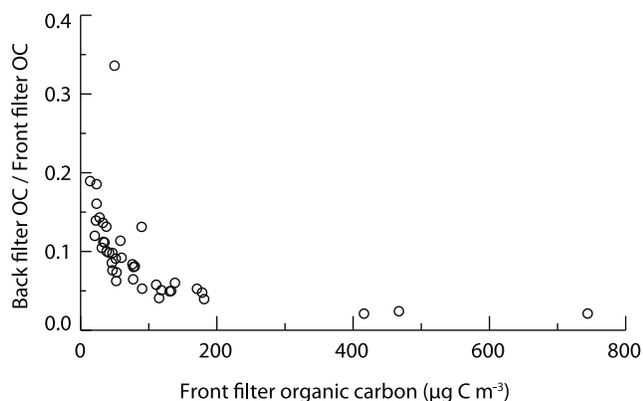


Figure 3. Organic carbon (OC) concentrations measured on the back IMPROVE quartz filter normalized by OC measured on the front IMPROVE quartz filter, as a function of front filter OC. Chamber burns only.

It is unclear which method provides a more accurate measure of the EC content of the aerosol. In the remainder of this work, we use the Hi-vol/NIOSH 5040/Sunset OC and EC measurements, simply because they are a more complete data set (available for both stack and chamber burns).

[27] Filter-based carbonaceous aerosol measurements are prone to artifacts caused by gas-phase adsorption onto filter fibers (positive artifact) and volatilization of the sampled particle phase organic material (negative artifact) [e.g., Kirchstetter *et al.*, 2001; Mader and Pankow, 2001; Turpin *et al.*, 1994]. During FLAME, the IMPROVE PM_{2.5} and IMPROVE PM₁₀ samplers collected aerosol using front and back quartz filters arranged in series. Ideally, the mass of OC (adsorbed gases) measured on the back filter equals the mass of OC measured on the front filter that was due to adsorbed gases. Overall, adsorption artifacts during FLAME appeared to be relatively small (Figure 3). At high OC concentrations (>100 µg m⁻³), when presumably more semivolatile material was in the particle phase, back filter OC was ~2–5% of the front filter OC. At lower OC concentrations (<50 µg m⁻³), when more semivolatile material should be in the gas phase, back filter OC approached 20% of the front filter OC, closer to the 20–50% values reported by Fine *et al.* [2001] and Lipsky and Robinson [2005, 2006]. In those studies, the aerosol samples were diluted to lower concentrations than we sampled during FLAME, which may have altered the partitioning of semivolatile species toward the gas phase.

3.7. Emitted and Consumed Mass Calculations

[28] For the canister and denuder measurements, the total emitted mass of each species was computed from the product of the excess mixing ratios and the sample volume. The canister and denuder samplers operated throughout each stack burn and represented fire-integrated emissions. The continuous measurements of CO and CO₂ during chamber burns showed that the concentrations of these species did not vary significantly after the chamber became well-mixed, within 30 min of ignition. The canisters were

used to capture a sample from the chamber approximately 60 min into the experiment.

[29] During the stack burns, filter and denuder samples were collected over multiple, replicate burns to ensure adequate concentrations for compositional analysis, particularly trace organic species [Sullivan *et al.*, 2008]. We usually sampled three replicate burns on a single filter during FLAME 1 and two replicate burns on a single filter during FLAME 2. In the calculation of emission factors for each aerosol species, we multiplied the mass concentrations of each species determined from the filter measurements by the total volume of air sampled through the stack. We calculated the mass of aerosol species emitted during the chamber burns by multiplying mass concentrations determined from filter measurements by the total volume of the chamber. This approach assumes that the emissions were well-mixed, and therefore the calculations of emission factors for chamber burns have higher uncertainty than those for stack burns.

[30] The mixing ratio measurements from the real-time gas analyzers were multiplied by the volume flux of air through the stack and integrated over the lifetime of the burn to obtain the total masses of CO, CO₂ and NO_x emitted during the stack burns. For chamber burns, we calculated the average gaseous-species mixing ratios for the period from 30 to 35 min following ignition, and multiplied by the chamber volume.

[31] We adjusted the total masses of CO and CO₂ emitted for burns that used the ethanol-coil ignition system by subtracting the mean of the total emissions for each species during the two ethanol-coil test burns (0.13 g CO, 12.5 g CO₂). In general, the mass of plant material burned was 5–10 times greater than the mass of ethanol consumed during the ignition procedures. Exceptions were burns featuring low fuel masses conducted during FLAME 2 chamber burns. Emission data for burns that used the propane torch ignition method were adjusted by subtracting the total torch emissions, which were determined by multiplying the time the torch was on by the species emission rate. Burns that required the torch to be applied to maintain combustion for a period greater than half of the total burn time were omitted from the analyses.

[32] The mass of dry biomass consumed (m_{consumed}), assuming the residual material contained no water, was calculated as

$$m_{\text{consumed}} = \frac{m_{\text{fuel}}}{1 + FM} - m_{\text{residual}}, \quad (1)$$

where FM is the fuel moisture fraction, m_{fuel} is the initial (wet) fuel mass and m_{residual} is the mass of ash and unburned fuel remaining. The carbon consumed (C_{consumed}) during each burn was calculated by multiplying m_{consumed} by x_c (section 3.9).

3.8. Modified Combustion Efficiency Calculation

[33] Since biomass burning emissions are known to depend on the combustion conditions, it is useful to include a measure of the combustion efficiency in reporting observations. We adopt the approach used in many prior studies [e.g., Yokelson *et al.*, 2008] and report the fire-integrated

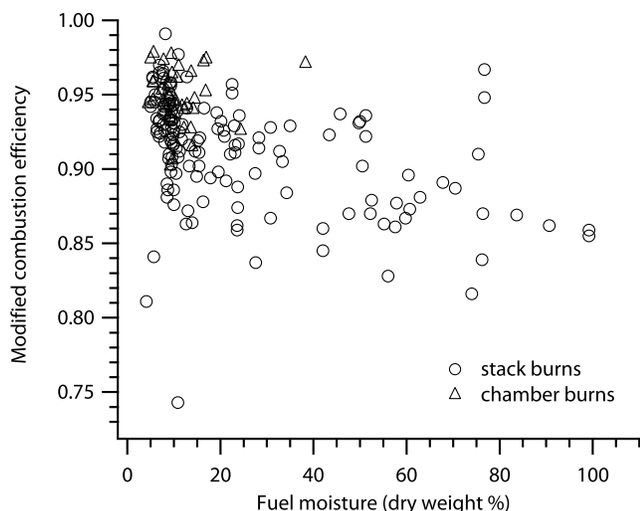


Figure 4. Fire-integrated modified combustion efficiency plotted as a function of fuel moisture (in dry weight %).

modified combustion efficiency (MCE), which depends on the molar ratio of the emitted CO and CO₂ [Ward and Radke, 1993],

$$MCE = \frac{\Delta[\text{CO}_2]}{\Delta[\text{CO}] + \Delta[\text{CO}_2]}, \quad (2)$$

where $\Delta[\text{CO}_2]$ and $\Delta[\text{CO}]$ are the fire-integrated excess molar mixing ratios of CO₂ and CO. To compute the excess quantities, we assumed the ambient concentrations of CO and CO₂ were equal to their mean values measured in the stack or chamber immediately prior to ignition (usually from 120 to 10 s before ignition). For stack burns, we determined the fire-integrated MCE for each burn by dividing the total mass of CO₂ (in g C) emitted by the net mass of CO₂ plus CO emitted, also in g C. For chamber burns, we computed the mean fire-integrated MCE during the 5-min period between 30 and 35 min following ignition, as was done for other gases (section 3.7). Auxiliary material Data Set S1 lists the fire-integrated MCE for each burn.

3.9. Emission Ratios and Emission Factors

[34] Fire-integrated emission factors were calculated using the carbon mass balance (CMB) approach [Ward and Radke, 1993], in which the concentrations of emitted carbon-containing species are a proxy for the mass of dry fuel consumed during the fire. The emission factor for species i emitted by a fuel with carbon mass fraction (x_c) of the dry fuel mass is given by

$$EF_i = \frac{m_i}{\Delta\text{CO} + \Delta\text{CO}_2 + \Delta\text{PM}_C + \Sigma(\Delta\text{HC})} x_c, \quad (3)$$

where m_i is the mass of species i emitted, PM_C is the mass of particulate-phase carbon and ΣHC is the sum of the total mass of C contained in gas-phase hydrocarbons, estimated during FLAME as the sum of the measured C_{1–4} hydrocarbons. We used the measured values of dry fuel mass x_c reported in Table 1 or assumed a value of 0.45 [Andreae and Merlet, 2001] in the absence of fuel carbon information. To report gas-phase emission factors on a burn-by-burn basis we

ignored the PM_C term in equation (3), but it was usually a small fraction of the carbon emissions [Lipsky and Robinson, 2006] and, together with the contribution from carbon-containing gases not measured, caused an overestimation of EF on the order of only 1–2% [Andreae and Merlet, 2001]. All emission factors reported here are in units of g species per kg dry fuel (denoted g kg⁻¹ fuel for simplicity), unless stated otherwise.

4. Results

4.1. Fire Behavior and Combustion Efficiency

[35] Fire-integrated MCE values ranged from approximately 0.75–0.95, but we also observed MCE values outside this range for burns in which we only sampled flaming or smoldering phase emissions (see auxiliary material Data Set S1). Our best estimate of the variability in fire-integrated MCE for a single fuel was derived from 15 replicate ponderosa pine needle litter burns with constant FM (9.9 ± 0.5%) and initial fuel mass (246 ± 6 g), for which we calculated fire-integrated MCE values ranging from 0.88 to 0.94 with a mean and standard deviation of 0.92 ± 0.02.

[36] In some cases, fuels with higher FM burned with lower MCE (Figure 4). For example, untreated ponderosa pine needles (FM ~60%) had a fire-integrated MCE of 0.86 whereas dry ponderosa pine needles (FM ~10%) had a fire-integrated MCE of 0.94. However, factors other than FM affected MCE. We observed larger MCE values when we increased the mass of fuel while holding fuel moisture constant during a series of ponderosa pine needle burns. Burning different plant components also resulted in differ-

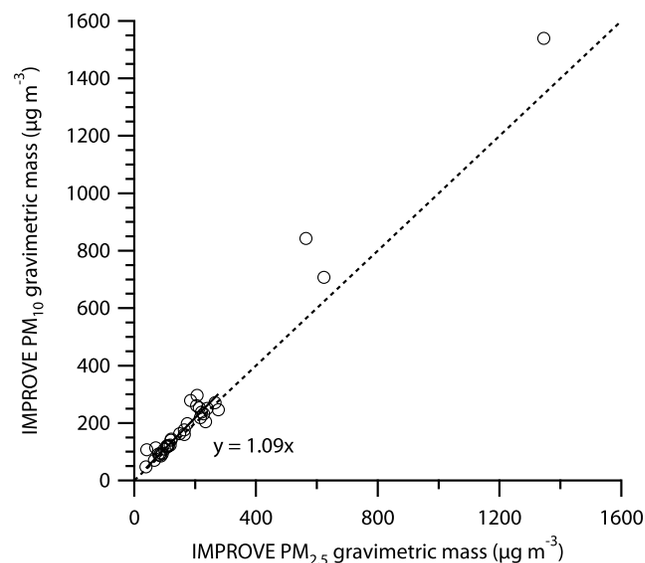


Figure 5. Gravimetrically determined mass concentrations of particles with aerodynamic diameters less than 10 µm (PM₁₀) compared to gravimetrically determined mass concentrations of particles with diameters less than 2.5 µm (PM_{2.5}) for IMPROVE filter samples obtained during chamber burns. Dashed line is the 1:1 line. Solid line gives the linear regression of PM₁₀ mass onto PM_{2.5} mass, forced through the origin, for all but the highest three concentration samples.

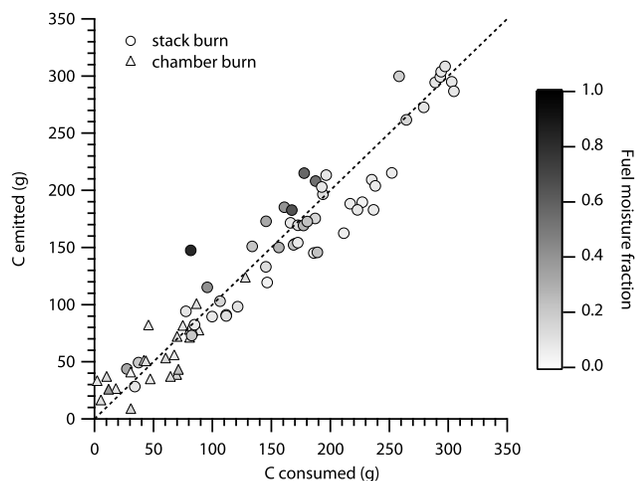


Figure 6. Carbon mass consumed versus carbon mass emitted during FLAME. Carbon mass consumed was calculated assuming the residual mass had zero water content. Carbon mass emitted consists of the sum of carbon monoxide, carbon dioxide, methane, C_{2-4} hydrocarbons, and particulate carbon. Points are shaded by fuel moisture to indicate samples where the assumption is less likely to be valid. Circles indicate stack burns, and triangles indicate chamber burns.

ent combustion behavior; we observed higher MCE for chamise and Douglas fir woody material compared to leaves and needles.

4.2. Total Particulate Emissions

[37] The gravimetric mass concentration data from the chamber burns confirmed that the PM_{10} mass concentrations were dominated by $PM_{2.5}$ mass concentrations (Figure 5). The PM_{10} to $PM_{2.5}$ mass ratio was 1.09, estimated from a zero-intercept linear regression of all but the three highest-concentration samples. The ratio increased to 1.16 if all samples were included in the regression. On average, aerosol emissions were dominated by carbon and TC made up almost 90% of reconstructed $PM_{2.5}$ mass emissions, which we computed by summing all identified aerosol species, as gravimetric data were only available for chamber burns,

$$\text{reconstructed } PM_{2.5} = \sum(\text{ionicspecies})_{URG} + EC + OC \times 1.5. \quad (4)$$

The rationale for the factor of 1.5 is discussed in section 4.3.4. We observed a large range in fire-integrated $PM_{2.5}$ emission factors ($1.9\text{--}82.1 \text{ g kg}^{-1}$ dry fuel). Since OC dominated $PM_{2.5}$ and its emissions were higher in smoldering combustion, the $PM_{2.5}$ EF also depended on MCE. Reid *et al.* [2005] estimated fine aerosol emission factors of $\sim 9 \text{ g kg}^{-1}$ fuel based on flaming combustion measurements, which they defined as $MCE > 0.9$, and $\sim 34 \text{ g kg}^{-1}$ fuel for smoldering combustion measurements ($MCE < 0.9$). Yokelson *et al.* [2008] obtained an average $EF_{PM_{2.5}}$ of 9.93 g kg^{-1} dry fuel in their laboratory studies of tropical fuels, similar to the recommendation of Reid *et al.* [2005], with variations between 2.17 and 16.61 g kg^{-1} fuel for various fuels that

had fire-integrated MCEs between 0.88 and 0.979. Ward and Hardy [1991] recommended $EF_{PM_{2.5}}$ of 10 g kg^{-1} fuel for cured grasses, 15 g kg^{-1} fuel for chaparral and palmetto/gallberry fires and $20\text{--}50 \text{ g kg}^{-1}$ fuel for long-needled conifer fires. In FLAME, the average $EF_{PM_{2.5}}$ for chaparral species was $11.6 \pm 15.1 \text{ g kg}^{-1}$ dry fuel; for palmetto, $11.4 \pm 10.5 \text{ g kg}^{-1}$ dry fuel; and for montane fuels (long-leaf conifers) $29.4 \pm 25.1 \text{ g kg}^{-1}$ dry fuel, on average, all very similar to previous recommendations.

4.3. Carbon Species

4.3.1. Total Carbon Mass Balance

[38] We calculated the mass of carbon emitted (C_{emitted}) during each burn by adding together carbon emitted in the form of CO_2 , CO , CH_4 , C_{2-4} hydrocarbons, and particle-phase OC and EC, for burns where all of these measurements were available. Figure 6 compares C_{emitted} to C_{consumed} , with the points coded by burn type and shaded by FM because the assumption of zero residual water content may not be valid for high moisture content fuels. The masses of carbon emitted and consumed were highly correlated ($r^2 = 0.96$) and close to the 1:1 line, indicating that emissions were effectively captured by the stack and could justifiably be assumed to be well-mixed in the chamber. On average, $89 \pm 5.7\%$ of the carbon was emitted in the form of CO_2 , followed by CO ($6.9 \pm 3.0\%$), OC ($2.3 \pm 2.5\%$), C_{2-4} hydrocarbons ($1.3 \pm 1.9\%$), CH_4 ($0.5 \pm 0.4\%$), and EC ($0.2 \pm 0.2\%$).

4.3.2. Carbon Monoxide and Carbon Dioxide

[39] We report fire-integrated emission factors for CO and CO_2 in auxiliary material Data Set S1 and in Table 2a give emission factors averaged for the plant species and ecosystem classifications described in section 2. The species and ecosystem data are the averages of all burns for that species or ecosystem type, so the numerical values depend on the number and variety of burns performed. The emission factors for many species were driven by the relative contributions from flaming and smoldering combustion during each burn, as expressed through fire-integrated MCE in this work, and the carbon abundance in the fuel. For example, Alaskan duff featured a strong contribution from smoldering combustion (average $MCE = 0.867 \pm 0.074$), but had a lower CO emission factor than several fuels with higher average MCE because it contained less carbon per unit mass (Table 1).

[40] The average EF_{CO_2} for montane fuels was $1552 \pm 150 \text{ g kg}^{-1}$ dry fuel (mean ± 1 standard deviation), near the $1569 \pm 131 \text{ g CO}_2 \text{ kg}^{-1}$ dry fuel recommended by Andreae and Merlet [2001] for extratropical forests. The EF_{CO_2} for rangeland fuels was somewhat lower ($1489 \pm 176 \text{ g kg}^{-1}$ dry fuel) and for coastal plain fuels was somewhat higher ($1632 \pm 150 \text{ g kg}^{-1}$), reflecting the different contributions from flaming and smoldering combustion quantified through the fire-integrated MCE. The average EF_{CO} for montane fuels was $92 \pm 34.1 \text{ g kg}^{-1}$ dry fuel, somewhat lower than the value recommended by Andreae and Merlet [2001] for extratropical forests ($107 \pm 37 \text{ g kg}^{-1}$ dry fuel). Rangeland and chaparral fuels had similar average EF_{CO} as montane fuels, but the average coastal plain value was lower ($78.0 \pm 27.7 \text{ g kg}^{-1}$ dry fuel), again reflecting different average contributions of flaming and smoldering combustion.

Table 2a. Gas-Phase Emission Factors for Individual Species and Ecosystem Groups^a

Species/Group	MCE	CO ₂	CO	CH ₄	C ₂ H ₂	C ₃ H ₆	NO	NO ₂	NH ₃	SO ₂
Montane	0.915 ± 0.033	1552 ± 150	92.0 ± 34.1	3.7 ± 2.7	5.7 ± 4.8	1.7 ± 1.2	1.5 ± 1.9	0.7 ± 0.9	1.7 ± 1.5	0.5 ± 0.4
Douglas fir	0.906 ± 0.036	1579 ± 193	106.8 ± 32.4	4.1 ± 3.8	5.8 ± 4.4	2.0 ± 1.6	3.8 ± 1.9	2.1 ± 1.0	1.6 ± 0.9	0.3 ± 0.2
Lodgepole pine	0.920 ± 0.035	1528 ± 106	84.6 ± 38.8	4.2 ± 2.5	8.3 ± 7.7		0.4 ± 0.2	0.4 ± 0.3	2.1 ± 2.3	0.6 ± 0.3
Montana grass	0.863 ± 0.062	1172 ± 228	115.3 ± 50.5	4.2	8.4					
Ponderosa pine	0.920 ± 0.026	1589 ± 85	88.4 ± 30.7	3.2 ± 2.0	4.4 ± 3.7	1.4 ± 0.4	0.9 ± 1.3	0.4 ± 0.2	1.6 ± 1.4	0.6 ± 0.5
Rangeland	0.905 ± 0.043	1489 ± 176	96.4 ± 38.2	3.3 ± 3.1	3.5 ± 3.0	1.5 ± 1.0	4.6 ± 2.0	0.3 ± 0.2	2.7 ± 2.2	0.6 ± 0.4
Juniper	0.956	1713	51	0.2	0.7		2.2	0.2	0.8	0.4
Rabbitbrush	0.935	1529	68	1.3	1.5	2.2	1.4	0.5	0.8	0.2
Sagebrush	0.889 ± 0.041	1437 ± 173	111.2 ± 34.6	4.6 ± 3.1	4.4 ± 3.0	1.3 ± 1.1	5.7 ± 0.7		4.0 ± 1.8	0.7 ± 0.4
Chaparral	0.909 ± 0.029	1538 ± 125	93.2 ± 24.1	2.5 ± 2.1	3.3 ± 2.1	1.4 ± 1.1	1.7 ± 2.2	0.5 ± 0.2	1.5 ± 1.3	0.4 ± 0.2
Ceanothus	0.913 ± 0.012	1623 ± 51	98.3 ± 11.6	1.7 ± 0.4	1.7 ± 0.6	0.7 ± 0.5	4.3 ± 3.9	1.1	1.4 ± 0.3	0.3 ± 0.2
Chamise	0.914 ± 0.030	1562 ± 112	86.1 ± 20.9	2.3 ± 1.6	3.4 ± 2.0	1.5 ± 1.2	1.7 ± 2.2	0.4 ± 0.1	1.2 ± 0.8	0.4 ± 0.3
Manzanita	0.899 ± 0.030	1471 ± 138	104.4 ± 28.9	3.8 ± 3.6	4.1 ± 3.0	1.9 ± 1.3	1.3 ± 1.8	0.5 ± 0.1	2.1 ± 2.2	0.4 ± 0.1
Coastal plain	0.930 ± 0.029	1632 ± 150	78.0 ± 27.7	2.7 ± 1.7	2.6 ± 2.2	1.1 ± 1.2	4.5 ± 2.4	0.7 ± 0.4	2.0 ± 1.8	0.9 ± 1.4
Black needlerush	0.891 ± 0.030	1538 ± 114	119.0 ± 28.3	5.4 ± 1.8	4.1 ± 1.7	0.7	3.8 ± 0.3		1.1	0.5
Common reed	0.957 ± 0.013	1656 ± 9	47.0 ± 15.6	1.6 ± 0.4	2.7 ± 0.0		8.1 ± 2.1		1.9	1.3
Gallberry	0.947 ± 0.004	1868 ± 5	66.0 ± 4.2	2.4 ± 0.2	1.7 ± 0.2	0.5 ± 0.0	7.3		1.3	0.4
Hickory	0.933 ± 0.005	1583 ± 24	72.0 ± 4.2	2.7 ± 0.5	1.9 ± 0.1	0.5 ± 0.0			2.6	0.7
Kudzu	0.857 ± 0.003	1096 ± 35	116.5 ± 0.7	4.8 ± 1.7	8.4 ± 1.0	2.3 ± 0.2	6.5 ± 1.2		6.6	1.1
Longleaf pine	0.944 ± 0.023	1659 ± 78	60.8 ± 27.6	2.1 ± 0.9	2.7 ± 3.4	1.6 ± 2.2	3.2 ± 1.5	1.3	3.6 ± 3.2	0.5 ± 0.4
Oak	0.943 ± 0.007	1622 ± 43	65.7 ± 3.5	1.7 ± 0.6	2.4 ± 0.4	0.6 ± 0.2	9.6		2.2	0.7
Palmetto	0.933 ± 0.018	1678 ± 65	75.9 ± 19.5	2.3 ± 1.6	1.5 ± 0.8	0.6 ± 0.4	2.9 ± 1.9	0.5 ± 0.2	0.9 ± 0.4	1.7 ± 2.6
Rhododendron	0.961	1783	46	1.8	1.2		3.9		1.3	0.1
Sawgrass	0.900 ± 0.008	1522 ± 16	107.0 ± 8.5	3.4 ± 0.0	2.0 ± 0.2		5.9 ± 0.6		1.8	0.7
Titi	0.942	1825	71	1.8	1.1		7.6			
Turkey oak	0.886 ± 0.006	1580 ± 31	129.5 ± 4.9	5.9 ± 1.2	4.3 ± 0.8	1.4 ± 0.3	6.3 ± 0.2		3.7	0.5
Wax myrtle	0.915 ± 0.013	1622 ± 61	95.7 ± 11.5	2.9 ± 0.9	3.7 ± 2.7	2.0 ± 2.3	3.6 ± 3.7	1	1.8 ± 0.0	0.3 ± 0.2
Wire grass	0.965 ± 0.007	1680 ± 9	43.0 ± 1.4	0.6 ± 0.2	0.4 ± 0.1	0.2	3.5 ± 0.4		0.4	0.8
Boreal forest	0.917 ± 0.068	1311 ± 325	70.6 ± 40.2	1.4 ± 0.9	1.7 ± 1.6	0.7 ± 0.6	3.3 ± 1.8	1.6 ± 1.1	1.4 ± 0.7	0.1 ± 0.1
Alaskan duff	0.867 ± 0.074	1034 ± 175	96.4 ± 43.0	2.3 ± 0.9	2.5 ± 2.7	1.1 ± 1.0	2.0 ± 0.7	1.0 ± 0.5	1.9 ± 0.6	0.0 ± 0.0
Black spruce	0.957 ± 0.012	1588 ± 125	44.8 ± 11.2	0.8 ± 0.4	1.3 ± 0.4	0.5 ± 0.3	3.9 ± 2.0	3.2	0.7 ± 0.5	0.2 ± 0.1
White spruce	0.971			1.7					1.7	0.1
Other	0.922 ± 0.035	1411 ± 82	84.4 ± 29.8	2.8 ± 2.8	2.0 ± 2.0	0.8 ± 0.7	2.2 ± 2.1	0.5 ± 0.1	0.9 ± 0.4	0.6 ± 0.5
Fern	0.943	1571	60	1.7	2.3	2.1	0.8	0.5	0.7	0
PR mixed woods	0.952			1.7	0.8	0.8				
Rice straw	0.911 ± 0.032	1394 ± 64	87.1 ± 30.3	3.6 ± 3.3	2.4 ± 2.4	0.5 ± 0.3	2.5 ± 2.2	0.4 ± 0.0	1.0 ± 0.5	0.7 ± 0.5
Sugarcane	0.977			0.8	0.9					

^aEmission factors are reported in g species kg⁻¹ dry fuel. Bold type indicates the average of all samples for each group. Blank entries are below detection limit or not calculated (see text for details).

4.3.3. Gas-Phase Hydrocarbons

[41] Fire-integrated emission factors for most of the measured hydrocarbon species were positively correlated with MCE, with r^2 ranging from 0.39 to 0.67. In Figure 7, we compare our results to the regressions reported by *Christian et al.* [2003] for emissions from African fuels burned at the FSL. The FLAME and *Christian et al.* [2003] regressions for CH₄ are in nearly perfect agreement. The two studies took place in the same facility, but examined different fuels, and used a different method to determine CH₄ concentrations (gas chromatography versus open path FTIR). *Yokelson et al.* [2003] measured slightly higher emission factors for CH₄ over African savanna fires, but obtained a similar slope. A number of FLAME samples fall on the *Yokelson et al.* [2003] regression, but it is unclear if this is just a coincidence or reflects a systematic difference in CH₄ emissions for different fuels or fire regimes. The FLAME emission factors for C₂H₂ and C₂H₄ were higher than the *Yokelson et al.* [2003] and *Christian et al.* [2003] regressions predicted, particularly for fire-integrated MCE values lower than 0.85, but both of those studies examined a narrower range of higher MCEs than those achieved in FLAME. The least squares fitting method used for the FLAME regressions was strongly influenced by the high emission factor values we observed at low MCE. Similar

comparisons between domestic and other fuels, and over a broader range of fire-integrated MCE, might be in order for other important emissions.

4.3.4. Carbonaceous Aerosols

[42] Elemental carbon emissions were associated with flaming-phase combustion, consistent with temperature and oxidant-dependent soot formation mechanisms. Figure 8 illustrates the relationship between fire-integrated MCE and EC/TC for emissions from two fuel classes during FLAME: needle and branch components of ponderosa pine (Figure 8a) and several chaparral and desert shrub fuels, including sagebrush, chamise, and manzanita (Figure 8b). EC/TC ratios were less than 10% for MCE values below ~0.93, and increased strongly for MCE > 0.93 for both fuel classes. The EC/TC ratio was ~0 for a sample collected during only the smoldering phase of the fire (MCE = 0.80) and 0.5 for a sample collected during the flaming phase (MCE = 0.99).

[43] The relationships in Figure 8 were similar to previous measurements for similar fuels. *Battye and Battye* [2002] summarized recommended EF derived from a number of airborne field studies reported in the gray literature. For Ponderosa pine, EC/TC ratios for flaming/smoldering combustion were 0.06/0.16; for chaparral species in smoldering combustion, 0.11, whereas flaming conditions yielded 0.11–0.22. Findings from prior laboratory studies are shown in

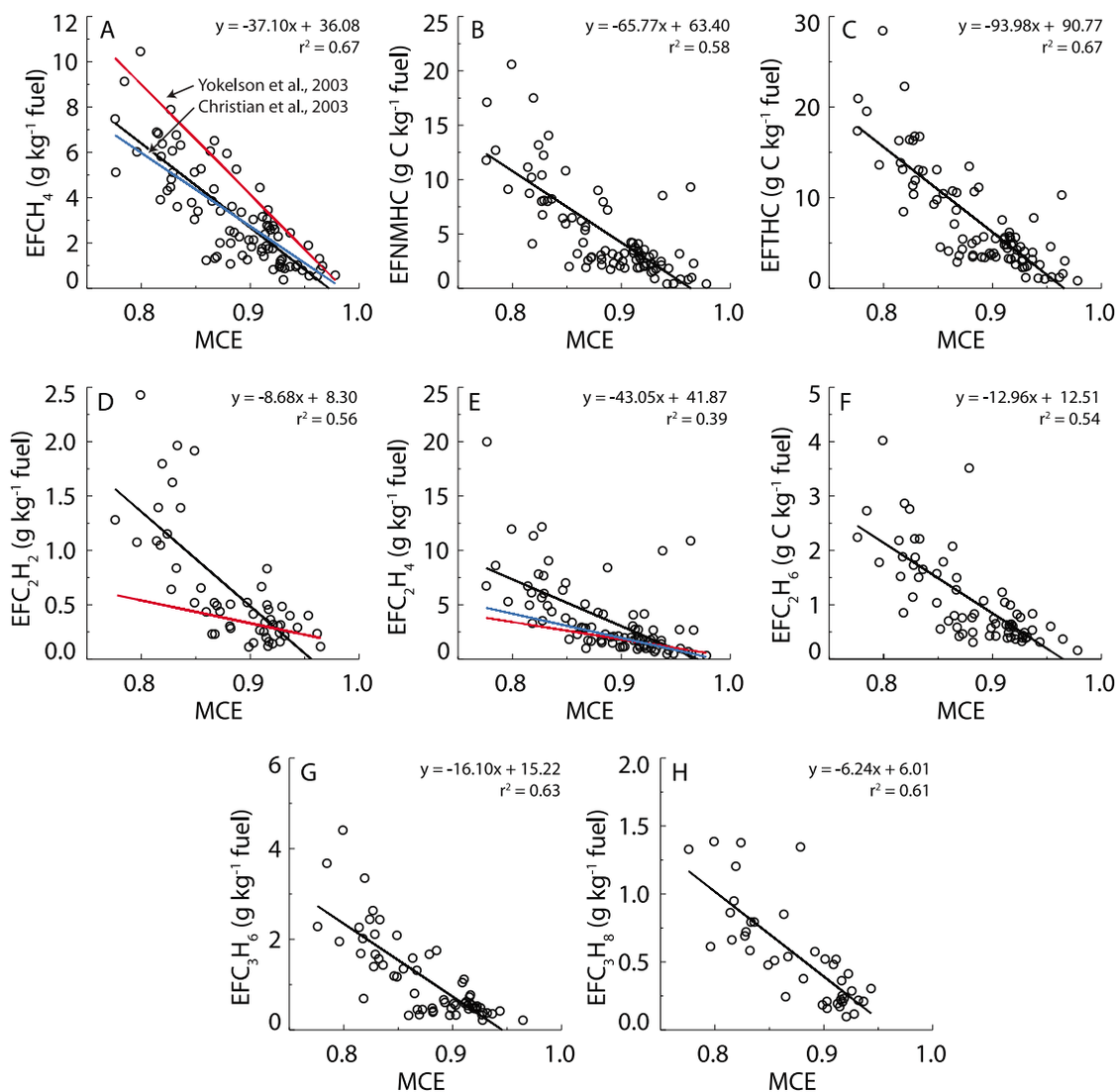


Figure 7. Fire-integrated emission factors for hydrocarbon gas species calculated from canister gas chromatography measurements as a function of fire-integrated modified combustion efficiency (MCE) for all tested fuels. Emission factors are shown for (a) methane (CH_4), (b) total nonmethane hydrocarbons (NMHC), (c) total hydrocarbons (THC), (d) acetylene (C_2H_2), (e) ethene (C_2H_4), (f) ethane (C_2H_6), (g) propene (C_3H_6), and (h) propane (C_3H_8). Black lines indicate the linear least squares regression of the emission factors onto MCE.

Figure 8 [Chen *et al.*, 2007; Hays *et al.*, 2002; Inuma *et al.*, 2007; Christian *et al.*, 2003]. Note that Hays *et al.* [2002] did not report fire-integrated MCE, so we estimated fire-integrated MCE from their reported time series of $\Delta[\text{CO}_2]$ and $\Delta[\text{CO}]$ mixing ratios, and Inuma *et al.* [2007] reported only the median and not burn-integrated values of $\Delta[\text{CO}]$ and $\Delta[\text{CO}_2]$. Further, different techniques were used to measure EC in the various studies. Nevertheless, at similar values of MCE, the various field and laboratory measurements are in general agreement. We note that the larger range of MCE accessed in the FLAME experiments enabled a better overall picture of the variations in emissions with MCE. For example, conditions with $\text{MCE} \sim 0.95$ are not frequently accessible during field studies since they are associated with the intense flaming phase of combustion, but our data showed that large fractions of EC can be emitted by chaparral species under

those conditions. This variability over a fire lifetime may be important in estimating the total emissions of EC to the atmosphere.

[44] The patterns in Figure 8 were not evident for all fuels. Several produced little or no EC when burned despite featuring a substantial flaming contribution and associated high MCE. These fuels, rice straw in particular, also produced particles with some of the highest inorganic mass fractions of total $\text{PM}_{2.5}$, so it is possible the two are linked. Inorganic salts may catalyze combustion of EC on the filter during the OC stages of the thermal optical analysis (TOA), but photoacoustic measurements of the aerosol made online during the burn showed the emissions were only weakly absorbing [Lewis *et al.*, 2008], confirming the lack of EC. In their microscopy analysis of aerosol emissions, Hopkins *et al.* [2007] identified a distinct category of fuels that featured

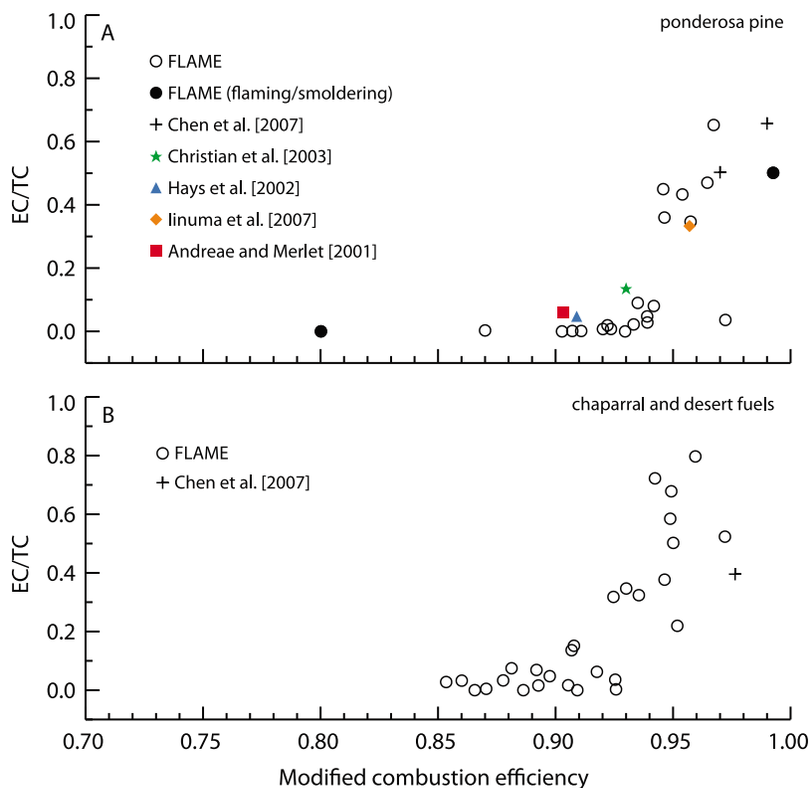


Figure 8. Elemental-to-total aerosol carbon (EC/TC) ratios observed for emissions from (a) ponderosa pine and (b) chaparral and desert shrub fuels versus fire-integrated modified combustion efficiency (MCE). Ponderosa pine data include needle, branch, needles and branches, needle litter, and duff burns. Samples collected during only flaming (high MCE) and smoldering (low MCE) combustion of ponderosa pine needles are indicated by the solid circles; all others are fire-integrated. Previously measured ratios from selected studies are also shown.

a strong flaming phase when burned, but produced a significant concentration of inorganic salts and had optical properties inconsistent with EC.

[45] Figure 9 shows fire-integrated emission factors for OC, EC, and TC for all tested fuels as a function of fire-integrated MCE. The data are also tabulated by plant species and ecosystem in Table 2a and for each burn in auxiliary material Data Sets S2 and S3. A factor of 1.5 was used to compute the total organic mass concentration, accounting for associated O, H, N, and other elements, from the measured C mass concentrations attributed to OC. The 1.5 factor was within the range of OM-to-OC factors of 1.4–1.8 for biomass burning aerosol recommended by Reid *et al.* [2005], and was validated for FLAME data from comparisons of reconstructed total aerosol mass concentrations with measured gravimetric mass concentrations (Levin *et al.*, manuscript in preparation, 2009).

[46] Organic carbon emission factors were negatively correlated with MCE ($r^2 = 0.36$), increasing, as expected, with increasing contributions from smoldering-phase combustion (Figure 9a). Emission factors ranged from $\sim 0.5 \text{ g C kg}^{-1}$ fuel at high MCE to $\sim 50 \text{ g C kg}^{-1}$ fuel at lower MCE values. Juniper, rabbitbrush, rhododendron and white spruce were examples of plants with low OC emission factors, with emissions dominated by flaming combustion, as reflected by the fire-integrated MCE. Examples of plants with high OC emission factors included “leafy” fuels such as kudzu, turkey

oak, sagebrush, and manzanita that had low fire-integrated MCE. The coastal plain category had the highest average OC emission factor ($12.4 \pm 12.0 \text{ g C kg}^{-1}$ fuel) and those in the chaparral category had the lowest ($6.6 \pm 10.1 \text{ g C kg}^{-1}$ fuel), but these averages do not account for the relative abundances of particular plants in the ecosystem. The range of OC emission factors reported in the literature is very large, even for single species, as we would expect given the sensitivity of emissions to combustion conditions. OC emission factors reported for ponderosa pine range from at least $3\text{--}30 \text{ g C kg}^{-1}$ [Hays *et al.*, 2002]. Andreae and Merlet [2001] suggest an OC emission factor for extratropical forest fires of $8.6\text{--}9.7 \text{ g C kg}^{-1}$ fuel, somewhat lower than the averages for montane fuels we report in Table 2b, but higher than the average for boreal species.

[47] Elemental carbon emission factors during FLAME ranged from 0 to 8 g C kg^{-1} fuel (Figure 9b). The significance of the relationship between EC and MCE was weaker ($r^2 = 0.09$) than that between OC and MCE. Rangeland and coastal plain species tended to have higher EC emission factors compared to fuels from other regions, but with considerable variability within the classifications. The study-average EC emission factor for montane species was $0.4 \pm 0.8 \text{ g C kg}^{-1}$ fuel compared to the literature average of 0.56 ± 0.19 reported by Andreae and Merlet [2001] for extratropical forests. The lower MCE in FLAME sagebrush burns, compared to those reported by Chen *et al.* [2007], led to averages of $0.63 \pm$

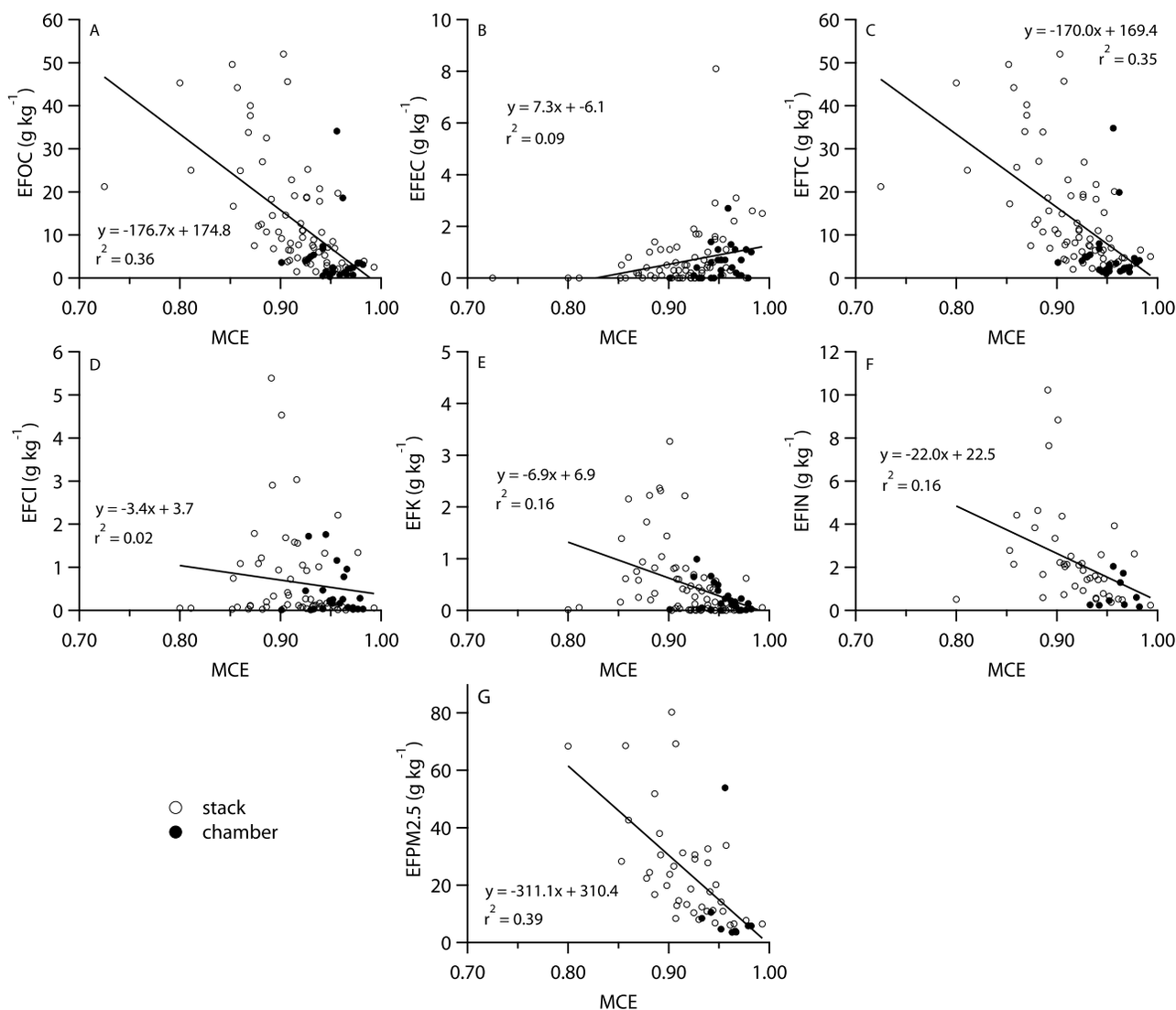


Figure 9. Fire-integrated aerosol emission factors (EF) as a function of fire-integrated modified combustion efficiency (MCE) for (a) organic carbon (OC), (b) elemental carbon (EC), (c) total aerosol carbon (TC), (d) chloride, (e) potassium, (f) total inorganic aerosol species, and (g) reconstructed PM_{2.5}. Black lines indicate the linear regression of EF onto MCE with coefficients and coefficient of variation indicated on the plot for each species.

0.42 g kg⁻¹ fuel compared with 1.4 g kg⁻¹ fuel in that earlier study. Several studies have reported EC emission factors for ponderosa pine [e.g., *Chen et al.*, 2007; *Christian et al.*, 2003; *Hays et al.*, 2002], ranging from 0.4 to 2.6 g kg⁻¹, compared to 0.48 ± 0.83 g kg⁻¹ in our study. *Ward and Hardy* [1991] give a range of emission factors for “graphitic carbon” of 0.46–1.18 g kg⁻¹ for fires burning in the Pacific northwest, a region with large populations of ponderosa pine.

[48] *de Gouw and Jimenez* [2009] recently compared emission ratios for organic aerosols from a number of biomass burning sources, and found they ranged from approximately 60 to 130 $\mu\text{g m}^{-3}$ (ppm ΔCO)⁻¹ for primary organic aerosol. The study average for FLAME was higher, at 180 ± 170 $\mu\text{g m}^{-3}$ (ppm ΔCO)⁻¹, closer to organic aerosol/ ΔCO ratios of 200 $\mu\text{g m}^{-3}$ (ppm ΔCO)⁻¹ in an aged urban/biomass burning plume near Mexico City reported by *DeCarlo et al.* [2008]. Recent work by *Grieshop et al.* [2009] showed that biomass burning emissions can be

oxidized and form secondary organic aerosol, leading to increases in the organic aerosol/ ΔCO ratio, but *Capes et al.* [2008] did not observe any ratio increase over fires in Africa despite other evidence of aging. The FLAME results showed that high organic aerosol/ ΔCO emission ratios can exist in fresh biomass burning emissions with a high level of variability, making it difficult to draw conclusions about the importance of primary and secondary sources of organic aerosol.

4.4. Nitrogen Emissions

4.4.1. Gas-Phase Nitrogen

[49] We compared the mass of NH₃ and NO_x emitted to the mass of N consumed in the burn, rather than to the N present in the fuel, to account for the N ash component. The NO_x measurements for FLAME 2 were estimated using measurements of NO and the mean ratio of NO₂:NO observed during FLAME 1 because an instrument malfunction

Table 2b. Aerosol-Phase Emission Factors by Ecosystem Species and Group^a

Species/Group	MCE	OC	EC	K ⁺	Na ⁺	NH ₄ ⁺	Cl ⁻	NO ₃ ⁻	SO ₄ ²⁻	PM _{2.5}
Montane	0.915 ± 0.033	18.4 ± 16.3	0.4 ± 0.8	0.1 ± 0.2		0.0 ± 0.1	0.1 ± 0.0	0.3 ± 0.7	0.2 ± 0.5	29.4 ± 25.1
Douglas fir	0.906 ± 0.036	26.0 ± 14.9	0.36 ± 0.75	0.07 ± 0.10		0.08 ± 0.10	0.03 ± 0.01	0.84 ± 1.17	0.62 ± 0.88	42.9 ± 22.9
Lodgepole pine	0.920 ± 0.035	11.3 ± 15.2	0.45 ± 0.70	0.19 ± 0.38		0.01 ± 0.01	0.06 ± 0.05	0.04 ± 0.04	0.08 ± 0.07	18.1 ± 23.1
Montana grass	0.863 ± 0.062									
Ponderosa pine	0.920 ± 0.026	17.6 ± 17.0	0.48 ± 0.83	0.03 ± 0.07		0.02 ± 0.02	0.06 ± 0.04	0.13 ± 0.27	0.11 ± 0.15	27.7 ± 26.0
Rangeland	0.905 ± 0.043	9.4 ± 8.1	1.2 ± 0.9	1.1 ± 0.8	0.1 ± 0.1	0.0 ± 0.0	1.2 ± 1.1	0.2 ± 0.3	0.5 ± 0.4	18.9 ± 13.9
Juniper	0.956	0.7	2.7	0.28		0	0.16	0.01	0.14	4.2
Rabbitbrush	0.935	0.5	1.4	0.66	0.02	0.01	0.47	0.02	0.39	3.4
Sagebrush	0.889 ± 0.041	15.3 ± 1.2	0.63 ± 0.42	1.50 ± 0.76	0.09 ± 0.09	0.04 ± 0.01	1.78 ± 1.08	0.40 ± 0.31	0.73 ± 0.34	29.0 ± 1.9
Chaparral	0.909 ± 0.029	6.6 ± 10.1	0.5 ± 0.4	0.5 ± 0.3	0.0 ± 0.0	0.0 ± 0.0	0.2 ± 0.1	0.1 ± 0.1	0.2 ± 0.1	11.6 ± 15.1
Ceanothus	0.913 ± 0.012	3.8 ± 0.1	0.35 ± 0.35	0.74 ± 0.12	0.00 ± 0.00	0.03 ± 0.00	0.43 ± 0.03	0.11 ± 0.12	0.31 ± 0.06	7.8 ± 1.2
Chamise	0.914 ± 0.030	3.2 ± 2.5	0.56 ± 0.48	0.42 ± 0.35	0.00 ± 0.01	0.01 ± 0.02	0.23 ± 0.09	0.10 ± 0.19	0.25 ± 0.11	6.5 ± 4.2
Manzanita	0.899 ± 0.030	14.8 ± 17.3	0.35 ± 0.31	0.38 ± 0.15	0.01 ± 0.02	0.03 ± 0.03	0.09 ± 0.02	0.07 ± 0.08	0.14 ± 0.04	23.5 ± 25.9
Coastal plain	0.930 ± 0.029	12.4 ± 12.0	0.9 ± 1.7	0.7 ± 0.9	0.1 ± 0.3	0.2 ± 0.1	1.3 ± 1.5	0.2 ± 0.1	0.3 ± 0.2	23.4 ± 18.7
Black needlerush	0.891 ± 0.030	18.3	0.3	2.36	1.16	0.51	5.39	0.16	0.61	38.4
Common reed	0.957 ± 0.013	19.7	0.4	0.14		0.27	2.21	0.6	0.33	36.2
Gallberry	0.947 ± 0.004	7.1	8.1	0.57		0.08	0.08	0.17	0.48	20.5
Hickory	0.933 ± 0.005	7.1	0.3	0.37		0.12	0.11	0.25	0.38	12.5
Kudzu	0.857 ± 0.003	44.2		0.61		0.1	0.07	0.49	0.66	70.5
Longleaf pine	0.944 ± 0.023	23.8 ± 8.9	0.93 ± 0.32	0.23 ± 0.04	0.09 ± 0.09	0.15 ± 0.12	0.78 ± 0.47	0.12 ± 0.03	0.14 ± 0.01	38.3 ± 13.6
Oak	0.943 ± 0.007	10.6	0.4	0.44		0.12	0.1	0.15	0.36	18.2
Palmetto	0.933 ± 0.018	5.0 ± 6.6	0.47 ± 0.34	0.59 ± 0.81	0.06 ± 0.07	0.26 ± 0.12	1.45 ± 0.83	0.08 ± 0.08	0.19 ± 0.14	11.4 ± 10.5
Rhododendron	0.961	2.1	0.2	0.08		0.01	0.07	0.04	0.04	3.7
Sawgrass	0.900 ± 0.008	9.2	1.1	3.27	0.24	0.13	4.53	0.16	0.44	24.6
Titi	0.942									
Turkey oak	0.886 ± 0.006	32.5	1.4	0.82		0.04	0.2	0.19	0.35	52.2
Wax myrtle	0.915 ± 0.013	6.3 ± 2.5	0.35 ± 0.07	0.80 ± 0.27	0.25 ± 0.10	0.10 ± 0.04	1.22 ± 0.70	0.10 ± 0.03	0.29 ± 0.02	12.2 ± 4.0
Wire grass	0.965 ± 0.007	3.5	0.3	0.03		0.02	0.09	0.14	0.16	6.4
Boreal forest	0.917 ± 0.068	7.8 ± 7.2	0.2 ± 0.4	0.1 ± 0.0		0.0 ± 0.0	0.1 ± 0.1	0.1 ± 0.0	0.1 ± 0.1	12.7 ± 11.3
Alaskan duff	0.867 ± 0.074	10.2 ± 10.0	0.00 ± 0.00	0.03 ± 0.02	0.00 ± 0.01	0.02 ± 0.02	0.03 ± 0.02	0.09 ± 0.01	0.09 ± 0.11	16.1 ± 15.9
Black spruce	0.957 ± 0.012	6.2 ± 2.8	0.60 ± 0.46	0.05 ± 0.03	0.01 ± 0.01	0.03 ± 0.03	0.07 ± 0.04	0.10 ± 0.07	0.08 ± 0.04	10.4 ± 4.2
White spruce	0.971	3.5		0.13		0.05	0.28	0.08	0.01	5.9
Other	0.922 ± 0.035	5.6 ± 3.5	0.1 ± 0.1	0.6 ± 0.3	0.0 ± 0.1	0.2 ± 0.2	1.3 ± 0.6	0.0 ± 0.0	0.1 ± 0.1	10.2 ± 6.6
Fern	0.943	2.2	0.1	0.13		0.03	0.25	0.02	0.05	3.9
PR mixed woods	0.952									
Rice straw	0.911 ± 0.032	6.2 ± 3.5	0.08 ± 0.15	0.69 ± 0.22	0.03 ± 0.07	0.26 ± 0.16	1.54 ± 0.34	0.04 ± 0.03	0.13 ± 0.10	11.8 ± 6.5
Sugarcane	0.977									

^aEmission factors are reported in g species kg⁻¹ dry fuel, except that OC and EC are g C kg⁻¹ dry fuel and PM_{2.5} was reconstructed from species measurements (equation (4)). Bold type indicates the average of all samples for each group. Blank values indicate that the measurement was below the detection limit or not calculated (see text for details).

prevented accurate measurement of NO₂. The N consumed by the burn was assumed to be equal to the product of the dry fuel N content and the dry mass consumed during the burn. Ammonia emissions represented approximately 21 ± 30% and nitrogen oxides represented 27 ± 26% of the N consumed, but NO_x emissions were much larger during FLAME 2 compared to FLAME 1. In FLAME 1, NH₃ and NO_x accounted for ~20% of the N consumed on average, whereas in FLAME 2 they represented ~50%. There was no strong difference in the average N contents for the fuels we burned during each of the studies, and the mass of fuel used in each burn was similar, so that fire size, as hypothesized by *Goode et al.* [1999], did not appear to be a factor. It is possible that the changes in the ignition method between the two studies may be responsible for the observed differences.

[50] Laboratory and field measurements have shown that NO_x is emitted primarily via flaming combustion and NH₃ is emitted primarily by smoldering combustion [*Goode et al.*, 2000; *Lobert et al.*, 1991; *Yokelson et al.*, 1996]. However, emission factors for individual nitrogen species are not strongly correlated with MCE and instead depend primarily on fuel nitrogen content [*Andreae and Merlet*, 2001; *Lobert et al.*, 1991; *Yokelson et al.*, 2008]. To account for the fuel N dependence, *Yokelson et al.* [1996] and *Goode et al.* [1999,

2000] compared molar ratios of NH₃ and NO_x to MCE. They showed that a linear relationship between NH₃/NO_x and MCE was consistent for fire emissions measured in the laboratory and field for a variety of fuels. Figure 10 compares the *Goode et al.* [2000] relationship between NH₃/NO_x molar ratios and MCE with FLAME observations and other recently published data. The FLAME data points are shaded according to the absolute NO_x mass emissions to illustrate increasing uncertainty in the molar NH₃/NO_x ratios calculated for low NO_x cases. A linear least squares regression to the high-NO_x data (defined as having absolute NO_x emissions greater than 0.6 g equivalent NO) indicated that NH₃ composed the majority of the identified N emissions below a fire-integrated MCE ~0.85. Most of the samples that deviated from the linear fit corresponded to burns with low NO_x emissions and high uncertainties in the calculated NH₃/NO_x molar ratios.

[51] NH₃/NO_x molar ratios during FLAME were about a factor of 2 lower than those reported and summarized by *Goode et al.* [2000] at similar MCE. *Goode et al.* [2000] treated all NO_x emissions as NO because NO₂ mixing ratios were below their instrument's detection limits. The high-NO_x FLAME data agreed with the *Goode et al.* [2000] fit if NH₃:NO molar ratios are considered. Several other field

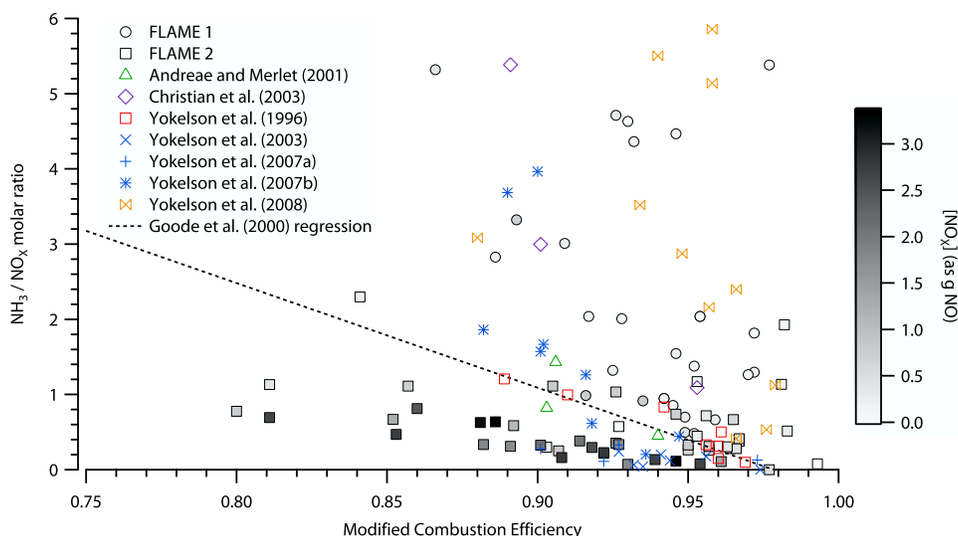


Figure 10. Molar ratios of NH_3 -to- NO_x emissions as a function of fire-integrated modified combustion efficiency (MCE) during FLAME and as reported for several other biomass burning field and laboratory experiments, as indicated in the legend. FLAME data are shaded to reflect the magnitude of the NO_x measurement and therefore reflect the confidence in the measured ratio. The dashed line indicates the fit provided by *Goode et al.* [2000] for several sets of laboratory and field biomass burning measurements. Note that this plot is truncated to better illustrate the majority of NH_3/NO_x data from our study and the literature. A maximum NH_3/NO_x ratio of ~ 12 at an MCE of 0.82 was reported by *Christian et al.* [2003].

measurements of NH_3 and NO_x from open-path and aircraft-based Fourier Transform Infrared spectrometry (FTIR) published this decade also deviated significantly from the *Goode et al.* [2000] fit, as shown in Figure 10. An improved description of NH_3/NO_x ratios in emissions may be important in estimates of global N budgets, as well as in source apportionment studies that rely on accurate profile information.

[52] We calculated emission factors for NO , NO_2 , and NH_3 following the same approach used to calculate CO , CO_2 and hydrocarbon emission factors (auxiliary material Data Set S1). Fire-integrated NO emission factors ranged from 0.04 to 9.6 g NO kg^{-1} dry fuel, with a study mean and standard deviation of $2.6 \pm 2.4 \text{ g NO kg}^{-1}$ dry fuel. There was a large difference between the average FLAME 1 EFNO ($0.7 \pm 0.5 \text{ g NO kg}^{-1}$) and the average FLAME 2 EFNO ($3.9 \pm 2.4 \text{ g NO kg}^{-1}$). This could have been due to the larger number of N-rich grasses and other plants we tested during FLAME 2. Average NO emission factors for species in the coastal plain and rangeland categories were almost 3 times higher than for montane and chaparral species and NH_3 emission factors were roughly 50% higher. The higher rangeland averages were due primarily to sagebrush, which had emission factors for NO and NH_3 of 5.7 ± 0.7 and $4.0 \pm 1.8 \text{ g kg}^{-1}$ fuel, respectively. The FLAME sagebrush averages were considerably higher than the EFNO of 2.94 g kg^{-1} and EF NH_3 of 0.19 g kg^{-1} reported by *Yokelson et al.* [1996].

[53] Nitric acid (HNO_3) concentrations measured using the denuder samplers were typically much lower than the other N-containing gas species we measured. The study average emission factor was $0.12 \pm 0.58 \text{ g HNO}_3 \text{ kg}^{-1}$ dry fuel, but the concentrations of HNO_3 were below the MDL for most of the samples. Nitric acid emissions were $\sim 5\%$ of the N emitted in the form of NO_x .

4.4.2. Particulate Nitrogen

[54] We measured particulate-phase nitrogen in the form of NH_4^+ , NO_3^- , and NO_2^- and found that these species generally accounted for only a small fraction of the fuel nitrogen as well as a small fraction of the total $\text{PM}_{2.5}$ mass. Nitrate emission factors ranged from 0.01 to 2.9 $\text{g NO}_3^- \text{ kg}^{-1}$ dry fuel, with a study-average value of $0.2 \pm 0.4 \text{ g kg}^{-1}$ dry fuel. The observations span the range previously reported in the literature [*Andreae et al.*, 1998; *Hays et al.*, 2002; *Hegg et al.*, 1987]. Including the particulate nitrogen species, we were able to identify between 10 and 50% of the original fuel nitrogen, consistent with the findings of *Lobert et al.* [1990] and *Kuhlbusch et al.* [1991]. The remaining fuel nitrogen was likely emitted in the form of N_2 , HCN , and nitrogen-containing organic species [*Yokelson et al.*, 2007a] or remained in the ash following the burn.

4.5. Sulfur Emissions

4.5.1. Sulfur Dioxide

[55] Sulfur dioxide emission factors ranged from approximately 0–7 $\text{g SO}_2 \text{ kg}^{-1}$ dry fuel. *Andreae and Merlet* [2001] recommended an SO_2 emission factor of $1.0 \text{ g SO}_2 \text{ kg}^{-1}$ dry fuel for extratropical forests. *Ferek et al.* [1998] observed SO_2 emission factors in the tropics ranging from roughly 0.2–1.5 $\text{g SO}_2 \text{ kg}^{-1}$ C burned, which corresponds to a range of roughly 0.1–0.7 $\text{g SO}_2 \text{ kg}^{-1}$ dry fuel assuming a fuel C fraction of 0.45. *Ferek et al.* [1998] noted that EF SO_2 increased weakly with MCE, but did not observe a strong correlation between MCE and EF SO_2 , which was also not observed in our data set.

4.5.2. Sulfate

[56] Sulfate emission factors ranged from 0.0 to 5.7 $\text{g SO}_4^{2-} \text{ kg}^{-1}$ dry fuel and were weakly correlated with MCE, increasing slightly with decreasing MCE. For savanna fires

in Africa, *Sinha et al.* [2003] observed sulfate emission factors on the order of $0.2 \text{ g SO}_4^{2-} \text{ kg}^{-1}$ dry fuel, whereas *Andreae et al.* [1998] reported $0.6 \text{ g SO}_4^{2-} \text{ kg}^{-1}$ dry fuel. Even higher SO_4^{2-} emission factors have been measured further from the source; e.g., the airborne data of *Andreae et al.* [1998] yielded 4–10 times higher SO_4^{2-} emission factors than did ground-based measurements closer to the fire. In our experiments, SO_2 was emitted at roughly 1–4 times the rate of SO_4^{2-} . If this emitted SO_2 is subsequently oxidized in the atmosphere to form SO_4^{2-} , the combined emission factors suggest an equivalent SO_4^{2-} yield of $0.6 \pm 1.4 \text{ g SO}_4^{2-} \text{ kg}^{-1}$ dry fuel.

4.6. Other Inorganic Species

4.6.1. Chlorine

[57] On average, chloride was the most abundant inorganic species in the aerosol during FLAME, accounting for $40 \pm 14\%$ of the soluble inorganic and $4.5 \pm 2.5\%$ of the reconstructed $\text{PM}_{2.5}$ mass concentrations. *Reid et al.* [2005] estimated Cl^- made up 2–5% of $\text{PM}_{2.5}$ in fresh biomass burning emissions and *Chen et al.* [2007] found that chloride accounted for 0.1–9.6% of $\text{PM}_{2.5}$ for several of the same fuels we burned. Emissions from several southeastern fuels burned during FLAME contained high mass fractions of chloride relative to other inorganic species. For example, chloride was $\sim 60\%$ of the inorganic emissions for a palmetto leaf (*Serenoa repens*) burn.

[58] Chloride emission factors ranged from 0.0 to 5.4 g kg^{-1} fuel (study average, $0.4 \pm 0.7 \text{ g kg}^{-1}$ fuel) and were not a function of MCE (Figure 9d). Previously reported EFCl include $\sim 0.0\text{--}3.2 \text{ g kg}^{-1}$ fuel [*Keene et al.*, 2006]; $0.0\text{--}1.8 \text{ g kg}^{-1}$ fuel [*Christian et al.*, 2003]; and $1\text{--}2 \text{ g kg}^{-1}$ fuel [*Andreae et al.*, 1998]. Several studies showed that roughly one third of fuel chlorine was emitted in the form of particulate matter for tropical and savannah fuels [*Christian et al.*, 2003; *Keene et al.*, 2006; *Yokelson et al.*, 2008]. Although we did not measure the fuel chlorine content, chloride mass fractions of total inorganics within fuel classes were relatively constant, indicating that fuel type and chlorine content were the major drivers of chloride emissions.

4.6.2. Potassium

[59] Excess (nonsoil and non-sea-salt) potassium has long been used as a tracer for biomass burning aerosol [*Andreae*, 1983]. It was the second-most abundant inorganic species measured during FLAME, making up $2.5 \pm 1.8\%$ of reconstructed $\text{PM}_{2.5}$ mass concentrations and $22 \pm 8\%$ of the inorganic mass. Potassium emission factors ranged from 0.0 to 3.3 g kg^{-1} fuel, with a study average of $0.3 \pm 0.5 \text{ g kg}^{-1}$ fuel (Figure 9e). *Christian et al.* [2003] reported EFK ranging from 0.02 to 1.29 g kg^{-1} for African savanna, Indonesian peat, and several wildland plant species and *Andreae and Merlet* [2001] provide literature-average values ranging from 0.08 to 0.41 g kg^{-1} fuel for extratropical forests. The higher values observed in FLAME were a result of the types of fuels burned. In particular, rangeland plant species had large EFK, along with many coastal plain fuels. Fire-integrated molar ratios of potassium to chloride and sulfate were consistent with K being in the form of predominantly KCl with a minor contribution from K_2SO_4 .

4.6.3. Other Species

[60] Sodium, calcium, magnesium and nitrite made up the remainder of the analyzed inorganic species in the emissions. The totals of all measured inorganic emission factors were only weakly correlated with MCE ($r^2 = 0.16$) (Figure 9f), as expected since fuel composition should play the largest role in emissions of inorganic aerosol species [*Christian et al.*, 2003; *Keene et al.*, 2006].

5. Discussion

[61] The dependencies of carbonaceous and inorganic emission factors on fuel and burn characteristics have implications for predictions of biomass burning impacts on climate, air quality, and visibility, because these are sensitive to the chemical composition of the aerosol. Estimates of smoke aerosol optical properties require accurate information regarding combustion conditions in order to estimate the relative abundances of EC and OC, which to a large extent determine the single scattering albedo. Emission factors for OC and $\text{PM}_{2.5}$ are stronger functions of combustion conditions, compared to EF for inorganic compounds, but depend only weakly on plant species. Lack of data over a broad range of MCE may result in biased estimates of fire-related aerosol amounts and properties. For example, if smoldering emissions are underestimated in current biomass burning inventories, then total $\text{PM}_{2.5}$ concentrations attributable to biomass burning are likely to be underestimated: (1) the emission factors for $\text{PM}_{2.5}$ increase with decreasing MCE; (2) emissions of most carbonaceous gas species increase with decreasing MCE, and it is likely that a fraction of these eventually form secondary organic aerosol; (3) as MCE decreases, more N is released in the form of NH_3 , which can readily convert to particulate-phase ammonium. Ongoing work examining time-resolved aerosol mass spectrometer (AMS) data obtained in the FLAME 2 studies is exploring the relationships between particulate matter emissions and fire phase more closely.

[62] There are limits to the usefulness of the MCE in capturing other effects of the fire. *Ward and Hardy* [1991] found that emission factors for total PM increased relative to $\text{PM}_{2.5}$ emissions as fire energy release rates increased. They attributed the increased PM emissions to increased turbulence for the larger fire, which lofted larger-sized PM, including ash and soil material. *Andreae et al.* [1998] observed increases in the Ca^{2+} and Mg^{2+} content of coarse mode aerosol over intense savanna fires, which they also attributed to the lofting of soil material by the turbulence in the fire. This lofting effect is not captured by the MCE, nor would the laboratory studies reproduce these soil emissions. Proxies for combustion behavior other than MCE may provide a more practical tool for linking laboratory measurements to the modeling of observed fires. For example, recent laboratory work by *Ichoku et al.* [2008] showed that fire radiative energy (FRE) measured by a thermal imaging system was strongly correlated with aerosol emission rates. This work could be extended to examine the relationships between FRE and individual gas- and particle-phase species. An advantage of FRE-based emission factors is that they can be applied to satellite measurements to develop more accurate emissions inventories.

[63] Source apportionment techniques attempt to separate fire-related particles from other sources and to apportion the fire-related aerosols retrospectively to various fire types such as wildland, prescribed, agricultural, and residential. Most apportionment studies have been conducted using chemical transport models, receptor models, and hybrids of the two. While chemical transport models require accurate emission inventories, a necessary component of which are accurate EF, receptor-type models require appropriate tracer species to apportion sampled aerosols to these various sources. The use of a subset of FLAME data (measurements of aerosol OC, water-soluble potassium, and levoglucosan, a smoke marker compound) to develop better source profiles for biomass burning aerosols is discussed in *Sullivan et al.* [2008]. The ratios of EF we report in the auxiliary material for various aerosol species can also be applied as source emission profiles. For example, *Park et al.* [2007] examined observed TC-to-nonsoil-potassium ratios across the IMPROVE network to investigate the contributions by biomass burning to annual U.S. aerosol concentrations. They estimated TC/K ratios near 10 for grassland and shrub fires in the south and ratios approaching 130 for fires in the north. We found similar ratios in the emissions from individual plant species from these regions, suggesting that our measured TC/K ratios could be used to estimate primary fire contributions to TC from the studied fuel types.

6. Conclusions

[64] We have reported fire-integrated emission factors and aerosol mass fractions for 33 predominantly North American wildland plant species. To our knowledge, many have not been previously studied in laboratory open burning experiments, including the chaparral species chamise, manzanita, and ceanothus, and species common to the southeastern United States (common reed, hickory, kudzu, needlegrass rush, rhododendron, cord grass, sawgrass, titi, and wax myrtle). These species frequently burn in wildland fires and prescribed burns near urban centers, so their emissions have important effects on urban air quality. We note here that the EF reported for EC, an aerosol component that plays a key role in radiative forcing, are up to a factor of 2 lower than those that would be obtained if an alternate analysis protocol were used to analyze the filters, as shown by our comparisons for a limited number of burns. The magnitude of the emission factor for EC remains a significant uncertainty in estimates of the climate impacts of biomass burning.

[65] To assist in the interpretation of our gas- and aerosol-phase measurements, we report the corresponding fire-integrated MCE. Our results are consistent with previous work that found carbonaceous gas- and particle-phase emissions depend more strongly on MCE than did the emissions of inorganic species, which depend most strongly on fuel type and composition [*Ward and Hardy*, 1991]. Combustion behavior still plays a role in the form of the inorganic emissions (e.g., NO_x versus NH_3), but the relationships between fire-integrated inorganic gas and particle emission factors and fire-integrated MCE are weak. The aerosol composition data provide a basis set for interpreting simultaneous measurements of aerosol optical and hygroscopic properties, CCN activity, and IN activity that were conducted during FLAME.

[66] The generally consistent relationships between laboratory- and field-derived EF that we found in this work support the integrated approach advocated by *Yokelson et al.* [2008] for the development of more comprehensive descriptions of EF for use in modeling. As those authors pointed out, different ranges of MCE are accessed in laboratory, airborne and ground-based sampling strategies, and capturing EF over a large measured range of MCE can be expected to enhance the accuracy of modeled emissions estimates. Several examples from their own work showed how combining sources of data led to insights on the variation of emissions with fire phase that were not obvious from measurements over a limited range of MCE. However, two caveats in combining such data are (1) the MCE and EF we measured in the laboratory are fire-integrated, whereas those measured in a field study may represent only a portion of the burn history and (2) the emissions in a small-scale laboratory fire do not fully reflect those in a true wildfire. Nevertheless, we have confirmed here that many EF for specific fuels are surprisingly consistent when interpreted through the corresponding MCE. These findings suggest value in continuing controlled laboratory studies of emissions from important fuel types that have also been observed in the field, combining the observations from various platforms and approaches to develop more robust, MCE-dependent emissions estimates. In particular, further work should revisit the relationship between NH_3/NO_x and MCE because our results confirm the large variation observed in previous measurements.

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