

Moisture effects on carbon and nitrogen emission from burning of wildland biomass

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Abstract. Carbon (C) and nitrogen (N) released from biomass burning have multiple effects on the Earth's biogeochemical cycle, climate change, and ecosystem. These effects depend on the relative abundances of C and N species emitted, which vary with fuel type and combustion conditions. This study systematically investigates the emission characteristics of biomass burning under different fuel moisture contents, through controlled burning experiments with biomass and soil samples collected from a typical alpine forest in North America. Fuel moisture in general lowers combustion efficiency, shortens flaming phase, and introduces prolonged smoldering before ignition. It increases emission factors of incompletely oxidized C and N species, such as carbon monoxide (CO) and ammonia (NH₃). Substantial particulate carbon and nitrogen (up to 4 times C in CO and 75% of N in NH₃) were also generated from high-moisture fuels, mainly associated with the pre-flame smoldering. This smoldering process emits particles that are larger and contain lower elemental carbon fractions than soot agglomerates commonly observed in flaming smoke. Hydrogen (H)/C ratio and optical properties of particulate matter from the high-moisture fuels show their resemblance to plant cellulose and brown carbon, respectively. These findings have implications for modeling biomass burning emissions and impacts.

1 Introduction

On a global scale, biomass burning is an important process mobilizing carbon (C) and nitrogen (N) from the biosphere to the atmosphere, with $2\text{--}5 \times 10^{15}$ g of C and $15\text{--}46 \times 10^{12}$ g of N burned and released each year (Crutzen and Andreae, 1990). C emissions in the form of greenhouse gases, including carbon dioxide (CO₂) and methane (CH₄), are known to contribute to climate change (IPCC, 2007). N is an essential nutrient for all living organisms, and its emissions in the form of nitrous oxide (N₂O), nitrogen oxides (NO_x), ammonia (NH₃), particulate nitrogen, etc. can be utilized by the living organisms and therefore impact the ecosystem (Vitousek et al., 1997). In addition, many of the C- and/or N-containing volatile and particulate components from biomass burning play vital roles in radiative forcing, atmospheric chemistry, and air pollution (Keil and Haywood, 2003; Kondo et al., 2004; Crouse et al., 2009). The fate (i.e., atmospheric lifetime) and effects of C and N from biomass burning depend on how they are distributed between gaseous and particulate phases and among various species. The dry deposition velocity of NH₃, for instance, is at least 10 times that of NO_x (Neirynek et al., 2007). N in NH₃ more likely remains close to where biomass burning occurs rather than being removed from the source region due to long-range transport.

Emission rates and compositions are sensitive to fuel type and combustion conditions (e.g., fuel load, moisture content, and wind). Flaming combustion involves a more complete oxidation, converting C and N to CO₂ and NO_x, respectively, than smoldering combustion (Koppmann et al., 2005; Chen et al., 2007). Particles are generated from both flaming



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and smoldering phases, but they differ in size, morphology, and optical properties (Reid et al., 2005a, b; Chen et al., 2006; Chakrabarty et al., 2006; McMeeking et al., 2009). Light-absorbing black carbon with a structure similar to soot agglomerates dominates in high temperature flames, while smoldering particles are whiter and more spherical in shape. Particle hygroscopicity increases with the inorganic fraction of particulate mass which depends more on fuel type than on combustion conditions (Carrico et al., 2010).

Laboratory experiments are useful for controlling and investigating individual parameters effecting the biomass burning emission. The Fire LAB at Missoula Experiments (FLAME I and II) conducted in 2006 and 2007 investigated a wide range of biomass types and loads in a laboratory combustion system (McMeeking et al., 2009). Fuels used during the FLAME study were either fresh or dried sufficiently to readily ignite. A systematic investigation of the fuel moisture effect on biomass burning was not part of FLAME I and II and is rarely reported in the literature. However, moisture contents of wildland biomes could vary corresponding to climate and weather conditions, substantially influencing emissions from wild fires and prescribed burns.

As part of the Tahoe Prescribed Fires and Nutrient Emission Experiment (TPFNEE), laboratory-controlled burning experiments were conducted with fuels prepared at different moisture levels. The experiments emphasized shrubs and downed materials that were subject to both prescribed and wild fires. This paper documents changes in emission factors of C and N species as a function of fuel moisture content. Based on continuous and time-integrated measurements, it also examines the emission mechanisms needed for process-based emission models. The Tahoe basin is a typical alpine forest in North America. Decades of fire suppression resulted in large biomass accumulation in this region. Prescribed burning is considered as an option for fuel reduction, and TPFNEE was tasked to provide a critical assessment on the environmental impact of such practices.

2 Methods

Biomass and soil samples were collected in Incline Village, Nevada, northeast of the Lake Tahoe Basin, in spring 2008 from two plots that were scheduled for prescribed burning by the United States Forest Service (USFS). Within each of the plots, downed materials in a $20 \times 20 \text{ cm}^2$ area, 5 cm deep bulk volume were picked up from the forest floor at 12 randomly selected coordinates for a total of 6–8 kilograms of materials. These materials were then segregated into: 1) litter (the surface layer which is not in an advanced stage of decomposition, typically consisting of freshly fallen and dried leaves, needles, twigs, stems, bark, and fruits); 2) duff (the layer dominated by partially to fully decaying leaves and branches); and 3) topsoil (mixture of mineral soil, humus, and/or degraded organic material). Topsoil was examined

because Murphy et al. (2006) found that wild fires nearly deplete C and N in soils. Common aboveground shrubs were also collected and separated in the laboratory into leaves and stems of different diameters. Acquired samples were stored in sealed polyethylene bags under refrigeration ($<4^\circ\text{C}$).

The field moisture content of each sample was quantified by drying a small portion of it at 85°C and measuring the weight loss after 24 h. Litter, duff, and soil showed moisture contents between 1–10% of dry mass, with median values of 3.5%, 5.0%, 3.8%, respectively. Moisture contents of aboveground shrub samples (i.e., Bitterbrush, Green Leaf Manzanita, Huckleberry Oak, and Squaw Carpet) were higher, ranging from 6.5–50.0% for leaves and 9.3–49.0% for stems. Fractions of C and N in the dried mass were determined by CHNS-O analysis (Thermo EA1110-CHNS-O elemental analyzer, Thermo Nicolet Corporation, Waltham, USA).

To prepare fuels for desired/different moisture contents, individual litter, duff, and soil samples from different locations were first homogenized and air dried for a week at 30% relative humidity. Calculated amounts of water were added into these samples to achieve fuels with moisture contents of 10 or 20% of dry mass. Shrub leaf and stem samples were also composited, and either air dried or soaked in water for 24 to 96 h. The fuel moisture content, determined from the weight losses of small fractions of samples, increased over time and gradually reached saturation (Fig. 1). Dry, 24-h soaked, and 96-h soaked biomass were used for the burning experiments. Table 1 shows the fuels and moisture levels examined in this study. The two wet moisture levels (II & III) simulated natural fuels of relatively high moisture contents. Two replicate burns were conducted for each fuel-moisture combination.

Figure 2 shows the combustion facility and measurement suite. Fuels were loaded on a uniformly-heated hotplate ($25 \times 25 \text{ cm}^2$) with adjustable temperatures up to 500°C . Fuels were weighed before and after the experiment. Fuel loads were 23–69 g per burn. The hotplate kept fuels warm throughout the experiment to simulate large scale burns where environmental temperatures could be much higher than those in the laboratory without a heater. The hotplate was located in a 60-cm diameter fire pit. Horizontal air flow was kept calm, and most of the smoke was vented through the chimney with an exhaust fan. Fuels were placed on the hotplate (maintained at 450°C) for ~ 30 s before starting an electric hot-air gun (Loof Lighter) to ignite them. Dry fuels were ignited within seconds, while it generally required application of hot air ($\sim 600^\circ\text{C}$) for 1–2 min to ignite high moisture fuels. Some wet fuels required additional use of the hot air gun for several minutes to sustain combustion after ignition. The hot air gun avoided interference from emissions of combustion-based igniter (e.g., propane torch).

The smoke sampling probe consisted of 2-inch (5.08 cm) diameter conductive tubing stretching from the top of chimney, about 1.5 m above the fuels, to a mixing plenum (Fig. 2). Plumes generally equilibrated with ambient temperatures in

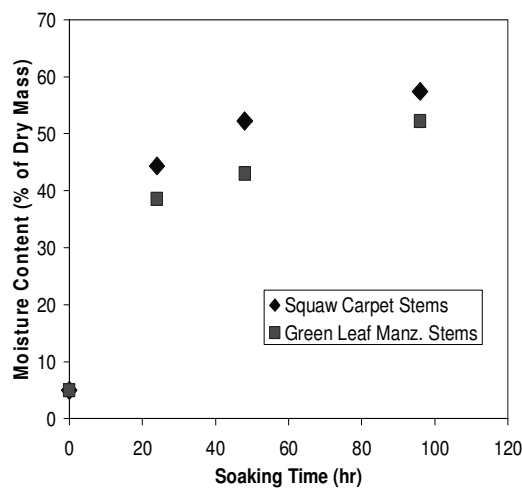


Fig. 1. Examples of fuel moisture content, starting air dried (0 h), as a function of time soaking in water. Excess water was drained before the measurement of moisture content.

the plenum. Through two size-cut cyclones, $PM_{2.5}$ (particles with aerodynamic diameter $<2.5 \mu m$) were sampled onto 1) a Teflon-membrane filter (for $PM_{2.5}$ mass and elements) with a calcium carbonate impregnated cellulosic backup filter (for sulfur dioxide [SO_2]) and 2) a quartz-fiber filter (for organic carbon [OC] and elemental carbon [EC] by the IMPROVE_A protocol (Chow et al., 2007a) and water-soluble ions) with citric acid impregnated cellulosic backup filter (for NH_3). The analytical methods for filter samples are the same as described in Chow (1995) and Chow et al. (2004). Lower quantifiable limits (LQLs) were established with dynamic blanks through the same sampling channels but in the absence of combustion. Measurement precision is $<10\%$ for concentrations greater than 10 times the LQLs. Additionally, portions of quartz-fiber filters were submitted to the CHNS-O analyzer for quantifying total C (TC), N (TN), and hydrogen (TH) in $PM_{2.5}$. TC by CHNS-O analysis agrees with OC + EC by IMPROVE_A within $\pm 8\%$.

An Electric Low Pressure Impactor (ELPI) was used to measure particle size and number concentrations in real time. The ELPI has 12 stages covering an aerodynamic size range of 10 nm to $10 \mu m$. Size retrieval followed the algorithm developed by Marjamäki et al. (2005), which is shown to give accurate size distributions for particles smaller than $3 \mu m$ (Pagels et al., 2005). Through Teflon tubing from the plenum and downstream of a Teflon particle filter (Fig. 1), several gaseous species were continuously monitored with a nondispersive infrared CO_2/H_2O analyzer (LI-840, LiCor Biosciences, Lincoln, NE), an extractive Fourier Transform Infrared (FTIR) Spectrometer (Midac, Costa Mesa, CA, for CO), and research-grade NO_x and NH_3 analyzers (TECO 42 and 17C, Thermo Scientific, Waltham, MA). These analyzers were calibrated against standards traceable to the National

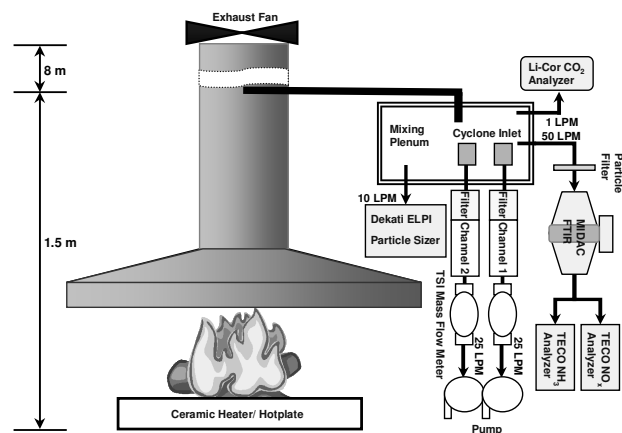


Fig. 2. Schematic of the controlled burning experiment configurations.

Institute of Standard and Technology before and after the experiments. The detection ranges (10-s average) were 0.001–10, 0.1–3000, and 1–10 000 ppmv for the TECO, Li-Cor, and Midac analyzers, respectively. The precision of gas measurements is estimated to be $<5\%$ if they are within the instrumental detection range. All our experiments met the condition.

Since the experimental setup sampled a small portion of smoke plumes, production efficiencies of gaseous and particulate species (per unit fuel consumption) were calculated using a carbon balance approach (Andreae and Merlet, 2001). The concentration ratio of a measured species over grand total carbon (GTC) released from a burn defines the emission factor (EF) of that species in gram per kilogram of carbon burned, which can be scaled to gram per kilogram of fuel consumed with known carbon content of the fuel. GTC includes C in CO_2 , CO, and $PM_{2.5}$. Volatile organic compounds (VOCs) including CH_4 were not measured, though under most circumstances VOCs are minor contributors, compared to CO_2 and CO, in the carbon budget of emissions ($<5\%$). Natural variability of EFs, according to replicate burns, usually exceeds the EF uncertainties estimated from the analytical precision. Table 1 shows the average and standard deviation of fuel-specific EFs, while Fig. 3 illustrates their dependence on the fuel moisture content. All leaf and stem burns have been combined in Fig. 3 to increase the number of replicate measurements.

3 Results and discussion

Dry and wet biomass fuels showed distinct fire behaviors. Dry fuels were ignited easily by the hot air and flames quickly spread over the fuel. Residues after the burn were small except for the duff composite, for which only half of the dry mass was combusted (Table 1). The duff sample likely contained substantial mineral material, as its carbon

Table 1. Time-integrated combustion efficiency (CE) and emission factors^a for major C and N species as well as PM_{2.5} from controlled burning experiments.

Fuel Type →		Downed Material					Aboveground Shrub			
Moisture/Parameter		Litter	Composite Duff	Soil	Bitterbrush		Manzanita		Squaw Carpet	
					Leaves	Stems	Leaves	Stems	Leaves	Stems
Moisture Level I (Dry ^e)	Carbon % ^b	50%	32%	3%	52%	48%	49%	48%	47%	50%
	Burned % ^c	92%	52%	9%	92%	81%	90%	98%	83%	93%
	CE	0.94±0.01	0.92±0.02	0.86±0.02	0.93±0.01	0.94±0.00	0.91±0.01	0.94±0.00	0.92±0.01	0.92±0.00
	CO ^f	126.2±12.7	140.6±27.3	297.3±48.2	112.6±30.3	127.4±8.8	113.2±43.5	128.4±6.7	144.5±22.3	162.1±8.9
	OC	2.5±0.2	(9.3±6.7)	14.9±4.4	9.1±1.8	6.4±2.0	21.2±7.9	5.5±0.6	11.3±0.2	4.7±0.1
	EC	4.6±0.3	5.7±0.1	0.7±0.2	15.4±2.7	2.9±0.3	15.9±3.9	2.0±0.4	8.1±0.8	2.5±0.5
	PM _{2.5}	8.1±0.9	(18.3±10.2)	N.D.	27.1±5.8	13.0±3.9	47.5±14.1	8.7±0.4	25.3±3.2	14.5±1.1
	NO _x ^f	5.9±0.3	4.9±0.9	0.9±0.2	9.1±2.8	9.1±1.0	7.1±3.0	5.4±1.0	7.7±1.5	10.5±0.8
	NH ₃	(1.3±1.1)	(3.7±2.4)	(11.5±6.1)	7.7±2.4	3.4±0.6	4.0±1.6	1.9±0.8	6.8±1.6	5.6±0.1
	NO ₃ ⁻	0.04±0.01	(0.03±0.02)	N.D.	0.05±0.01	0.12±0.00	0.16±0.04	0.12±0.00	0.07±0.00	0.09±0.02
	NH ₄ ⁺	0.05±0.01	0.06±0.03	N.D.	0.04±0.00	0.04±0.01	0.04±0.00	0.03±0.00	0.05±0.00	0.08±0.00
	OPN	(0.24±0.19)	(0.44±0.35)	(0.31±0.23)	0.40±0.11	0.31±0.12	0.47±0.21	0.09±0.03	0.45±0.04	0.28±0.01
	Moisture Level II	Moisture ^d	10%	10%	10%	73%	39%	40%	39%	48%
Burned % ^c		72%	47%	10%	88%	66%	73%	53%	69%	92%
CE		0.79±0.06	0.70±0.07	0.80±0.06	0.88±0.01	0.82±0.06	0.72±0.03	0.86±0.11	0.67±0.09	0.86±0.06
CO ^f		253.2±59.8	444.9±68.1	431.4±143.8	113.2±42.4	261.3±44.0	(97.5±76.7)	107.1±2.4	(77.7±57.7)	(108.1±96.0)
OC		99.8±34.9	110.9±41.5	17.5±1.0	66.1±9.9	(66.6±36.1)	236.0±62.4	(62.8±64.1)	292.2±115.1	92.6±18.4
EC		5.2±0.3	1.0±0.1	1.5±0.5	(7.3±4.0)	3.7±0.7	5.6±1.1	(2.3±1.2)	8.5±3.2	2.6±0.3
PM _{2.5}		160.5±56.2	143.9±62.2	N.D.	96.9±18.3	(119.3±73.7)	214.9±84.1	(94.9±91.3)	270.8±68.4	109.6±32.7
NO _x ^f		6.4±1.5	6.7±1.4	2.8±0.8	9.9±4.0	13.0±3.1	(2.2±2.9)	2.9±1.2	(4.0±5.4)	(10.7±7.9)
NH ₃		4.3±1.7	(26.7±14.0)	24.7±6.6	10.2±1.5	10.7±0.9	10.0±0.5	3.8±0.5	11.9±4.1	11.0±5.4
NO ₃ ⁻		0.17±0.00	0.13±0.01	N.D.	0.23±0.02	0.20±0.02	0.08±0.03	0.08±0.01	0.23±0.04	0.20±0.01
NH ₄ ⁺		0.24±0.04	0.15±0.01	N.D.	0.10±0.00	(0.03±0.02)	0.31±0.09	(0.05±0.06)	0.28±0.11	0.06±0.01
OPN		1.62±0.59	3.57±1.50	0.99±0.26	2.03±0.05	1.39±0.68	4.4±1.42	(1.07±1.33)	7.70±3.28	2.17±0.46
Moisture Level III		Moisture ^d	20%	20%	20%	84%	57%	60%	52%	66%
	Burned % ^c	80%	45%	9%	86%	78%	68%	92%	65%	90%
	CE	0.74±0.10	0.69±0.07	0.82±0.01	0.51±0.07	0.84±0.04	0.62±0.09	0.87±0.00	0.72±0.01	0.88±0.08
	CO ^f	313.6±105.8	407.6±29.2	357.7±15.9	215.6±31.1	216.8±60.4	134.2±43.5	188.5±25.1	142.6±22.3	(133.1±102.6)
	OC	116.6±48.8	131.1±59.2	22.1±0.8	393.6±54.1	67.8±11.1	315.3±111.1	43.1±5.8	210.4±20.2	(61.5±39.7)
	EC	(7.6±4.9)	1.1±0.2	2.3±0.9	5.3±0.5	3.6±0.1	7.7±1.3	2.6±1.0	4.8±0.0	2.7±0.7
	PM _{2.5}	179.9±54.7	208.9±56.2	N.D.	768.7±43.2	99.5±4.4	493.8±231.9	61.5±3.8	387.2±31.1	129.1±43.5
	NO _x ^f	8.4±2.5	3.8±0.2	5.4±0.1	9.6±2.0	7.5±0.5	(2.7±2.0)	4.7±0.5	6.4±2.2	9.1±2.3
	NH ₃	5.3±0.4	17.7±5.9	(24.5±18.0)	19.1±1.4	10.9±1.8	10.1±0.4	3.8±0.0	9.5±0.8	(9.7±4.9)
	NO ₃ ⁻	0.20±0.05	0.09±0.03	N.D.	0.28±0.11	0.18±0.02	0.12±0.00	0.11±0.01	0.14±0.03	(0.13±0.08)
	NH ₄ ⁺	0.30±0.11	0.15±0.06	N.D.	0.32±0.11	0.06±0.01	0.42±0.14	0.07±0.01	0.18±0.00	(0.05±0.03)
	OPN	1.59±0.56	3.36±1.40	0.60±0.07	11.83±0.53	1.59±0.39	4.32±1.52	0.66±0.04	4.43±0.36	(1.33±0.76)

^a In g/kgC (g per kilogram of carbon burned). Values are based on the average and standard deviation of two replicate burns. N.D. indicates the species are below detection limit. Emission factors with higher variability (average/standard deviation <2) are shown in bracket.

^b Percentage of carbon in dry fuels.

^c Combined percentage of dry fuel burned.

^d Fuel moisture content in percentage of dry mass.

^e Moisture content <5%.

^f Not filter-based measurements.

content (32%) was lower than the nominal value (i.e., 45–50%) for biomass (McMeeking et al., 2009). It took much longer for wet fuels to ignite when they were heated by the same hot air. Smoke was visible before ignition. Flaming periods were relatively short, followed by prolonged smoldering combustion. Wet fuels left more and variable amounts of unburnable residues than dry fuels. For soil, flames were absent throughout the experiments, and only 9–10% of the dry mass was consumed, regardless of moisture content.

Particles from each individual burn were collected on a single set of Teflon and quartz-fiber filters that yielded burn-average chemical compositions. For consistency, all continuous measurements were baseline subtracted and integrated over the filter sampling period. NH₃ from the TECO 17C correlated reasonably with those of filters ($r = 0.83$), but the TECO 17C values were lower by an average of 21% and as much as 80%. This possibly results from the loss of NH₃ in the Teflon sampling tubing and to the pre-analyzer particle filter (e.g., Mukhtar et al., 2003). EF calculations

(i.e., Table 1) were therefore based on filter measurements which deployed much shorter/larger diameter tubing from the plenum. CO₂, CO, and NO_x adsorption on Teflon surface are generally negligible.

3.1 Time-integrated carbon measurement

Combustion efficiency (CE), defined as the fraction of C emissions in the form of CO₂, best indicates the relative importance of flaming (high CE) and smoldering (low CE) phases (Sinha et al., 2004; Janhäll et al., 2010). CE was >0.9 for all dry fuels (except soil) and was <0.9 for all wet fuels in this study (Table 1). In general, wet duff/leaves produced lower CEs than wet litter/stems, though there are no apparent differences in CEs between the two wet moisture levels (II and III). CO often constitutes most of the non-CO₂ carbon from biomass burning (Yokelson et al., 1996; Andreae and Merlet, 2001; McMeeking et al., 2009). For fuels with high moisture contents, however, particulate carbon approached or exceeded the carbon in CO emissions. The sum of OC and EC (moisture level III, Table 1) exceeds 2–4 times the carbon from CO for three stem burns and is 57–113% of CO carbon for the rest of burns except soil.

Thermal/optical analysis indicates that most of the particulate carbon is OC. The increasing PM_{2.5} and OC EFs with fuel moisture content are clear for leaf combustion (and to a less degree for litter, duff, and stem, Fig. 3). This likely reflects an enhanced primary process converting plant material to OC, since the experimental setup does not allow plumes to age sufficiently for secondary OC formation. Soil contains little biomass, and the influence of soil moisture on OC emissions is limited. Although only leaf and duff burns show EC EFs to decrease with increasing fuel moisture content, the EC fraction in TC ranges from 0.01 to 0.68 with all the highest values found from burning dry biomass. This is consistent with EC being generated from flaming combustion that intensifies with dry fuels. From such burns, Charkrabarty et al. (2006) identified soot-based particles similar to those from diesel engines.

The TH/TC ratio increases with the OC fraction in TC (Fig. 4), which corroborates the dominance of hydrocarbon and elemental C in the (operationally-defined) OC and EC, respectively. For 17 biomass burns in which OC/TC > 0.97 (i.e., low to no EC), the TH/TC ratio averages at 0.13 ± 0.02. Considering TH/TC ratios of 0.167, 0.139, and 0.108 for sugar [(CH₂O)_n], cellulose [(C₆H₁₀O₅)_n], and lignin [(C₁₀H₁₃O₃)_n], respectively, the OC may result from decomposition of cellulose, which is the most abundant organic polymer in biomass (Gani and Naruse, 2007).

Particles from wet fuel combustion often appear yellowish to brownish on filters and in water extracts, in contrast to a black appearance from dry fuels (Andreae and Gelencser, 2006). A two-wavelength (370 and 880 nm) filter transmittance instrument (OT21, Magee Scientific Co, Berkeley, CA) was used to examine the loaded Teflon filters for wavelength-

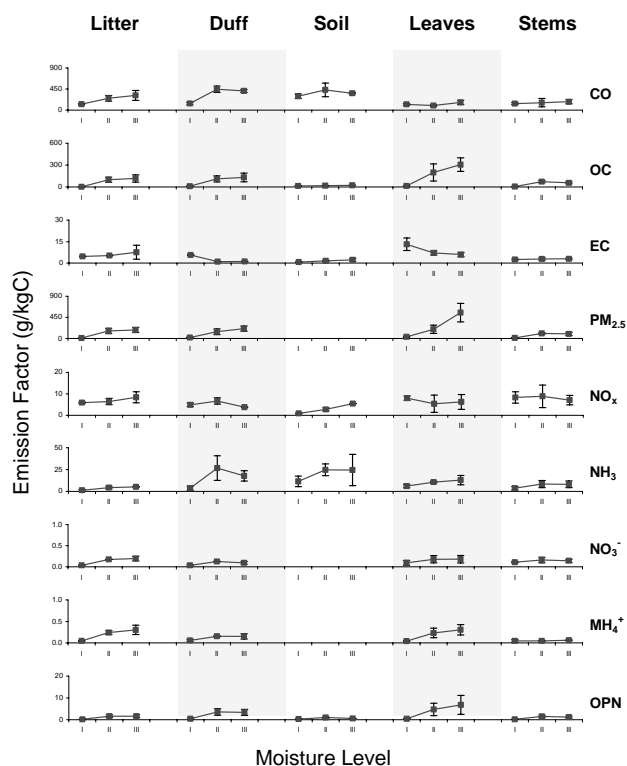


Fig. 3. Emission factors, by fuel type, as a function of fuel moisture level, based on data presented in Table 1 (all the leaf and stem data have been combined).

dependent absorption of particles. The absorption exponent (AE) was calculated by

$$AE = -\frac{\ln(ATN_{370}) - \ln(ATN_{880})}{\ln(370) - \ln(880)} \quad (1)$$

where ATN is the filter attenuation used as a surrogate for light absorption (Moosmüller et al., 2009). The increasing AE with higher OC/TC ratio (Fig. 4) shows that OC absorption is skewed toward shorter wavelengths than EC absorption, a principle characteristic of brown carbon (BrC). Ultraviolet (UV) absorption by such BrC can be comparable to that by black carbon (Kirchstetter et al., 2004). Though levoglucosan (C₆H₁₀O₅) is regarded as the main product from cellulosic decomposition during biomass burning and has been used as a marker for apportioning biomass burning contributions to ambient PM_{2.5} (Simoneit et al., 1999; Rinehart et al., 2006; Chow et al., 2007b), it is colorless. Formation of BrC from biomass burning must involve not only thermal decomposition but also oxidation and/or other reactions in the plumes.

3.2 Time-integrated nitrogen measurement

Lobert et al. (1990) suggested that NO_x accounts for most of fuel N detected in biomass burning plumes, followed by

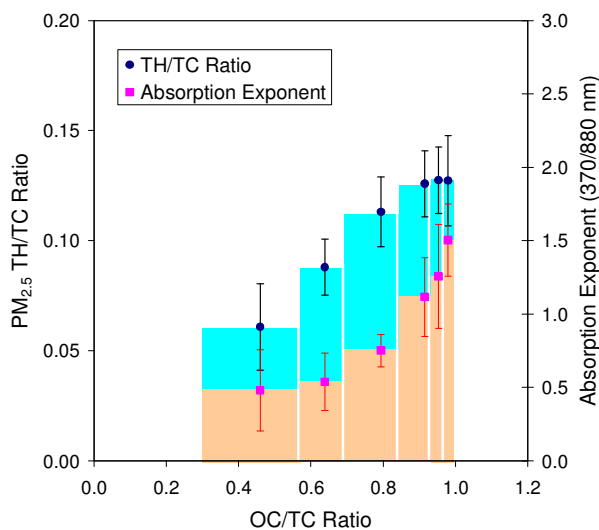


Fig. 4. Conditional means of TH/TC ratio and absorption exponent of biomass-burning $\text{PM}_{2.5}$ as a function of OC fraction in TC. The symbols correspond to six groups of samples with OC/TC ratio of 1) 0.3–0.575, 2) 0.575–0.7, 3) 0.7–0.85, 4) 0.85–0.94, 5) 0.94–0.97, and 6) 0.97–1. Soil combustion samples were excluded from this analysis.

NH_3 , hydrogen cyanide (HCN), N_2O , and nitriles. This is the case for some dry biomass (i.e., litter and stems) burns in this study (Table 1). In other cases NH_3 carries the most nitrogen, while N_2O is below the detection limit and HCN not quantified. Particulate N, which was not previously addressed in Lobert et al. (1990), is substantial from wet fuel combustion emissions. Only a small fraction of the particulate N can be attributable to water soluble NO_3^- and NH_4^+ . Other particulate nitrogen (OPN, particulate N minus N from NO_3^- and NH_4^+) EFs increase with fuel moisture content (Fig. 3) and are particularly high ($>2 \text{ g/kgC}$ and up to 75% of N in NH_3) when burning wet duff and leaves (see moisture level II and III in Table 1). The fact that OPN and to a lesser extent NH_3 EFs are anti-correlated with CE for all fuel types, including soil, indicates that they are favored products during smoldering combustion. This is not the case for NO_x .

It is possible that some N-containing gases, such as NH_3 , mixed with $\text{PM}_{2.5}$ in plumes and/or on filters and was measured as OPN. NH_3 adsorbed on quartz-fiber filters should be detected as NH_4^+ under the current procedure (i.e., water extraction/automated colorimetry) while adsorption of other gases seems minor compared to the amounts of OPN observed. In addition, OPN is highly correlated with OC ($r=0.93$; slope=0.022) for all biomass burns. This suggests that most OPN forms, along with OC, from the decomposition or pyrolysis of plant material and is in the form of organic nitrogen.

The nitrogen balance is evaluated by scaling N/GTC ratios in the combustion products to those in the fuels (Fig. 5). Bet-

ter closure for wet fuels results, for the most part, from higher NH_3 and OPN emissions. Nearly 100% of N was recovered from the wet litter composite and bitterbrush leaf burns. In other burns, however, substantial N was missing. Besides HCN and nitriles, Lobert et al. (1990) suggested that up to 50% of biomass N could end up becoming molecular nitrogen (N_2), which cannot be detected in open fires due to the large N_2 fraction of the ambient air. Combustion experiments conducted in a closed chamber supplied with helium/oxygen air showed dominant N_2 emissions from biomass burning, particularly during flaming combustion (Kuhlbusch et al., 1991). The mechanism of N_2 formation is unclear, but such a process is inhibited during the relatively low-temperature smoldering phase, leaving N available for other species.

3.3 Time-resolved measurement

Continuous data provide additional insights into the biomass combustion process. Temporally-resolved dry- and wet-fuel combustion emissions are compared in Fig. 6a–b using Manzanita leaves as an example. NH_3 concentrations in Fig. 6 (from TECO 17C) may be biased low as the averaged NH_3 for these two burns are $\sim 10\%$ lower than those measured on filters. They are only discussed qualitatively below.

The flaming phase for wet Manzanita leaves, consistent with the period with highly evaluated CO_2 concentrations, was brief ($\sim 1 \text{ min}$) relative to that of dry Manzanita leaves (3–4 min), though most of the CO_2 and NO_x were emitted during this phase. High temperatures in the flames provide sufficient thermal energy to break up plant organic matter into small fragments, producing CH_4 , VOCs, CO, NH_3 , etc., which are subsequently oxidized to CO_2 and NO_x . Dry fuels generate more intense flames, and therefore CO_2 and NO_x dominate the C and N species (except N_2) in the smoke. From the end of flaming phase through the smoldering phase, thermal energy decreases and higher fractions of CO, VOCs, and NH_3 are observed (e.g., Fig. 6a and Chen et al., 2007). For wet fuels, NH_3 could be the dominant N species throughout the burn (e.g., Fig. 6b).

To investigate the emission of particle mass and the dominant size fraction, real-time ELPI particle size and number measurements were converted to particle volume concentration ($\mu\text{m}^3/\text{cm}^3$) assuming spherical particles. The burn-averaged particle volume concentration correlates well with $\text{PM}_{2.5}$ mass measured on Teflon filters ($r=0.90$, across all the samples) with a mass/volume slope of $1.01 \pm 0.08 \text{ g/cm}^3$.

For the two Manzanita leaf burns in Fig. 6, the time-resolved mean particle volume (V_p in μm^3) was calculated from the ratio of particle volume and number concentrations. From V_p the particle volume mean (aerodynamic) diameter (D_v) was determined:

$$D_v = \left(\frac{6V_p}{\pi} \right)^{1/3} \quad (2)$$

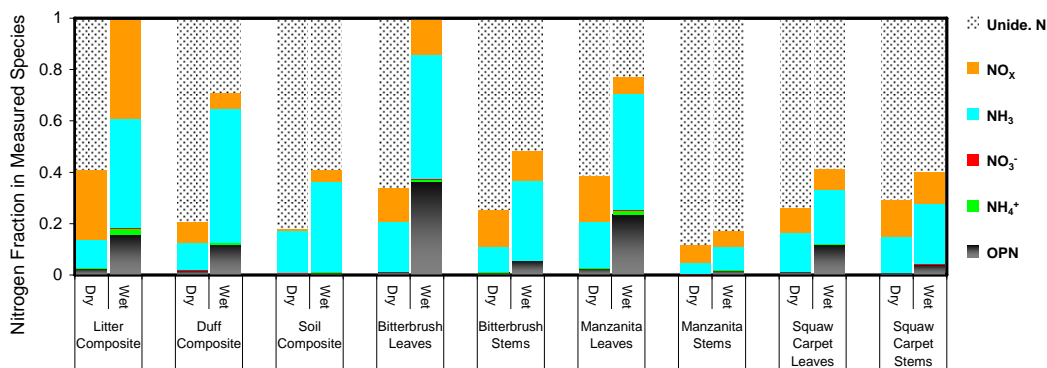


Fig. 5. Nitrogen balance with respect to fuel nitrogen content, determined from N/C ratios in smoke plumes and in fuels. Unidentified N (Unide. N) is the fuel N that is not accounted for by measured species. Dry and wet fuels correspond to the Moisture Level I and III in Table 1.

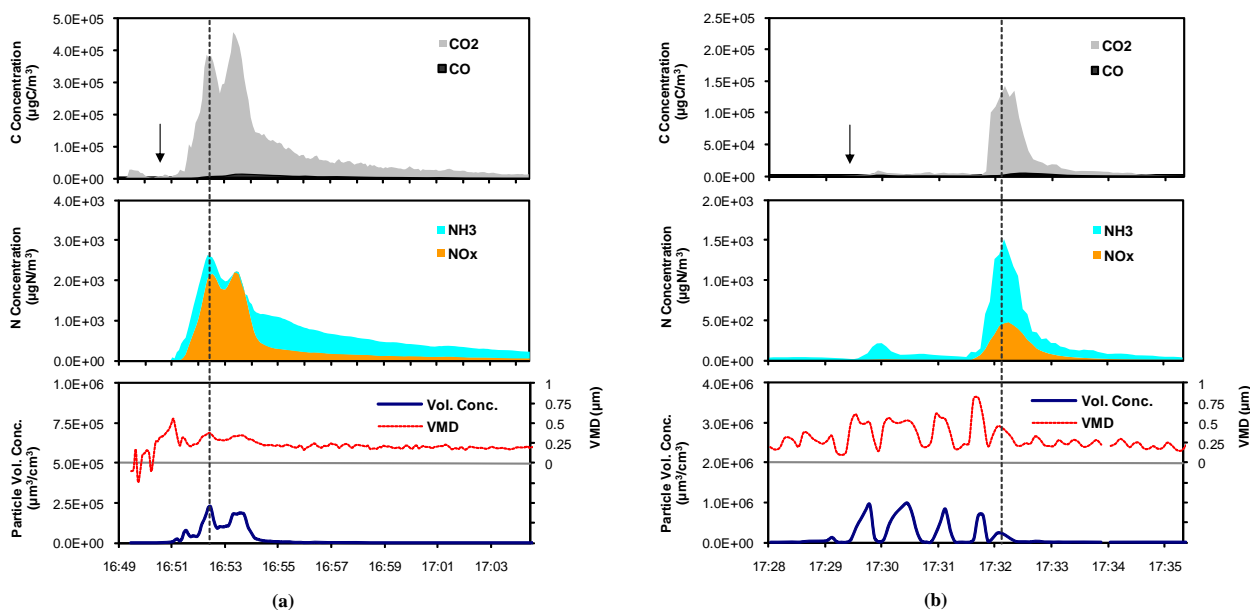


Fig. 6. Time series of gaseous (stacked area) and particulate concentrations during the burn of (a) dry and (b) wet Manzanita leaves. Particle size is represented by the volume mean diameter (VMD, i.e., D_v in the text). Arrows mark the start of hot-air igniter, and vertical dashed lines align the emission peaks.

Soot particles that mostly consist of EC are expected to form in the flames, and indeed peaks of particle volume concentration track those of CO_2 well in Fig. 6a. D_v at the peaks is estimated to be 0.34–0.38 μm . Larger D_v up to 0.56 μm are observed right before the flames start, which could be related to the initial breakdown of plant organic matter. These large particles, however, do not appear to account for substantial $\text{PM}_{2.5}$ volume or mass from the dry fuel combustion. Reid et al. (2005) and Guyon et al. (2003) reported typical values of D_v ranging from 0.25–0.3 μm for fresh smoke, but with some values as high as 0.5 μm .

Figure 6b shows NH_3 emissions, in the absence of NO_x emission, during a prolonged heating/smoldering period before the wet Manzanita leaves were ignited. Unlike Fig. 6a, the ELPI recorded intense particle emissions during this pre-flame period, and this explains the much higher OC and OPN EFs from the wet fuels. D_v that correspond to each of the pre-flame peaks range from 0.56 μm to 0.81 μm , with the largest D_v occurring right before the flame starts. These particles possibly consist of polyaccharides and/or its derivatives from the decomposition of cellulose. There is a smaller ELPI peak corresponding with the major CO_2 hump (flaming phase). Particle size at this peak appears to be smaller, i.e., $D_v=0.45 \mu\text{m}$, but is more consistent with flaming soot particles observed in Fig. 6a.

4 Conclusions

To study the moisture effect on biomass burning emission, litter, duff, soil, and aboveground shrub vegetation had been collected from an alpine forest, prepared in the laboratory for three moisture levels (dry, partially wetted, and fully wetted), and submitted to laboratory-controlled combustion experiments. These experiments show that fuel moisture lowers the overall combustion efficiency, shortens the flaming phase, and prolongs smoldering period before flames start. EFs of CO, OC, NH₃, and OPN increase with the fuel moisture content; the effect is generally larger for plant leaves and duff materials than for stems, litter, or soil. For high moisture leaves, emitted particulate OC can be as much as 4 times C in CO and OPN can be up to 75% of N in NH₃. OPN and OC are found to highly correlate ($r=0.93$).

CO₂ and CO emissions are mostly associated with flaming combustion and post-flame smoldering, respectively, and so are NO_x (flaming) and NH₃ (smoldering). N₂ emissions could be important especially during the flaming phase but cannot be quantified in open fires. The pre-flame smoldering of wet biomass mainly emits particles. These particles, with relatively large volume mean diameters (e.g., >0.55 μm), most likely consist of OC and OPN from decomposition of plant materials. It appears that higher thermal energy in flames is required to break the plant materials further into simpler C and N molecules, facilitating a full oxidation to CO₂ and NO_x. Particulate matter emissions associated with flaming combustion are relatively minor and smaller in size (e.g., 0.34–0.45 μm).

OC emitted from biomass burning, particularly with wet fuels, have TH/TC ratios similar to that of cellulose, the principle component of plant cells. However, the OC shows characteristics of BrC, i.e., absorbing UV much more strongly than cellulose. Though it is possible that polyaccharides and other derivatives of cellulose dominate the OC mass, how BrC is formed and its chemical nature warrant further investigations.

This study confirms that fuel moisture content is a critical factor controlling the C and N partitioning in biomass burning emissions. Since high-moisture fuels lead to substantial NH₃ and OPN emissions that have higher deposition rates, prescribed burning during wet seasons (e.g., spring in Lake Tahoe) would mitigate nutrient loss from long-range transport but increase the potential of nutrient deposition into the lake water. Distinct particle properties between low- and high-moisture fuel combustion imply different radiative effects of forest fires in dry and wet seasons. Moisture-dependent EFs and emission mechanisms represent an important step towards establishing accurate biomass burning emission inventories and source functions for global and/or regional climate models.

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