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aerosols in an aged
prescribed fire plume**

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Organic composition of carbonaceous aerosols in an aged prescribed fire plume

B. Yan¹, M. Zheng¹, Y. T. Hu², S. Lee¹, H. K. Kim², and A. G. Russell²

¹School of Earth and Atmospheric Sciences, Georgia Institute of Technology, Atlanta, GA, USA

²School of Civil and Environmental Engineering, Georgia Institute of Technology, Atlanta, GA, USA

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Correspondence to: A. G. Russell (ted.russell@ce.gatech.edu)

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Aged smoke from a prescribed fire (dominated by conifers) impacted Atlanta, GA on 28 February 2007 and dramatically increased hourly ambient concentrations of PM_{2.5} and organic carbon (OC) up to 140 and 72 $\mu\text{g m}^{-3}$, respectively. It was estimated that over 1 million residents were exposed to the smoky air lasting from the late afternoon to midnight. To better understand the processes impacting the aging of fire plumes, a detailed chemical speciation of carbonaceous aerosols was conducted by gas chromatography/mass spectrometry (GC/MS) analysis. Ambient concentrations of many organic species (levoglucosan, resin acids, retene, *n*-alkanes, *n*-alkanoic acids) associated with wood burning emission were significantly elevated on the event day. Levoglucosan increased by a factor of 10, while hopanes, steranes, cholesterol and major polycyclic aromatic hydrocarbons (PAHs) did not show obvious increases. Strong odd over even carbon number predominance was found for *n*-alkanes versus even over odd predominance for *n*-alkanoic acids. Alteration of resin acids during transport from burning sites to monitors is suggested by the observations. Our study also suggests that large quantities of biogenic volatile organic compounds (VOCs) and semivolatile organic compounds (SVOCs) were released both as products of combustion and unburned vegetation heated by the fire. Higher leaf temperature can stimulate biogenic VOC and SVOC emissions, which enhanced formation of secondary organic aerosols (SOA) in the atmosphere. This is supported by elevated ambient concentrations of secondary organic tracers (dicarboxylic acids, 2-methyltetrols, pinonic acid and pinic acid). An approximate source profile was built for the aged fire plume to help better understand evolution of wood smoke emission and can be used for source apportionment.

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1 Introduction

Wildland fire (wild fire and prescribed burning) is estimated to contribute about 20% of total fine particulate matter (PM_{2.5}) emissions in the United States (EPA, 2000). In 2006, a total of 96 385 wildland fires were reported to burn 39 958 square kilometers, 125% above the 10-year average (NIFC, 2007). Among these forests, 11 010 square kilometers were treated with prescribed fires, which is 1 659 square kilometers above last year's total and is the second highest since 1998 (NIFC, 2007). Such large and increasing emission contributions are of concern to air quality managers, particularly in areas near or above the applicable air quality standards. For example, the prescribed fires on 28 February 2007 in Georgia and later the Georgia-Florida wildfires lasting from April through May severely impacted Atlanta, Georgia with thick wood smoke (Hu et al., 2007¹; Lee et al., 2007). During such events, hourly concentrations of PM_{2.5} increased by over 100 μg m⁻³.

Previous studies have shown that the major component of PM_{2.5} from forest burning events is organic carbon (OC), accounting for 30–70% of PM_{2.5} mass (Nopmongcol et al., 2007; Robinson et al., 2004; Ward et al., 2006). However, detailed data speciating OC in PM_{2.5} impacted by wildland fires is sparse, and even less data is available for aged plumes. During the 28 February 2007 event, OC reached 72 μg m⁻³ at 06:00 p.m. and contributed approximately 51% of the ambient PM_{2.5} in Atlanta, GA (Lee et al., 2007). This smoke event provides an opportunity to characterize OC in such carbonaceous aerosols, further understand processes impacting the aging of fire plumes, and estimate the composition of prescribed fire-derived PM_{2.5} for source apportionment studies.

Numerous organic compounds have been identified and quantified from ambi-

¹Hu, Y. T., Odman, M. T., Chang, M. E., Jackson, W., Lee, S., Edgerton, E. S., Baumann, K., and Russell, A. G.: Simulation of Air Quality Impacts from Prescribed Fire on an Urban Area: Analysis of the 28 February 2007, Environ. Sci. Technol., in review, 2007.

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ent PM_{2.5} samples including *n*-alkanes, hopanes, steranes, alkanolic acids, alkanedioic acids, PAHs, resin acids, and others (syringols, levoglucosan, cholesterol, 2-methyltetrols, etc.). Some compounds are reasonably unique tracers for certain sources and are widely used to track specific sources of carbonaceous aerosols. Similar sets of organic species have also been measured for source emissions. Together, they can be used to quantify source impacts on ambient PM_{2.5}.

Source impacts from biomass burning are usually traced through a few organic tracers including levoglucosan, resin acids, syringols and retene. As a pyrolysis product of cellulose in wood biopolymers, levoglucosan has been considered a particularly useful molecular marker of biomass burning (Simoneit et al., 1999). With its large emission abundance and reasonable thermal stability in the atmosphere, levoglucosan is frequently used to assess air quality impacts from biomass burning (Fraser and Lakshmanan, 2000; Schauer and Cass, 2000). Resin acids are thermal alteration products of coniferous wood resins and emitted exclusively from softwood burning (various pines, firs, etc.) (Rogge et al., 1998; Simoneit et al., 1993; Standley and Simoneit, 1994). In contrast, hardwood combustion produces much higher quantities of syringols (Hawthorne et al., 1988; Hawthorne et al., 1989). Although PAHs are emitted from multiple combustion processes of fuels (biomass, natural gas, diesel and gasoline) and ubiquitous in the atmosphere, retene, a thermal alteration of abietane compounds (resin diterpenoids), is considered as an organic tracer specific for coniferous wood burning (Ramdahl, 1983).

Other primary sources can be also linked to some specific organic tracers. Hopanes and steranes are emitted from both gasoline-powered vehicle and diesel-powered vehicle (Simoneit, 1985; Zielinska et al., 2004). They have been widely used as molecular markers of vehicular emissions in source apportionment of PM_{2.5} and OC (Fraser et al., 2003b; Schauer et al., 1996; Zheng et al., 2002, 2006, 2007). Cholesterol, found in animal fats and oils, is thought as an excellent molecular marker of meat cooking emission (Rogge et al., 1991). Vegetative detritus emissions are characterized by high-molecular weight *n*-alkanes with pronounced odd over even carbon number

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predominance (Rogge et al., 1993a).

In addition to primary components of PM_{2.5}, secondary organic aerosol (SOA) formation can result from gaseous emissions of isoprenoids (isoprene and monoterpene)(Claeys et al., 2004a; Kavouras et al., 1998). A few biogenic SOA species have been identified and quantified including 2-methyltetrols (oxidation products of isoprene), pinonic acid and pinic acid (oxidation products of monoterpene) (Claeys et al., 2004a, b; Yu et al., 1999a, b). Although dicarboxylic acids (alkanedioic acids and dicarboxylic aromatic acids) can be emitted from various primary sources (mobile emission, meat cooking, etc.), previous studies suggested that atmospheric photochemical formation is probably the main source of these dicarboxylic acids (Fine et al., 2004b; Fraser et al., 2003a; Schauer et al., 2002).

Here, we collected PM_{2.5} filter samples before, during and after the 28 February 2007 prescribed fire episode impacting Atlanta, GA and analyzed organic composition of carbonaceous aerosols using GC/MS. We further capitalize on the quantified organic tracers to better understand the evolution of wood smoke from wildland fires.

2 Method description

2.1 Ambient sampling

Daily PM_{2.5} samples were collected on 47 mm quartz fiber filters with particulate composition monitors (PCM) at the Assessment of Spatial Aerosol Composition in Atlanta (ASACA) sites (Butler et al., 2003). These sites are located in the Atlanta metro area about 80 km downwind from the 28 February 2007 prescribed fires and were impacted directly by the smoke plume. The 24-h daily samples were analyzed for organic carbon/elemental carbon (OC/EC), metals and ions using thermal optical transmittance (TOT), X-ray fluorescence (XRF) and ion chromatography (IC), respectively. Details of ambient sampling, chemical measurements and particle compositions are described elsewhere (Baumann et al., 2003; Lee et al., 2005, 2007).

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2.2 Organic speciation

Due to the limitation of low air volumes sampled, the OC mass on a single PCM filter is usually not enough for organic tracer analysis. Therefore, three composite PCM samples (named “Before_Fire”, “Event” and “After_Fire”) were prepared as below. “Before_Fire” was composed of 6 PCM filters from three ASACA sites for the two days before the smoke day. “Event” was composed of 3 PCM filters collected at the same sites on 28 February, the day most directly impacted. “After_Fire” was composed of 7 PCM filters from the same sites within the three days after the smoke day. The three composite samples along with a composite field blank were analyzed for organic compounds in $PM_{2.5}$ using a standardized method described elsewhere (Nolte et al., 2002; Zheng et al., 2002, 2006). Briefly, each filter composite was spiked with deuterated internal standard (IS) mixtures and then successively extracted using hexane and benzene/isopropanol (2:1, v/v). After filtering, extracts were concentrated with rotary evaporation followed by blowdown under pure nitrogen. Half of each concentrated extract was then derivatized with diazomethane to convert organic acids to their methyl esters. These methylated extracts were analyzed by GC/MS along with authentic standards. To quantify a few polar organic compounds (levoglucosan, cholesterol, 2-methyltetrols), underivatized remains of concentrated extracts were silylated with BSTFA (N, O-bis(trimethylsilyl)acetamide) to convert polar compounds to trimethylsilyl (TMS) derivatives. After one hour reaction under $70^{\circ}C$, these silylated extracts were analyzed using GC/MS along with authentic standards.

3 Results and discussion

Our results show that the observed 24-h average OC concentration jumped from 3.5 to $17.7 \mu g m^{-3}$ on the event day, accounting for more than 60% (when converted to organic matter by a factor of 1.4) of the total 24-h average $PM_{2.5}$ mass of $37 \mu g m^{-3}$. Major sources of carbonaceous aerosols are traced using variations of their associated

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organic tracers.

3.1 Organic tracers of biomass burning

During the smoke, large increases were observed for biomass burning tracers (levoglucosan, resin acids, retene, etc.). Levoglucosan was detected in all ambient samples as the most abundant organic compound (Fig. 1). Before and after the smoke event, the observed levoglucosan concentrations were 114 and 145 ng m⁻³, respectively. On the event day, the levoglucosan concentration increased dramatically to 1210 ng m⁻³ and contributed 7% of the total OC, suggesting that the wood burning emission impact was 10 times higher on the event day than the non-smoke days. Along with levoglucosan, concentrations of resin acids also increased, especially dehydroabietic acid and 7-oxodehydroabietic acid, increasing to 42 and 19 ng m⁻³, approximately 9 and 23 times higher respectively than the levels before the burning day. Unlike levoglucosan, their ambient concentrations remained elevated after the fires (Fig. 2). To elucidate the processes occurring during transport, comparisons between wood burning source emissions and ambient data are conducted here. Ratios of major resin acids to levoglucosan were calculated and compared for both ambient data and a few source emissions from prominent softwood species in the southern United States (Table 1). In the softwood source emissions, abietic acid and dehydroabietic acid generally account for the majority of resin acids, about 58% and 32% respectively on average, while 7-oxodehydroabietic acid is minor. However, dehydroabietic acid and 7-oxodehydroabietic acid constituted a major fraction of the observed resin acids, about 65% and 29% respectively, in the ambient sample impacted by the fires. This comparison provides evidence that dehydroabietic acid and 7-oxodehydroabietic acid are being formed from other resin acids (i.e. diterpenoids) during the 3–4 h (around 80 km travel distance) transport from the burning sites to the monitors. Note that further alterations of dehydroabietic acid and 7-oxodehydroabietic acid exist, leading to their ratios to levoglucosan being variable over time during transport. Previous studies have also proposed that dehydroabietic acid and 7-oxodehydroabietic acid can be formed through

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oxidation processes of other resin acids (i.e. abietic acids), resulting in accumulated concentrations in the atmosphere (Oros and Simoneit, 2001; Rogge et al., 1993b). Although most PAHs did not show significant elevation in ambient concentrations during the event day, retene, a potential softwood burning tracer, increased by around 12 and 7 times, respectively compared to before and after the fire, and predominated among all PAHs (Fig. 2).

n-alkanes and *n*-alkanoic acids identified in this study ranged from C₁₇ to C₃₆ and from C₁₄ to C₃₀, respectively (Fig. 3). Generally, *n*-alkanes are associated with plant wax and fossil fuel contributions, depending on carbon numbers. To approximately compare source impacts from plant wax versus fossil fuel combustion, the carbon preference index (CPI) measuring carbon number predominance in homologous compound series is calculated (Mazurek and Simoneit, 1984; Oros et al., 2006). In the samples before and after the fire, *n*-alkanes only show a slight odd carbon number predominance with CPIs of 1.4 and 1.7, respectively. However, *n*-alkanes exhibited distinctly strong odd carbon number predominance (CPI=3.1, carbon number maximum C_{max}=29) on the event day, reflecting a major contribution from plant waxes. Likewise, *n*-alkanoic acids had a strong even carbon number predominance on the event day (CPI=5.7, C_{max}=24). The concentration of even-over-odd carbon number series on the event day was 50 ng m⁻³, much larger than those before and after the event (12 and 10 ng m⁻³, respectively). Thus, a dominant contribution from plant waxes is suggested. These compounds are considered natural products of epicuticular waxes and internal lipid substances in leaf surfaces and emitted as vegetative detritus or through direct thermal volatilization (Rogge et al., 1993a; Simoneit, 2002). Higher leaf temperatures presumably lead to volatilization of these high-molecular weight organic compounds, which can then condense into the particle phase in the atmosphere. This result indicates that such waxes are not unique tracers for vegetative detritus when wildland fires also significantly impact air quality.

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3.2 Other primary organic tracers

A few non-biomass organic molecular markers were also analyzed as indicators of other major primary sources. Vehicular engine exhaust and meat cooking are the two other major sources of primary organic carbon in urban areas. The before, during and after concentrations of hopanes and steranes did not show significant variation, i.e. 1.7, 1.7 and 1.0 ng m⁻³, indicating relatively consistent source contributions from vehicular emissions in the Atlanta area (Fig. 4). Cholesterol, an organic tracer of meat cooking, did not exhibit major variation as well (Fig. 5). Thus, it is clear that vehicular sources and meat cooking emissions are not responsible for the large increase in OC, which accounts for 81% of the total OC, on the event day, and that the greatly increased carbonaceous aerosol concentrations during the smoke episode are from prescribed fires. The source apportionment results calculated using organic molecular marker-based chemical mass balance (CMB-MM) model also indicate significant increases of prescribed burning emissions on the event day, but not for other major primary sources (i.e. vehicular source and meat cooking) (Lee et al., 2007). In addition, about 43% of the total observed OC can not be explained by primary source contributions, suggesting that secondary organic aerosol (SOA) might be a main contributor to the increased OC.

3.3 Secondary organic tracers

Our results for secondary organic tracers indicate that additional biogenic SOA were formed during the 28 February 2007 fire episode. 2-methyltetrols, secondary organic products of isoprene, were detected only on the smoke plume day with a concentration of 0.8 ng m⁻³ (Fig. 5). Pinonic acid and pinic acid, photo-oxidation products of monoterpenes, increased from 2.9 and 1.2 ng m⁻³ to 5.0 and 3.1 ng m⁻³, respectively, from the day prior to the fire through the smoke day (Fig. 5). Additionally, elevated dicarboxylic acids (alkanedioic acids and dicarboxylic aromatic acids) provided supportive evidence of increased SOA formation in the atmosphere. They are further

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considered here products of biogenic emissions enhanced by the fire since significant increases were not found in anthropogenic emissions (i.e. vehicular source and meat cooking). The sums of dicarboxylic acids were 7.5, 51, 8.4 ng m⁻³ before, during and after the event, respectively. There are three possible hypotheses relating to the large increases in these dicarboxylic acids. The most direct one is that precursors of these compounds are associated with prescribed fire emissions. As stated previously, emissions of biogenic VOC and SVOC (isoprenes, terpenes, sesquiterpenes, alcohols, esters, carbonyls, acids, etc.) would be enhanced by either wood combustion process or increasing leaf temperature. Emissions of isoprenoids (isoprene and monoterpene) have been observed to be higher during forest fires due to increased temperatures (Alessio et al., 2004). A previous study also suggests that leaf temperatures rising from 25 to 35°C would increase biogenic VOC and SVOC emissions by 4 and 1.5 times from isoprene-emitting deciduous trees and terpene-emitting conifers, respectively (Lamb et al., 1987). The second hypothesis is that the dramatic increase in OC mass can enhance formation of SOA due to increased partitioning of these diacids to the particle phase. However, unless there is a significant chemical affinity between OC and the compounds, such a shift is not expected to be so large (less than 10% of enhancement in SOA) (Nopmongcol et al., 2007; Odum et al., 1996). The third hypothesis is that enhanced photo-oxidation occurred. The observed increase of ozone in the plume is indicative that this effect might exist, but would be limited since ozone increased about 50% (from around 60 to 90 ppb), significantly less than the increase in dicarboxylic acids (8 times higher). Therefore, among the three hypotheses above, the evidence strongly supports that biomass burning is a major contributor to precursors of dicarboxylic acids. Results of air quality modeling and measurement of water soluble organic carbon (WSOC) provide further supporting evidence (Hu et al., 2007¹; Lee et al., 2007).

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3.4 Source profiles for an aged plume

Capturing the fire event provides information to assess the source composition profiles of an aged prescribed fire plume. Impacted by the fire plume, OC increased by $14.2 \mu\text{g m}^{-3}$, accounting for 81% of total observed OC on the event day while $\text{PM}_{2.5}$ increased by $23.9 \mu\text{g m}^{-3}$, accounting for 64% of total observed $\text{PM}_{2.5}$. Organic markers of other major primary emissions (vehicular source and meat cooking) did not show significant variation in Atlanta, GA during the episode. Thus, the large increases in OC, $\text{PM}_{2.5}$ and associated chemical species on the event day are considered a direct contribution from prescribed fire emissions. Here, two approximate source composition profiles were developed for the aged biomass burning plume by considering differences between the before_event day (non-fire event impacted) and the event day, designated “aged_plume profile” and “primary_plume profile”. “aged_plume profile” is constituted by fractions of individual chemical species (increased concentrations) in the total increased (fire-caused total) $\text{PM}_{2.5}$ mass, i.e. $23.9 \mu\text{g m}^{-3}$, on the event day (Table 2). This profile contains chemical compositions of the aged plume, but is not the primary $\text{PM}_{2.5}$ source profile due to large quantities of SOA in aerosol from the fire emissions. “primary_plume profile” is then calculated using fractions of individual chemical species (increased concentrations) in the fire-caused primary $\text{PM}_{2.5}$ mass where the estimated fire-caused secondary organic carbon (SOC) was subtracted from the fire-caused total $\text{PM}_{2.5}$ mass on the event day (Table 2). The fire-caused SOC on the event was estimated based on the CMB-MM modeling results and measured EC/OC ratios. Briefly, the CMB-MM apportionment results indicate that prescribed fires contributed approximately 1.1 and $8.7 \mu\text{g m}^{-3}$ to the total primary OC before and on the event day, respectively (Lee et al., 2007). The difference, $7.6 \mu\text{g m}^{-3}$, is attributed to the plumes. Results here find that the total, primary plus secondary, impact of the plumes on OC is $14.2 \mu\text{g m}^{-3}$, suggesting that $6.6 \mu\text{g m}^{-3}$ (47% of the fire-caused total OC) comes from enhanced SOC. A comparable SOC fraction in the fire-caused total OC was also estimated using the EC/OC ratio method. In the aged plume, the fire-caused EC/OC ratio

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is 0.039 while a value of 0.065 has been measured for prescribed burning emissions in Georgia (Lee et al., 2005). The lower EC/OC ratio suggests SOA formation in the aged plume. Using those ratios, SOC is calculated to account for 40% of the fire-caused total OC. Averaging the two, 44% of fire-caused total OC is taken as SOC during the event, and the fire-caused primary OC (POC_{fc}) is then estimated by:

$$POC_{fc} = OC(\text{fire-caused}) \times (1 - f_{SOC}) = [OC(\text{event}) - OC(\text{before-event})] \times (1 - f_{SOC}), \quad (1)$$

where f_{SOC} is the estimated SOC fraction in the fire-caused total OC. Similarly, the amount of fire-caused primary $PM_{2.5}$ (PPM_{fc}) is calculated as:

$$PPM_{fc} = [PM(\text{event}) - PM(\text{before-event})] - OC(\text{fire-caused}) \times f_{SOC}. \quad (2)$$

Finally, fractions of individual species (f_i) in fire-caused primary $PM_{2.5}$ are found as:

$$f_i = \frac{C_i(\text{event}) - C_i(\text{before-event})}{PPM_{fc}} \quad (3)$$

where C_i is the ambient concentration of individual chemical species in $PM_{2.5}$ that are viewed as being dominated by primary emissions. In addition, an overall uncertainty of each chemical species in the two plume source profiles was calculated by propagating uncertainties associated with the observed OC, estimated SOC fraction and measured chemical species (i.e. organic compounds, EC, ions and trace metals).

Our study indicates that the “primary plume profile” derived from the fire event is comparable with the prescribed burning emission profile measured by Lee et al. (2005) for some organic compounds, but significantly different for others (Table 2 and Fig. 6). In the aged fire plume, levoglucosan accounts for 14% of primary OC, comparable to 9.5% in the profile of Lee et al. (2005). Most *n*-alkanes also show comparable levels for the two source profiles. Hopanes and steranes were not significant in either set. The ratio of water-soluble potassium (K^+)/OC in the “primary plume profile” is 0.0067, lower than the values in Lee et al. (0.011) and in Fine et al. (0.012 on average), but comparable to the average ratio of 0.0058 from foliar fuel combustion (Fine et al., 2004a; Hays et

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al., 2002; Lee et al., 2005). However, significant differences were found for the fractions of PAHs, resin acids, *n*-alkanoic acid and dicarboxylic acids, suggesting aging of fire smoke after a 3–4 h travel distance. Lee et al. (2005) measured higher abundances for many PAHs and resin acids, implying direct combustion is the main contributor of these compounds and significant chemical alteration occurs during transport, especially for resin acids. In contrast, the aged plume has distinctly higher dicarboxylic acids (alkanedioic acids and dicarboxylic aromatic acids) fractions. Enhanced SOA formation in the atmosphere from the fire is suggested by these secondary indicators. The profile comparison between prescribed fire source emission and aged plume is useful to understand evolution of wood smoke during transport and to assess the organic species applied in CMB-MM modeling. Impacts from aging process of smoke plume should be considered in source apportionment methodology using CMB model, i.e. selection of fitting species.

4 Conclusions

Large increases in PM_{2.5} and OC in Atlanta caused by an aged prescribed fire plume through direct burning emissions and formation of secondary organic aerosols was captured by a system of monitors. Organic tracers of biomass burning, levoglucosan, resin acids and retene, exhibited sharp increases in concentrations during the episode. Observed resin acids and retene indicated predominant softwood (conifers) burning. Increases in resin acids were accompanied by high levels of dehydroabietic acids and 7-oxodehydroabietic acids, mainly formed during transport. Carbon number predominance for *n*-alkanes (odd over even) and *n*-alkanoic acids (even over odd) suggested that emissions from heat-exposed vegetation are important as one of major sources for increasing OC, especially secondary OC. Secondary organic tracers were observed with significantly higher ambient concentrations, providing further support for biogenic SOA formation. These data allows estimating source profiles for aged fire plumes, which can be used for improving source apportionment

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of wood burning by CMB modeling <http://www.atmos-chem-phys-discuss.net/7/18015/2007/acpd-7-18015-2007-supplement.pdf>.

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Table 1. Ratios of major resin acids to levoglucosan in source emissions and ambient samples.

Compounds	Softwood Emissions					Ambient Samples		
	Loblolly Pine ^a	Slash Pine ^a	White pine ^b	Hemlock ^b	Balsam fir ^b	Before_Fire	Smoke	After_Fire
Pimaric acid	0.069	0.029	0.008	0.001	0.001	0.001	0.000	0.002
Sandaracopimaric acid	0.013	0.011	0.026	0.002	0.001	0.003	0.001	0.011
Dehydroabietic acid	0.339	0.141	0.149	0.017	0.028	0.041	0.034	0.248
Abietic acid	0.801	0.056	0.391	0.021	0.240	0.001	0.000	0.002
Abieta-6,8,11,13,15-pentae-18-oic acid	0.007	0.003	0.005	0.000	0.002	0.000	0.000	0.000
Abieta-8,11,13,15-tetraen-18-oic acid	0.021	0.000	0.010	0.001	0.003	0.001	0.001	0.001
7-Oxodehydroabietic acid	0.009	0.004	0.005	0.001	0.000	0.007	0.015	0.042

^a Softwood species tested by Fine et al. (Fine et al., 2002);

^b Softwood species tested by Fine et al. (Fine et al., 2001).

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Table 2. Source composition profiles from the aged plume on the event and the previous prescribed burning emission (ng/μg PM_{2.5}).

Compound	Aged Plume ^a	PPM _{fc} ^b	Lee et al. ^c	Compound	Aged Plume ^a	PPM _{fc} ^b	Lee et al. ^c
	Fraction ^d ±std ^e	Fraction±std	Fraction±std		Fraction±std	Fraction±std	Fraction±std
<i>n</i> -alkanes							
Tetracosane	0.0375±0.0208	0.0508±0.0284	0.0836±0.0528	Triacotane	0.0451±0.0116	0.0610±0.0164	0.1168±0.0715
Pentacosane	0.0706±0.0294	0.0956±0.0404	0.1506±0.0954	Hentriacotane	0.2561±0.0731	0.3467±0.1020	0.1737±0.1054
Hexacosane	0.0225±0.0159	0.0305±0.0217	0.1374±0.0913	Dotriacotane	0.0292±0.0075	0.0395±0.0106	0.0000±0.0001
Heptacosane	0.1728±0.0497	0.2339±0.0693	0.1505±0.1137	Tritriacotane	0.0527±0.0156	0.0713±0.0217	0.0542±0.0480
Octacosane	0.0470±0.0171	0.0637±0.0236	0.0695±0.0416	Tetrtiacotane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
Nonacosane	0.3115±0.0881	0.4217±0.1231	0.4860±0.2950				
Branch-alkanes							
iso-Nonacosane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	iso-Hentriacotane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
anteiso-Triacotane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001				
Hopanes							
17α(H)-21β(H)-29-Norhopane	0.0022±0.0051	0.0029±0.0069	0.0000±0.0001	22R,17α(H),21β(H)-Homohopane	0.0018±0.0021	0.0025±0.0029	0.0000±0.0001
17α(H)-21β(H)-Hopane	0.0016±0.0047	0.0021±0.0064	0.0000±0.0001	22S,17α(H),21β(H)-Bishomohopane	0.0009±0.0013	0.0012±0.0017	0.0000±0.0001
22,29,30-Trisnormehopane	0.0022±0.0020	0.0029±0.0028	0.0000±0.0001	22R,17α(H),21β(H)-Bishomohopane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
22,29,30-Trisnorhopane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	22S,17α(H),21β(H)-Trishomohopane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
22S,17α(H),21β(H)-Homohopane	0.0028±0.0028	0.0038±0.0038	0.0000±0.0001	22R,17α(H),21β(H)-Trishomohopane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
Steranes							
20S,R-5α(H),14β(H),17β(H)-Cholestanes	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	20S,R-5α(H),14β(H),17β(H)-Ergostanes	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
20R-5α(H),14α(H),17α(H)-Cholestane	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	20S,R-5α(H),14β(H),17β(H)-Sitostanes	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
PAHs							
Fluoranthene	0.0000±0.0001	0.0000±0.0001	0.0539±0.0310	Benzo(j)fluoranthene	0.0004±0.0002	0.0006±0.0003	0.0077±0.0049
Acephenanthrylene	0.0009±0.0004	0.0012±0.0005	0.0168±0.0105	Benzo(e)pyrene	0.0039±0.0024	0.0053±0.0033	0.0300±0.0168
Pyrene	0.0000±0.0001	0.0000±0.0001	0.0643±0.0363	Benzo(a)pyrene	0.0006±0.0016	0.0008±0.0022	0.0179±0.0100
Retene	0.0284±0.0077	0.0385±0.0108	0.2102±0.1267	Perylene	0.0000±0.0001	0.0000±0.0001	0.0023±0.0015
Benzo(ghi)fluoranthene	0.0018±0.0018	0.0025±0.0025	0.0585±0.0517	Indeno(1,2,3-cd)fluoranthene	0.0024±0.0011	0.0033±0.0015	0.0000±0.0001
Cyclopenta(cd)pyrene	0.0000±0.0001	0.0000±0.0001	0.1740±0.1058	Indeno(1,2,3-cd)pyrene	0.0028±0.0025	0.0038±0.0034	0.0444±0.0265
Benzo(a)anthracene	0.0006±0.0017	0.0008±0.0023	0.0482±0.0268	Picene	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001
Chrysene/Triphenylene	0.0071±0.0041	0.0097±0.0056	0.0589±0.0332	Benzo(ghi)perylene	0.0026±0.0041	0.0036±0.0056	0.0215±0.0120

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Table 2. (Continued.)

Compound	Aged Plume ^a	PPM _{TC} ^b	Lee et al. ^c	Compound	Aged Plume ^a	PPM _{TC} ^b	Lee et al. ^c
	Fraction ^d ± std ^o	Fraction ± std	Fraction ± std		Fraction ± std	Fraction ± std	Fraction ± std
PAHs							
Benzo(b)fluoranthene	0.0043±0.0028	0.0059±0.0039	0.0304±0.0170	Coronene	0.0039±0.0037	0.0053±0.0051	0.0044±0.0031
Benzo(k)fluoranthene	0.0028±0.0019	0.0038±0.0026	0.0245±0.0136				
Resin Acids							
8,15-Pimaredienoic acid	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	Abietic acid	0.0112±0.0043	0.0152±0.0059	0.0000±0.0001
Pimaric acid	0.0185±0.0055	0.0251±0.0077	1.4856±0.8256	Abieta-6,8,11,13,15-pentae-18-oic acid	0.0088±0.0025	0.0120 ± 0.0036	0.0000±0.0001
Sandaracopimaric acid	0.0558±0.0169	0.0756±0.0235	0.0000±0.0001	Abieta-8,11,13,15-tetraen-18-oic acid	0.0263±0.0072	0.0356±0.0101	0.0000±0.0001
Isopimaric acid	0.0000±0.0001	0.0000±0.0001	1.7830±0.9994	7-Oxodehydroabietic acid	0.7430±0.1978	1.0058±0.2773	0.0000±0.0001
Dehydroabietic acid	1.5456±0.4331	2.0923±0.6051	20.073±11.450				
Aromatic Acids							
1,2-Benzenedicarboxylic acid	0.1185±0.0359	0.1604±0.0499	0.0088±0.0062	1,3-Benzenedicarboxylic acid	0.0286±0.0112	0.0387±0.0154	0.0000±0.0001
1,4-Benzenedicarboxylic acid	0.0436±0.0207	0.0591±0.0284	0.0054±0.0047				
Alkanoic Acids							
Tetradecanoic acid	0.0741±0.0248	0.1004±0.0343	1.2089±0.7432	Tricosanoic acid	0.1450±0.0392	0.1963±0.0548	0.3941±0.2273
Pentadecanoic acid	0.0359±0.0135	0.0486±0.0187	0.3838±0.2446	Tetracosanoic acid	0.5537±0.1509	0.7496±0.2112	2.9835±1.7399
Hexadecanoic acid	0.2784±0.1320	0.3769±0.1808	4.0548±2.3375	Pentacosanoic acid	0.0488±0.0133	0.0660±0.0186	0.2308±0.1332
Heptadecanoic acid	0.0207±0.0077	0.0280±0.0106	0.1494±0.0907	Hexacosanoic acid	0.3011±0.0811	0.4077±0.1136	2.2120±1.2819
Octadecanoic acid	0.1343±0.0642	0.1818±0.0879	1.3508±0.8111	Heptacosanoic acid	0.0172±0.0046	0.0233±0.0064	0.0850±0.0506
Nonadecanoic acid	0.0262±0.0075	0.0355±0.0105	0.2941±0.1870	Octacosanoic acid	0.1031±0.0276	0.1395±0.0387	0.5953±0.3501
Eicosanoic acid	0.1761±0.0487	0.2363±0.0681	0.7198±0.4371	Nonacosanoic acid	0.0216±0.0056	0.0292±0.0078	0.1014±0.0647
Heneicosanoic acid	0.0740±0.0202	0.1002±0.0283	0.1739±0.1009	Triacontanoic acid	0.1119±0.0299	0.1516±0.0419	0.4837±0.3137
Docosanoic acid	0.3794±0.1031	0.5136±0.1444	1.0757±0.6382				
Alkenoic Acids							
9-Hexadecenoic acid	0.0000±0.0001	0.0000±0.0001	0.3058±0.2272	9-Octadecenoic acid	0.0000±0.0001	0.0000±0.0001	1.4690±0.8808
9,12-Octadecanedienoic acid	0.0000±0.0001	0.0000±0.0001	1.3574±0.9208				
Alkanedioic Acids							
Propanedioic acid	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	Hexanedioic acid	0.0882±0.0264	0.1195±0.0367	0.0159±0.0132
Methylpropanedioic acid	0.0000±0.0001	0.0000±0.0001	0.0000±0.0001	Heptanedioic acid	0.0000±0.0001	0.0000±0.0001	0.0180±0.0159
Butanedioic acid	0.8009±0.2267	1.0842±0.3165	0.2368±0.1533	Octanedioic acid	0.1487±0.0409	0.2013±0.0572	0.0358±0.0280

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Table 2. (Continued).

Compound	Aged Plume ^a	PPM _{TC} ^b	Fraction \pm std ^d	Lee et al. ^c	Compound	Aged Plume ^a	PPM _{TC} ^b	Fraction \pm std ^d	Lee et al. ^c
Alkanedioic Acids									
Methylbutanedioic acid	0.1716 \pm 0.0470	0.2323 \pm 0.0658	0.0000 \pm 0.0001	0.0000 \pm 0.0001	Nonanedioic acid	0.2986 \pm 0.0835	0.4043 \pm 0.1167	0.1558 \pm 0.1006	
Pentanedioic acid	0.1198 \pm 0.0347	0.1622 \pm 0.0484	0.0489 \pm 0.0317						
Others									
Nonanal	0.0125 \pm 0.0088	0.0170 \pm 0.0120	0.0000 \pm 0.0001	0.0000 \pm 0.0001	Benz(de)anthracen-7-one	0.0036 \pm 0.0023	0.0049 \pm 0.0032	0.0000 \pm 0.0001	
Sinapyl aldehyde	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	3,5-Dimethoxy-4-hydroxyacetophenone	0.0275 \pm 0.0071	0.0372 \pm 0.0100	0.0000 \pm 0.0001	
Acetonylsyringol	0.0581 \pm 0.0181	0.0786 \pm 0.0252	0.0000 \pm 0.0001	0.0000 \pm 0.0001	Levoglucosan	45.911 \pm 12.692	62.151 \pm 17.746	57.087 \pm 32.377	
Coniferyl aldehyde	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	Cholesterol	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	
Propionylsyringol	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.0000 \pm 0.0001					
OC	593.85 \pm 180.15	450.19 \pm 250.55	602.50 \pm 185.20	Cl ⁻	0.4355 \pm 0.0898	0.5895 \pm 0.1287	5.2700 \pm 2.8900		
EC	22.962 \pm 12.421	31.085 \pm 16.961	39.200 \pm 11.300	NH ₄ ⁺	13.119 \pm 4.4281	17.760 \pm 6.1272	1.0700 \pm 1.0800		
Al	0.0000 \pm 0.0001	0.0000 \pm 0.0001	0.2290 \pm 0.4260	NO ₃ ⁻	19.310 \pm 5.1844	26.141 \pm 7.2624	4.4000 \pm 2.9900		
Si	0.2889 \pm 0.2276	0.3911 \pm 0.3094	0.1860 \pm 0.2580	SO ₄ ²⁻	13.800 \pm 8.5722	18.682 \pm 11.680	2.4500 \pm 1.1200		
K ⁺	2.2261 \pm 0.5044	3.0135 \pm 0.7159	6.4900 \pm 4.3500						

^a source composition profiles where individual chemical species are normalized to the fire-caused total PM_{2.5} mass on the event day;

^b source composition profiles where individual chemical species are normalized to the fire-caused primary PM_{2.5} mass estimated on the event day;

^c source composition profiles from the prescribed burning emission in Georgia (Lee et al., 2005);

^d fraction of chemical species in the associated PM_{2.5} mass (ng/ μ g PM_{2.5});

^e standard deviation.

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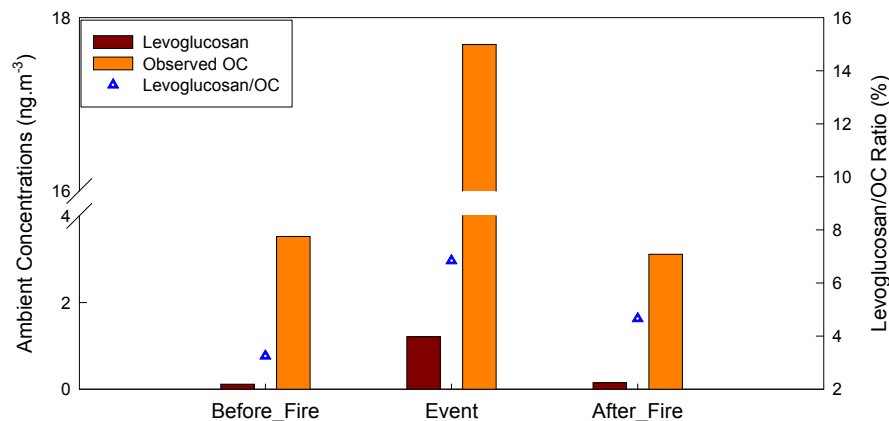


Fig. 1. 24-h average ambient concentrations of OC and levoglucosan observed before, during and after the event day at Atlanta area, GA.

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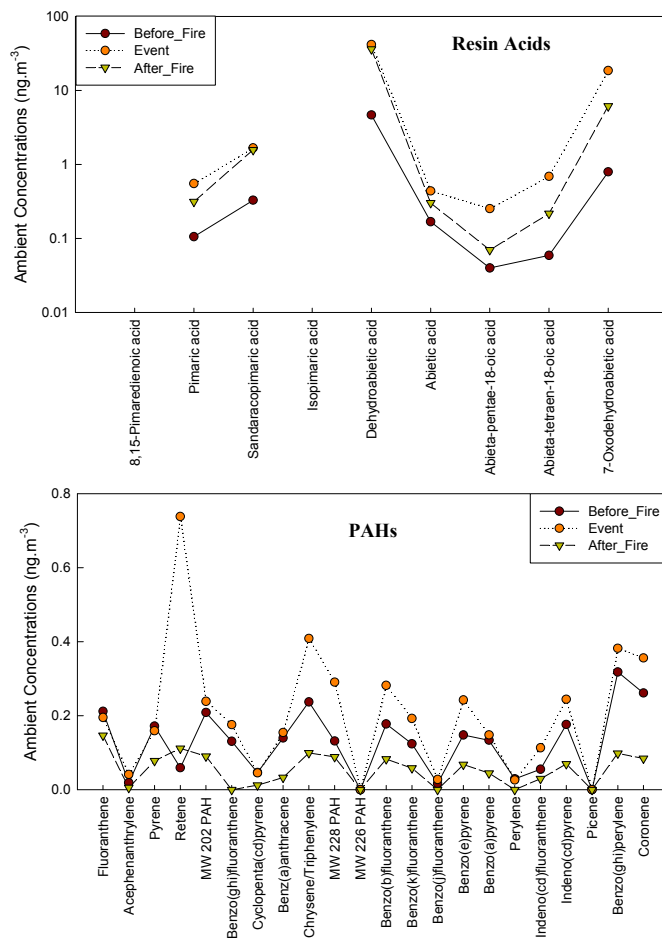


Fig. 2. 24-h average concentrations of resin acids and PAHs observed before, during and after the event day at Atlanta area, GA.

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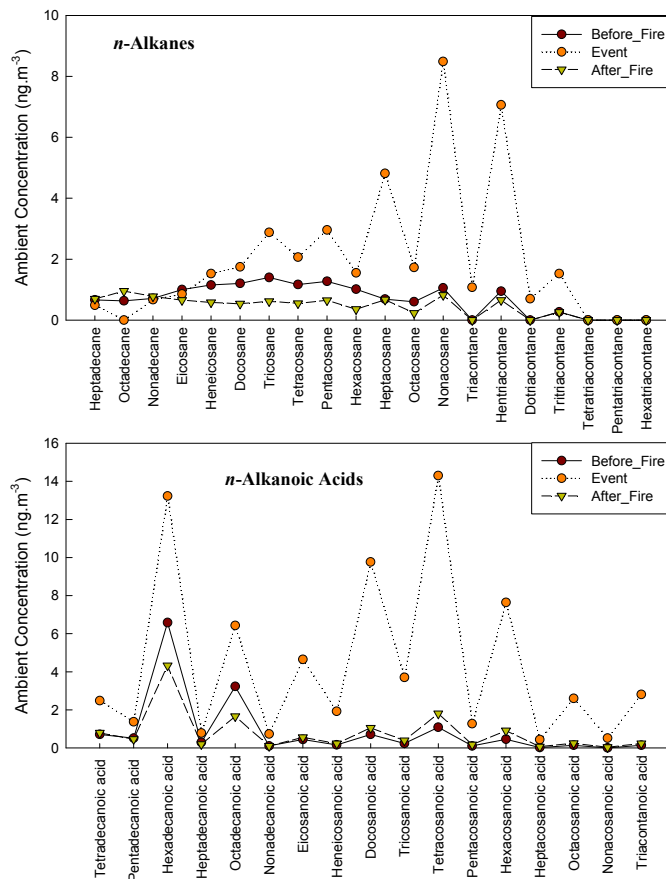


Fig. 3. 24-h average concentrations of *n*-alkanes and *n*-alkanoic acids observed before, during and after the event day at Atlanta area, GA.

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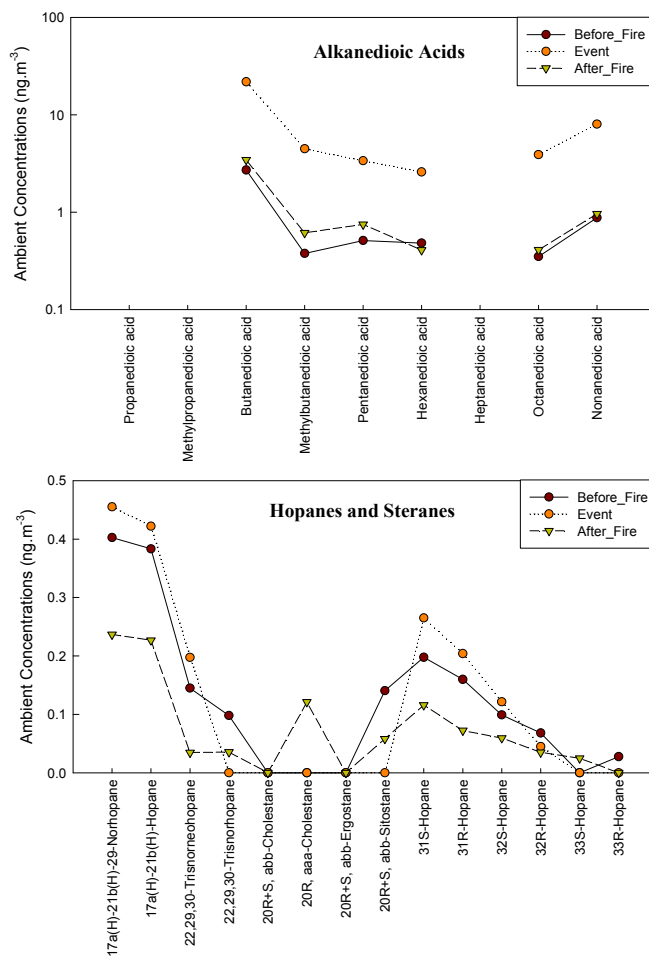


Fig. 4. 24-h average concentrations of alkanedioic acids, hopanes and steranes observed before, during and after the event day at Atlanta area, GA.
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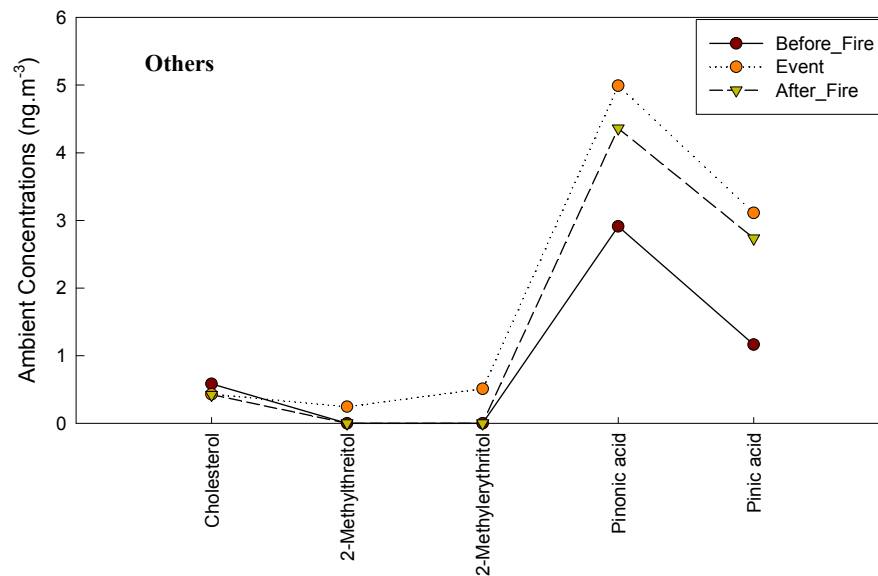


Fig. 5. 24-h average concentrations of other compounds (cholesterol, 2-methyltetrols, pinonic acid and pinic acid) observed before, during and after the event day at Atlanta area, GA.

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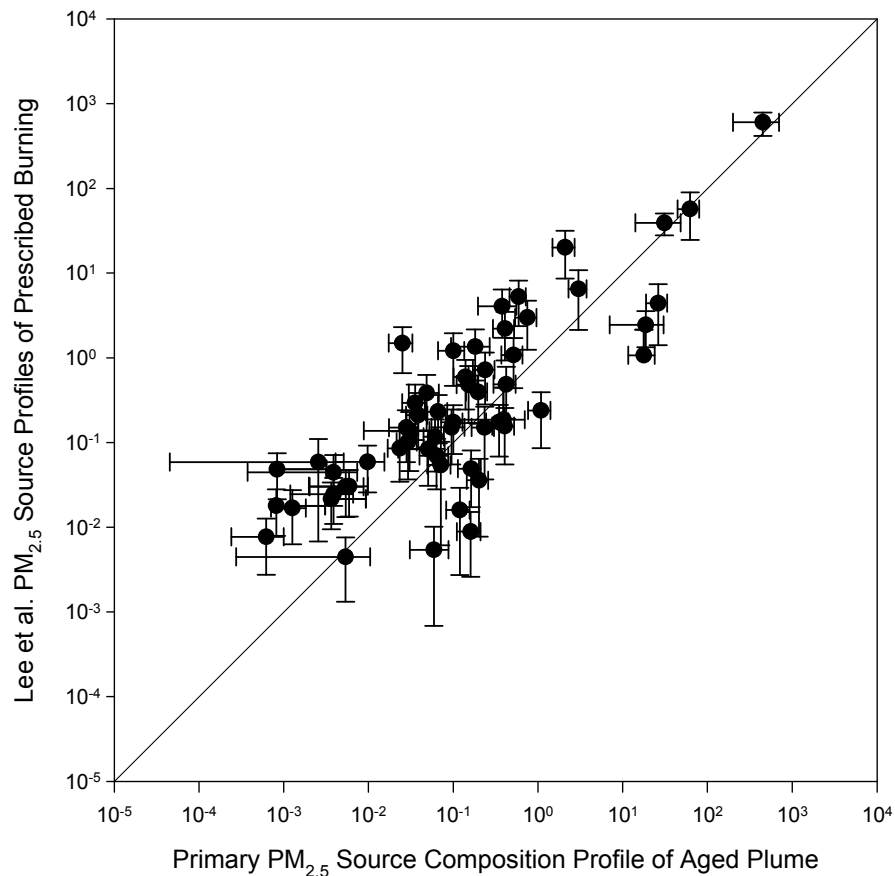


Fig. 6. Comparison between source composition profiles developed from the aged biomass burning plume on the event and the prescribed burning emission measured by Lee et al. (2005). The profiles are composed of individual species fractions, normalized to primary PM_{2.5} mass. A log-log scale is used.

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