Response to Reviewer #1

We thank Reviewer #1 for their helpful comments and suggestions. The original comments are provided below in gray, and our responses, with specific revisions, noted in **bold** font.

Aerosol composition measurements from an Aerodyne AMS and single-particle ATOFMS are presented for approximately 10 days at a rural ground site in Northern Michigan. Observations and HYSPLIT back trajectories suggest the site was influenced by a mixture of "background", wildfire, and urban sources. Secondary organic aerosol was the dominant aerosol species by mass regardless of source region, and the majority of particles throughout the campaign were identified as having originated from biomass burning. While the ubiquitous nature of biomass burning in the region is interesting, the paper might benefit from additional analysis to overcome the shortfalls of having such a short time series. The paper is certainly appropriate for ACP and should be published after satisfying the following critiques.

Major Comments:

What metric is being used to separate the wildfire and "regional background" time periods? Figure 1 would suggest that ozone and PM2.5 mass/number concentrations are of similar magnitude between the two sources. Likewise, the relative compositional breakdown is identical in Figure 3. I believe it is a bit misleading to label one of these times as "regional background" when they both seem equally influenced by diffuse smoke. If the authors have used a quantitative metric for isolating these sources, please highlight this method in the revised text. The similarity in back trajectories does not seem sufficient from Figure S2. I suggest only presenting "urban" and "background" sources.

Wildfire and regional background periods were differentiated by the presence of smoke at the field site as determined by the NOAA HMS smoke maps, shown in Figure 4 and described on P11 L2-4. We clarified this during the regional background section (Section 3.2) on P9 L13-15, which now states "This remote background air mass period was differentiated from the wildfire influenced periods (Section 3.3) based on the lack of smoke impacting the site, as indicated by NOAA Smoke and Fire products (Figure 4)." While average ozone is similar, average PM_{2.5} mass and number concentrations are significantly different between wildfire and regional background time periods as indicated by a t-test (P11 L 7-10). In addition, chemically-resolved ATOFMS mass concentrations (Figure 5) show lower biomass burning aerosol mass concentrations during the regional background period, compared to the rest of the study, when smoke influence was detected by the NOAA HMS smoke maps.

Ozone mixing ratios are quite low throughout the measurement period, as low as 10-20 ppb during background and wildfire periods. This seems anomalous for summer- time North America. Please provide more context/validation for these measurements, especially with respect to your conclusion that the region was frequently influenced by transported smoke.

The range in ozone concentrations (9-63 ppb) are similar to previous measurements conducted at UMBS during background periods (~10-90 ppb) (Cooper et al., 2001); this statement was added to P10 L2-3. While ozone increased significantly with biomass burning influence during the first wildfire period, ozone levels were only slightly above background

during the second period, which also featured lower biomass burning aerosol concentrations (Figures 1 and 5). It should be noted that previous studies have shown that ozone does not always increase within wildfire plumes (P11 L17-19) (Jaffe and Wigder, 2012).

Your result of persistent biomass burning influence is somewhat in contrast to previous reports that biogenic emissions play a role in SOA production (and thus aerosol mass concentrations), e.g., Sheesley et al. (2004). Has something fundamentally changed in the transport pathways to the region, or can these differences be attributed to the VERY short duration of observations presented here?

Summer 2014 was unique in that it was one of the worst (largest) Canadian wildfire seasons in the past 25 years (P15 L17-19); therefore, increased influence from biomass burning would be expected. Wildfire intensity and frequency are increasing due to climate change (Gillett et al., 2004; Knorr et al., 2016; Liu et al., 2010; Veira et al., 2016). We added a statement noting this on P16 L 10-13, which states "The observed levels of biomass burning aerosol influence are attributed to the abnormally active Canadian wildfire season of 2014, compared to previous typical summers in northern Michigan with primarily biogenic SOA influence (Sheesley et al., 2004)."

The AMS and ATOFMS have very different sampling capabilities with respect to particle size. This is evident in the vastly different mass concentrations presented in Figure 3 and Figure 5. Can you comment on comparisons between the two measurements, in the context of measured size distributions shown in Figure S4. Do you think the number-based prevalence of biomass burning is consistent with AMS results given that you are only assessing the tail of the mass distribution?

The NOAA HMS smoke maps show satellite-based detection of smoke over the field site (Figure 4), and this is consistent with the abundance of biomass burning aerosols detected by the ATOFMS. We now indicate the AMS and ATOFMS size ranges in Figure S4, which shows the measured aerosol size distributions, to illustrate the overlap between the instruments and with the aerosol population. The mode diameter of biomass burning aerosols upon emission is typically between 100-180 nm (Reid et al., 2005); therefore, based on the full mass-based size distributions in Figure S4, it would be expected that the majority of the mass corresponds to biomass burning particles coated with SOA. Also, Figure 5 shows that biomass burning aerosols dominated the 0.5-2.0 μ m aerosol mass, as measured by ATOFMS.

Rationale for this work is attributed to model under-prediction of OC in the Great Lakes region (i.e., in the abstract line-4), but the source of SOA is not entirely clear from the paper as written. The real question that I don't think is answered with this study is whether the precursor organic material contributing to the significant SOA mass coating biomass burning cores originated from the fires or from more-local biogenic sources encountered during transport. Can AMS (or ATOFMS) mass spectra help assess the SOA source? AMS source apportionment often utilizes the positive matrix factorization (PMF) technique to answer this sort of question and might provide very useful insight into SOA formation processes that could inform the model/measurement disconnect.

As stated in the response to Reviewer #1, while we agree that additional HR - AMS analysis would further support our findings, such analyses are unfortunately beyond the scope of what is feasible at this current time due to changes in personnel appointments following the field measurements. Therefore, we are unable to further apportion the SOA using the AMS data. However, we added discussion (P12 L17-19) of a previous SOA source apportionment study by Sheesley et al. (2004) who used molecular tracers, showing that the majority of summertime SOA in northern Michigan is from biogenic sources. Given the significant wildfire influence during our study, it is plausible that wildfire VOC precursors also contributed to SOA production.

The buildup of urban emissions culminating in, presumably, a frontal passage at night on 7/22 is an interesting aspect of the sampling period. During this time, it seems that sulfate aerosol is potentially being produced during transport and, as you state, aerosol properties like hygroscopicity and CCN activity are likely significantly altered. Can the authors comment on the meteorological conditions conducive to this sulfate production, and whether situations like this might be an important moderator of aerosol chemistry in the region?

We added the following sentences to P13 L12-14, "Stagnant air (wind speeds of ~ 2 m/s) led to the buildup of the urban-influenced PM, which peaked on July 22, as shown in Figure 1. The passing of a cold front, along with precipitation and a change in wind direction, led to a sudden decrease in PM concentrations late on July 22 (Figure 1)."

Minor Comments

8/2: It is not clear at all from the N and M size distributions that this statement is true. Figure S4 shows the smallest dM/dlogD peak for urban cases, opposite of the conclusion in the text. Please revise and be more specific.

Thank you for pointing this out. Upon further investigation we determined that the labels for the urban and wildfire periods were reversed and labeled incorrectly in this figure. Figure S4 (as well as Figure S3) are now labeled correctly and support the conclusions within the text.

8/25: The presence of Na and Ca attributed to Great Lakes or seawater sources is interesting. I understand that the additional manuscript with likely cover these details, but it might be useful to state here whether ratios are consistent with sea-salt or continental dust. Further, soil dust is often lofted into the atmosphere in conjunction with biomass burning, is there any evidence for this during smoke transport? Whether this dust was coated with SOA or sulfates would also be a nice result, based on ATOFMS measurements.

The ratios and other mass spectral ion peaks were not consistent with mineral dust, as stated on P8 L26-P9 L2.

10/3: Does the AMS mass spectrum support your hypothesis that these background particles are mostly SOA coated on a biomass burning core? m/z=60 has typically been used to quantify organic aerosols from biomass burning, can a lack of m/z=60 contribution be used to support the fact that most of the organics are not from combustion?

We now show a plot of AMS m/z 60 and 73 vs time as Figure S6, showing that biomass burning organic aerosol contributed to the OA measured and supporting the ATOFMS results in Figure 5.

13/5: Given the source region for the biomass burning emissions, fires were more likely from agricultural burning. Were there any notable differences that could be highlighted in the mass spectrum or chemical properties of these particles compared to wildfires?

Thank you for this suggestion. We now specify that the likely sources were an agricultural fire in Missouri and a forest fire in Alabama. As shown in Figure 6, a larger number fraction (58%) of the biomass burning particles detected during the urban air mass influence (agricultural burning influence) contained ammonium, compared to the Canadian wildfire periods (39%, by number). This is consistent with the AMS ammonium mass concentration being the highest of the study during this period and is attributed to agricultural ammonia emissions. This is described on P14 L12-14.

References

- Burling, I., Yokelson, R.J., Griffith, D.W., Johnson, T.J., Veres, P., Roberts, J., Warneke, C., Urbanski, S., Reardon, J., Weise, D., 2010. Laboratory measurements of trace gas emissions from biomass burning of fuel types from the southeastern and southwestern United States. Atmos. Chem. Phys. 10, 11115-11130.
- Cooper, O., Moody, J., Thornberry, T., Town, M., Carroll, M., 2001. PROPHET 1998 meteorological overview and air mass classification. J. Geophys. Res-Atmos. 106, 24289-24299.
- DeBell, L.J., Talbot, R.W., Dibb, J.E., Munger, J.W., Fischer, E.V., Frolking, S.E., 2004. A major regional air pollution event in the northeastern United States caused by extensive forest fires in Quebec, Canada. J. Geophys. Res-Atmos. 109, D19305.
- Gillett, N., Weaver, A., Zwiers, F., Flannigan, M., 2004. Detecting the effect of climate change on Canadian forest fires. Geophys. Res. Lett. 31, L18211.
- Jaffe, D.A., Wigder, N.L., 2012. Ozone production from wildfires: A critical review. Atmos. Environ. 51, 1-10.
- Jimenez, J., Canagaratna, M., Donahue, N., Prevot, A., Zhang, Q., Kroll, J.H., DeCarlo, P.F., Allan, J.D., Coe, H., Ng, N., 2009. Evolution of organic aerosols in the atmosphere. Science 326, 1525-1529.
- Knorr, W., Jiang, L., Arneth, A., 2016. Climate, CO2 and human population impacts on global wildfire emissions. Biogeosciences 13, 267-282.
- Liu, Y., Stanturf, J., Goodrick, S., 2010. Trends in global wildfire potential in a changing climate. Forest. Ecol. Manag. 259, 685-697.
- Pratt, K.A., Heymsfield, A.J., Twohy, C.H., Murphy, S.M., DeMott, P.J., Hudson, J.G., Subramanian, R., Wang, Z., Seinfeld, J.H., Prather, K.A., 2010. In situ chemical characterization of aged biomass-burning aerosols impacting cold wave clouds. J. Atmos. Sci. 67, 2451-2468.

- Reid, J., Koppmann, R., Eck, T., Eleuterio, D., 2005. A review of biomass burning emissions part II: intensive physical properties of biomass burning particles. Atmos. Chem. Phys. 5, 799-825.
- Seinfeld, J.H., Pandis, S.N., 2016. Atmospheric chemistry and physics: from air pollution to climate change. John Wiley & Sons, Hoboken, New Jersey.
- Sheesley, R.J., Schauer, J.J., Bean, E., Kenski, D., 2004. Trends in secondary organic aerosol at a remote site in Michigan's upper peninsula. Environ. Sci. Technol. 38, 6491-6500.
- Stockwell, C., Yokelson, R., Kreidenweis, S., Robinson, A., DeMott, P., Sullivan, R., Reardon, J., Ryan, K., Griffith, D., Stevens, L., 2014. Trace gas emissions from combustion of peat, crop residue, domestic biofuels, grasses, and other fuels: configuration and Fourier transform infrared (FTIR) component of the fourth Fire Lab at Missoula Experiment (FLAME-4). Atmos. Chem. Phys., 9727.
- VanReken, T., Mwaniki, G., Wallace, H., Pressley, S., Erickson, M., Jobson, B., Lamb, B., 2015. Influence of air mass origin on aerosol properties at a remote Michigan forest site. Atmos. Environ. 107, 35-43.
- Veira, A., Lasslop, G., Kloster, S., 2016. Wildfires in a warmer climate: Emission fluxes, emission heights, and black carbon concentrations in 2090–2099. J. Geophys. Res-Atmos. 121, 3195-3223.

Ubiquitous Influence of Wildfire Emissions and Secondary Organic Aerosol on Summertime Atmospheric Aerosol in the Forested Great Lakes Region

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- 20 Abstract. Long-range aerosol transport affects locations hundreds of kilometers from the point of emission, leading to distant particle sources influencing rural environments that have few major local sources. Source apportionment was conducted using real-time aerosol chemistry measurements made in July 2014 at the forested University of Michigan Biological Station near Pellston, Michigan, a site representative of the remote forested Great Lakes region. Size-resolved chemical composition of individual 0.5 2.0 µm particles was measured using an aerosol time-of-flight mass spectrometer (ATOFMS), and non-refractory aerosol mass less than 1 µm (PM₁) was measured by a high resolution aerosol mass spectrometer (HR-AMS). The field site was also influenced by air masses transporting Canadian wildfire emissions and urban pollution from Milwaukee and Chicago. During wildfire influenced periods, 0.5 2.0 µm particles were primarily aged biomass burning particles (88% by
- 30 number). These particles were heavily coated with secondary organic aerosol (SOA) formed during

transport, with organics (average O/C ratio of 0.8) contributing 89% of the PM₁ mass. During urbaninfluenced periods, organic carbon, elemental carbon/organic carbon, and aged biomass burning particles were identified, with inorganic secondary species (ammonium, sulfate, and nitrate) contributing 41% of the PM₁ mass, indicative of atmospheric processing. With current models under-predicting organic carbon (OC) in this region and biomass burning being the largest combustion contributor to SOA by mass, these results highlight the importance for regional chemical transport models to accurately predict the impact of long-range transported particles on air quality in the upper Midwest United States, particularly considering increasing intensity and frequency of Canadian wildfires.

10 1 Introduction

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Atmospheric particulate matter less than 2.5 µm in diameter (PM_{2.5}) has significant impacts on air quality, climate, and human health (Calvo et al., 2013; Pöschl and Shiraiwa, 2015). Atmospheric particles directly affect climate by scattering incoming solar radiation and indirectly by acting as cloud condensation (CCN) and ice nuclei (IN) (IPCC, 2013). Increased levels of PM_{2.5} are also linked to increased health risks, particularly respiratory and cardiovascular diseases (Brook et al., 2004; Pope and Dockery, 2006). Particles can impact areas hundreds of kilometers from their sources through long-range transport, with residence times of up to two weeks depending on particle size and chemical composition (Uno et al., 2009). Determining the impact of the long-range transported particles, as well as how they are transformed in the atmosphere during transport, is a critical topic to accurately predict their air quality

- 20 and climate effects (Ault et al., 2011; Creamean et al., 2013). During transport, particles undergo heterogeneous reactions and gas-particle partitioning, aging the particles and leading to primary particles (e.g., soot) to become becoming internally mixed with secondary species, including water, ammonium, nitrate, sulfate, and oxidized organic carbon, thus changing the chemical composition of individual particles (Moffet and Prather, 2009; Riemer and West, 2013). These aging processes are particularly
- 25 important since chemical composition is directly related to particle properties, including reactivity, hygroscopicity, toxicity, scattering, and absorption properties (Brook et al., 2004; Pöschl, 2005; Calvo et al., 2013; Fierce et al., 2016). Particle properties also differ based on the distribution of chemical species, or mixing state, within a population of particles whether various chemical species are contained within

a single particle (internally mixed) or within different particles (externally mixed). Particle mixing state representation in models is particularly important (Bauer et al., 2013), especially for predicting aerosol impacts on the climate (Matsui et al., 2013; Fierce et al., 2016).

- Long-range transport of atmospheric particles can contribute to both remote and populated locations being out of compliance with air quality regulations (National Research Council and National Academies, 2010). For example, elevated aerosol mass and ozone in Europe, eastern Canada, and northeastern United States has been attributed to transported Canadian wildfire emissions (Forster et al., 2001; Colarco et al., 2004; Müller et al., 2005; Wang et al., 2010b; Dutkiewicz et al., 2011; Miller et al., 2011; Dempsey, 2013; Kang et al., 2014; Dreessen et al., 2016). A multi-day exceedance of the National
- 10 Ambient Air Quality Standard for ozone in Maryland during the summer of 2015 was attributed to Canadian wildfire emissions (Dreessen et al., 2016). Similarly, elevated PM_{2.5} observed in New York and Wisconsin has been attributed to Ohio River Valley emissions. Transported pollutants can impact biogenic secondary organic aerosol (SOA) formation in remote locations (Carlton et al., 2010; Emanuelsson et al., 2013; Xu et al., 2015; Rattanavaraha et al., 2016). Finally, prior and on-going studies
- 15 through the IMPROVE program in rural locations throughout North America have investigated both transported and local contributions to the aerosol populations (Hand et al., 2011). Uncertainty in the contributions of long-range aerosols and limited measurements in remote areas can lead to inaccuracies in modeling of aerosol source contributions.

Relatively few studies have chemically characterized atmospheric aerosols in the rural Great Lakes region of the United States (Sheesley et al., 2004; Kim et al., 2005; Kim et al., 2007; Zhang et al., 2009; Jeong et al., 2011; Sjostedt et al., 2011; Kundu and Stone, 2014; Bullard et al., 2017). Except for the major metropolitan areas of Detroit (MI), Chicago (IL), Minneapolis (MN), and Milwaukee (WI), much of the land is characterized by rural agricultural areas and remote forests without significant anthropogenic emissions. A study in the upper peninsula of Michigan conducted by Sheesley et al. (2004)

25 observed major contributions from secondary organic aerosol from both biogenic and anthropogenic volatile organic compound (VOC) oxidation in the summer. Studies across rural Illinois and Ohio found major atmospheric contributions from secondary sulfate, nitrate, and organic carbon, consistent with aerosol aging during transport (Kim et al., 2005; Kim et al., 2007; Zhang et al., 2009), though these

locations were much less forested than the more northern Great Lakes regions. Kundu and Stone (2014) measured composition and sources at rural locations in Iowa, identifying major PM mass contributions from biomass burning, combustion, and dust. Jeong et al. (2011)-and, Sjostedt et al. (2011), and Slowik et al. (2011) identified contributions from secondary organic aerosol, elemental carbon, and dust in rural Harrow, Ontario, downwind of Detroit and Windsor. The scarcity of measurement data in the rural Great Lakes region provides limited opportunities for model evaluation and requires assumptions of background primary aerosol.

In remote regions, there are challenges in distinguishing and identifying primary and secondary aerosol sources, particularly for bulk methods (Pratt and Prather, 2012). Single-particle mass

- 10 spectrometry allows the identification of particle sources through comparisons with source 'fingerprints' and particle aging through characterization of individual particle chemical mixing state (Pratt and Prather, 2009, 2012). Therefore, to apportion the sources of the aerosol population influencing remote northern Michigan, single particle mass spectrometry measurements were conducted during July 2014 at the University of Michigan Biological Station (UMBS) near Pellston, MI. In this study, individual particle
- 15 chemical composition, measured in real-time using single-particle mass spectrometry, was used to identify the sources and secondary processing of transported particles at UMBS. In addition, high resolution aerosol mass spectrometry (HR-AMS) measured chemically-resolved mass concentrations of non-refractory aerosol (organics, sulfate, nitrate, ammonium, and chloride) to provide complementary mass-based characterization of the transported particles at UMBS.

20 2 Methods

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2.1 Field Site and Instrumentation

Atmospheric measurements were conducted from July 13-24, 2014 at the University of Michigan Biological Station (UMBS) near Pellston, MI, a 10,000-acre, remote, forested location with little local pollution (Carroll et al., 2001). The closest major cities are Milwaukee (370 kilometers southwest), Detroit (385 kilometers south), and Chicago (466 kilometers southwest). Instrumentation was located

within a laboratory at the base of the Program for Research on Oxidants: Photochemistry, Emissions, and Transport (PROPHET) tower, a 30-meter tall sampling tower (45°33'31"N, 84°42'52"W) (Carroll et al., 2001). Air was sampled from 34 m above ground level (~14 m above the forest canopy) through foam-insulated 1.09-cm I.D. copper tubing at a flow rate of 9.25 L min⁻¹ (laminar) with a residence time of 15 s. This tubing was connected to a shared sampling manifold at the base of the tower, allowing individual instruments to each have a dedicated sampling line while limiting particle loss.

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An aerosol time-of-flight mass spectrometer (ATOFMS model 3800, TSI, Inc., Shoreview, MN) (Gard et al., 1997; Dall'Osto et al., 2004), described briefly below, was used to measure the size and chemical composition of individual atmospheric particles ranging from $0.5 - 2.0 \mu m$ in vacuum

- 10 aerodynamic diameter (d_{va}) (Section 2.2). An Aerodyne high resolution aerosol mass spectrometer (HR-AMS) (DeCarlo et al., 2006) measured chemically-resolved mass concentrations of non-refractory fine particulate material (nominal vacuum aerodynamic diameter range of $0.05 - 1.0 \mu m$) from July 15–24, 2014. Concentrations for major composition classes (organics, sulfate, nitrate, ammonium, and chloride) are reported here. O/C ratios were calculated throughout the study using the methods described by
- 15 Canagaratna et al. (2015). The operation of the HR-AMS followed standard practice as described elsewhere (Jayne et al., 2000; Allan et al., 2003; Jimenez et al., 2003; Allan et al., 2004); the sampling resolution for the UMBS observations was 2.5 min. Calibrations for instrument flow rate, particle sizing and transmission, and ionization efficiency were conducted during the study following documented procedures (Jimenez and DeCarlo, 2017). Data were analyzed using SQUIRREL (version 1.60) and the
- 20 high resolution analysis software tool PIKA (version 1.20) (Sueper, 2010), with the concentrations corrected based on the estimated composition-dependent collection efficiency (Middlebrook et al., 2012). Additional instrumentation included an ozone analyzer (Thermo Scientific model 49), a scanning mobility particle sizer spectrometer (SMPS, TSI model 3936) with a sheath flow rate of 4 L/min and an aerosol flow rate of 0.4 L/min for measuring size-resolved number concentrations of mobility diameter particles
- 25 12-600 nm, and an aerodynamic particle sizer spectrometer (APS, TSI model 3321) for measuring sizeresolved number concentrations of 0.5-19 μm aerodynamic diameter particles. SMPS and APS size distributions were merged to give a continuous aerosol distribution from 0.01-2.5 μm (aerodynamic

diameter) using previously established methods (Khlystov et al., 2004), assuming a density of 1.5 g cm⁻³ and shape factor of 1.

2.2 Aerosol Time-of-Flight Mass Spectrometer (ATOFMS)

- Using the ATOFMS, 11,430 individual atmospheric particles ranging from 0.5 2.0 μm in dva were chemically analyzed from July 13–24, 2014. The design and operation of the ATOFMS has been described in detail elsewhere (Dall'Osto et al., 2004; Su et al., 2004). Briefly, particles are focused through an aerodynamic lens system and optically detected by two 532 nm continuous wave lasers spaced 6 cm apart. Particle aerodynamic diameter is determined from particle velocity, which was calibrated using monodispersed spherical polystyrene latex spheres (0.4 – 2.5 μm, Polysciences, Inc.) of known diameter and density. Particles are individually desorbed and ionized by a 266 nm Nd:YAG laser that was operated at ~1.2 mJ, and the resulting ions enter a dual-polarity reflectron time-of-flight mass spectrometer. Positive and negative ion mass spectra corresponding to the same individual particles are collected. Mass spectral peak lists for individual particles were generated using TSI MS-Analyze software.
- The individual particle mass spectra were analyzed using YAADA (yaada.org), a software toolkit for MATLAB. Particles were clustered in YAADA using the ART-2a algorithm with a vigilance factor of 0.80 and a learning rate of 0.05 for 20 iterations (Song et al., 1999). The top 50 clusters were manually classified into five particle types described in section 3.1. These top 50 clusters contained 92% of the 11,430 particle mass spectra collected and are the focus of the manuscript. Particle identification was based on characteristic ATOFMS mass spectral signatures previously described (Silva et al., 1999; Pastor et al., 2003; Qin et al., 2012). The errors associated with number fractions for each particle type were
- calculated using binomial statistics.

To obtain chemically-resolved number and mass concentrations for 0.5-2.0 μ m particles, ATOFMS particle counts were scaled with the APS size-resolved particle number concentration data using the method of Qin et al. (2006) to account for size-dependent particle transmission in the inlet.

25 Briefly, ratios of APS number concentration to ATOFMS non-scaled number concentration were calculated every three hours for each individual size bin defined by the APS for use as a scaling factor.

This scaling factor was then multiplied by the corresponding ATOFMS number concentration, providing size and chemically-resolved particle number concentrations for each of the four particle types. These number concentrations were then converted to mass concentrations using assumed spherical shape and compositionally-specific densities. The following densities were applied for the four particle types: 1.5 g

5 cm⁻³ for biomass burning, 1.5 g cm⁻³ for salts, and 1.25 g cm⁻³ for organic carbon-sulfate (OC-sulfate) and elemental carbon/organic carbon – sulfate (ECOC-sulfate) particles (Spencer et al., 2007; Moffet et al., 2008).

3 Results and Discussion

3.1 Overview

- 10 The UMBS campaign (July 13-24, 2014) was characterized by air masses from three primary directions: north, northwest, and southwest (Figures S1 and S2), representative of periods observed during previous UMBS summer studies (Cooper et al., 2001; VanReken et al., 2015). Analysis of NOAA HYSPLIT backward air mass trajectories showed four distinct air mass time periods (Figure S2). From July 13–15, air primarily came from northwestern Canada. From July 15–17, the wind shifted and came from directly
- 15 north crossing over Lake Superior and Lake Michigan before arriving at the field site. In contrast, from July 17–22 the air came mainly from south-southwest of the field site, crossing over the major metropolitan areas of Chicago and Milwaukee followed by Lake Michigan. Finally, from July 23–24, air came from the north-northwest of the field site, crossing Lake Superior and Lake Michigan from northern Canada (Figure S2). During summer 2009, VanReken et al. (2015) found that 60% of the air masses came
- from north/northwest of UMBS, similar to this study (57%). Air came from southern polluted regions 43% of the time during our study, compared to 29% during July-August 2009 (VanReken et al., 2015).

Total PM_{2.5} number, PM_{2.5} mass, and ozone concentrations ranged from 143 to 6,031 particles cm⁻³ (average \pm standard deviation: 1,822 \pm 1,181 particles cm⁻³), 1 to 43 µg/m³ (average \pm standard deviation: 8 \pm 8 µg/m³) and 9 to 63 ppb (average \pm standard deviation: 32 \pm 14 ppb), respectively (Figure

25 1). Maximum concentrations were detected when the air arrived from the southwestern urban areas, and

the minimum values were observed for air masses from the north during remote air transport (Figures S3 and S4). Previously, VanReken et al. (2015) observed an 85% increase in particle number concentration when air originating from these southwestern urban areas impacted UMBS. These results suggest a wide range of sources affecting the field site, which were directly observed by the ATOFMS. Here, we examine the influences of wildfires (Section 3.3) and urban pollution (Section 3.4) on summertime aerosol chemical composition, compared to remote background (Section 3.2), at UMBS.

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Major individual particle types observed by ATOFMS included biomass burning, organic carbonsulfate (OC-sulfate), and elemental carbon/organic carbon-sulfate (ECOC-sulfate) (Figure 2). Biomass burning particles were characterized by intense peaks at m/z 39 (K⁺) and -97 (HSO₄⁻), as well as less

- 10 intense peaks at m/z 12 (C⁺), 18 (NH₄⁺), and 27 (C₂H₃⁺) (Pratt et al., 2010). Biomass burning particles also contained a peak at m/z 43 (C₂H₃O⁺), a marker for oxidized OC on particles, which is addressed further in section 3.3. Biomass burning was the most prominent particle type, comprising ~80% of submicron (0.5 – 1.0 µm) and ~50% of supermicron (1 – 2 µm) particles, by number, throughout the study, with number fraction varying according to the level of influence from wildfires. OC-sulfate
- 15 particles contributed ~7%, by number, to submicron $(0.5 1.0 \ \mu\text{m})$ particles and ~8%, by number, to supermicron $(1.0 - 2.0 \ \mu\text{m})$ particles and were characterized by intense peaks at m/z 27 (C₂H₃⁺), 39 (C₃H₃⁺/K⁺), +/-43 (C₂H₃O^{+/-}), and -97 (HSO₄⁻). OC-sulfate particles can originate from a variety of sources including primary vehicular emissions (Toner et al., 2008) and secondary organic sources (Pratt and Prather, 2009). The intense m/z 43 (most intense OC-sulfate particle ion peak) is indicative of
- significant SOA coatings on combustion particles, including biomass burning (Pratt and Prather, 2009). ECOC-sulfate particles, characterized by C_n^+ fragment peaks, observed at m/z 12 (C⁺), 24 (C₂⁺), 36 (C₃⁺), 48 (C₄⁺), etc., as well as markers at m/z 27 (C₂H₃⁺), 18 (NH₄⁺), and -97 (HSO₄⁻), are attributed to vehicular emissions (Toner et al., 2006; Toner et al., 2008) and contributed ~5%, by number, to both sub- and supermicron particles with the majority observed on July 22 during an urban-influenced air mass. In
- 25 addition to the previously mentioned combustion and secondary particles, Na and Ca salts internally mixed with nitrate were episodically detected, primarily during July 16–18 and July 24–25. <u>Based on elemental ratios and established mass spectral fingerprints</u>, these salts <u>may have</u> originated from the Great Lakes (<u>Axson et al., 2016</u>)(Axson et al., 2016; May et al., 2018) and/or seawater, <u>rather than mineral</u>

dust (Sullivan et al., 2007; Ault et al., 2011; Fitzgerald et al., 2015), and are the focus of an upcoming manuscript. For each of the discussed particle types, we present the chemical mixing state by reporting the number percentage of particles within each particle type that contain a mass spectral marker corresponding to each secondary aerosol chemical species of interest, including sulfate (HSO₄⁻, m/z -97), nitrate (NO₂⁻, m/z -46, and/or NO₃⁻, m/z -62), ammonium (NH₄⁺, m/z 18), and oxidized OC (C₃H₂O⁻, m/z -43, or C₃H₂O⁺, m/z 43) (Oin et al., 2012).

 PM_1 mass measured by the HR-AMS was on average 73% organics (7.8 μ g/m³) throughout the study, with a substantial contribution from oxidized organics as determined by an average HR-AMS O/C

10 2008; Qin et al., 2012). O/C ratios between 0.6 – 1 are commonly associated with low volatility oxidized organic aerosol (LV-OOA) that has undergone extensive aging (Jimenez et al., 2009), consistent with the single-particle observation that SOA coated the major particle types. In addition, the ammonium balance of predicted ammonium versus measured ammonium throughout the study (Figure S5) shows a slight deficit in measured ammonium, typically indicative of acidic aerosol or the presence of organic

ratio of 0.84 and through the ATOFMS oxidized organic carbon ion marker m/z 43, C₂H₃O⁺ (Aiken et al.,

15 nitrates/sulfates (Farmer et al., 2010). Also consistent with atmospheric processing during long-range transport, 92% of all 0.5 – 2.0 μm particles, by number, were measured by the ATOFMS to be internally mixed with secondary species, including sulfate (HSO₄⁻, *m/z* -97), nitrate (NO₂⁻, *m/z* -46 and/or NO₃⁻, *m/z* -62), ammonium (NH₄⁺, *m/z* 18), and/or oxidized OC (C₃H₂O⁻, *m/z* -43 or C₃H₂O⁺, *m/z* 43) (Qin et al., 2012). On average, sulfate comprised 20% (2.2 μg/m³) of the total PM₁ mass measured by HR-AMS.

20 3.2 Remote Background Air Mass Influence

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From July 15-17, air arrived at UMBS originating from rural northern Canada. This remote background air mass period was differentiated from the wildfire influenced periods (Section 3.3) based on the lack of smoke impacting the site, as indicated by NOAA Smoke and Fire products (Figure 4). The average $PM_{2.5}$ number concentration was 903 ± 499 particles cm⁻³ (range of 143 - 2163 particles cm⁻³,

Figure 1), and the average PM_{2.5} mass concentration was $1.9 \pm 0.4 \,\mu\text{g/m}^3$ with a particle number mode of 82 nm (Figure 1 and S3), comparable the average particle number ($1630 \pm 1280 \,\text{cm}^{-3}$) and mode ($73 \,\text{cm}^{-3}$) and mod

21 nm) observed by VanReken et al. (2015) during background air mass influence at UMBS in summer 2009. The average ozone concentration was 17 ± 6 ppb (Figure 1)-, similar to previous measurements at UMBS during background air mass influence (Cooper et al., 2001). Despite the With a lack of direct wildfire influence (Figure 4), $61 \pm 1\%$ of the 0.5 – 2.0 µm particles, by number, were classified by ATOFMS as aged biomass burning aerosols, relatively similar to the background biomass burning particle influence reported by Hudson et al. (2004) and Pratt et al. (2010) for the United States free troposphere (33-52%, by number). Biomass burning particles were internally mixed with oxidized OC $(80 \pm 2\%)$, by number) or mixed with sulfate $(85 \pm 2\%)$. Nitrate was internally mixed with $8 \pm 2\%$, by number of biomass burning particles and $33 \pm 3\%$, by number, of OC-sulfate particles. It is likely that, 10 while the observed biomass burning particles have a small potassium-rich (biomass burning) core, they are primarily SOA by mass (Pratt and Prather, 2009; Moffet et al., 2010) (Section 3.3). The HR-AMS showed average PM₁ organic mass concentrations of 4.4 μ g/m³, with minimal contribution from sulfate $(0.3 \ \mu g/m^3)$, as well as nitrate and ammonium (both less than 0.1 $\mu g/m^3$ on average) (Figure 3).

- The significant internal mixing of oxidized OC combined with the significant organic mass loading (average HR-AMS O/C ratio of 0.9) is consistent with high SOA mass on the particles (Aiken et 15 al., 2008). Previous studies in rural and forested environments found similarly high O/C ratios during periods of non-polluted air and attributed this to regional SOA formation (Jimenez et al., 2009; Sun et al., 2009; Raatikainen et al., 2010; Sjostedt et al., 2011). There was a notable spike in O/C ratio on July 15 – 16 to 1.2, indicative very highly oxidized organics. O/C ratios of this magnitude have previously been 20 observed at the remote Whistler Mountain, where organic aerosol O/C ratios up to ~1.3 where observed during organic aerosol accumulation events (Sun et al., 2009). Sheesley et al. (2004) found that SOA, primarily biogenic-derived, contributed over 90% of the total organic carbon mass observed during the summer at the Seney National Wildlife Refuge in northern Michigan, located 120 km northwest of UMBS. Notably, ultrafine particle growth was observed at UMBS on July 16 during this high O/C ratio spike (Gunsch et al., 2017).(Gunsch et al., 2018). The air arriving during this period was not under the
- 25 influence of wildfires (Section 3.3) or urban areas (Section 3.4), and is therefore expected to be representative of remote background conditions.

3.3 Wildfire Influence

	From July 13-15 and July 24 mid-day through July 25, the NOAA Hazard Mapping System	
	(HMS) Smoke Product (Rolph et al., 2009) indicated that smoke plumes originating from wildfires within	
	the Northwest Territories (Canada) directly influenced UMBS (Figure 4). According to the Canadian	
5	Interagency Forest Fire Centre, over 5,500 km ² of land burned within the Northwest Territories during	
	July 2014 (CIFFC, 2014). Canadian wildfires are a major source of global PM _{2.5} , with estimates of ~1.6	
	Tg yr ⁻¹ emitted to the atmosphere (Wiedinmyer et al., 2006). Average PM _{2.5} number and concentrations	Formatted: Superscript
	during these two wildfire influence periods were statistically higher (t-test, $\alpha = 0.05$) at 1400 ± 800	
	particles cm ⁻³ (range of 147 – 4832 particles cm ⁻³ , Figure 1) and 5.4 \pm 2.6 $\mu g/m^3$ (range of 1.3 – 10.5	
10	μ g/m ³ , Figure 1), respectively, compared to the background period (Section 3.2). The particle <u>number</u>	
	mode during wildfire influence was 80 ± 46 nm, similar to the background periods air mass period (mode	
	of 82 ± 37 nm) (Figure S3), Ozone was also elevated during July 13-15 reaching as high as 35 ppb,	Formatted: Font: Bold
	compared to an average of 10 ppb during the background period (Figure 1). During these periods, the air	
	masses did not pass over any major urban areas (Figure S2), making ozone production within the smoke	
15	plume during transport the likely source (Jaffe and Wigder, 2012). Ozone did not increase during the July	
	24 smoke plume, staying near the average for the study (25 \pm 12 ppb) with a concentration of 26 \pm 3 ppb	
	(Figure 1). While an ozone increase is often observed for aged wildfire plumes, an increase does not	
	always occur during wildfire influence, such as when low NO_x levels within plumes, potentially due to	
	smoldering combustion, limitlimiting the production of ozone (Jaffe and Wigder, 2012).	
20	During the wildfire influenced periods, 88 \pm 1% of the measured 0.5 – 2.0 μm particles, by	
	number, were biomass burning particles, with an average mass concentration of 0.42 $\mu g/m^3$ (Figure 5)	
	and a maximum of 0.80 μ g/m ³ occurring during the early afternoon of July 14 when the heaviest wildfire	
	smoke was reported by the NOAA smoke product (Figure 4A). Minor contributions of OC-sulfate	
	particles (8 \pm 1% by number) were also measured. The OC-sulfate particle mass spectra (Figure 2B)	
25	showed that 75 \pm 5%, by number, contained potassium (K ⁺ , m/z 39), suggesting that these were highly	
	aged biomass burning particles coated by SOA such that the typical biomass burning mass spectral	

These OC-sulfate particles featured a dominant intense m/z 43 (C₂H₃O⁺) ion peak, indicating that these particles were heavily coated with SOA. During the afternoon event on July 24, PM₁ organic mass concentrations measured by HR-AMS nearly doubled from 2.5 ± 0.1 µg/m³ before the event to 4.5 ± 0.3 µg/m³ during the event (Figure 3), accounting for ~90% of the total PM₁ mass concentration. The HR-AMS O/C ratio was 0.8 during wildfire periods, consistent with biomass burning particles heavily coated with SOA (Aiken et al., 2008), as also observed by 95 ± 1%, by number, of the biomass burning and OC-

sulfate particles, measured by ATOFMS during these periods, featuring the oxidized OC ion marker (m/z 43, C₂H₃O⁺) (Figure 6). Freshly emitted biomass burning has a O/C ratio of ~0.2, which can increase to ~0.6 in only a few hours as <u>the emissions undergo photochemical aging and oxidized material condenses</u>

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- 10 onto the particles (Grieshop et al., 2009; Pratt et al., 2011)(Grieshop et al., 2009; Pratt et al., 2011; Liu et al., 2016; Zhou et al., 2017). While AMS levoglucosan ion markers (*m*/*z* 60 (C₂H₄O₂⁺) and 73 (C₃H₅O₂⁺)) (Alfarra et al., 2007) were observed, consistent with the ATOFMS observation of biomass burning particles, the levels were low (Figure S6), due to the expected degradation during atmospheric transport (Hennigan et al., 2010). As tThe wildfire air masses measured during the present study were transported
- 15 <u>48 72 h over multiple days</u> over Canadian forests, suggesting the accumulation of both biomass burning SOA and biogenic SOA, from condensation of monoterpene oxidation products (Slowik et al., 2010), likely contributed to the observed O/C ratio of 0.8 at UMBS. Previous studies conducted by Slowik et al. (2010) and Sheesley et al. (2004) observed monoterpene-derived SOA within summertime air masses passing over these forest. Therefore, it is likely that the oxidation (both gas- and aqueous-phase) of both
- 20 biomass burning and biogenic VOCs (Lee et al., 2011), contributed to the observed elevated O/C ratio of 0.8 1.1 at UMBS.

During transport of the biomass burning aerosols, accumulation of sulfate also occurred, with 97 \pm 1%, by number, of biomass burning particles internally mixed with sulfate (*m/z* -97, HSO₄⁻) (Figure 6). HR-AMS measured PM₁ sulfate also increased from less than 0.1 µg/m³ to 2 µg/m³ after mid-day July 24

25 (Figure 3). IncreasesSO₂ has been previously shown to be emitted from wildfires (e.g. Burling et al., 2010; Stockwell et al., 2014), and increases in particulate sulfate mass have been observed during wildfire plume aging through cloud processing (DeBell et al., 2004; Pratt et al., 2010). In comparison, the HR-

AMS measured limited amounts of PM₁ ammonium (~2% of total mass, 0.2 μ g/m³) during the wildfire event on July 24 (Figure 3). However, ammonium was internally mixed in 38 ± 2%, by number, of biomass burning and 68 ± 2%, by number, of OC-sulfate particles (Figure 6). This result indicates that while ammonium was present within many particles, it was a minor fraction of the particle mass. Nitrate was also internally mixed with 43 ± 2% of biomass burning particles, by number, and 17 ± 2%, by number, of OC-sulfate particles (Figure 6), and the HR-AMS only measured ~1% of PM₁ mass to be nitrate (0.06 μ g/m³). Therefore, it is likely that the ammonium was present in the form of ammonium sulfate internally mixed with biomass burning and OC-sulfate.

3.4 Urban Air Mass Influence

- From July 17-22, UMBS was influenced by air masses from the southwest, passing over the major metropolitan areas of Chicago and Milwaukee before arriving at the site (Figure S2) after transport times of 24-36 hours. Stagnant air (wind speeds of ~ 2 m/s) led to the buildup of the urban-influenced PM, which peaked on July 22, as shown in Figure 1. The passing of a cold front, along with precipitation and a change in wind direction, led to a sudden decrease in PM concentration late on July 22 (Figure 1). The average ozone concentration was elevated at an average of 41 ± 12 ppb similar to previous measurements by Cooper et al. (2001) at UMBS when under the direct influence of urban pollution (Figure 1). The PM_{2.5} number and mass concentration for this period were 2,700 ± 900 particles cm⁻³ (range of 414 6,031
 - particles cm⁻³) and $14 \pm 8 \ \mu g/m^3$ (range of $2 43 \ \mu g/m^3$), respectively, the highest for the study (Figure 1). The particle mode of 69 ± 29 nm was also the smallest of the study (Figure S3) <u>due to contributions</u>
- from combustion emissions, typically less than 50 nm (Seinfeld and Pandis, 2016), which likely grew to the observed sizes due to the condensation of secondary species during transport. A similar mode of 84 ± 18 nm was observed by VanReken et al. (2015) at UMBS during summer 2009 urban air mass influence. VanReken et al. (2015) also previously observed the highest particle number concentrations (3,000 ± 1,300 particles cm⁻³) at UMBS during the influence of southern air masses. Wildfire smokeSmoke influence was present during this period as shown by the NOAA smoke product (Figure 4C). However,
- unlike during the previous periods, this smoke originated mainly from the southern United States-(active

fires were located in Tennessee, Arkansas, and Missouri), and were from agricultural and forest fires (active agricultural fires were located in Missouri, with a forest fire in Arkansas). The HR-AMS showed levoglucosan ion markers (m/z 60 ($C_2H_4O_2^+$) and 73 ($C_3H_5O_2^+$)) (Alfarra et al., 2007) (Figure S6), consistent with the satellite-observed smoke influence. Biomass burning particles measured by ATOFMS

- 5 steadily increased in mass concentration throughout this period (Figure 5), with a notable spike in the mass concentration on July 22 observed in both the submicron (0.5-1.0 μ m particles: 2.3 μ g/m³) and supermicron (0.3 μ g/m³) size ranges (1.0-2.0 μ m particles: Figure 5). Overall, during urban influence, biomass burning particles accounted for 72 ± 2% of the particles, by number, and ~30% of the total mass concentration (Figure 5). The biomass burning particles were aged, as shown by internal mixtures of
- sulfate (88 ± 2%, by number), oxidized OC (92 ± 1%, by number), ammonium (58 ± 2%, by number), and nitrate (30 ± 2%, by number) (Figure 6). The greatest internal mixing with ammonium was observed during this period.Notably, a higher number fraction of the biomass burning particles during this period were internally mixed with ammonium, compared to the biomass burning particles detected during the Canadian wildfire influence. The HR-AMS also measured the highest average ammonium mass
 concentration during this period of 1.6 µg/m³, accounting for 10% of the total PM₁ particle mass (Figure

3). Agricultural activities, both crop and livestock, located to the south and southwest of the field site (Stephen and Aneja, 2008; Paulot et al., 2014) may be the source of the elevated ammonium levels.

ECOC-sulfate and OC-sulfate particles comprised the second most prominent particle types measured by ATOFMS during this urban-influenced period at 12 ± 1% and 9 ± 1% of the submicron (0.5 20 - 1.0 µm) particles, by number, and an average of 0.08 µg/m³ and 0.03 µg/m³, respectively (Figure 5). The influence of urban vehicular combustion resulted in the increased levels of measured ECOC-sulfate particles (Toner et al., 2006; Toner et al., 2008), compared to non-urban influenced periods (2 ± 1% by number). HR-AMS PM₁ mass concentrations (Figure 3) showed increased organic mass during urban influence with an average mass concentration of 9.7 µg/m³ (Figure 3), likely due to a mixture of biomass

25 burning, anthropogenic, and biogenic organic aerosol. The average HR-AMS O/C ratio during the urban period was the lowest of the study (0.78), likely due to increased contributions from hydrocarbon-like organic aerosol from urban vehicle combustion emissions (Aiken et al., 2008), in contrast to primarily oxidized organic aerosol during regional background periods (Jimenez et al., 2009). An increase in less oxidized organic aerosol was similarly observed in rural Ontario when the site was influenced by urban air masses from Detroit, compared to remote air masses (Sjostedt et al., 2011). The ECOC-sulfate and OC-sulfate particles were highly aged, with ~75%, by number, of each particle type internally mixed with ammonium, consistent with particle aging during transport (Figure 6C). Ammonium (1.6 μ g/m³) and sulfate (4.9 μ g/m³) comprised over 40% of the total PM₁ mass measured by the HR-AMS during these periods, likely in the form of ammonium sulfate (Figure 3). Urban influenced air masses had the highest mass concentration of sulfate (up to 10 μ g/m³) measured throughout the study. In contrast, there was little presence of nitrate internally mixed in the ECOC-sulfate (4 ± 2%, by number) and OC-sulfate (19 ± 5%, by number) particles (Figure 6). Finally, while, and nitrate only comprised 1% (0.2 μ g/m³) of the total PM. mass concentration during from the urban influence (Figure 3), this was the highest nitrate

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PM₁ mass concentration <u>during</u> from the urban influence (Figure 3), this was the highest nitrate contribution during the study, similar previous rural eastern United States studies (Jimenez et al., 2009).

4 Conclusions

Source apportionment of atmospheric particles in the summertime was conducted at the forested University of Michigan Biological Station, located in remote northern Michigan. The field site was impacted by air masses from three distinct areas: remote background, northwestern Canada, and southwestern urban areas. July 2014 was one of the most active burning seasons for the Northwest Territories in over two decades with a total of 10,643 km² of land burned, significantly more than the tenyear (1,944 km²) and twenty-five year (2,423 km²) averages (CIFFC, 2014). The increased wildfire activity noticeably impacted northern Michigan, as the presence of biomass burning particles was ubiquitous throughout the study and made up the majority of measured particle number and mass concentrations. While air also came from urban areas southwest of UMBS, aged biomass burning particles dominated particle number concentrations due to wildfire influences from the southern United States. Due to the urban influence, these air masses had the highest mass contributions of sulfate (over 50 times

25 the background) detected during the entire study. The accumulation of soluble secondary species,

including sulfate and nitrate, increases the CCN ability of biomass burning particles (Furutani et al., 2008; Petters et al., 2009; Wang et al., 2010a), illustrating the importance of transported wildfire emissions.

While biomass burning particles were the most dominant particle core detected, SOA was a major contributor to particle mass during the study. On average, the HR-AMS organic aerosol O/C ratio was

- 5 0.84, indicative of highly oxidized organic carbon (Aiken et al., 2008). During remote background periods, internal mixing of oxidized OC combined with the significant PM₁ organic mass loading is indicative of the high mass loading of biogenic SOA in the forested region (Sheesley et al., 2004). During wildfire-influenced air masses, organics contributed ~90% to the PM₁ mass, with SOA internally mixed with biomass burning and OC-sulfate particles, indicating that SOA from both biogenic VOC oxidation
- 10 and wildfire combustion is a major source of OC in the region. <u>The observed levels of biomass burning</u> aerosol influence are attributed to the abnormally active Canadian wildfire season of 2014, compared to previous typical summers in northern Michigan with primarily biogenic SOA influence (Sheesley et al., 2004). Models under-predict OC in this region, and Jathar et al. (2014) indicates that on a national level, models predict biomass burning is the largest combustion contributor to SOA by mass, consistent with
- 15 the significant influence of wildfires during this work.

Modeling studies have called for further investigations of wildfire emissions and areas they affect in order to reduce uncertainty within models due to limited data, particularly when modeling interactions between wildfire plumes and urban emissions. Wildfire plume ozone production can lead to areas far from the original source to be out of compliance with regulatory standards, demonstrating the importance

- 20 to be able to accurately model ozone production (Hu et al., 2008; Jaffe and Wigder, 2012; Lu et al., 2016). Also, as described here, particles aged through transport show internal mixtures of nitrate, sulfate and oxidized organics, which can lead to increased CCN activity (Furutani et al., 2008). With wildfires expected to increase in both intensity and frequency due to climate change (Gillett et al., 2004; Liu et al., 2010; Knorr et al., 2016; Veira et al., 2016), the contributions of long-range transported biomass burning
- 25 emissions to the upper Midwest US atmosphere are expected to increase, such that air quality modeling efforts will need to supplement their existing emissions to account for the expected increase in wildfire emissions (Smith and Mueller, 2010).

Competing Interests. The authors declare that they have no competing financial interests.

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References

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- Aiken, A. C., Decarlo, P. F., Kroll, J. H., Worsnop, D. R., Huffman, J. A., Docherty, K. S., Ulbrich, I. M., Mohr, C., Kimmel, J. R., and Sueper, D.: O/C and OM/OC ratios of primary, secondary, and ambient organic aerosols with high-resolution time-of-flight aerosol mass spectrometry, Environ. Sci. Technol., 42, 4478-4485, 2008.
- Alfarra, M. R., Prevot, A. S., Szidat, S., Sandradewi, J., Weimer, S., Lanz, V. A., Schreiber, D., Mohr, M., and Baltensperger, U.: Identification of the mass spectral signature of organic aerosols from wood burning emissions, Environ. Sci. Technol., 41, 5770-5777, 2007.
- Allan, J. D., Jimenez, J. L., Williams, P. I., Alfarra, M. R., Bower, K. N., Jayne, J. T., Coe, H., and Worsnop,
 D. R.: Quantitative sampling using an Aerodyne aerosol mass spectrometer 1. Techniques of data interpretation and error analysis, J. Geophys. Res-Atmos., 108, 4090, 2003.
 - Allan, J. D., Delia, A. E., Coe, H., Bower, K. N., Alfarra, M. R., Jimenez, J. L., Middlebrook, A. M., Drewnick, F., Onasch, T. B., and Canagaratna, M. R.: A generalised method for the extraction of chemically resolved mass spectra from Aerodyne aerosol mass spectrometer data, J. Aerosol. Sci., 35, 909-922, 2004.
 - Ault, A. P., Williams, C. R., White, A. B., Neiman, P. J., Creamean, J. M., Gaston, C. J., Ralph, F. M., and Prather, K. A.: Detection of Asian dust in California orographic precipitation, J. Geophys. Res-Atmos., 116, 2011.
- Axson, J. L., May, N. W., Colón-Bernal, I. D., Pratt, K. A., and Ault, A. P.: Lake Spray Aerosol: A Chemical
 Signature from Individual Ambient Particles, Environ. Sci. Technol., 50, 9835 9845, 2016.
 - Bauer, S. E., Ault, A., and Prather, K. A.: Evaluation of aerosol mixing state classes in the GISS modelE - MATRIX climate model using single - particle mass spectrometry measurements, J. Geophys. Res-Atmos., 118, 9834-9844, 2013.
 - Brook, R. D., Franklin, B., Cascio, W., Hong, Y., Howard, G., Lipsett, M., Luepker, R., Mittleman, M.,
- 25 Samet, J., and Smith, S. C.: Air pollution and cardiovascular disease, Circulation, 109, 2655-2671, 2004.
 - Bullard, R. L., Singh, A., Anderson, S. M., Lehmann, C. M., and Stanier, C. O.: 10-Month characterization of the aerosol number size distribution and related air quality and meteorology at the Bondville, IL Midwestern background site, Atmos. Environ., 2017.
- 30 Burling, I., Yokelson, R. J., Griffith, D. W., Johnson, T. J., Veres, P., Roberts, J., Warneke, C., Urbanski, S., Reardon, J., and Weise, D.: Laboratory measurements of trace gas emissions from biomass burning of fuel types from the southeastern and southwestern United States, Atmos. Chem. Phys., 10, 11115-11130, 2010.
- Calvo, A., Alves, C., Castro, A., Pont, V., Vicente, A., and Fraile, R.: Research on aerosol sources and
 chemical composition: past, current and emerging issues, Atmospheric Research, 120, 1-28, 2013.
 - Canagaratna, M., Jimenez, J., Kroll, J., Chen, Q., Kessler, S., Massoli, P., Hildebrandt Ruiz, L., Fortner, E., Williams, L., and Wilson, K.: Elemental ratio measurements of organic compounds using aerosol mass spectrometry: characterization, improved calibration, and implications, Atmos. Chem. Phys., 15, 253-272, 2015.

- Carlton, A. G., Pinder, R. W., Bhave, P. V., and Pouliot, G. A.: To what extent can biogenic SOA be controlled?, Environ. Sci. Technol., 44, 3376-3380, 2010.
- Carroll, M. A., Bertman, S. B., and Shepson, P. B.: Overview of the Program for Research on Oxidants: Photochemistry, Emissions, and Transport (PROPHET) summer 1998 measurements intensive, J. Geophys. Res-Atmos., 106, 24275-24288, 2001.
- Centre, C. I. F. F.:Canadian Interagency Forest Fire Centre, National Wildland Fire Situation Report: http://www.ciffc.ca, Accessed July 2016.,

- Colarco, P., Schoeberl, M., Doddridge, B., Marufu, L., Torres, O., and Welton, E.: Transport of smoke from Canadian forest fires to the surface near Washington, DC: Injection height, entrainment, and optical properties, J. Geophys. Res-Atmos., 109, D06203, 2004.
- Cooper, O., Moody, J., Thornberry, T., Town, M., and Carroll, M.: PROPHET 1998 meteorological overview and air mass classification, J. Geophys. Res-Atmos., 106, 24289-24299, 2001.
- Creamean, J. M., Suski, K. J., Rosenfeld, D., Cazorla, A., DeMott, P. J., Sullivan, R. C., White, A. B., Ralph,
 F. M., Minnis, P., and Comstock, J. M.: Dust and biological aerosols from the Sahara and Asia
 influence precipitation in the western US, Science, 339, 1572-1578, 2013.
- Dall'Osto, M., Beddows, D., Kinnersley, R. P., Harrison, R. M., Donovan, R. J., and Heal, M. R.: Characterization of individual airborne particles by using aerosol time - of - flight mass spectrometry at Mace Head, Ireland, J. Geophys. Res-Atmos., 109, 2004.
- DeBell, L. J., Talbot, R. W., Dibb, J. E., Munger, J. W., Fischer, E. V., and Frolking, S. E.: A major regional air pollution event in the northeastern United States caused by extensive forest fires in Quebec, Canada, J. Geophys. Res-Atmos., 109, D19305, 2004.
 - DeCarlo, P. F., Kimmel, J. R., Trimborn, A., Northway, M. J., Jayne, J. T., Aiken, A. C., Gonin, M., Fuhrer, K., Horvath, T., and Docherty, K. S.: Field-deployable, high-resolution, time-of-flight aerosol mass spectrometer, Anal. Chem., 78, 8281-8289, 2006.
- 25 Dempsey, F.: Forest fire effects on air quality in Ontario: Evaluation of several recent examples, B. Am. Meteorol. Soc., 94, 1059-1064, 2013.
 - Dreessen, J., Sullivan, J., and Delgado, R.: Observations and Impacts of Transported Canadian Wildfire Smoke on Ozone and Aerosol Air Quality in the Maryland Region on 9-12 June, 2015, J. Air Waste Manage. Assoc., 66, 842-862, 2016.
- 30 Dutkiewicz, V. A., Husain, L., Roychowdhury, U. K., and Demerjian, K. L.: Impact of Canadian wildfire smoke on air quality at two rural sites in NY State, Atmos. Environ., 45, 2028-2033, 2011.
- Emanuelsson, E. U., Hallquist, M., Kristensen, K., Glasius, M., Bohn, B., Fuchs, H., Kammer, B., Kiendler-Scharr, A., Nehr, S., and Rubach, F.: Formation of anthropogenic secondary organic aerosol (SOA) and its influence on biogenic SOA properties, Atmos. Chem. Phys., 13, 2837-2855, 2013.
 - Farmer, D., Matsunaga, A., Docherty, K., Surratt, J., Seinfeld, J., Ziemann, P., and Jimenez, J.: Response of an aerosol mass spectrometer to organonitrates and organosulfates and implications for atmospheric chemistry, Proc. Natl. Acad. Sci., 107, 6670-6675, 2010.
- Fierce, L., Bond, T. C., Bauer, S. E., Mena, F., and Riemer, N.: Black carbon absorption at the global scale is affected by particle-scale diversity in composition, Nat. Commun., 7, 2016.

- Fitzgerald, E., Ault, A. P., Zauscher, M. D., Mayol-Bracero, O. L., and Prather, K. A.: Comparison of the mixing state of long-range transported Asian and African mineral dust, Atmos. Environ., 115, 19-25, 2015.
- Forster, C., Wandinger, U., Wotawa, G., James, P., Mattis, I., Althausen, D., Simmonds, P., O'Doherty, S., Jennings, S. G., and Kleefeld, C.: Transport of boreal forest fire emissions from Canada to Europe, J. Geophys. Res-Atmos., 106, 22887-22906, 2001.

- Furutani, H., Dall'osto, M., Roberts, G. C., and Prather, K. A.: Assessment of the relative importance of atmospheric aging on CCN activity derived from field observations, Atmos. Environ., 42, 3130-3142, 2008.
- 10 Gard, E., Mayer, J. E., Morrical, B. D., Dienes, T., Fergenson, D. P., and Prather, K. A.: Real-time analysis of individual atmospheric aerosol particles: Design and performance of a portable ATOFMS, Anal. Chem., 69, 4083-4091, 1997.
 - Gillett, N., Weaver, A., Zwiers, F., and Flannigan, M.: Detecting the effect of climate change on Canadian forest fires, Geophys. Res. Lett., 31, L18211, 2004.
- 15 Grieshop, A., Donahue, N., and Robinson, A.: Laboratory investigation of photochemical oxidation of organic aerosol from wood fires 2: analysis of aerosol mass spectrometer data, Atmos. Chem. Phys., 9, 2227-2240, 2009.
- Gunsch, M. J., Schmidt, S., Gardner, D. J., Bondy, A. L., May, N., Bertman, S. B., Pratt, K. A., and Ault, A.
 P.: Particle Growth in an Isoprene-Rich Forest: Influences of Urban, Wildfire, and Biogenic
 Precursors, <u>SubmittedAccepted</u> to Atmospheric Environment, <u>20172018</u>.
- Hennigan, C. J., Sullivan, A. P., Collett, J. L., and Robinson, A. L.: Levoglucosan stability in biomass burning particles exposed to hydroxyl radicals, Geophys. Res. Lett., 37, 2010.
 - Hu, Y., 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 fires on an urban area, Environ. Sci. Technol., 42, 3676-3682, 2008.
- Hudson, P. K., Murphy, D. M., Cziczo, D. J., Thomson, D. S., De Gouw, J. A., Warneke, C., Holloway, J., Jost, H. J., and Hübler, G.: Biomass - burning particle measurements: Characteristic composition and chemical processing, J. Geophys. Res-Atmos., 109, D23S27, 2004.
- IPCC: IPCC, 2013: climate change 2013: the physical science basis. Contribution of working group I
 to the fifth assessment report of the intergovernmental panel on climate change, edited by:
 Stocker, T., Qin, D., Plattner, G., Tignor, M., Allen, S., Boschung, J., Nauels, A., Xia, Y., Bex, B.,
 and Midgley, B., Cambridge University Press, 2013.
 - Jaffe, D. A., and Wigder, N. L.: Ozone production from wildfires: A critical review, Atmos. Environ., 51, 1-10, 2012.
- 35 Jathar, S. H., Gordon, T. D., Hennigan, C. J., Pye, H. O., Pouliot, G., Adams, P. J., Donahue, N. M., and Robinson, A. L.: Unspeciated organic emissions from combustion sources and their influence on the secondary organic aerosol budget in the United States, Proc. Natl. Acad. Sci., 111, 10473-10478, 2014.

- Jayne, J. T., Leard, D. C., Zhang, X., Davidovits, P., Smith, K. A., Kolb, C. E., and Worsnop, D. R.: Development of an aerosol mass spectrometer for size and composition analysis of submicron particles, Aerosol. Sci. Technol., 33, 49-70, 2000.
- Jeong, C.-H., McGuire, M. L., Godri, K. J., Slowik, J. G., Rehbein, P., and Evans, G.: Quantification of aerosol chemical composition using continuous single particle measurements, Atmos. Chem. Phys., 11, 7027-7044, 2011.

35

- Jimenez, J., Canagaratna, M., Donahue, N., Prevot, A., Zhang, Q., Kroll, J. H., DeCarlo, P. F., Allan, J. D., Coe, H., and Ng, N.: Evolution of organic aerosols in the atmosphere, Science, 326, 1525-1529, 2009.
- 10 Jimenez, J., and DeCarlo, P.:Field ToF-AMS Operation: <u>http://www.ciffe.ea</u>, Accessed September 2017.,<u>http://cires1.colorado.edu/jimenez-group/wiki/index.php/Field_ToF-AMS_Operation</u>, Accessed September 2017.,
- Jimenez, J. L., Jayne, J. T., Shi, Q., Kolb, C. E., Worsnop, D. R., Yourshaw, I., Seinfeld, J. H., Flagan, R. C., Zhang, X., and Smith, K. A.: Ambient aerosol sampling using the aerodyne aerosol mass spectrometer, J. Geophys. Res-Atmos., 108, 8425, 2003.
- Kang, C.-M., Gold, D., and Koutrakis, P.: Downwind O 3 and PM 2.5 speciation during the wildfires in 2002 and 2010, Atmos. Environ., 95, 511-519, 2014.
- Khlystov, A., Stanier, C., and Pandis, S.: An algorithm for combining electrical mobility and aerodynamic size distributions data when measuring ambient aerosol, Aerosol. Sci.
 Technol., 38, 229-238, 2004.
- Kim, E., Hopke, P. K., Kenski, D. M., and Koerber, M.: Sources of fine particles in a rural midwestern US area, Environ. Sci. Technol., 39, 4953-4960, 2005.
 - Kim, M., Deshpande, S. R., and Crist, K. C.: Source apportionment of fine particulate matter (PM2.5) at a rural Ohio River Valley site, Atmos. Environ., 41, 9231-9243, 2007.
- 25 Knorr, W., Jiang, L., and Arneth, A.: Climate, CO2 and human population impacts on global wildfire emissions, Biogeosciences, 13, 267-282, 2016.
 - Kundu, S., and Stone, E. A.: Composition and sources of fine particulate matter across urban and rural sites in the Midwestern United States, Env. Sci. Process. Impact, 16, 1360-1370, 2014.
- Lee, A. K., Herckes, P., Leaitch, W., Macdonald, A., and Abbatt, J.: Aqueous OH oxidation of ambient
 organic aerosol and cloud water organics: Formation of highly oxidized products, Geophys. Res. Lett., 38, 2011.
 - Liu, X., Zhang, Y., Huey, L., Yokelson, R., Wang, Y., Jimenez, J., Campuzano Jost, P., Beyersdorf, A., Blake, D., and Choi, Y.: Agricultural fires in the southeastern US during SEAC4RS: Emissions of trace gases and particles and evolution of ozone, reactive nitrogen, and organic aerosol, J. Geophys. Res-Atmos., 121, 7383-7414, 2016.
 - Liu, Y., Stanturf, J., and Goodrick, S.: Trends in global wildfire potential in a changing climate, Forest. Ecol. Manag., 259, 685-697, 2010.
 - Lu, X., Zhang, L., Yue, X., Zhang, J., Jaffe, D. A., Stohl, A., Zhao, Y., and Shao, J.: Wildfire influences on the variability and trend of summer surface ozone in the mountainous western United States, Atmos. Chem. Phys., 16, 14687-14702, 2016.

- Matsui, H., Koike, M., Kondo, Y., Moteki, N., Fast, J. D., and Zaveri, R. A.: Development and validation of a black carbon mixing state resolved three dimensional model: Aging processes and radiative impact, J. Geophys. Res-Atmos., 118, 2304-2326, 2013.
- May, N. W., Olson, N. E., Panas, M., Axson, J. L., Tirella, P. S., Kirpes, R. M., Craig, R. L., Gunsch, M. J., China, S., Laskin, A., Ault, A. P., and Pratt, K. A.: Aerosol Emissions from Great Lakes Harmful Algal Blooms, Environ. Sci. Technol., 52, 397-405, 10.1021/acs.est.7b03609, 2018.

15

- Middlebrook, A. M., Bahreini, R., Jimenez, J. L., and Canagaratna, M. R.: Evaluation of compositiondependent collection efficiencies for the aerodyne aerosol mass spectrometer using field data, Aerosol. Sci. Technol., 46, 258-271, 2012.
- 10 Miller, D. J., Sun, K., Zondlo, M. A., Kanter, D., Dubovik, O., Welton, E. J., Winker, D. M., and Ginoux, P.: Assessing boreal forest fire smoke aerosol impacts on US air quality: A case study using multiple data sets, J. Geophys. Res-Atmos., 116, 2011.
 - Moffet, R. C., Qin, X., Rebotier, T., Furutani, H., and Prather, K. A.: Chemically segregated optical and microphysical properties of ambient aerosols measured in a single particle mass spectrometer, J. Geophys. Res-Atmos., 113, D12213, 2008.
- Moffet, R. C., and Prather, K. A.: In-situ measurements of the mixing state and optical properties of soot with implications for radiative forcing estimates, Proc. Natl. Acad. Sci., 106, 11872-11877, 2009.
- Moffet, R. C., Henn, T. R., Tivanski, A. V., Hopkins, R. J., Desyaterik, Y., Kilcoyne, A., Tyliszczak, T., Fast,
 J., Barnard, J., and Shutthanandan, V.: Microscopic characterization of carbonaceous aerosol particle aging in the outflow from Mexico City, Atmos. Chem. Phys., 10, 961-976, 2010.
- Müller, D., Mattis, I., Wandinger, U., Ansmann, A., Althausen, D., and Stohl, A.: Raman lidar observations of aged Siberian and Canadian forest fire smoke in the free troposphere over Germany in 2003: Microphysical particle characterization, J. Geophys. Res-Atmos., 110, 2005.
 - National Research Council, and National Academies: Global sources of local pollution: an assessment of long-range transport of key air pollutants to and from the United States, xiii, 234 p., National Academies Press, Washington, D.C., xiii, 234 p. pp., 2010.
- Pastor, S. H., Allen, J. O., Hughes, L. S., Bhave, P., Cass, G. R., and Prather, K. A.: Ambient single particle
 analysis in Riverside, California by aerosol time-of-flight mass spectrometry during the
 SCOS97-NARSTO, Atmos. Environ., 37, 239-258, 2003.
- Paulot, F., Jacob, D. J., Pinder, R., Bash, J., Travis, K., and Henze, D.: Ammonia emissions in the United States, European Union, and China derived by high resolution inversion of ammonium wet deposition data: Interpretation with a new agricultural emissions inventory (MASAGE NH3), I. Geophys. Res-Atmos., 119, 4343-4364, 2014.
 - Petters, M. D., Carrico, C. M., Kreidenweis, S. M., Prenni, A. J., DeMott, P. J., Collett, J. L., and Moosmueller, H.: Cloud condensation nucleation activity of biomass burning aerosol, J. Geophys. Res-Atmos., 114, 2009.

Pope, C. A., and Dockery, D. W.: Health effects of fine particulate air pollution: lines that connect, J. 40 Air Waste Manage. Assoc., 56, 709-742, 2006. Pöschl, U.: Atmospheric Aerosols: Composition, Transformation, Climate and Health Effects, Angew. Chen. Int. Ed., 44, 7520 - 7540, 2005.

Pöschl, U., and Shiraiwa, M.: Multiphase chemistry at the atmosphere–biosphere interface influencing climate and public health in the anthropocene, Chem. Rev., 115, 4440-4475, 2015.

5

- Pratt, K., Murphy, S., Subramanian, R., DeMott, P., Kok, G., Campos, T., Rogers, D., Prenni, A., Heymsfield, A., and Seinfeld, J.: Flight-based chemical characterization of biomass burning aerosols within two prescribed burn smoke plumes, Atmos. Chem. Phys., 11, 12549-12565, 2011.
- 10 Pratt, K. A., and Prather, K. A.: Real-time, single-particle volatility, size, and chemical composition measurements of aged urban aerosols, Environ. Sci. Technol., 43, 8276-8282, 2009.
 - Pratt, K. A., Heymsfield, A. J., Twohy, C. H., Murphy, S. M., DeMott, P. J., Hudson, J. G., Subramanian, R., Wang, Z., Seinfeld, J. H., and Prather, K. A.: In situ chemical characterization of aged biomass-burning aerosols impacting cold wave clouds, J. Atmos. Sci., 67, 2451-2468, 2010.
- 15 Pratt, K. A., and Prather, K. A.: Mass spectrometry of atmospheric aerosols—Recent developments and applications. Part II: On - line mass spectrometry techniques, Mass Spectrom. Rev., 31, 17-48, 2012.
 - Qin, X., Bhave, P. V., and Prather, K. A.: Comparison of two methods for obtaining quantitative mass concentrations from aerosol time-of-flight mass spectrometry measurements, Anal. Chem., 78, 6169-6178, 2006.
 - Qin, X., Pratt, K. A., Shields, L. G., Toner, S. M., and Prather, K. A.: Seasonal comparisons of singleparticle chemical mixing state in Riverside, CA, Atmos. Environ., 59, 587-596, 2012.
- Raatikainen, T., Vaattovaara, P., Tiitta, P., Miettinen, P., Rautiainen, J., Ehn, M., Kulmala, M., Laaksonen, A., and Worsnop, D. R.: Physicochemical properties and origin of organic groups detected in boreal forest using an aerosol mass spectrometer, Atmos. Chem. Phys., 10, 2063-2077, 2010.
 - Rattanavaraha, W., Chu, K., Budisulistiorini, S. H., Riva, M., Lin, Y.-H., Edgerton, E. S., Baumann, K., Shaw, S. L., Guo, H., and King, L.: Assessing the impact of anthropogenic pollution on isoprene-derived secondary organic aerosol formation in PM 2.5 collected from the
- 30 Birmingham, Alabama, ground site during the 2013 Southern Oxidant and Aerosol Study, Atmos. Chem. Phys., 16, 4897-4914, 2016.
 - Riemer, N., and West, M.: Quantifying aerosol mixing state with entropy and diversity measures, Atmos. Chem. Phys., 13, 11423-11439, 2013.
- Rolph, G. D., Draxler, R. R., Stein, A. F., Taylor, A., Ruminski, M. G., Kondragunta, S., Zeng, J., Huang,
 H.-C., Manikin, G., and McQueen, J. T.: Description and verification of the NOAA smoke
 - forecasting system: the 2007 fire season, Weather Forecast., 24, 361-378, 2009. <u>Seinfeld, J. H., and Pandis, S. N.: Atmospheric chemistry and physics: from air pollution to climate</u> change, John Wiley & Sons, Hoboken, New Jersey, 2016.
- Sheesley, R. J., Schauer, J. J., Bean, E., and Kenski, D.: Trends in secondary organic aerosol at a remote site in Michigan's upper peninsula, Environ. Sci. Technol., 38, 6491-6500, 2004.
 - 23

- Silva, P. J., Liu, D.-Y., Noble, C. A., and Prather, K. A.: Size and chemical characterization of individual particles resulting from biomass burning of local Southern California species, Environ. Sci. Technol., 33, 3068-3076, 1999.
- Sjostedt, S., Slowik, J., Brook, J., Chang, R.-W., Mihele, C., Stroud, C., Vlasenko, A., and Abbatt, J.: Diurnally resolved particulate and VOC measurements at a rural site: indication of significant biogenic secondary organic aerosol formation, Atmos. Chem. Phys., 11, 5745-5760, 2011.

10

25

- Slowik, J., Stroud, C., Bottenheim, J., Brickell, P., Chang, R.-W., Liggio, J., Makar, P., Martin, R., Moran, M., and Shantz, N.: Characterization of a large biogenic secondary organic aerosol event from eastern Canadian forests, Atmos. Chem. Phys., 10, 2825-2845, 2010.
- Slowik, J., Brook, J., Chang, R.-W., Evans, G., Hayden, K., Jeong, C.-H., Li, S.-M., Liggio, J., Liu, P., and McGuire, M.: Photochemical processing of organic aerosol at nearby continental sites: contrast between urban plumes and regional aerosol, Atmos. Chem. Phys., 11, 2991-3006, 2011.
- 15 Smith, S., and Mueller, S.: Modeling natural emissions in the Community Multiscale Air Quality (CMAQ) Model–I: building an emissions data base, Atmos. Chem. Phys., 10, 4931-4952, 2010.
 - Song, X.-H., Hopke, P. K., Fergenson, D. P., and Prather, K. A.: Classification of single particles analyzed by ATOFMS using an artificial neural network, ART-2A, Anal. Chem., 71, 860-865, 1999.
- 20 Spencer, M. T., Shields, L. G., and Prather, K. A.: Simultaneous measurement of the effective density and chemical composition of ambient aerosol particles, Environ. Sci. Technol., 41, 1303-1309, 2007.
 - Stephen, K., and Aneja, V. P.: Trends in agricultural ammonia emissions and ammonium concentrations in precipitation over the Southeast and Midwest United States, Atmos. Environ., 42, 3238-3252, 2008.
- Stockwell, C., Yokelson, R., Kreidenweis, S., Robinson, A., DeMott, P., Sullivan, R., Reardon, J., Ryan, K., Griffith, D., and Stevens, L.: Trace gas emissions from combustion of peat, crop residue, domestic biofuels, grasses, and other fuels: configuration and Fourier transform infrared (FTIR) component of the fourth Fire Lab at Missoula Experiment (FLAME-4), Atmos. Chem.
 30 Phys., 9727, 2014.
 - Su, Y., Sipin, M. F., Furutani, H., and Prather, K. A.: Development and characterization of an aerosol time-of-flight mass spectrometer with increased detection efficiency, Anal. Chem., 76, 712-719, 2004.
- Sueper, D.: ToF-AMS analysis software, Available at: <u>http://cires1.colorado.edu/jimenez-</u> 35 <u>group/ToFAMSResources/ToFSoftware/index.html</u>, 2010.
- Sullivan, R., Guazzotti, S., Sodeman, D., and Prather, K.: Direct observations of the atmospheric processing of Asian mineral dust, Atmos. Chem. Phys., 7, 1213-1236, 2007.
 - Sun, Y., Zhang, Q., Macdonald, A. M., Hayden, K., Li, S. M., Liggio, J., Liu, P. S. K., Anlauf, K. G., Leaitch, W. R., Steffen, A., Cubison, M., Worsnop, D. R., van Donkelaar, A., and Martin, R. V.: Sizeresolved aerosol chemistry on Whistler Mountain, Canada with a high-resolution aerosol
 - 24

mass spectrometer during INTEX-B, Atmos. Chem. Phys., 9, 3095-3111, 10.5194/acp-9-3095-2009, 2009.

- Toner, S. M., Sodeman, D. A., and Prather, K. A.: Single particle characterization of ultrafine and accumulation mode particles from heavy duty diesel vehicles using aerosol time-of-flight mass spectrometry, Environ. Sci. Technol., 40, 3912-3921, 2006.
- Toner, S. M., Shields, L. G., Sodeman, D. A., and Prather, K. A.: Using mass spectral source signatures to apportion exhaust particles from gasoline and diesel powered vehicles in a freeway study using UF-ATOFMS, Atmos. Environ., 42, 568-581, 2008.

5

20

- Uno, I., Eguchi, K., Yumimoto, K., Takemura, T., Shimizu, A., Uematsu, M., Liu, Z., Wang, Z., Hara, Y.,
 and Sugimoto, N.: Asian dust transported one full circuit around the globe, Nat. Geosci., 2,
 557-560, 2009.
 - VanReken, T., Mwaniki, G., Wallace, H., Pressley, S., Erickson, M., Jobson, B., and Lamb, B.: Influence of air mass origin on aerosol properties at a remote Michigan forest site, Atmos. Environ., 107, 35-43, 2015.
- 15 Veira, A., Lasslop, G., and Kloster, S.: Wildfires in a warmer climate: Emission fluxes, emission heights, and black carbon concentrations in 2090–2099, J. Geophys. Res-Atmos., 121, 3195-3223, 2016.
 - Wang, J., Cubison, M., Aiken, A., Jimenez, J., and Collins, D.: The importance of aerosol mixing state and size-resolved composition on CCN concentration and the variation of the importance with atmospheric aging of aerosols, Atmos. Chem. Phys., 10, 7267-7283, 2010a.
 - Wang, Y., Huang, J., Zananski, T. J., Hopke, P. K., and Holsen, T. M.: Impacts of the Canadian forest fires on atmospheric mercury and carbonaceous particles in northern New York, Environ. Sci. Technol., 44, 8435-8440, 2010b.
- Wiedinmyer, C., Quayle, B., Geron, C., Belote, A., McKenzie, D., Zhang, X., O'Neill, S., and Wynne, K. K.:
 Estimating emissions from fires in North America for air quality modeling, Atmos. Environ., 40, 3419-3432, 2006.
 - Xu, L., Guo, H., Boyd, C. M., Klein, M., Bougiatioti, A., Cerully, K. M., Hite, J. R., Isaacman-VanWertz, G., Kreisberg, N. M., and Knote, C.: Effects of anthropogenic emissions on aerosol formation from isoprene and monoterpenes in the southeastern United States, Proc. Natl. Acad. Sci., 112, 37-42, 2015.
 - Zhang, Y., Sheesley, R. J., Schauer, J. J., Lewandowski, M., Jaoui, M., Offenberg, J. H., Kleindienst, T. E., and Edney, E. O.: Source apportionment of primary and secondary organic aerosols using positive matrix factorization (PMF) of molecular markers, Atmos. Environ., 43, 5567-5574, 2009.
- 35 Zhou, S., Collier, S., Jaffe, D. A., Briggs, N. L., Hee, J., Sedlacek III, A. J., Kleinman, L., Onasch, T. B., and Zhang, Q.: Regional influence of wildfires on aerosol chemistry in the western US and insights into atmospheric aging of biomass burning organic aerosol, Atmos. Chem. Phys., 17, 2477-2493, 2017.

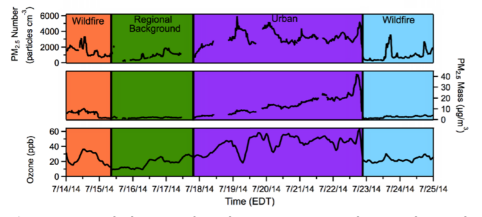


Figure 1. Time-resolved PM_{2.5} number and mass concentrations and ozone mole ratios during the different periods of air mass influence. Periods without data are due to instrument down time. Colors of the different time periods correspond to the colors of the corresponding HYSPLIT backward air mass trajectories in Figure S2.

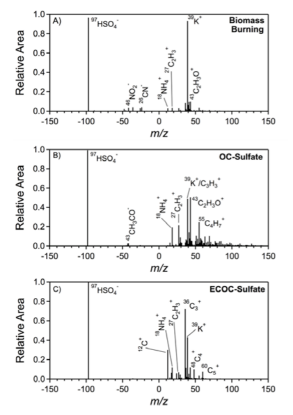


Figure 2. Average positive and negative ion single-particle mass spectra (ATOFMS), with characteristic peaks labeled, for the dominant aged combustion particle types observed: (A) biomass burning, (B) OC-sulfate, and (C) ECOC-sulfate.

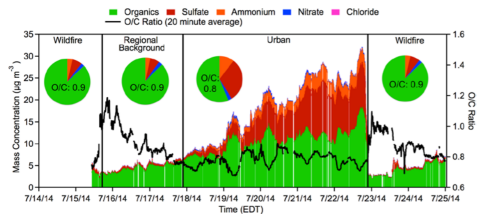


Figure 3. PM_1 non-refractory chemically speciated mass concentrations, as well as O/C ratios (20 min averages), measured by HR-AMS. Periods of influence are notated and separated by solid vertical lines. Pie charts represent the average mass fractions for each air mass period, with average O/C ratio inset.

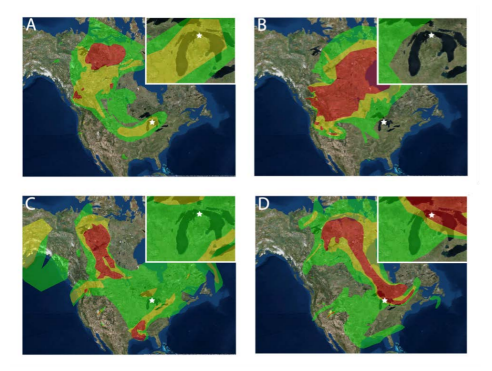


Figure 4. Representative NOAA HMS smoke maps for four representative days during the time periods of different air influence: (A) July 14, wildfire influence; (B) July 16, remote background influence; (C) July 21, Urban influence; (D) July 24, wildfire influence. Inset enlarges the state of Michigan to clearly display smoke influence on the field site, shown as a star.

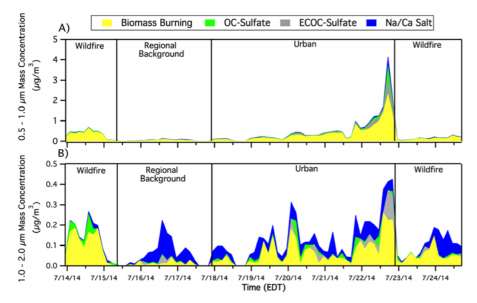
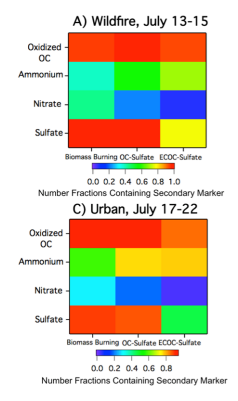
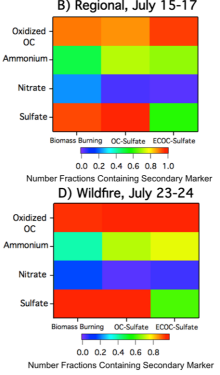


Figure 5. Three hour binned mass concentrations of (A) $0.5 - 1.0 \mu m$ and (B) $1.0 - 2.0 \mu m$ particle types, as measured by ATOFMS. Gaps in the data correspond to periods when APS data were not available for scaling.





B) Regional, July 15-17

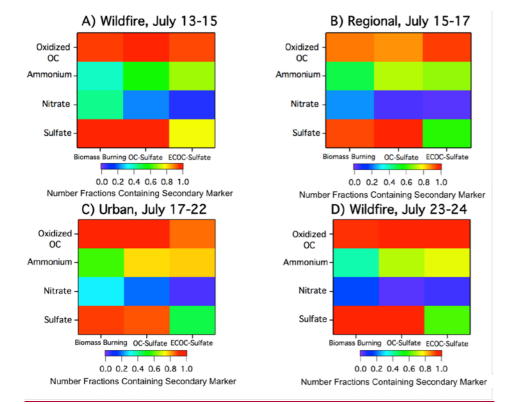


Figure 6. Number fractions of individual biomass burning, OC-sulfate, and ECOC-sulfate particles that were internally mixed with secondary species, as determined by ATOFMS ion markers, including oxidized OC ($C_2H_3O^+$, m/z 43), ammonium (NH_4^+ , m/z 18), nitrate (NO_2^- , m/z -46, and/or NO_3^- , m/z -62), and sulfate (HSO_4^- , m/z -97). Since a given particle can contain more than one secondary species, the number fractions can add to greater than one for a given particle type. Chemical mixing states are provided here for the four air mass time periods: (A) wildfire influence from July 13-15, (B) clean air from northern Canada from July 15-17, (C) mix of wildfire and urban influences from July 17-22, (D) mix of clean air and Canadian wildfires from July 23-24.

Supplemental Information for Ubiquitous Influence of Wildfire Emissions and Secondary Organic Aerosol on Summertime Atmospheric Aerosol in the Forested Great Lakes Region

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1 Supporting Measurements

Meteorological data (Figure S1), including wind direction, wind speed, relative humidity, and temperature, were collected by a Vaisala WXT510 weather sensor located at the top of the PROPHET tower. Variations in meteorological conditions throughout the study, and average meteorological conditions during each period of influence, are discussed in the main text. In order to determine the origin of the influential air masses (Figure S2), backward air mass trajectories were calculated using the NOAA Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) Model (Stein et al., 2015). A final altitude of 500 m AGL was used for the field site, with each trajectory modeling the proceeding 72 h. During each of the three influence air mass locations, median, was well as 25th and 75th percentile aerosol

10 number, mass, and size distributions were calculated based on SMPS measurements (Figures S3 and S4). Predicted NH₄⁺ (Figure S4) was calculated using the methods described by Sueper (2010). Though these calculations indicate the aerosol is likely acidic, there are caveats associated with these calculations, as outlined by Hennigan et al. (2015). Therefore, the pH cannot be reliably calculated beyond a qualitative indication of whether or not the aerosol is acidic.

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References

- Hennigan, C.J., Izumi, J., Sullivan, A.P., Weber, R.J., Nenes, A., 2015. A critical evaluation of proxy methods used to estimate the acidity of atmospheric particles. Atmos. Chem. Phys. 15, 2775-2790.
- Stein, A., Draxler, R., Rolph, G., Stunder, B., Cohen, M., Ngan, F., 2015. NOAA's HYSPLIT atmospheric transport and dispersion modeling system. B. Am. Meteorol. Soc. 96, 2059-2077.
 - Sueper, D., 2010. ToF-AMS analysis software. Available at: http://cires1.colorado.edu/jimenezgroup/ToFAMSResources/ToFSoftware/index.html.
- VanReken, T., Mwaniki, G., Wallace, H., Pressley, S., Erickson, M., Jobson, B., and Lamb, B.: Influence of air mass origin on aerosol properties at a remote Michigan forest site, Atmos. Environ., 107, 35-43, 2015.

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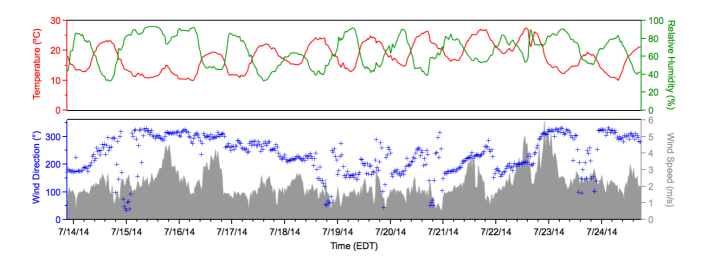


Figure S1. Meteorological conditions measured from a height of ~30 m at the UMBS PROPHET Tower.

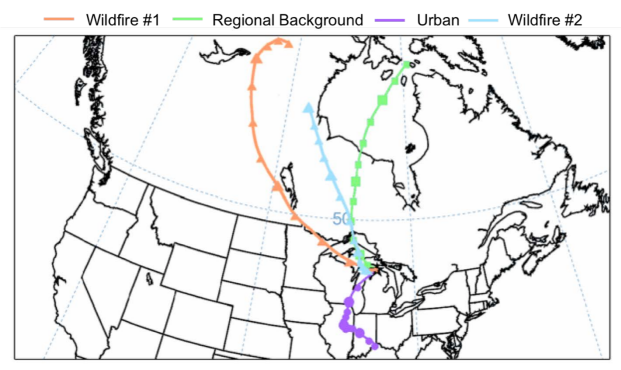


Figure S2. Representative 72 h HYSPLIT back trajectories with a final altitude of 500 m for the four air mass influences, with markers indicating 6 h intervals. Trajectory start times were: Wildfire #1: 7/14/2014 07:00 EDT, Regional Background: 7/17/2014 07:00 EDT, Urban: 7/21/2014 07:00 EDT, Wildfire #2: 7/24/14 07:00 EDT. Colors correspond to the air mass of influence indicated in Figure 3.

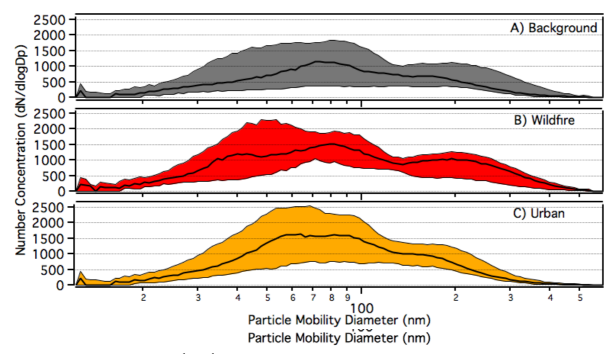


Figure S3. Median and 25th/75th percentiles of size-resolved particle number concentration distributions, as measured by SMPS, during the three air mass periods of interest: (A) Background, (B) Wildfire, and (C) Urban. For comparison, particle size distributions by air mass origin at UMBS in summer 2009 were previously discussed in detail by VanReken et al. (2015).

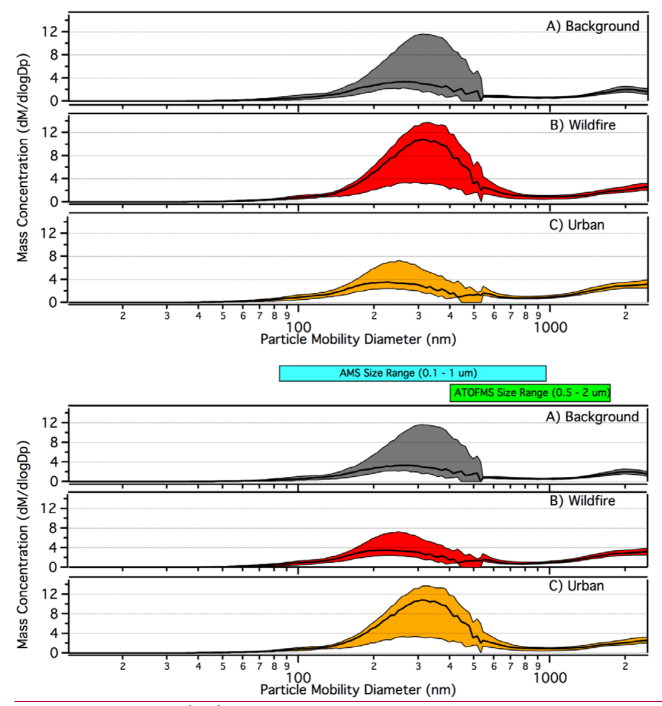


Figure S4. Median and 25th/75th percentiles of size-resolved particle mass distributions, as measured by SMPS and APS (assuming a density of 1.5 g cm⁻³), during the three air mass periods of interest: (A) Background, (B) Wildfire, and (C) Urban. For comparison, the size range of measurements (converted

from vacuum aerodynamic diameter to mobility diameter) made by the AMS (blue) and ATOFMS (green) throughout the study are notated above the figure.

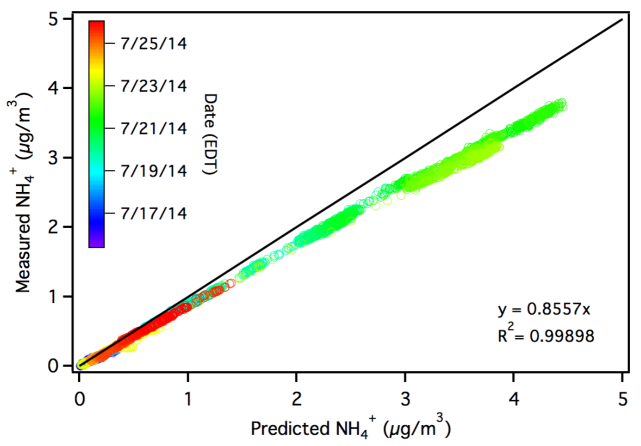


Figure S5. Ammonium balance calculated from predicted ammonium versus measured ammonium from the HR-AMS, following the method of Sueper (2010). A 1:1 line is showed in black for reference.

