Organic tracers of fine aerosol particles in central Alaska: summertime composition and sources

Dhananjay Kumar Deshmukh1, M. Mozammel Haque1,2, Yongwon Kim3, and Kimitaka Kawamura1

1Chubu Institute for Advanced Studies, Chubu University, Kasugai 487-8501, Japan
2Yale-NUIST Center on Atmospheric Environment, School of Applied Meteorology, Nanjing University of Information Science and Technology, Nanjing 210044, China
3International Arctic Research Center, University of Alaska Fairbanks, Fairbanks 99775, USA

Correspondence: Kimitaka Kawamura (kkawamura@isc.chubu.ac.jp)

Received: 27 March 2019 – Discussion started: 21 May 2019
Revised: 8 October 2019 – Accepted: 14 October 2019 – Published: 21 November 2019

Abstract. PM2.5 aerosols were collected at Fairbanks (64.51° N and 147.51° W) in central Alaska during the summer of 2009 and analyzed for organic tracer compounds using a gas chromatograph–mass spectrometer. The organic compounds were grouped into 14 classes based on their functional groups and sources. Concentrations of the total organics measured ranged from 113 to 1664 ng m$^{-3}$ (avg 535 ng m$^{-3}$). Anhydrosugars (avg 186 ng m$^{-3}$) and $n$-alkanoic acids (avg 185 ng m$^{-3}$) were 2 major classes among the 14 compound classes. The similar temporal trends and strong positive correlations among anhydrosugars and $n$-alkanoic acids demonstrated that biomass burning (BB) is the major source of organic aerosols (OAs) in central Alaska. The dominance of higher molecular weight $n$-alkanoic acids over lower molecular weight homologs and their carbon preference index (5.6–9.8) confirmed that they were mostly emitted from plant waxes during BB in central Alaska. The concentration ratios of levoglucosan to mannosan denoted that softwood is the main biomass burned. The rainfall event distinctly enhanced the levels of mannitol and arabitol due to the growth of fungi and active discharge of fungal spores in the subarctic region. Molecular compositions of biogenic secondary organic aerosol (BSOA) tracers inferred that isoprene is a crucial precursor of BSOA over central Alaska. Our results suggest forest fires and plant emissions to be the crucial factors controlling the levels and molecular composition of OAs in central Alaska. We propose that PM$_{2.5}$ laden with OAs derived in central Alaska may significantly impact the air quality and climate in the Arctic via long-range atmospheric transport.

1 Introduction

Atmospheric aerosols can absorb and scatter solar radiation and alter the radiative forcing of the atmosphere (Seinfeld and Pandis, 1998; Wilkening et al., 2000). Fine aerosol particles have a diameter size close to the wavelengths of visible lights and thus are expected to have a stronger climatic impact than coarse particles (Kanakidou et al., 2005). They can also be transported far away from the source regions, and thus their climatic and environmental effects are delocalized compared to the emission areas. Aerosol particles that are hydrophilic can act as cloud condensation nuclei (CCN) and have an indirect climatic effect through modification of cloud properties (Novakov and Penner, 1993; Novakov and Corri-gan, 1996).

Organic aerosols (OAs) that are comprised of thousands of organic compounds contribute about 20 % to 50 % of the total mass of fine particles in the continental mid-latitudinal atmosphere (Saxena and Hildemann, 1996), whereas it is around 90 % in tropical forest areas (Crutzen and Andreae, 1990; Andreae and Rosenfeld, 2008). They are derived from anthropogenic and natural sources. They can alter the physical and chemical properties of atmospheric particles depending on the meteorological conditions. OAs are highlighted for the past decade because they are related to the changes in global and regional climate and chemical composition of the atmosphere as well as public health. Primary organic aerosols (POA) are directly emitted as particulate forms, whereas secondary organic aerosols (SOA) refer to particulate organic matters that are transformed to the aerosol phase via gas-phase oxidation of organic precursors. Emissions of POA...
particles and SOA precursors can be released from numerous sources near the ground surface and subsequently mixed in the boundary layer and to a lesser extent in the free troposphere. The dry depositional removal of OAs mainly depends on the sizes of the aerosol particles.

The molecular composition of OAs can be used as a tracer to better understand the sources and formation pathways. Advances were made during the last decade to better understand the formation of OAs and their precursors in the atmosphere. On a global scale, the emission of biogenic volatile organic compounds (VOCs) is 1 order of magnitude higher than that of anthropogenic VOCs (Seinfeld and Pandis, 1998). It is notable that biogenic VOCs are comprised of unsaturated hydrocarbons with double bonds and are more reactive towards the atmospheric oxidants such as hydroxyl (OH) radicals and ozone ($\text{O}_3$) than anthropogenic VOCs that are largely comprised of aromatic hydrocarbons. This specific feature of biogenic VOCs further enhances their significance as a conceivable supplier to the global burden of OAs in the atmosphere. Laboratory and chamber experiments have also documented that biogenic VOCs are the potential precursor for SOA formation in the atmosphere (Kavouras et al., 1998; Jaoui et al., 2007).

Although early Arctic explorers had noticed atmospheric haze (Nordenskiold, 1883), the remote Arctic atmosphere was believed to be extremely clean. Pilots flying over the North American Arctic in the 1950s observed widespread haze that could be seen every winter and early spring (Mitchell, 1957). It took until the 1970s for scientists to realize that the haze was air pollution transported from the middle latitudes (Barrie, 1986). Over the past 3 decades there has been much research on the climate consequences of this pollution that is also present in summer. Surface air temperature has increased more than the global average over the past few decades and is predicted to warm by about 5°C over a large part of the Arctic by the end of the 21st century (IPCC, 2001). The Arctic atmosphere is considered a unique natural laboratory for photochemical reactions and transformations during the polar sunrise (Kawamura et al., 1996). Arctic atmosphere is influenced by marine-derived OAs from the Arctic Ocean as well as continentally derived OAs and their precursors from mid-latitudes in Eurasia or North America (Stohl et al., 2006; Law and Stohl, 2007).

Previous analyses have reported a substantial contribution of summertime boreal forest fires to the chemical composition of aerosol over the Arctic (Iziomon et al., 2006; Kaplan and New, 2006; Stohl et al., 2006). French et al. (2003) proposed that wildfire contributed a substantial amount of carbon-based gas from 1950 to 1999 in the atmosphere of the boreal region of Alaska. Based on the modeling and in situ observations of black carbon (BC) and soot during the FROSTFIRE campaign, Kim et al. (2005) revealed that BC and soot particles of 0.4 to 10 µm in radius can be transported to the Arctic and the whole area of Alaska in a very short time. The results of Kaplan and New (2006) delivered strong evidence that high-latitude ecosystems are sensitive to climate change due to the increase in concentrations of greenhouse gases. Iziomon et al. (2006) examined summertime aerosols based on column-integrated and surface aerosol measurements at Borrow in the North Slope of Alaska between 1998 and 2003. They noticed high loadings of aerosols at least 8 d each summer and demonstrated that the pollution events with the highest aerosol loadings were associated with smoke from wildfires in northwest Canada. Stohl et al. (2006) explored the impact of boreal forest fire emissions on the light-absorbing aerosol levels at the Barrow Arctic station. They proposed that boreal forest fires could result in elevated concentrations of light-absorbing aerosols throughout the entire Arctic with an impact on the radiation transmission of the Arctic atmosphere.

The results of Hegg et al. (2009) and Warneke et al. (2009) confirm that biomass burning (BB) causes a more efficient transport and deposition of BC aerosol in Arctic snow, causing a strong climate forcing at high latitudes. Based on in situ measurements in the Arctic and a transport model of carbon monoxide (CO), Warneke et al. (2010) proposed that BB plumes transported to the Arctic in spring in 2008 more than doubled the Arctic atmospheric burden in other seasons. The results of Ward et al. (2012) based on chemical mass balance modeling revealed that wood smoke was the major source of PM$_{2.5}$ particles, mainly during the winter months at several locations in Fairbanks. Ward et al. (2012) and Wang and Hopke (2014) demonstrate that Arctic air pollution could be so severe that the city of Fairbanks has been labeled a serious non-attainment area by the United States Environmental Protection Agency. Biogenic emissions from boreal forest largely increase during the summertime growing season. The year-round measurements conducted at Fairbanks by Haque et al. (2016) have shown that SOA derived from biogenic VOC emissions dominated the organic chemical composition of total suspended particles during summer in central Alaska. They found higher contributions of isoprene oxidation products than monoterpene and sesquiterpene oxidation products to SOA formation in summer due to the higher isoprene emissions and high levels of oxidants. They estimated isoprene-derived secondary organic carbon (SOC) approximately 5 times higher than SOA derived from monoterpene and nearly 2 times higher than sesquiterpene-derived SOA in central Alaska. Because climate change is generally proceeding fastest at the high latitudes (Serreze et al., 2000; Hinzman et al., 2005), there is an increasing demand for better understanding of the chemical compositions and sources of OAs in the Arctic atmosphere.

We collected PM$_{2.5}$ samples during the summer of 2009 at Fairbanks in central Alaska. The samples were analyzed for several organic tracer compounds to characterize OAs in the North American subarctic region. This paper discusses the molecular compositions of various organic compound classes and the factors controlling temporal changes in their concentrations in central Alaska. We also discuss the sources...
of organic compounds detected and the secondary formation processes as well as atmospheric implications for the burden of OAs in the Arctic and subarctic atmosphere.

2 Methodology

2.1 Description of the sampling area

Alaska is located in a subarctic zone. Fairbanks is the largest city in central Alaska. It is situated in the central Tanana Valley connecting the Chena River near the Tanana River. The location of the sampling site in Fairbanks (64.51° N and 147.51° W) and its surroundings are shown in Fig. 1. The altitude of the sampling location is 136 m above sea level. The total area of Fairbanks is nearly 85 km², with a population of 31,500. The sampling site is located at the downwind side of Fairbanks, where a forest is very close to the campus of the University of Alaska Fairbanks. The highest levels of atmospheric aerosol burden in the United States have been recorded in Fairbanks (Ward et al., 2012). The National Emission Inventory database pointed out that forest fires and combustion of fossil fuels are the two critical sources of air pollution in Fairbanks (Shakya and Peltier, 2013; Ware et al., 2013).

2.2 Sample collection

Atmospheric particle samples of sizes less than 2.5 µm in diameter (PM$_{2.5}$) were collected on the rooftop of the International Arctic Research Center building of the University of Alaska Fairbanks during the summer season from 5 June to 21 September in 2009 when a forest fire was active in the region. The collection of samples was performed using a low-volume air sampler model, URG-2000-39EH (USA), with a flow rate of 16.7 L min$^{-1}$. PM$_{2.5}$ particles were retained on a quartz fiber filter of 47 mm in diameter that was precombusted at 450°C for 6 h. The sampler was operated for 3 to several days to get enough aerosol particles on the filter to detect trace organic species with very low concentrations. We collected 13 samples (Alaska 01 to 13) and three field blanks during the campaign. The samples and field blank filters were individually placed in a pre-heated glass vial with a Teflon-lined screw cap. We stored the aerosol samples in a dark room at −20°C to prevent the samples from microbial degradation and loss of semivolatile organic compounds.

2.3 Analysis of organic tracers

We analyzed the samples for organic compounds using a gas chromatograph–mass spectrometer (GC–MS) system: a Hewlett-Packard (HP) model 6890 GC coupled to an HP model 5973 mass-selective detector. A 5.0 cm² filter area of each aerosol sample was extracted with a 10 mL dichloromethane (CH$_2$Cl$_2$) and methanol (CH$_3$OH) mixture (2 : 1) through ultrasonication (10 min × 3). The solvent extracts were filtered through a Pasteur pipet packed with pre-combusted (450°C for 6 h) quartz wool to remove particles and filter debris. The extracts were concentrated by a rotary evaporator and then dried under a stream of pure nitrogen gas. The hydroxyl (OH) and carboxyl (COOH) groups of organic compounds in the extracts were derivatized to trimethylsilyl ethers and esters, respectively, by the reaction with 50 µL N,O-bis(trimethylsilyl)trifluoroacetamide including 1% trimethylsilyl chloride and 10 µL pyridine at 70°C for 3 h (Schauer et al., 1996; Simoneit et al., 2004a). n-Hexane containing 1.43 ng µL$^{-1}$ of a C$_{13}$ n-alkane internal standard (40 µL) was added into the derivatives before injection of the sample into a GC–MS.

The separation of compounds was performed on a 30 m long DB-5MS fused silica capillary column (0.25 mm inner diameter and 0.25 µm film thickness). Helium was used as a carrier gas at a flow rate of 1.0 mL min$^{-1}$. The GC oven temperature was programmed from 50°C for 2 min to 120°C at 30°C min$^{-1}$ and then 300°C at 6°C min$^{-1}$ with a final isotherm held at 300°C for 16 min. The sample was injected on a splitless mode with an injector temperature of 280°C. The mass detection was conducted at 70 eV on an electron ionization mode with a scan range of 50 to 650 Dalton. The organic compounds were determined by the comparison of the GC retention times and mass fragmentation patterns of a sample with those of authentic standards and National Institute of Standards and Technology library data. The mass spectral data were acquired and processed using HP Chemstation software. The GC–MS relative response factor of each compound was calculated using authentic standards or surrogate compounds. The recoveries of authentic standards or surrogates were above 80% for target compounds. The data reported here were not corrected for recoveries. The relative standard deviation of the measurements based on duplicate analyses was within 10%. The field blank filters were analyzed by the procedure described above. The target compounds were not detected in the blank filters.

2.4 Meteorology and air mass trajectories

Figure 2 shows temporal changes in daily average meteorological parameters at the campaign site. The daily mean temperature was in a range of 2.0 to 33°C with an average of 13.9°C, whereas the daily average relative humidity ranged from 19% to 99% with a mean of 63%. The mean wind speed was 5.2 km h$^{-1}$ and the total rainfall was 122 mm during the sampling period. The 5 d air mass backward trajectories at the height of 500 m above the ground level were computed from the Hybrid Single Particle Lagrangian Integrated Trajectory model (Draxler and Rolph, 2013). The air mass backward trajectories arriving over the observation site during the collection of aerosol samples is presented in Fig. 3.
Figure 1. The location of the sampling site at Fairbanks in central Alaska (64.51° N and 147.51° W) and its surrounding regions. The map was downloaded from Map data © 2018 Google.

Figure 2. The daily average variations of meteorological parameters from 5 June to 21 September 2009 at the observation site in central Alaska.

3 Results and discussion

3.1 Overview of the molecular composition of organic aerosols

A total of 96 organic compounds were detected in PM$_{2.5}$ samples collected at Fairbanks during the sampling period. We grouped them into 14 compound classes as listed in Table 1 together with the mean concentrations and ranges. Figure 4 shows the chemical compositions of OAs in individual samples (Alaska 01 to 13). The levels of all the quantified organic compounds in Alaskan samples ranged from 113 to 1664 ng m$^{-3}$ (avg 535 ng m$^{-3}$) with the predominance of anhydrosugars (avg 186 ng m$^{-3}$) and $n$-alkanoic acids (avg 185 ng m$^{-3}$). Anhydrosugars are produced by pyrolysis of cellulose and hemicellulose, followed by the subsequent emission into the atmosphere, and are widely used as specific tracers of BB (Simoneit et al., 1999; Sang et al., 2013). $n$-Alkanoic acids are derived directly from the surface of plant leaves and marine phytoplankton as well as BB and meat cooking (Kawamura and Gagosian, 1987; Rogge et al., 1993; Fine et al., 2001). Sugar alcohols were de-
tected in ample amounts in three samples collected during the end of the campaign. We also detected a substantial amount of isoprene-derived SOA tracers and \( n \)-alkanols in Alaskan samples, while the concentrations of other compound classes are relatively low.

### 3.2 Anhydrosugars and lignin and resin products: tracers of biomass burning

The combustion of biopolymers including cellulose and hemicellulose as well as lignin and suberin produces several organic molecules that have been recognized as important source tracers of OAs. Anhydrosugars and lignin and resin acids are specific tracers of BB among the pyrolysis products of biopolymers (Simoneit et al., 1999). The pyrolysis of cellulose and hemicellulose produces anhydrosugars such as levoglucosan and its isomers mannosan and galactosan (Simoneit et al., 1999; Engling et al., 2009). Figure 5a presents molecular distributions of anhydrosugars in Alaskan PM\(_{2.5}\) samples. Levoglucosan is the dominant anhydrosugar, followed by mannosan and galactosan in Alaskan samples. Their concentrations ranged from 23 to 463 ng m\(^{-3}\) (avg 125 ng m\(^{-3}\)), 4.1 to 180 ng m\(^{-3}\) (avg 36 ng m\(^{-3}\)) and 3.5 to 106 ng m\(^{-3}\) (avg 26 ng m\(^{-3}\)), respectively.

Because 90% of levoglucosan exists in the atmospheric particles with an aerodynamic diameter less than 2 \( \mu \text{m} \) (Graham et al., 2000; Deshmukh et al., 2012), it is reasonable to compare the levoglucosan concentrations of Alaskan PM\(_{2.5}\) samples with those reported in PM\(_{10}\) and TSP during summer or the BB season. We found that the concentration levels of levoglucosan in central Alaska are substantially higher than those from the Bering Sea (10 ng m\(^{-3}\)) and Arctic Ocean (5.2 ng m\(^{-3}\)) (Hu et al., 2013), Chichijima (0.24 ng m\(^{-3}\)) and Okinawa (0.57 ng m\(^{-3}\)) islands in the western North Pacific (Verma et al., 2015; Zhu et al., 2015), northern Japan (7.8 ng m\(^{-3}\)) (Agarwal et al., 2010), and Mt. Everest (47.2 ng m\(^{-3}\)) (Cong et al., 2015). They are comparable to those reported at the urban site of Chennai in India (avg 111 ng m\(^{-3}\)) (Fu et al., 2010) but lower than those reported at the rural site of Lumbini in Nepal (avg 771 ng m\(^{-3}\)) (Wan et al., 2017), the forest site of Rondônia in Brazil (avg 1180 ng m\(^{-3}\)) (Graham et al., 2012).
et al., 2002), and Chiang Mai in Thailand (avg 1222 ng m\(^{-3}\)) (Thepnuan et al., 2019).

The emission strength of BB products and their long-range atmospheric transport influence the atmospheric levels of anhydrosugars. The backward trajectories reveal that air masses mostly came from the ocean during the campaign (Fig. 3). This result shows that anhydrosugars present in the Alaskan aerosols were mainly associated with the local and regional BB during the campaign. The higher level of levoglucosan in Fairbanks than other sites in the Arctic implies a possible effect of BB on the air quality and climate in the Arctic region. Stocks et al. (2000) and Grell et al. (2011) proposed that the frequency of boreal forest fires recently increased in summer due to global warming. Figure 6a\textendash}c show the temporal trends of anhydrosugars in the Alaskan aerosols. The levels of anhydrosugars expressively alter during the campaign period. The lower levoglucosan levels were found at the beginning of the campaign, whereas they became very high (241 to 50 ng m\(^{-3}\)) on 4–23 July (Fig. 6a). Another peak of levoglucosan was found on 30 July to 4 August (169 ng m\(^{-3}\)). The concentrations of levoglucosan decreased towards the end of the campaign (23 to 50 ng m\(^{-3}\)). Forest fire smokes were seen during 4–23 July and 30 July to 4 August over central Alaska. This observation demonstrates that levoglucosan levels became high due to the local forest fire in central Alaska. mannosan and galactosan presented similar temporal variations to levoglucosan (Fig. 6b and c). The chemical reaction of anhydrosugars with OH radicals could also influence their concentrations in the atmosphere. Although previous studies have reported that levoglucosan can remain stable in the atmosphere for around 10 d with no substantial degradation (Fraser and Lakshmanan, 2000; Schkolnik and Rudich, 2006), recent findings (Hoffmann et al., 2009; Hennigan et al., 2010; Gensch et al., 2018) reported significant chemical reactivity of levoglucosan and have raised a question over the stability of levoglucosan in the atmosphere. Hennigan et al. (2010) carried out a smog chamber experiment and reported the lifetime of atmospheric levoglucosan to be 0.7 to 2.2 d when exposed to 10^6 molecules of OH cm\(^{-3}\). This lifetime is within the range of 0.5 to 3.4 d predicted by Hoffmann et al. (2009) using the Spectral Aerosol Cloud Chemistry Interaction Model. Lai et al. (2014) found that the atmospheric lifetime of levoglucosan ranged from 1.2 to 3.9 d by the control experiment integrating OH in a flow reactor under different environmental conditions and different mixing states. Nevertheless, Bai et al. (2013) reported an atmospheric lifetime of levoglucosan to be 26 d when exposed to an OH level of 2 \times 10^6 molecules cm\(^{-3}\) that is much longer than other predictions.

It is notable from the above discussion that the degradation of levoglucosan is mostly induced by photochemical aging via oxidation by OH radicals during long-range transport. Therefore, the degradation of levoglucosan could be insignificant if the receptor site is close to the source region. As discussed previously, anhydrosugars detected in Alaskan aerosols during the campaign originated from local and regional BB, and we consider that the degradation of anhydrosugars may not be important for explaining the low levels of BB tracers in the samples collected at the beginning and end of the campaign. The low concentrations of anhydrosugars during the beginning and end of the campaign might be caused by the decreased emission rate of BB tracers due to lower BB activities in the source region. Wet deposition may be another reason to lower the level of anhydrosugars in aerosol samples collected at the beginning and end of the campaign because we observed rainfall especially on 5 June to 3 July and 6 August to 17 September in Fairbanks (Fig. 2). Although the concentrations of both mannosan and galactosan are much lower than levoglucosan (Fig. 5a), we observed strong positive correlations \((r = 0.94–0.97)\) among

### Table 1. Concentrations (ng m\(^{-3}\)) of organic tracer compound classes detected in PM\(_{2.5}\) aerosols from central Alaska.

<table>
<thead>
<tr>
<th>Compound classes</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Mean</th>
<th>Median</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anhydrosugars</td>
<td>31</td>
<td>749</td>
<td>186</td>
<td>69</td>
<td>217</td>
</tr>
<tr>
<td>Lignin acids</td>
<td>0.5</td>
<td>15</td>
<td>3.8</td>
<td>2.2</td>
<td>4.2</td>
</tr>
<tr>
<td>Resin acid</td>
<td>0.9</td>
<td>19</td>
<td>6.1</td>
<td>5.1</td>
<td>4.7</td>
</tr>
<tr>
<td>n-Alkanes</td>
<td>0.5</td>
<td>77</td>
<td>24</td>
<td>21</td>
<td>23</td>
</tr>
<tr>
<td>n-Alkanols</td>
<td>5.3</td>
<td>119</td>
<td>46</td>
<td>29</td>
<td>38</td>
</tr>
<tr>
<td>n-Alkanoic acids</td>
<td>9.2</td>
<td>562</td>
<td>185</td>
<td>82</td>
<td>209</td>
</tr>
<tr>
<td>Primary sugars</td>
<td>0.3</td>
<td>44</td>
<td>13</td>
<td>12</td>
<td>11</td>
</tr>
<tr>
<td>Sugar alcohols</td>
<td>1.0</td>
<td>24</td>
<td>14</td>
<td>18</td>
<td>7.4</td>
</tr>
<tr>
<td>Phthalate esters</td>
<td>0.4</td>
<td>6.6</td>
<td>1.7</td>
<td>0.9</td>
<td>1.8</td>
</tr>
<tr>
<td>Aromatic acid</td>
<td>0.1</td>
<td>0.9</td>
<td>0.3</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>Polyacids</td>
<td>1.2</td>
<td>10</td>
<td>3.3</td>
<td>2.6</td>
<td>2.5</td>
</tr>
<tr>
<td>Isoprene oxidation products</td>
<td>2.0</td>
<td>142</td>
<td>41</td>
<td>20</td>
<td>43</td>
</tr>
<tr>
<td>Monoterpene oxidation products</td>
<td>1.0</td>
<td>36</td>
<td>9.2</td>
<td>7.0</td>
<td>8.7</td>
</tr>
<tr>
<td>Sesquiterpene oxidation products</td>
<td>0.1</td>
<td>3.4</td>
<td>0.9</td>
<td>0.3</td>
<td>1.0</td>
</tr>
<tr>
<td>Sum of all</td>
<td>113</td>
<td>1664</td>
<td>535</td>
<td>251</td>
<td>517</td>
</tr>
</tbody>
</table>
Figure 6. Temporal changes in the concentrations of biomass burning tracers and other organic compounds in the Alaskan aerosols.

Figure 6. Temporal changes in the concentrations of biomass burning tracers and other organic compounds in the Alaskan aerosols.

Table 2. Statistical summary of correlations among the organic tracers in PM2.5 aerosols from central Alaska.

<table>
<thead>
<tr>
<th>Linear regression</th>
<th>Correlation coefficient</th>
<th>Significance of correlation at ( P ) value &lt;0.05</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levoglucosan vs. mannosan</td>
<td>0.97</td>
<td>Significant</td>
</tr>
<tr>
<td>Levoglucosan vs. galactosan</td>
<td>0.94</td>
<td>Significant</td>
</tr>
<tr>
<td>Mannosan vs. galactosan</td>
<td>0.95</td>
<td>Significant</td>
</tr>
<tr>
<td>Glucose vs. fructose</td>
<td>0.91</td>
<td>Significant</td>
</tr>
<tr>
<td>Glucose vs. sucrose</td>
<td>0.82</td>
<td>Significant</td>
</tr>
<tr>
<td>Fructose vs. sucrose</td>
<td>0.94</td>
<td>Significant</td>
</tr>
<tr>
<td>Arabitol vs. mannitol</td>
<td>0.95</td>
<td>Significant</td>
</tr>
<tr>
<td>Trehalose vs. arabitol</td>
<td>0.85</td>
<td>Significant</td>
</tr>
<tr>
<td>Trehalose vs. mannitol</td>
<td>0.74</td>
<td>Significant</td>
</tr>
<tr>
<td>DEP(^a) vs. DBP(^b)</td>
<td>0.85</td>
<td>Significant</td>
</tr>
<tr>
<td>DEP(^a) vs. DiBP(^c)</td>
<td>0.87</td>
<td>Significant</td>
</tr>
<tr>
<td>DEP(^a) vs. DEHP(^d)</td>
<td>0.71</td>
<td>Significant</td>
</tr>
<tr>
<td>DBP(^b) vs. DiBP(^c)</td>
<td>0.81</td>
<td>Significant</td>
</tr>
<tr>
<td>DBP(^b) vs. DEHP(^d)</td>
<td>0.88</td>
<td>Significant</td>
</tr>
<tr>
<td>DiBP(^b) vs. DEHP(^d)</td>
<td>0.75</td>
<td>Significant</td>
</tr>
<tr>
<td>C(_5)-Alkene triols vs. 2-Methyltetrols</td>
<td>0.97</td>
<td>Significant</td>
</tr>
<tr>
<td>Glyceric acid vs. tartaric acid</td>
<td>0.84</td>
<td>Significant</td>
</tr>
<tr>
<td>Glyceric acid vs. citric acid</td>
<td>0.67</td>
<td>Significant</td>
</tr>
<tr>
<td>Tartaric acid vs. citric acid</td>
<td>0.87</td>
<td>Significant</td>
</tr>
<tr>
<td>Benzoic acid vs. glyceric acid</td>
<td>0.53</td>
<td>Not significant</td>
</tr>
<tr>
<td>Benzoic acid vs. tartaric acid</td>
<td>0.39</td>
<td>Not significant</td>
</tr>
<tr>
<td>Benzoic acid vs. citric acid</td>
<td>0.17</td>
<td>Not significant</td>
</tr>
<tr>
<td>Glyceric acid vs. isoprene SOA tracer</td>
<td>0.78</td>
<td>Significant</td>
</tr>
<tr>
<td>Tartaric acid vs. isoprene SOA tracer</td>
<td>0.75</td>
<td>Significant</td>
</tr>
<tr>
<td>Citric acid vs. isoprene SOA tracer</td>
<td>0.67</td>
<td>Significant</td>
</tr>
</tbody>
</table>

\( \text{a Diethyl phthalate. } \text{b Dibutyl phthalate. } \text{c Diisobutyl phthalate. } \text{d Diethylhexyl phthalate.} \)

these tracers (Table 2). This result indicates that they might have originated from similar types of biomass via the burning in central Alaska.

Levoglucosan (L) is largely produced by thermal decomposition of cellulose, while mannosan (M) is mainly a pyrolysis product of hemicellulose (Simoneit et al., 1999). Klemm et al. (2005) found that hardwood contains almost 55% to 65% of cellulose and 20% to 30% of hemicellulose. Accordingly, in a laboratory chamber analysis, Schmidl et al. (2008) found L/M ratios of nearly 2.5 to 3.9 for softwood burning and around 14 to 15 for hardwood burning. It is worth using the L/M ratio to identify the relative contribution of biomass types: hardwood vs. softwood in central Alaska. The L/M ratios in Alaskan samples ranged from 2.2 to 6.8 (avg 4.6), which are much lower than the ratios found in smoke samples derived from the burning of hardwoods but almost similar to the ratios found in samples derived from the burning of softwoods. The L/M ratios found in Alaskan aerosol samples are also much lower than the ratios reported for the samples derived from burning of rice straw (12.3 to 55.0) (Sheesley et al., 2003; Sullivan et al., 2008; Engling et al., 2009), cereal straw (55.7) (Zhang et al., 2007), wheat straw (12.7) and corn straw (19.5) (Cheng et al., 2013).

Fine et al. (2001, 2002, 2004) presented the concentrations of BB tracers in fine particles derived from the burning of several wood species collected from the United States. Based on their data, we calculated the L/M ratios to be 3.4 to 6.7 for softwood burning and 10.7 to 83.4 for hardwood burning. Our values from the Alaskan aerosol (2.2 to 6.8) are well within the range of L/M ratios for softwood burning from the United States. The ratios in Alaskan aerosol samples are comparable to those for marine aerosols collected from the Arctic Ocean (avg 3.5) (Fu et al., 2013), South China Sea (6.4) and western North Pacific (avg 4.6) (Fu et al., 2011). Our values are also similar to those found in aerosol samples collected at Montana in the USA (4.6) (Ward et al., 2006), the Vienna (4.1 to 6.4) and Salzburg (5.4 to 5.7) sites in Austria.
Burning of lignin produces phenolic compounds such as 4-hydroxybenzoic (4-HBA), vanillic and syringic acids, whereas dehydroabietic acid (DHAA) is a specific pyrolysis product of resin present in the bark surface and needle leaves and woody tissues of conifers (Simoneit et al., 1993). We detected 4-HBA and DHAA in the Alaskan aerosols, although their concentrations were much lower than BB tracers produced from cellulose and hemicellulose burning (Fig. 5b). The concentrations of 4-HBA and vanillic acid ranged from 0.4 to 6.4 ng m\(^{-3}\) (avg 1.7 ng m\(^{-3}\)) and from 0.1 to 8.6 ng m\(^{-3}\) (avg 1.8 ng m\(^{-3}\)), respectively, whereas those of syringic acid ranged from 0.02 to 1.1 ng m\(^{-3}\) (avg 0.2 ng m\(^{-3}\)). Shakya et al. (2011) and Myers-Pigg et al. (2016) reported that syringic to vanillic acid ratios for the burning of woody and non-woody angiosperm range from 0.1 to 2.4, whereas the ratios of softwood are 0.01 to 0.24. The concentration ratio of syringic to vanillic acid can therefore be used as a marker to distinguish the type of vegetation burned. We found that syringic to vanillic acid ratios in Fairbanks aerosols ranged from 0.02 to 0.5 (avg 0.2), suggesting that softwood is the more important biomass burned in central Alaska during the campaign. This conclusion is consistent with the observation on the L/M ratios as discussed above. The temporal variation of 4-HBA is very similar to that of anhydro sugars, whereas vanillic and syringic acids presented rather similar temporal trends to DHAA in Alaskan aerosols (Fig. 6d–g). Simoneit et al. (1993) proposed that the emission of DHAA is different than those of lignin pyrolysis products (Fig. 5b). This result suggests that the burning of conifer is a common source of OAs in central Alaska.

### 3.3 Lipids: tracers of leaf waxes and marine sources

Series of lipid class compounds, including \(n\)-alkanes (C\(_{21}\) to C\(_{33}\)), \(n\)-alkanols (C\(_{8}\) to C\(_{30}\)), and \(n\)-alkanoic acids (C\(_{12}\) to C\(_{18}\)), were detected in Alaskan aerosols. \(n\)-Alkanoic acids are the major lipid class compounds in Alaskan aerosols (avg 185 ng m\(^{-3}\)), which is several times higher than those of \(n\)-alkanols (avg 46 ng m\(^{-3}\)) and \(n\)-alkanes (avg 24 ng m\(^{-3}\)) (Table 1). Figure 7a–c show the average molecular distributions of lipid compounds in Alaskan aerosols. The molecular distribution of \(n\)-alkanes is characterized by an odd-carbon-number predominance with maxima at heptacosane (C\(_{27}\): avg 6.8 ng m\(^{-3}\)). Low molecular weight (LMW) \(n\)-alkanes are dominant in particles derived from fossil fuel combustion, whereas those derived from leaf waxes are enriched with high molecular weight (HMW) \(n\)-alkanes (Rogge et al., 1993; Hays et al., 2005; Wang et al., 2009). A remarkable feature in the molecular signature of \(n\)-alkanes is the presence of only the HMW species (C\(_{21}\) to C\(_{33}\)) in Alaskan aerosols. This molecular signature in the \(n\)-alkanes of the PM\(_{2.5}\) samples suggests that leaf waxes are the major source of \(n\)-alkanes in central Alaska, with no significant contribution from fossil fuel combustion. This feature is different from the result of marine aerosols collected over the Arctic Ocean (Fu et al., 2013), in which \(n\)-alkanes were mostly of fossil fuel origin. We conclude that fossil fuel combustion is not an important source of OAs over central Alaska during the summer campaign. This remark is consistent with the fact that the fossil fuel biomarkers such as hopanes and steranes (Ding et al., 2009; Wang et al., 2009) were not detected in the Alaskan samples. These results demonstrate that biogenic \(n\)-alkanes emitted from boreal forest fires largely overwhelmed fossil fuel combustion-derived \(n\)-alkanes in central Alaska.

The carbon preference index (CPI) is a powerful tool to characterize the anthropogenic vs. biogenic sources of lipid compounds (Simoneit et al., 1991; Kawamura et al., 2003). The CPI value of \(n\)-alkanes in fossil fuel emission is usually...
close to 1, while it is more than 5 for leaf waxes (Peltzer and Gagosian, 1989). The calculated CPI values of n-alkanes in the Alaskan aerosols are in the range of 5.2 to 9.9, with an average of 6.6. These values are significantly higher than those reported in urban aerosols from megacities in China (0.9 to 1.8) (Wang et al., 2006), India (1.2 to 2.3) (Fu et al., 2010) and Japan (1.1 to 2.8) (Kawamura et al., 1994), where aerosol particles were seriously affected by fossil fuel combustion. The broader range of CPI values was found in aerosol particle samples collected over Mt. Tai (1.1 to 8.0) (Fu et al., 2008) and the western North Pacific (1.8 to 15) (Kawamura et al., 2003), where the input of plant waxes overwhelms the contribution of fossil fuel combustion. Together with these assessments our results strongly infer that n-alkanes over the Alaskan atmosphere mainly originated from plant leaf waxes. The wax covering the external surface of a plant leaf is composed of a mixture of long-chain aliphatic compounds. Kolattukudy (1976) found that odd-carbon-number n-alkanes (C_{25} to C_{33}) are one of the most abundant compound classes in the leaf wax. Simonet et al. (1991) considered the excess of odd homologs minus the neighboring even homologs to be the abundance of plant-derived n-alkanes in atmospheric samples. The contribution of estimated plant-derived n-alkanes to the total n-alkanes ranged from 53% to 70% (avg 61%), implying that leaf wax is a major source of n-alkanes in the Alaskan aerosols.

The average molecular characteristics of n-alkanols and n-alkanoic acids displayed even-carbon-number predominance (Fig. 7b and c). n-Alkanol presented maxima at docosanol (C_{22}: avg 9.2 ng m^{-3}), whereas n-alkanoic acids demonstrated a peak at tetracosanoic acid (C_{24}: avg 63 ng m^{-3}). Microbes and marine phytoplankton are the sources of LMW n-alkanols and n-alkanoic acids, while their HMW homologs are specifically derived from higher plant waxes (Kawamura et al., 2003; Wang and Kawamura, 2005). Simonet (2002) has proposed that BB also emits a large extent of n-alkanols and n-alkanoic acids into the atmosphere. The average concentrations of HMW n-alkanols (C_{21} to C_{30}: 31 ng m^{-3}) and HMW n-alkanoic acids (C_{21} to C_{32}: 122 ng m^{-3}) are twice as high as those of LMW n-alkanols (C_{8} to C_{20}: 15 ng m^{-3}) and LMW n-alkanoic acids (C_{12} to C_{20}: 58 ng m^{-3}) in the Alaskan aerosols. This result shows that locally derived plant waxes might be the source of HMW n-alkanols and n-alkanoic acids in central Alaska. The CPI values of n-alkanols and n-alkanoic acids are in the range of 3.0 to 10 (avg 6.2) and 5.6 to 9.8 (avg 7.9), respectively, suggesting a large contribution of plant waxes to lipid class compounds in central Alaskan aerosols.

The concentrations of n-alkanes and n-alkanols slightly decreased from the 5–12 June to late June samples (25 June to 4 July) and then dramatically increased in the 4–6 July sample (Fig. 6h and i). The concentration peaks of n-alkanes and n-alkanols were also observed in the sample of 14–23 July, whereas their concentrations constantly decreased from 30 July to the end of the campaign. The levels of n-alkanoic acids were low at the beginning of the campaign and then increased drastically in the 4–6 July sample and remained high in the two samples collected on 6–23 July (Fig. 6j). Concentrations of n-alkanoic acids decreased from 30 July to 21 September. Fascinatingly, the temporal variations of lipid class compounds were similar to those of anhydrosugars (Fig. 6a–c and h–j). Figure 8a–c show the linear regression analysis of lipid compounds with levoglucosan. We found strong correlations (r = 0.90–0.96) of lipid compounds with levoglucosan in Alaskan aerosols. These results suggest that forest fires significantly control the atmospheric levels of lipids in central Alaska via the evaporative ablation of leaf waxes of terrestrial plants.

We also detected unsaturated n-alkanoic acids in Alaskan aerosol samples. Oleic (C_{18:1}) and linoleic (C_{18:2}) acids are major constituents of the cell membranes in terrestrial plants. They are released into the atmosphere directly from the leaf surface by wind action (Yokouchi and Ambe, 1986; Noureddini and Kanabur, 1999). Fine et al. (2001) and Hays et al. (2005) proposed that BB also emits significant amounts of C_{18:1} and C_{18:2} into the atmosphere. They are subjected to photochemical oxidation in the atmosphere. C_{18:1} and C_{18:2} are more reactive due to a double bond than C_{18:0} in the atmosphere with oxidants such as OH radicals and O_3. The ratio of C_{18:1}+C_{18:2} to octadecanoic acid (C_{18:0}) is thus used as an indicator of photochemical processing of OAs (Kawamura and Gagosian, 1987). The ratios ranged from 0.03 to 0.3, with an average of 0.2 in Alaskan aerosols. Because average concentrations of C_{18:1} (0.9 ng m^{-3}) and C_{18:2} (0.5 ng m^{-3}) in Alaskan samples are significantly lower than that of C_{18:0} (10 ng m^{-3}), C_{18:1} and C_{18:2} may be rapidly degraded in the atmosphere by photochemical oxidations.

### 3.4 Sugar compounds: tracers of primary biological particles

Nine sugar compounds were detected in Alaskan aerosol samples, with five primary sugars and four sugar alcohols (Fig. 9). The concentrations of primary sugars were in the range of 0.3 to 44 ng m^{-3} (avg 13 ng m^{-3}), whereas those of sugar alcohols ranged from 1.0 to 24 ng m^{-3} (avg 14 ng m^{-3}). The concentrations of total sugar compounds ranged from 1.3 to 62 ng m^{-3} (27 ng m^{-3}), in which sugar alcohols contributed more to the total sugars (avg 54.2%) than primary sugars (avg 45.8%) in Alaskan aerosols. Primary sugars are abundantly present in vascular plants. They are produced during the photosynthetic process in leaves and then accumulated in growing plants (Medeiros et al., 2006). Figure 9a presents the average molecular distributions of primary sugars in Alaskan aerosols. Primary sugars are characterized by the predominance of glucose in Alaskan samples, with a concentration range of 0.1 to 19 ng m^{-3} (avg 6.8 ng m^{-3}), followed by trehalose (avg 2.6 ng m^{-3}). Although sucrose (avg 1.6 ng m^{-3}) and fructose (avg 1.3 ng m^{-3}) are not abundant (Fig. 9a), glucose...
Correlations of organic compounds with the biomass burning tracer levoglucosan in the Alaskan aerosol samples showed strong positive correlations with fructose ($r = 0.91$) and sucrose ($r = 0.82$) (Table 2). Fructose also presented a strong correlation with sucrose ($r = 0.94$) (Table 2). These correlations indicate their similar source and origin in the atmosphere of central Alaska.

Glucose and fructose are carbohydrates enriched in tree barks as well as branches and leaves (Medeiros et al., 2006; Li et al., 2016). They are present in plant nectars and fruits as well as pollen and fern spores (Baker et al., 1998; Graham et al., 2002). Dust and BB-derived particles have also been reported as major sources of glucose and fructose in the atmosphere (Nolte et al., 2001; Rogge et al., 2007). Sucrose is produced in plant leaves and distributed to several portions of the plant body (Jia et al., 2010). Sucrose has also been reported in airborne pollen grains produced from blooming plants (Pacini, 2000), surface soil and associated microbiota (Simoneit et al., 2004b) and dehydrated plant materials (Ma et al., 2009). We found that glucose shows moderate correlation ($r = 0.48$) with levoglucosan (Fig. 8d). Shafizadeh and Fu (1973) documented that glucose is a minor product of cellulose pyrolysis. The predominance of glucose among primary sugars together with a moderate correlation with levoglucosan suggests that pyrolysis of cellulose and hemicellulose is not the source of glucose in central Alaska. Pullman and Buchanan (2008) found that soluble carbohydrates such as glucose are a major component of conifers, where it can be stored in a large amount as deposited or dissolved free molecules. The temporal trend of glucose showed a peak in the sample collected during 14–23 July (Fig. 6k). Interestingly, the same sample showed a high loading of DHAA that is a unique tracer of the burning of conifer trees (Fig. 6g). This result suggests that the burning of conifer plants is the source of glucose in central Alaska.

Trehalose is a well-known constituent of microbes and fungal spores as well as plant species and suspended soil particles (Graham et al., 2003; Medeiros et al., 2006). The levels of trehalose stayed constant from 5 June to 14 July, dramatically decreased on 23 July to 8 August and then increased towards the end of the campaign when rainfall occurred in central Alaska (Figs. 2 and 6m). This result shows that the major source of trehalose might be the fungi in the surface soil of central Alaska that were emitted after the rainfall event. Terrestrial plants and marine phytoplankton as well as soil dust particles and associated microorganisms release xylose into the atmosphere (Cowie and Hedges, 1984). Although xylose is a minor primary sugar in Alaskan aerosols (avg 1.1 ng m$^{-3}$), its temporal trend is very similar to that of anhydrosugars (Fig. 6a–c and n). This result together with a strong positive correlation of xylose with levoglucosan ($r = 0.92$) implies its BB origin in central Alaska (Fig. 8g). This finding is similar to that of Sullivan et al. (2011), who documented that atmospheric levels of xylose in the Midwestern United States were attributed to BB emission.

Sugar alcohols presented the predominance of arabitol (avg 6.6 ng m$^{-3}$) and mannitol (avg 6.2 ng m$^{-3}$) (Fig. 9b). The concentration levels of erythritol (avg 1.0 ng m$^{-3}$) and inositol (avg 0.3 ng m$^{-3}$) are much lower than those of arabitol and mannitol in Alaskan aerosols. Arabitol and mannitol concentrations were higher during the beginning and end of the campaign than those during the middle of the campaign (Fig. 6o and p). We found that arabitol and mannitol are strongly correlated ($r = 0.95$), implying their similar source in the Alaskan aerosols (Table 2). The major sources of arabitol and mannitol are airborne fungal spores (Pashynska et al., 2002; Bauer et al., 2008). Debris from mature leaves has also been proposed as a source of arabitol and mannitol in the forest areas (Pashynska et al., 2002; Zhang et al., 2010). Guasco et al. (2013) and Prather et al. (2013) proposed that bubble bursting of seawater contributes bacteria and dissolves organic species along with sea salts to aerosol particles. Although air masses mostly originated from the ocean (Fig. 3), the altitude of most of the air masses dropped at several places and went on to Fairbanks by maintaining low height. Therefore, we presume a negligible input of ma-
The ambient concentrations of total phthalate esters ranged from 0.4 to 6.6 ng m$^{-3}$ (avg 1.7 ng m$^{-3}$), which are slightly higher than those from the North Sea to the high Arctic (0.4 to 1.0 ng m$^{-3}$) (Xie et al., 2007), comparable to or slightly lower than those observed in the North Pacific (0.7 to 4.48 ng m$^{-3}$) (Atlas and Giam, 1981), Great Lakes (0.1 to 10 ng m$^{-3}$) (Eisenreich et al., 1981), and Canadian High Arctic (0.28 to 11 ng m$^{-3}$) (Fu et al., 2009), but much lower than those reported in Sweden (0.5 to 127 ng m$^{-3}$) (Thuren and Larsson, 1990), mountainous aerosols (9.6 to 985 ng m$^{-3}$) (Fu et al., 2008) and urban aerosols from megacities in India and China (62 to 2200 ng m$^{-3}$) (Wang et al., 2006; Fu et al., 2010). Figure 10 shows the average molecular distributions of phthalate esters in Alaskan aerosols. We found DEP (avg 0.8 ng m$^{-3}$) to be a dominant phthalate, followed by DBP and DEHP (avg 0.4 ng m$^{-3}$), whereas DiBP was less abundant (avg 0.2 ng m$^{-3}$). The predominance of DEP among phthalate esters in Alaskan aerosol is different than those found in marine aerosol from the Arctic Ocean (Xie et al., 2007; Fu et al., 2013) and urban aerosols from India and China (Wang et al., 2006; Fu et al., 2010), where DEHP was the dominant species. We found similar temporal variations with significant positive correlations among detected phthalate esters ($r = 0.71–0.88$) (Fig. 11a–d and Table 2), suggesting that they have similar sources in central Alaska.

3.5 Phthalate esters: tracers of plastic burning

Phthalates are widely used as a plasticizer in synthetic polymers and as a softener in polyvinylchloride (Thuren and Larsson, 1990; Wang et al., 2006). They can be emitted into the atmosphere by evaporation from polymers because they are not chemically bonded (Staples et al., 1997). The compositions of phthalate esters are the subject of scientific discussion and public concern due to their potential carcinogenic and endocrine-disrupting properties (Sidhu et al., 2005; Swan et al., 2005). We detected four phthalate esters in Alaskan aerosols, including diethyl phthalate (DEP), dibutyl phthalate (DBP), diisobutyl phthalate (DiBP) and diethylhexyl phthalate (DEHP).

The ambient concentrations of total phthalate esters ranged from 0.4 to 6.6 ng m$^{-3}$ (avg 1.7 ng m$^{-3}$), which are slightly higher than those from the North Sea to the high Arctic (0.4 to 1.0 ng m$^{-3}$) (Xie et al., 2007), comparable to or slightly lower than those observed in the North Pacific (0.7 to 4.48 ng m$^{-3}$) (Atlas and Giam, 1981), Great Lakes (0.1 to 10 ng m$^{-3}$) (Eisenreich et al., 1981), and Canadian High Arctic (0.28 to 11 ng m$^{-3}$) (Fu et al., 2009), but much lower than those reported in Sweden (0.5 to 127 ng m$^{-3}$) (Thuren and Larsson, 1990), mountainous aerosols (9.6 to 985 ng m$^{-3}$) (Fu et al., 2008) and urban aerosols from megacities in India and China (62 to 2200 ng m$^{-3}$) (Wang et al., 2006; Fu et al., 2010). Figure 10 shows the average molecular distributions of phthalate esters in Alaskan aerosols. We found DEP (avg 0.8 ng m$^{-3}$) to be a dominant phthalate, followed by DBP and DEHP (avg 0.4 ng m$^{-3}$), whereas DiBP was less abundant (avg 0.2 ng m$^{-3}$). The predominance of DEP among phthalate esters in Alaskan aerosol is different than those found in marine aerosol from the Arctic Ocean (Xie et al., 2007; Fu et al., 2013) and urban aerosols from India and China (Wang et al., 2006; Fu et al., 2010), where DEHP was the dominant species. We found similar temporal variations with significant positive correlations among detected phthalate esters ($r = 0.71–0.88$) (Fig. 11a–d and Table 2), suggesting that they have similar sources in central Alaska.
Molecular distributions of phthalate esters in PM$_{2.5}$ aerosols collected in central Alaska. See Fig. 5 for the description of the box-and-whisker diagram.

SOA is a crucial component of the atmosphere that has an impact on the radiation budget directly by scattering sunlight and indirectly by acting as CCN (Kanakidou et al., 2005; Carlton et al., 2009). Isoprene has conjugated double bonds, and thus it is more reactive towards oxidants such as O$_3$ and NO$_x$, resulting in various intermediates and stable products via a series of oxidative reactions in the atmosphere. We detected six organic compounds, including 2-methylglyceric acid (2-MGA), three C$_5$-alkene triols and two 2-methyltetrols (2-MTLs), as isoprene-SOA tracers in the Alaskan aerosols. Their total concentrations ranged from 2.0 to 142 ng m$^{-3}$ (avg 41 ng m$^{-3}$), which are significantly higher than those reported over the North Pacific (0.11 to 0.48 ng m$^{-3}$) (Fu et al., 2011), Canadian High Arctic (avg 0.30 ng m$^{-3}$) (Fu et al., 2009), North Pacific to Arctic (avg 0.62 ng m$^{-3}$) (Ding et al., 2013), western North Pacific (0.05 to 7.22 ng m$^{-3}$) (Zhu et al., 2016), a forest site in western Germany (avg 20.5 ng m$^{-3}$) (Kourtchev et al., 2008a), and Mumbai in India (avg 1.1 ng m$^{-3}$) (Fu et al., 2016), but lower than those in Mt. Changbai (22 to 280 ng m$^{-3}$) (Wang et al., 2008) and Mt. Fuji (avg 69 ng m$^{-3}$) (Fu et al., 2014), Research Triangle Park in the USA (19.9 to 384 ng m$^{-3}$) (Lewandowski et al., 2007) and several sites in China (8.65 to 554 ng m$^{-3}$) (Ding et al., 2014).

Molecular compositions of isoprene-SOA tracers are characterized by the predominance of C$_5$-alkene triols (avg 20 ng m$^{-3}$) and 2-MTLs (avg 19 ng m$^{-3}$) in Alaskan aerosols (Fig. 12). Surratt et al. (2010) proposed that C$_5$-alkene triols and 2-MTLs are higher-generation products from the photooxidation of epoxydiols of isoprene under low-NO$_x$ conditions. C$_5$-Alkene triols were strongly correlated with 2-MTLs ($r = 0.97$) in the Alaskan aerosols (Table 2). The abundant co-presence of C$_5$-alkene triols and 2-MTLs suggests their similar sources and formation pathways in central Alaska. 2-Methylerythritol (avg 13 ng m$^{-3}$) is twice as abundant as 2-methylthreitol (avg 5.9 ng m$^{-3}$), similar to previous studies (Ion et al., 2005; Cahill et al., 2006). The concentration of 2-MGA is significantly lower (avg 2.2 ng m$^{-3}$) than C$_5$-alkene triols and 2-MTLs (Fig. 12). Surratt et al. (2006) suggested that 2-MGA is produced by the oxidation of methacrolein and methacrylic acid and has been detected as an important gas-phase intermediate in the SOA formation from isoprene under high-NO$_x$ conditions. Temporal variations of isoprene-SOA tracers were very similar to each other in Alaskan aerosols (Fig. 11e–g). Their concentrations increased in the sample of 12–25 June to 6–14 July and decreased in the sample of 14–23 July and 23–30 July. They increased significantly in the 30 July to 4 Aug...
The concentration ratio of PA to PNA of about 1 in the Southeastern United States is higher than those in Canada and other sites in Europe (Kavouras and Stephanou, 2002). PA is the initial photooxidation product of β-caryophyllene, which is hydroxylated in leaves and wood because of low volatility and then processed into β-caryophyllinic acid (Jaoui et al., 2007). The concentration ratio of β-caryophyllene to PNA is about 1 in the southeast of China (Ding et al., 2014) and Arctic Ocean (avg 0.017 ng m$^{-3}$), which are higher than those from the Canadian High Arctic (avg 0.02 to 0.22 ng m$^{-3}$) (Fu et al., 2009, 2013) but lower than those reported at several sites in China (0.17 to 17.4 ng m$^{-3}$) (Ding et al., 2014) and Research Triangle Park in the USA (5.9 to 25 ng m$^{-3}$) (Lewandowski et al., 2007). The temporal variation of β-caryophyllinic acid is very different from those of isoprene and monoterpene-SOA tracers detected in Alaskan aerosols (Fig. 11l). Akagi et al. (2011) reported that biogenic VOCs and monoterpene-SOA tracers detected in Alaskan aerosols showed a high level of β-caryophyllinic acid in the atmosphere (Jaoui et al., 2007).

The much lower vapor pressure of PA than that of PNA. However, this pattern is different from those found in summertime aerosols at the summit of Mt. Tai in China (Fu et al., 2008) and other sites in Europe (Kavouras and Stephanou, 2002) and North America (Cahill et al., 2006), in which PNA is more abundant than PA.

PA and PNA are the initial photooxidation products of monoterpene-SOA. It can be further photooxidized to 3-MBTCA in the atmosphere (Gómez-González et al., 2012). The concentration ratio of 3-MBTCA to PA+PNA can therefore be used to differentiate the fresh and photochemically processed monoterpene-SOA tracers in the atmosphere. We found the ratios to be 0.1 to 1.4 (avg 0.5), demonstrating that monoterpene-SOA are mostly fresh in central Alaska. The ratios of 3-HGA to 3-MBTCA are mostly lower when they are produced from α-pinene as compared to β-pinene. Lewandowski et al. (2013) documented a greater contribution of α-pinene to monoterpene-SOA tracers based on the 3-HGA / 3-MBTCA ratio of about 1 in the southeastern United States than those observed in California (1.8 to 3.8). Ding et al. (2014) also suggested α-pinene as the major precursor in China based on the low ratios in Hefei (1.16) and Qianyazhou (0.75). We found the ratios ranged from 0.5 to 2.0 in Alaskan aerosols (avg 1.0), indicating that α-pinene mainly contributes to monoterpene-SOA tracers. 3-HGA and 3-MBTCA have somewhat similar temporal variations, although the patterns are different from PA and PNA (Fig. 11b–k), which showed very similar variations in Alaskan aerosols.

Sesquiterpenes are BVOCs with high reactivity and relatively low vapor pressure (Duhl et al., 2008). β-Caryophyllene is the dominant sesquiterpene. The ozonolysis or photooxidation of β-caryophyllene produces β-caryophyllinic acid in the atmosphere (Jaoui et al., 2007). Concentrations of β-caryophyllinic acid in Alaskan aerosols ranged from 0.1 to 3.4 ng m$^{-3}$ (avg 0.9 ng m$^{-3}$), which are higher than those from the Canadian High Arctic (avg 0.02 to 0.22 ng m$^{-3}$) and Arctic Ocean (avg 0.017 ng m$^{-3}$) (Fu et al., 2009, 2013) but lower than those reported at several sites in China (0.17 to 17.4 ng m$^{-3}$) (Ding et al., 2014) and Research Triangle Park in the USA (5.9 to 25 ng m$^{-3}$) (Lewandowski et al., 2007). The temporal variation of β-caryophyllinic acid is very different from those of isoprene and monoterpene-SOA tracers detected in Alaskan aerosols (Fig. 11l). Akagi et al. (2011) reported that biogenic VOCs could also be emitted from biomass burning. Our result showed a high level of β-caryophyllinic acid in the samples that were affected by BB in central Alaska. Ciccioni et al. (2014) proposed that sesquiterpenes could be accumulated in leaves and wood because of low volatility and then abundantly emitted upon heating. The temporal trend variation of β-caryophyllinic acid is similar to those of anhydro sugars (Figs. 6a–c and 11l). Interestingly, we found a strong correlation ($r = 0.98$) of β-caryophyllinic acid with levoglucosan (Fig. 8j), again indicating that forest fire largely con-
tributes to the formation of β-caryophyllinic acid in central Alaska.

3.7 Aromatic acids and polyacids: tracers of SOA

We detected benzoic acid in the Alaskan aerosol with the concentration range of 0.1 to 0.9 ng m\(^{-3}\) (avg 0.3 ng m\(^{-3}\)). Benzoic acid is produced from several anthropogenic sources. It is a primary pollutant in the automobile emission and smokes derived from burning of biomass and biofuels (Rogge et al., 1993; Kawamura et al., 2002). It is also a secondary product of photochemical degradation of toluene emitted from anthropogenic sources (Suh et al., 2003). It can play an important role in enhancing the new particle formation in the atmosphere (Zhang et al., 2004). The temporal variation of benzoic acid is similar to anhydrosugars detected in Alaskan samples (Figs. 6a–c and 11m). We also found a strong positive correlation \(r = 0.95\) of benzoic acid with levoglucosan (Fig. 8k), demonstrating that BB is the source of benzoic acid in central Alaska.

Polyacids are also the secondary photooxidation products of atmospheric organic precursors (Wang et al., 2007; Fu et al., 2012). Concentrations of total polyacids ranged from 1.2 to 10 ng m\(^{-3}\) (avg 3.3 ng m\(^{-3}\)), among which glyceric acid (avg 1.6 ng m\(^{-3}\)) was dominant (Fig. 13). Significant positive correlations were found among all polyacids \((r = 0.67–0.87)\) in Alaskan aerosols (Table 2). These results imply that polyacids may have similar sources or formation pathways in central Alaska. We found that polyacids showed no significant correlations with benzoic acid \((r = 0.17–0.53)\), which is mostly of BB origin in Alaskan samples, as discussed above (Table 2). These correlations and different temporal trends of benzoic acid and polyacids suggest that forest fires are not the main source of polyacids in the Alaskan samples (Fig. 11m–p). This remark is further supported by the insignificant correlations of polyacids with levoglucosan \((r = 0.29–0.47)\) (Fig. 8l–n). Claey s et al. (2004) suggested that SOA tracers such as tartaric acid are produced by the photochemical oxidation of isoprene. Interestingly, significant positive correlations \((r = 0.67–0.78)\) of polyacids were found with total isoprene-SOA tracers detected in Alaskan samples (Table 2), suggesting that they may be produced by photooxidation of isoprene in the Alaskan atmosphere.

3.8 Contributions of different compound classes to aerosol organic carbon

The contributions of each compound class to organic carbon (OC) in the Alaskan aerosols are given in Table 3. BB tracers accounted for 1.35 % to 8.35 % (avg 4.40 %) of OC. The contribution of anhydrosugars to OC was substantially higher (avg 4.26 %) than that of lignin and resin acids (avg 0.14 %). This value from Fairbanks is notably higher than those reported in aerosol samples collected from a round-the-world cruise (avg 0.15 %) (Fu et al., 2011), Gosan, Jeju Island in Korea (avg 0.29 %), Sapporo (avg 0.44 %) and Chichijima (avg 0.06 %) in Japan (Simoneit et al., 2004a), and Chennai in India (avg 0.59 %) (Fu et al., 2010). The lipid compound classes in Fairbanks samples accounted for 1.16 to 45.8 % (avg 12.4 %) of OC. n-Alkanoic acids contributed on average 7.48 % (0.67 to 15.9 %), which is much higher than those estimated in samples of the round-the-world cruise (avg 0.82 %), Sapporo (avg 0.62 %) and Chichijima (avg 0.78 %) (Simoneit et al., 2004a; Fu et al., 2011). The tracers of primary biological aerosol particles accounted for on average 0.85 % (0.17 % to 1.50 %), among which comparable contributions of primary sugars (avg 0.39 %) and sugar alcohols (avg 0.46 %) to OC were found in Alaskan aerosols. Plastic burning tracers accounted for 0.02 to 1.07 % of OC (avg 0.14 %), which is lower than those from the Sapporo (avg 1.1 %) and Chichijima samples (avg 1.2 %) (Simoneit et al., 2004a) and tropical samples from India (avg 4.50 %) (Fu et al., 2010). Biogenic SOA tracers contributed 0.18 to 3.99 % of OC (avg 1.66 %), among which the contribution of isoprene-derived SOA tracers was high (avg 1.28 %), followed by monoterpene (avg 0.34 %) and sesquiterpene (avg 0.03 %) SOA tracers. Other SOA tracers with minor contribution to OC include polyacids (avg 0.08 %) and aromatic acids (avg 0.02 %).

With the consideration of water-soluble dicarboxylic acids and related polar compounds measured in the same sample sets as reported in Deshmukh et al. (2018), the total organic compounds identified in the Alaskan aerosols accounted for 6.37 % to 59.2 % with a mean of 21.4 % of OC. This result indicates that a substantial fraction of OAs studied in the Alaskan site can be identified at a molecular level.

4 Conclusions and implications

We identified 96 organic compounds in PM\(_{2.5}\) samples collected at Fairbanks in central Alaska during the summer campaign in 2009. Concentrations of total organic compounds ranged from 113 to 1664 ng m\(^{-3}\) (avg 535 ng m\(^{-3}\)). The most abundant compound classes in the Alaskan aerosol are anhydrosugars (avg 186 ng m\(^{-3}\)) and n-alkanoic acids (avg 185 ng m\(^{-3}\)). The temporal variations of anhydrosugars dramatically changed during the campaign, showing peaks during BB events. The similar temporal trends of lipids and strong correlations with levoglucosan demonstrated that local forest fires likely control the atmospheric levels of OAs in central Alaska. The concentration ratios of levoglucosan to mannose (2.2 to 6.8) and syringic to vanillic acid (0.02 to 0.5) suggest that burning of softwood is a common source of OAs. The higher levels of HMW n-alkanoic acids and n-alkanols than their LMW homologs together with high CPI values of n-alkanes (5.2 to 9.9), n-alkanols (3.0 to 10) and n-alkanoic acids (5.6 to 9.8) further suggest that they were emitted by the thermal ablation of plant waxes during forest fires in central Alaska. The temporal patterns of mannitol and
Table 3. Contributions (\%) of individual organic compound classes to organic carbon (OC) in PM$_{2.5}$ aerosols from central Alaska$^a$.

<table>
<thead>
<tr>
<th>Compound classes</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Mean</th>
<th>Median</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biomass burning tracers</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anhydrosugars</td>
<td>1.32</td>
<td>8.12</td>
<td>4.26</td>
<td>3.64</td>
<td>2.13</td>
</tr>
<tr>
<td>Lignin and resin acids$^b$</td>
<td>0.03</td>
<td>0.51</td>
<td>0.14</td>
<td>0.11</td>
<td>0.13</td>
</tr>
<tr>
<td>Subtotal</td>
<td>1.35</td>
<td>8.35</td>
<td>4.40</td>
<td>3.71</td>
<td>2.24</td>
</tr>
<tr>
<td><strong>Lipid compounds</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$n$-Alkanes</td>
<td>0.05</td>
<td>8.53</td>
<td>1.55</td>
<td>0.98</td>
<td>2.19</td>
</tr>
<tr>
<td>$n$-Alkanols</td>
<td>0.40</td>
<td>21.3</td>
<td>3.32</td>
<td>1.82</td>
<td>5.47</td>
</tr>
<tr>
<td>$n$-Alkanoic acids</td>
<td>0.67</td>
<td>15.9</td>
<td>7.48</td>
<td>6.71</td>
<td>4.80</td>
</tr>
<tr>
<td>Subtotal</td>
<td>1.16</td>
<td>45.8</td>
<td>12.4</td>
<td>9.20</td>
<td>11.3</td>
</tr>
<tr>
<td><strong>Primary biological aerosols</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Primary sugars</td>
<td>0.05</td>
<td>0.85</td>
<td>0.39</td>
<td>0.50</td>
<td>0.26</td>
</tr>
<tr>
<td>Sugar alcohols</td>
<td>0.07</td>
<td>0.95</td>
<td>0.46</td>
<td>0.33</td>
<td>0.33</td>
</tr>
<tr>
<td>Subtotal</td>
<td>0.17</td>
<td>1.50</td>
<td>0.85</td>
<td>0.74</td>
<td>0.56</td>
</tr>
<tr>
<td>Phthalate esters</td>
<td>0.02</td>
<td>1.07</td>
<td>0.14</td>
<td>0.05</td>
<td>0.28</td>
</tr>
<tr>
<td>Aromatic acid</td>
<td>0.01</td>
<td>0.09</td>
<td>0.02</td>
<td>0.01</td>
<td>0.02</td>
</tr>
<tr>
<td>Polyacids</td>
<td>0.02</td>
<td>0.25</td>
<td>0.08</td>
<td>0.09</td>
<td>0.06</td>
</tr>
<tr>
<td><strong>Biogenic SOA tracers</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isoprene oxidation products</td>
<td>0.07</td>
<td>3.20</td>
<td>1.28</td>
<td>0.83</td>
<td>1.12</td>
</tr>
<tr>
<td>Monoterpen epoxidation products</td>
<td>0.07</td>
<td>0.75</td>
<td>0.34</td>
<td>0.34</td>
<td>0.20</td>
</tr>
<tr>
<td>Sesquiterpe epoxidation products</td>
<td>0.02</td>
<td>0.04</td>
<td>0.03</td>
<td>0.03</td>
<td>0.01</td>
</tr>
<tr>
<td>Subtotal</td>
<td>0.18</td>
<td>3.99</td>
<td>1.66</td>
<td>1.22</td>
<td>1.29</td>
</tr>
<tr>
<td>Dicarboxylic acids and related compounds</td>
<td>1.15</td>
<td>2.97</td>
<td>1.90</td>
<td>1.87</td>
<td>0.58</td>
</tr>
<tr>
<td>Total detected organic compounds</td>
<td>6.37</td>
<td>59.2</td>
<td>21.4</td>
<td>16.9</td>
<td>13.8</td>
</tr>
</tbody>
</table>

$^a$ All the organic compounds quantified were converted to carbon contents and then divided by OC. See Deshmukh et al. (2018) for OC and dicarboxylic acids and related compounds. $^b$ The results of lignin and resin acids were combined due to the very low contribution of resin acid to OC.

arabitol suggested that the rainfall played an important role in enhancing their levels in central Alaska. The molecular compositions of phthalate esters showed that diethyl phthalate is a commonly used plasticizer in central Alaska. The molecular composition of biogenic SOA tracers with a predominance of isoprene-SOA tracers (avg 41 ng m$^{-3}$) suggested that isoprene is a crucial precursor of SOA over central Alaska. Our results provide valuable information to better understand the compositions of OAs and their sources and formation pathways in the subarctic atmosphere.

The Arctic is a critical region on the Earth, with a significant warming and high sensitivity to climate forcing due to a strong effect on an albedo–sea ice feedback system. Our results confirmed that forest fires and plant emissions are important factors controlling the organic chemical composition of fine aerosol particles in central Alaska. It is worthwhile therefore to note from the above discussion that Fairbanks exemplifies many of the problems of pollution in the Arctic regions. The local and regional BB episodes in the warmer season enhanced the atmospheric levels of OAs in central Alaska. Because the residence time of fine particles is relatively long in the atmosphere, we propose that OAs of PM$_{2.5}$ at Fairbanks can be subjected to long-range transport to the Arctic causing a significant influence on the air quality and climate in the Arctic region. Although we studied the aerosol samples collected in 2009, further research is needed to characterize the seasonal and interannual trends of OAs using more recent aerosol samples to better evaluate their current impact in the Arctic atmosphere.

**Data availability.** The data set of this paper is given in Table S1 in the Supplement.

**Supplement.** The supplement related to this article is available online at: https://doi.org/10.5194/acp-19-14009-2019-supplement.
Author contributions. KK designed the research. YK collected the aerosol samples. DKD and MMH analyzed the samples for organic tracer compounds. DKD evaluated the data and wrote the paper under the supervision of KK. All the authors contributed to discussing results and commenting on the paper.

Competing interests. The authors declare that they have no conflict of interest.

Acknowledgements. We acknowledge financial support from the Japan Society for the Promotion of Science (JSPS). This work was partially conducted under the Japan Agency for Marine-Earth Science and Technology – International Arctic Research Center (JAMSTEC-IARC) collaboration project with funding from JAMSTEC. We appreciate the financial support from a JSPS fellowship to Dhananjay Kumar Deshmukh (PU17907) and the Special Grant Program for International Students (SGPIS) of Hokkaido University, Sapporo, Japan, for the financial support to M. Mozammel Haque. We thank Bang-Yong Lee for his support during the sample collection.

Financial support. This research has been supported by the Japan Society for the Promotion of Science (grant no. 24221001) and National Research Foundation of Korea (grant nos. NRF-2016M1A5A1901769 and KOPRI-PN-19081).

Review statement. This paper was edited by Maria Kanakidou and reviewed by three anonymous referees.

References


Ding, L. C., Fu, K., Wang, D. K. W., Dann, T., and Austin, C. C.: A new direct thermal desorption-GC/MS method: organic spe-


D. K. Deshmukh et al.: Organic tracers of fine aerosol particles in central Alaska


Kourtchev, I., Warnke, J., Maenhaut, W., Hoffmann, T., and Claeys, M.: Polar organic marker compounds in PM$_{2.5}$ aerosol from a mixed forest site in western Germany, Chemosphere, 73, 1308–1314, 2008a.


Thepnuan, D., Chantara, S., Lee, C. T., Lin, N. H., and Tsai, Y. I.: Molecular markers for biomass burning associated with the characterization of PM_{2.5} and component sources during dry season


