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Author(s)	Pokhrel, Ambarish; Kawamura, Kimitaka; Ono, Kaori; Seki, Osamu; Fu, Pingqing; Matoba, Sumio; Shiraiwa, Takayuki
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4	since 1660s: Implication for climate change variability in the North Pacific Rim
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19 Abstract

20 Monoterpene and isoprene secondary organic aerosol (SOA) tracers are reported for the 21 first time in an Alaskan ice core to better understand the biological source strength before and 22 after the industrial revolution in the Northern Hemisphere. We found significantly high 23 concentrations of monoterpene- and isoprene-SOA tracers (e.g., pinic, pinonic, and 2methylglyceric acids, 2-methylthreitol and 2-methylerythritol) in the ice core, which show 24 25 historical trends with good correlation to each other since 1660s. They show positive 26 correlations with sugar compounds (e.g., mannitol, fructose, glucose, inositol and sucrose), and 27 anti-correlations with α -dicarbonyls (glyoxal and methylglyoxal) and fatty acids (e.g., C₁₈₋₁) in 28 the same ice core. These results suggest similar sources and transport pathways for 29 monoterpene- and isoprene-SOA tracers. In addition, we found that concentrations of C₅-30 alkene triols (e.g., 3-methyl-2,3,4-trihydroxy-1-butene, *cis*-2-methyl 1,3,4-trihydroxy-1-butene 31 and *trans*-2-methyl-1,3,4-trihydroxy-1-butene) in the ice core have increased after the Great Pacific Climate Shift (late 1970s). They show positive correlations with α -dicarbonyls and 32 33 fatty acids (e.g., $C_{18:1}$) in the ice core, suggesting that enhanced oceanic emissions of biogenic 34 organic compounds through the marine boundary layer are recorded in the ice core from 35 Alaska. Photochemical oxidation process for these monoterpene- and isoprene-/sesquiterpene-36 SOA tracers are suggested to be linked with the periodicity of multi-decadal climate 37 oscillations and retreat of sea ice in the Northern Hemisphere. 38

Keywords: Monoterpenes, isoprene, SOA tracers, ice core, Alaska, multi-decadal climate
oscillation, climate change

41 **1. Introduction**

42 Organic aerosols (OA) are an important fraction of atmospheric fine particles. They can 43 alter the earch radiation budget directly by scattering sunlight, and indirectly by acting as cloud 44 condensation nuclei (CCN) and ice nuclei (IN) (Aalto et al., 2001; Kanakidou et al., 2005). OA are derived from anthropogenic and biogenic sources via primary emissions and secondary 45 46 photochemical oxidaiton of various precursors (Claeys et al., 2004; Guenther et al., 2006; 47 *Kunwar and Kawamura 2014a, b*). The former can be classified as primary organic aerosols 48 (POA) whereas the latter as secondary organic aerosols (SOA). POA are emitted from plant 49 debris, fungal spore, fossil fuel combustion, biomass burning, and soil particles (Guenther et 50 al., 2006; Fu et al., 2014) whereas SOA are photochemically produced by homogeneous 51 (Claevs et al., 2004) and heterogeneous (Limbeck et al., 2003) oxidations of biogenic volatile 52 organic compounds (BVOCs) and anthropogenic VOCs. 53 Globally, annual emissions of BVOCs are estimated to be 1150 TgC/yr, where isoprene (C_5H_8) and monoterpenes $(C_{10}H_{16})$ consist 44%, and 11%, respectively. In contrast, emissions 54 55 of anthropogenic VOCs (110 TgC/yr) are one order of magnitude smaller than those of BVOC; 56 contribution of aromatic hydrocarbon is estimated to be only 13% of total anthropogenic VOCs 57 (Guenther et al. 1995, references therein). Particularly, terrestrial and oceanic phytoplankton's isoprene is the largest source of VOCs, contributing ~50% of total global emission of 309–706 58 59 TgC/yr (Acosta Navarro et al., 2014, references therein). In contrast, α -/ β -pinenes can 60 contribute 35% of total global emissions of BVOCs (Griffin et al., 1999). These BVOCs could 61 contribute significantly to the formation of SOA in the troposphere. 62 Isoprene- and monoterpene-oxidation products have been studied using field- (Claeys 63 et al., 2004; Fu et al., 2011) and laboratory-based measurements (Claevs et al., 2007; 64 Hallquist et al., 2009; Surratt et al., 2010; Noziere et al., 2015). Being different from the 65 previous assumption made by Limbeck et al. (2003), Claeves et al. (2004) first reported the

66 presence of 2-methyltetrols in the Amazonian aerosols as oxidation products of isoprene. 67 Based upon the laboratory work, the formation of SOA from isoprene is further confirmed 68 (Surratt et al., 2010). Laboratory experiments showed that the oxidation products of isoprene 69 (e.g., C₅-alkene triols and 2-methyltetrols) are enhanced in the presence of NO_x, SO₂ and/or 70 H₂SO₄ (Surratt et al., 2006, 2010). Moreover, in-cloud aqueous phase oxidation of isoprene is 71 suggested to significantly contribute to the SOA formation (Carlton et al., 2009; Volkamer et 72 al., 2009; Hallquist et al., 2009). Thus, these studies (i.e. field and laboratory measurement) 73 provide insight on the sources, spatial and temporal distributions and molecular evolution of 74 SOA.

McNeill et al. (2012) reported that organic compounds in snow/glaciers and/or in ice particles could be biological in origin and deposited from the atmosphere. Isoprene- and monoterpene-oxidation products are rarely studied in ice and snow, although biomass burning products and photooxidation products (i.e., dicarboxylic acids) of various organic precursors have been reported (*Kawamura et al., 2001; 2012a,b*). However, studies of monoterpene- and isoprene-SOA in snow and glacier ice remain poorly explored.

81 Here, we studied monoterpene- and isoprene-SOA tracers in an ice core from Alaska. 82 Our objectives are to obtain the temporal profiles of monoterpene- and isoprene-SOA tracers 83 that are derived from atmospheric photooxidation of isoprene and α/β -pinenes to better 84 understand the historical changes in biological source strength of BVOCs. In this study, we 85 report for the first time monoterpene- and isoprene-SOA tracers from mountain glaciers in 86 Alaska to better understand both POA and SOA cycles in the North Pacific Rim.

87

88 2. Samples and Analytical Procedures

89 2.1. Site description

We collected a ~180 m deep ice core on the saddle of Aurora Peak in 2008 AD, which
is located in the southeast of Fairbanks (63.52°N; 146.54°W, elevation: 2,825 m) (Figure 1).

92 The ice core sample was cut into \sim 50 cm-long pieces and directly transported to the laboratory 93 of the Institute of Low Temperature Science (ILTS), Sapporo, Japan and stored in a dark cold 94 room at -20°C until analysis. The ice core ages were determined by annual counting of 95 hydrogen isotopes (δD) and Na⁺ seasonal cycles and the age control was provided by reference horizons of tritium peaks in 1963 and 1964 (Tsushima et al., 2015). The bottom of the sample 96 97 was estimated to be 343 years old, i.e., 1665 AD (Tsushima et al., 2015). 10-day air mass 98 backward trajectory based on Lagrangian tracking method for 1992 – 2002 has been already 99 reported for Alaskan regions in the troposphere (>300 hPa) (Yasunari and Yamazaki, 2009).

100 2.2. Chemical analysis

101 Total numbers of ice core sections (50 cm long) were 147, which means that sampling 102 frequency is ~40% of the 180 m deep ice core. This sampling frequency is much better than 103 that (8%) of the previous ice core studies in Greenland (Kawamura et al., 2001). Roughly one 104 quarter of each section was used after removing the surface 5-10 mm of the ice section. These 105 ice core samples (ca. 150 mL) were first molten and then transferred to a pear shape flask (300 106 ml) and concentrated to almost dryness using a rotary evaporator under a vacuum. The total 107 dissolved and particulate organic matters were extracted with a mixture of CH₂Cl₂/CH₃OH 108 (2:1) using ultrasonic bath. The extracts were transferred to a 1.5 mL glass vial, dried under 109 nitrogen stream and then derivatized with 99% N, O-bis-(trimethylsilyl) trifluoroacetamide 110 (BSTFA) + 1% trimethylchlorosilane (TMCS) and pyridine at 70 °C for three hours (Fu et al., 111 2010a, b; Kawamura et al., 2012b; Zhu et al., 2015).

The derivatized fractions were diluted with n-hexane containing an internal standard (*n*-C₁₃ alkane) prior to the analysis with a gas chromatograph/mass spectrometer (GC/MS) installed with a capillary column (HP-5MS, 30 m × 0.32 mm I.D. × 0.25 μ m film thickness) and split/splitless injector (*Kawamura et al., 2012b*). Monoterpene- and isoprene-SOA tracers (Table 1) were determined using GC/MS method which has been reported elsewhere (*Fu et al.,* 117 2014). Triplicate analysis of real ice core sample was also conducted. Analytical errors were 118 less than 8%. Laboratory blank using pure water (150 ml), i.e., Milli-Q water, showed blank 119 levels of less than 3% of real ice core samples. All the data reported here are corrected for the 120 laboratory blank. We believe that the preservation conditions of organic compounds in the ice 121 core is fairly good and thus post-depositional microbial changes in the signals are not 122 important because no ice lense was found in the ice core studied. However, we cannot exclude 123 the posssibility that some oxidants deposited in snow may react with organic species, which 124 may decompose and/or produce some organics.

125

126 **3. Results and Discussion**

127 3.1. Historical changes of monoterpene-SOA tracers

128 Table 1 presents ice core concentrations of monoterpene- and isoprene-SOA tracers 129 since 1660s. Pinic and pinonic acids are detected as α/β -pinene oxidation products. These 130 acids are observed in smog chamber experiments (Glasius et al., 2000), and are produced from 131 photooxidation of α/β -pinene in the presence of O₃ and OH radicals (*Hoffman et al.*, 1997). 132 Pinic acid showed sporadic peaks in 1750 (concentration, 427 ng/L), 1786 (529), 1870 (406), 133 1875 (484), 1875, 1880 (3001), 1913 (630), 1973 (853), 1993 (352), 1998 (295) and 2005 134 (430) compared to its average concentration $(157\pm148 \text{ ng/L})$. Similar spikes were detected for 135 pinonic acid, except for 1980s and 1990s (Figure 2a and b). The periods for the peaks can be 136 considered as significant monoterpene oxidation periods corresponding to the formation of 137 SOA in the North Pacific region. 138 Interestingly, average concentration of pinic acid (157 ng/L) is more than double of that

of pinonic acid (70.6 ng/L), which contradicts the trend reported for aerosol studies from Mt.
Tai, East China (*Fu et al., 2010a*), tropical India (*Fu et al., 2010b*), central Greece (*Kavouras et al., 1999*), Sierra Nevada Mountains of California (*Cahill et al., 2006*) and North Carolina

142 (Bhat et al., 2007). It should be noted that vapor pressure of pinonic acid is twice higher than 143 that of pinic acid (Fu et al., 2010b). Coniferous forest is common in southern Alaska, being 144 similar to Research Triangle Park, USA and Germany (e.g., Kleindienst et al., 2007; Plewka et 145 al., 2006). A good correlation between pinonic and pinic acids (R = 0.93) in the ice core 146 indicates a similar sources and formation pathways and/or similar atmospheric fate in southern 147 Alaska for 1660s-1980s (Figure 4a). Interestingly, these compounds show a decrease during 148 the 1800-1850s and after the 1920s (except for 1973) with sporadic peaks during the 1990s, 149 suggesting that source and/or source region could be shifted to this sampling site. 150 We found significant levels of 3-hydroxyglutaric acid (3-HGA), whose historical 151 trends are completely different (Figure 2c) with pinic and pinonic acids. Smog chamber 152 experiment shows that 3-HGA is an oxidation product of α -pinene under UV irradiation in the 153 presence of NO_x (*Claeys et al., 2007*). Average concentration of 3-HGA (22.4±41 ng/L) is 154 several times lower than pinic and pinonic acids (Table1). It should be noted that we didn't 155 detect any β-caryophyllinic acid and related compounds. For instance, many aerosol studies showed higher concentrations of β -caryophyllinic acid in late winter (e.g., Fu et al., 2010b). 156 157 Correlations (R) of 3-HGA with pinic and pinonic acids are 0.46 and 0.46, respectively. These 158 results suggest that 3-HGA involves other processes than pinic and pinonic acids over the 159 southern Alaska.

160 3.2. Historical changes of isoprene-SOA tracers

161 We detected six isoprene-SOA tracers (Table 1 and Figure 3). Isoprene is readily

162 oxidized in the atmosphere by OH, O₃, NO_x, SO₂ and H₂SO₄ (*Carlton et al, 2009; Surratt et al.,*

163 2010). There are many key oxidation products of isoprene via methacrolein (a key as an

164 intermediate in SOA fromation) in ambient aereosl has been found (*Noziere et al., 2015;*

165 Hallquist et al., 2009 and references therein). Some isoprene oxidation products have less

166 vapor pressures and thus stay in particle phase, leading to the formation of SOA (Hoffmann et

167 al., 1997). 2-Methylglyceric acid can be formed by the oxidation of methacrolein and 168 methacrylic acid from two major gas-phase oxidation products of isoprene (Surrat et al., 2007; 169 *Hu et al.*, 2008), which can be altered and/or associated with land sea breeze atmospheric 170 circulation (Claevs et al., 2007; Fu et al., 2010b and references therein). 171 2-Methylglyceric acid showed higher concentrations in 1750 (74.8 ng/L), 1786 (175), 1870 (83.4), 1875 (93.1), 1880 (73.1), 1913 (166), 1973 (144), 1977 (67.3), 1993 (230), 1998 172 173 (65.9), 1999 (71.8) and 2005 (282), whose concentrations are much higher than its average 174 (35.6±48 ng/L). These periods are consistent with those of 2-methylthreitol (ave. 349±383 ng/L) and 2-methylerythritol (692±709 ng/L) as shown in Figure 3a, b and c. Interestingly, 175 176 historical trends are similar to those of pinic and pinonic acids and isoprene-SOA tracers 177 (Figure 2a, b, and 3a, b, c) before 1990s. Correlation coefficients (R) of pinonic and pinic acids 178 with 2-methylglyceric acid are 0.71 and 0.83, respectively whereas that of 2-methylglyceric 179 acid and 3-HGA is 0.67 (Figure 4b, c and d). Correlation coefficients of pinonic and pinic 180 acids with 2-methylthreitol are 0.73 and 0.77, respectively (Figure 5a, b). In addition, those of 181 pinonic and pinic acids with 2-methylerythritol are 0.71 and 0.75 (Figure 5c, d) and correlation 182 coefficients (R) of 2-methylglyceric with 2-methylthreitol and 2-methylerythritol are 0.82 and 183 0.67, respectively (not shown in figure).

184 The source strength of these SOA tracers is somewhat higher before the industrial 185 revolution than in recent years in the southern Alaskan ice core (Figure 3a-c). It should be 186 noted that 3-methyl-1,2,3-butanetricarboxylic acid (MBTCA) was reported during smog 187 chamber experiments of α -pinene with UV irradiation in the presence of NO_x (*Claevs et al.*, 188 2007). But we didn't detect MBTCA from the 1660's till 2008, suggesting that anthropogenic 189 contributions are insignificant for the last 3 centuries. In contrast, C5-alkene triols showed 190 similar historical trends each other (Figure 3d,e and f), with somewhat different trends to other 191 SOA tracers (e.g., Figure 3a, b and c). Particularly, 3-methyl-2,3,4-trihydroxy-1-butene

showed higher concentrations in 1786 (13.8 ng/L), 1947 (12.0), 1973 (30.2), 1989 (41.1), 1993 (93.5), 1998 (23.0), 1999 (31.7), 2004 (35.1), 2005 (140.7) and 2006 (46.4) compared to its average concentration (6.99±17 ng/L-ice). These periods are also consistent with those for the reaction products of *cis* and *trans* hydroxy alkenes (e.g., Figure 3d, e and f). Correlation coefficients (R) of 3-methyl-2,3,4-trihydroxy-1-butene with pinonic and pinic acids in the ice core are 0.51 and 0.63, whereas those of cis 2-methy-1 1,3,4-trihydroxy-1butene with pinonic and pinic acids are 0.49 and 0.52, respectively.

199 Moreover, correlations of 2-methylglyceric acid with 3-methyl-2,3,4-trihydroxy-1-200 butene, trans-2-methyl-1,3,4-trihydroxy-1-butene and cis 2-methyl 1,3,4-trihydroxy-1-butene 201 are 0.83, 0.77 and 0.75, respectively. Correlations of 2-methylthreitol to 2-methylerythritol, 3-202 methyl-2,3,4-trihydroxy-1-butene, cis 2-methyl 1,3,4-trihydroxy-1-butene and trans-2-methyl-203 1,3,4-trihydroxy-1-butene are positive (R=0.87, 0.65, 0.54 and 0.57, respectively). Hence, we 204 consider that atmospheric fate could be strongly involved with local or regional meteorological 205 conditions with an insignificant influence of anthropogenic activities. The most prevalent SOA 206 precursors on a global scale are terpenoids emitted from vegetations (*Claevs et al., 2007*; 207 Faiola et. al., 2014). It should be noted that more than 100 organic compounds including many 208 numbers of SOA precursors are emitted from leaf litter and/or soil, although they are less 209 significant than direct emission from plant leaves (Faiola et. al., 2014). SOA formation in the 210 forest environment (i.e., from leaf litter and/or forest soil) is also important in spring to fall 211 (Faiola et. al., 2014). For instance, rate of particle formation was high during spring from soil 212 and leaf litter in a boreal forest of southern Finland (Makela et al., 2000; Bigg et al., 2004). In 213 addition, emissions of SOA precursors (i.e., terpenoids) are high during late spring and autumn 214 in boreal pine forest floor (e.g., Aaltonen et al., 2011 and references therein). These results 215 suggest the additional sources for higher spikes in an ice core.

216 3.3. Responses of monoterpene- and isoprene-SOA tracers to tropospheric temperature

217 We found that monoterpene- (Figure 2a-b) and isoprene-SOA tracers (Figure 3a-c) 218 inversely correlate with marine biogenic tracers, i.e., even carbon numbered fatty acids (e.g., 219 C_{16:0}, C_{18:0} and C_{18:1}) (*Pokhrel et al., 2015*). In contrast, those SOA tracers show a positive 220 correlation with sugar compounds (e.g., mannitol, fructose, glucose, inositol and sucrose) from 221 the same ice core, suggesting the continental sources (unpublished data, Pokhrel and 222 Kawamura, 2015). Hence, emissions of isoprene and monoterpenes from vegetations could be 223 controlled by biological activity of the ocean and continents in different years, which could be 224 linked to histrorical variations of thermohaline circulation (Boyce et al., 2010 and references 225 *therein*).

226 Reconstructed air temperatures from the Gulf of Alaska (GOA) using tree ring showed 227 a sharp drop in June to September since 1800 to1875 except for 1830s (e.g., Wilson et al., 228 2007b), whose historical trend is similar to lower concentrations of monoterpene- and 229 isoprene-SOA tracers (Figure 2a-b and Figure 3a-c). It should be noted that sampling sites for 230 the tree ring data are located close to coastal range of the GOA and/or south of the Brooks 231 Range at about 68 °N. This GOA temperature expresses a strong consistency with multi-232 decadal climate oscillation and short-term atmospheric activities (e.g., ENSO/El-Nino and/or Aleutian Low). Interestingly, reconstructed Northern Hemisphere annual temperature since 233 234 1671 based on high-latitude tree-ring data from North America (Jacoby and D'Arrigo, 1989) is 235 also similar histrorical trend (except for few decades). In addition, tree-ring growth data and 236 inferred temperature variability at the North American Arctic tree line (D'Arrigo et al., 2009) 237 are also consistent with the trends of ice core SOA tracers.

Based on 10 day backward trajectory analyses for the years of 1992-2002, *Yasunari and Yamazaki* (2009) reported that Alaskan regions can receive significant air parcels from
adjacent North Pacific Regions, East Asia, Eastern Russia, Siberia, the Okhotsk and Bering
Seas, higher latitudes of Alaskan regions, the Gulf of Alaska, Japan, Canada, the Arctic Ocean

and Europe in the troposphere (>300 hPa). These areas are represented by extra tropical
Northern Hemispheric temperature (ENHT). ENHT is the robust temperature trends for the
Northern Hemisphere, which is calculated with more than 25 different sampling sites from tree
ring proxy records of the world. ENHT shows the positive correlations (R) with the ambient
temperatures in European Alps (0.67), western Siberia (0.61), Mongolia (0.70), Nepal (0.49),
Northern Yukon (0.60), Wrangell Mountains (0.60), British Columbia (0.77), Idaho (0.41) and
Northern Quebec (0.42) (e.g., *Wilson et al., 2007a*).

249 Lower concentrations of monoterpene- and isoprene-SOA tracers (Figures 2 and 3) 250 during this period are further supported by historical trend of ENHT (Figure 3h). For instance, 251 decreasing trends for all these SOA tracers showed positive correlations with decreasing 252 historical trends of ENHT during 1800-1860. Particularly, correlation coefficients (R) of each 253 11 points running mean (11-RM) of pinonic, pinic and 3-HGA with 11-RM of ENHT are 0.84, 254 0.86 and 0.54, respectively. Similarly, correlation coefficients (R) of 11-RM of each isoprene-SOA tracers (e.g., Figure 3a - f) with 11-RM of ENHT are 0.84, 0.95, 0.94, 0.82, 0.89, and 255 256 0.67, respectively. It should be noted that NH temperature departure and reconstructed solar 257 irradiance are well correlated with many climate periodicity and non-periodicity cycles in the 258 NH, which are reported elsewhere (Wang et al., 2012; Lean et al., 2005 and references 259 *therein*).

Aleutian Low (AL) can easily alter air parcel flow and heat exchange between the Arctic and mid-latitudinal region (*Wang et al., 2012*). *Mantua et al.* (1997) pointed out that the AL was significantly correlated with the Pacific Decadal Oscillation (PDO), which can be described as a long-lived El Niño-like pattern of Pacific climate variability. For instance, the warm phase of PDO corresponds to the strong AL, while the cold phase of PDO corresponds to the weak AL. The AL would have mixed continental and marine air mass, and more drive prevailing winds from the northwestern Pacific regions to northeastward in North America, 267 which can intensify the positive North Pacific Index (NPI) over American west (Trouet and 268 Taylor, 2010). It represents the region of 30°N-65°N, 160°E-140°W for decadal variations in 269 the atmospheric circulation (https://climatedataguide.ucar.edu). Moreover, on a decadal scale, 270 the AL has been consistently strong and drifted eastward since 1970s (Wang et al, 2012). 271 These monoterpene- and isoprene-SOA tracers (Figure 2a-b and Figure 3a-c, respectively) showed decreased trends since 1920s, which are similar to the trend of NPI. 272 273 Historical trends of C₅-alkene triols (Figure 3d-f) showed anti-correlation with those of other 274 compounds. Particularly, large-scale atmospheric circulation (or relatively local) is associated 275 with the AL system in the Bering Sea (or Alaskan regions) (Sasaki and Minobe, 2005). If this 276 is the case, it is important to understand the historical variations of atmospheric circulation 277 over the Bering Sea, Gulf of Alaska and Alaskan regions. The climatic oscillations can 278 represent the atmospheric transport in the lower latitudes. The effect of climatic oscillations 279 can be observed using NPI (Trouet and Taylor, 2010). For instance, relations between 7-RM of 280 C₅-alkene triols, i.e., 3-methyl-2,3,4-trihydroxy-1-butene, cis 2-methyl 1,3,4-trihydroxy-1-281 butene and trans-2-methyl-1,3,4-trihydroxy-1-butene (Figure 3d-f), anti-correlate with 7-RM 282 of NPI since 1899-2007 (R = 0.31, 0.32 and 0.29, respectively). These results suggest that 283 decadal variability (or atmospheric stability) may be involved with the alteration in the 284 concentrations of SOA tracers. *Tsushima et al.* (2015) reported that concentrations of Na^+ in the same ice core 285

increased after the 1970s. This increase could be resulted from changes in the position and/or strength of winter storms in the Gulf of Alaska, which is associated with the PDO shift in 1976. We obtained higher concentrations of all the compounds around 1975 (Figures 2 and 3), being similar to Na⁺ cycle of same ice core, rapid change of sea ice concentrations (SIC) and temporal wind pattern (TW) over the Bering Sea (*Sasaki and Minobe, 2005*). Moreover, historical trends C₅-alkene triols (Figure 3d-f) are similar to changes of SIC and TW over the Bering Sea for spring season after 1950s. It should be noted that SOA formation in forest
environment is important during spring and fall season compared to summer season (*Faiola et al., 2014*). *Makela et al.* (2000) and *Bigg et al.* (2004) reported that particle formation rate was
high during spring season in the boreal forest of southern Finland. SOA precursors and/or
terpenoids are also high during late spring and autumn in boreal pine forest floor (*Aaltonen et al., 2011 and references therein*).

298 In contrast, all other compounds in the ice core (Figure 2a-c and 3a-c) exhibit similar 299 trends with wintertime SIC and TW over the Bering Sea during the same periods. These results 300 indicate that different pattern of wind circulations can alter the temporal trends of these 301 compounds in ice core, further suggesting that the first case is linked with local circulation 302 over the North Pacific Rim compared to regional wind circulation for the latter case. These 303 results may suggest that climatic variations could be recorded in the ice core profile of SOA 304 tracers. Similarly, we can explain another process for vaiable concentrations of SOA tracers 305 since 1660s. For instance, we found lower concentrations of SOA-tracers around 1650-1700s 306 and 1800-1850s, which are known as cooling period of Maunder and Dalton minima and a part 307 of Little Ice Age (Figure 3h, i), respectively (Krivova, et al., 2010), suggesting lower emission 308 of isoprene and monoterpenes during the cold periods. Our ice core records of SOA tracers can 309 be further supported by air-sea and oceanic heat fluxes of the northern Gulf of Alaska (Janout 310 et al., 2013). Net heat flux and/or heat flux anomalies in winter are significantly increased 311 during 1975-2010 (except around 1998) compared to 1950-1975 (Janout et al., 2013), which is 312 consistent with historical trends of C₅-alkene triols.

These considerations suggest that concentrations of SOA tracers are controlled by temperature, pressure and wind field of the Gulf of Alaska and/or Northern North Pacific Rim. These three parameters alter the atmospheric transport (e.g., the AL can alter the atmospheric transport), which are correlated with ENHT and NPI elsewhere (NPI has negative correlation 317 with C₅-alkenes) (Wilson et al., 2007a,b; Wang et al., 2012 and references therein). We found 318 that C₅-alkene triols positively correlate with ENHT during the 1750s to 1980s. Particularly, 319 correlation coefficients (R) between 5 points running mean (5-RM) of C₅-alkene triols (Figure 320 3d, e and f) with ENHT are 0.73, 0.80 and 0.78 for individual species (3-methyl-2,3,4-321 trihydroxy-1-butene, cis 2-methyl 1,3,4-trihydroxy-1-butene and trans-2-methyl-1,3,4-322 trihydroxy-1-butene), respectively. It is likely that BVOC emissios could be associated with 323 type and density of vegetation in landscape, temperature and radiation, having a significant 324 impact on SOA formation (Pokhrel, 2015).

325 In addition, C₅-alkene triols have increasing trends of the concentrations since 1920s, 326 which is similar to sea surface temperature (SST) over the past century (Boyce et al., 2010 and 327 references therein). Hence, these increasing trends of C₅-alkene triols (see Figure 3d-f) could 328 be involved with climatic and oceanographic variability. Particularly, an increased SST over 329 the past century since 1920s (except for few points) is similar to historical trends of SOA 330 tracers, suggesting that an enhanced ocean warming could cause a restructuring of marine 331 ecosystems and/or likely ocean circulation (Boyce et al., 2010; and references therein), 332 followed by intensified production of SOA in southern Alaska. In the same ways, we may 333 explain other historical trends since 1660s, which could be altered by complex mechanisms of 334 thermohaline circulation and productivity of marine biota, which are reported elsewere (Boyce 335 et al., 2010; and references therein).

336 Moreover, C₅-alkene triols have positive correlations with α -dicarbonyls (i.e., glyoxal

and methylglyoxal) and even-carbon numbered low molecular weight fatty acids (e.g., C_{16:0}

and $C_{18:0}$) including unsaturated fatty acids ($C_{18:1}$), which are originated from marine

339 phytoplankton (*Pokhrel et al., 2015*). Increased historical trends of C₅-alkenes after 1920s are

340 somewhat similar to the reconstructed solar total irradiance (Figure 3i) (Krivova et al., 2010;

341 Ball et al., 2012). In addition, historical concentrations of C₅-alkene triols are

changed/increased significantly after the Great Pacific Climate Shift. In contrast, other compounds (Figure 2a-c and 3a-c) show negative correlation with α-dicarbonyls and fatty acids including oleic acid (e.g., $C_{16:0}$, $C_{18:0}$ and $C_{18:1}$). But they have positive correlations with sugar compounds (e.g., mannitol, fructose, glucose, inositol and sucrose) from same ice core sample (unpublished data, Pokhrel and Kawamura, 2015). These results all suggest that an increased biogenic emission capacity and/or source could be changed, being recorded in a clear historical profile of dual sources; marine phytoplankton and terrestrial plants.

349

350 4. Summary and Conclusions

We report for the first time historical records of monoterpene- and isoprene-derived secondary organic aerosols (SOA) tracers in glacier ice core from Aurora Peak in Alaska since 1660s, which could be derived from dual biogenic sources (marine and continental). Positive correlations of monoterpene tracers with 2-methylglyceric acid and 2-methyltetrols are found to be strong, indicating the shared source regions and transport. In contrast, correlations of monoterpene tracers with C₅-alkene triols are weak, suggesting the different source regions and transport pathways.

358 Ice core concentrations of monoterpene- and isoprene-SOA tracers showed lower levels 359 during 1800-1860, which are similar to temperature trends of the Gulf of Alaska and showed a 360 positive correlation with extra-tropical Northern Hemisphere temperature (ENHT). In addition, 361 C_5 -alkene triols showed positive correlations with α -dicarbonyls, ENHT and fatty acids, and 362 their historical trends are increased after the Great Pacific Climate Shift. In contrast, all other 363 compounds (i.e., monoterpene-SOA tracters) have negative correlation with α -dicarbonyls and 364 fatty acids including oleic acid $(C_{18:1})$. But they have positive correlations with sugar 365 compounds (e.g., mannitol, fructose, glucose, inositol and sucrose) in the same ice core.

366 These results suggest that source could be changed significantly after the Great Pacific

367 Climate Shift and exhibit a clear historical dual sources likely to be marine boundary layer.

368 This study reveals that ice core records of SOA tracers and atmospheric oxidizing capability or

- 369 emission of monoterpenes and isoprene are associated with multi-decadal climate oscillation
- 370 periodicity in the North Pacific Rim, which are associated with the Northern Hemispheric
- 371 temperature signals.
- 372

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- 562 Atmos. Chem. Phys. 15, 1–15. doi: 10.5194/acp-15-1-2015. 563
- 564

565 Table 1. Concentrations of monoterpene and isoprene secondary organic tracers in ice core

566 samples from southeast Alaska in 1665 - 2008.

	Concentrations (ng/L)			
Name	Ave.	Min.	Max.	SD
α/β-Pinene-SOA tracers				
Pinic acid	157	BDL	853	148
Pinonic acid	70.6	BDL	300	68
3-HGA	22.4	BDL	234	41
Isoprene-SOA tracers				
2-methylglyceric acid	35.6	BDL	282	48
2-methylthreitol	349	BDL	1740	383
2-methylerythritol	692	BDL	3509	709
3-methyl-2,3,4-trihydroxy-1-butene	6.99	BDL	141	17
cis-2-methyl-1,3,4-trihydroxy-1-butene	23.1	BDL	384	52
trans-2-methyl-1,3,4-trihydroxy-1-butene	36.5	BDL	786	98
BDL= Below detection limit (0.001 ng/g-ice)				





571 Figure 1. Map showing the geographical location of Aurora Peak of Alaska, from which 180-

- 572 meter long ice core sample was drilled in 2008 (*Pokhrel et al., 2014*).

- ___



Figure 2. Concentration changes of (a) pinic acid, (b) pinonic acid (c) 3-hydroxyglutaric acid
(3-HGA) and (d) average concentration changes of monoterpene tracers (a)-(c) in the Alaska
ice core records collected from Aurora peak.



Figure 3. Concentration changes of (a) 2-methylglyceric (b) 2-methylthreitol (c) 2-methylerythritol (d) cis 2methyl 1,3,4-trihydroxy-1-butene (e) 3-methyl-2,3,4-trihydroxy-1-butene (f) trans-2-methyl-1,3,4-trihydroxy-1butene and (g) total annual average concentrations of all these isoprene SOA tracers (a – f) in the Alaska ice core
records collected from Aurora peak (h) extra tropical Northern Hemispheric temperature (ENHT) departure, e.g.,
Wilson et al. (2007a) and (i) 7 years running mean of reconstructed solar total irradiance (STI) from IPCC AR5
based on Krivova et al. (2010) and Ball et al. (2012).



595

596 Figure 4. Correlation between the concentrations of (a) pinonic and pinic (b) pinonic acid and

- 597 2-methylglyceric (e) pinic acid and 2-methylglyceric and (f) 2-methylglyceric and 3-
- 598 hydroxyglutaric acid (3-HGA) in the Alaska ice core records collected from saddle of Aurora
- 599 Peak.





Figure 5. Correlation between the concentrations of (a) pinonic and 2-methylthreitol (b) pinic
and 2-methylthreitol, (c) pinonic and 2-methylerythritol, and (d) pinic and 2-methylerythritol
in the Alaskan ice core records collected from saddle of Aurora Peak.

605 Highlights

- 606 1. Monoterpene-SOA tracers in south Alaskan ice core are associated with continental sources.
- 607 (80 with space)
- 608 2. Isoprene-SOA tracers in the ice core are derived from both continental and marine sources.
- 609 (75 with space)
- 610 3. Monoterpene- and isoprene-SOA tracers are linked with climate oscillations of Aleutian Low.
- 611 (80 with space)