

To
Prof. Jason Surratt
Editor
Atmospheric Chemistry and Physics (ACP)

RE: Submission of revised manuscript entitle “**Thirteen years of observations on primary sugars and sugar alcohols over remote Chichijima Island in the western North Pacific**” for publication in Atmospheric Chemistry and Physics

Reference: MS No.: acp-2017-480

Dear Prof. Surratt,

Thanks for the decision letter on our manuscript. We have revised the manuscript meticulously following the referees' comments and suggestions. The authors greatly appreciate the critical and useful comments by anonymous referee's that significantly improved the quality of our manuscript. Our responses and changes are included in the response letter as blue color. The changes in the revised manuscript are highlighted in yellow. Please find the response letter and revised manuscript.

We believe that revised manuscript is acceptable for publication in ACP.

Thanking you.

Sincerely yours,

Kimitaka Kawamura
Professor of Chemistry,
Chubu Institute for Advanced Studies,
Chubu University, Kasugai 487-8501,
Japan
E-mail: kkawamura@isc.chubu.ac.jp

Authors' responses to the comments Referee # 1

General:

In general this paper reports valuable measurements for a number of sugars and sugar alcohols over a remote site in the middle of the open ocean over a long time period and conducts rational discussions for their seasonal and temporal trends, variation due to transportation and meteorological parameters, and source apportionment. The chemical, statistical, and other supplementary techniques for sample and data analyses are handled quite properly in order to provide and support their observations and conclusions.

As another anonymous referee mentioned, the English writing of the manuscripts should be further carefully checked and improved.

Overall, I think the paper is of decent quality for ACP to accept for publication after minor revision.

Authors are thankful to the reviewer for the positive recommendation. We carefully checked and improved English writing of through out the manuscripts.

Individual Questions:

1. Although this might be complex and not so feasible, is there a better way to clearly present the types or contributions of parameters (i.e., local emission, wind, oceanic emission, etc) that dominate or influence the observed concentrations in different seasons?

Response: Authors are thankful to the reviewer for suggestion. We studied the atmospheric conditions over Chichijima Island and found that the aerosol composition of the Chichijima Island is strongly influenced by the atmospheric circulation in the western North Pacific with first westerly winds and second trade winds. The westerly winds transport continental air mass from East Asia during winter/spring, whereas the trade winds transport clean air mass from the central Pacific Ocean during summer/autumn. In our study we have discussed both local sources and long-range transport of SCs as follows:

1. The seasonal variation of sucrose in Chichijima aerosols was suggested with multiple sources, including both long-range transported and local emission. "The seasonal mean concentrations of sucrose are almost equal during spring (8.80 ± 18.0 ng m⁻³), summer (7.31 ± 11.5 ng m⁻³) and winter (6.60 ± 13.1 ng m⁻³), except for autumn (2.76 ± 4.35 ng m⁻³) (Table 1). The similar seasonal distributions suggest multiple sources of sucrose in Chichijima aerosols." The locally emitted pollen by flowering plant influenced the springtime sucrose concentration. The long-range transport of pollen from East Asia, Mongolia and Russian For East under the influence of westerly winds significantly contributes to the sucrose concentrations in the aerosols collected over Chichijima Island. (Please see lines 360-363 in the revised MS).

2. The contributions of arabitol and mannitol in the Chichijima aerosols were locally originated by the fungal and microbial metabolic activities under the favorable meteorological conditions (RH and temperature). "Consequently, bacteria and fungi associated with bioaerosols grow extensively during summer/autumn, when the climate conditions (i.e., higher RH and temperature) are favorable for their metabolic activities (Morris et al., 2004)." (Please see lines 532-534 in the revised MS).

3. The results of decadal study suggested an increased long-range air mass transport of bioaerosols from East Asia to the western North Pacific for the last decade. "Accordingly, an increased transport of bioaerosols for the last decade may have caused a drastic increase in the concentrations of sugar alcohols during P-II and P-III compared to P-I period over

the western North Pacific.” (Please see lines 535-537 in the revised MS).

4. The trade winds transport clean oceanic air mass from the central Pacific Ocean over Chichijima during summer/autumn. The higher concentrations of arabitol, mannitol, glucose, trehalose and erythritol are observed in Chichijima aerosols during summer/autumn. Although we also tried to discuss the oceanic sources for those SCs from the Pacific Ocean but due to the lack of sufficient data set and recent literature, we could not clarify either those sources are authentic or not. “Several studies have described the occurrence of fungi in marine environment.....Therefore, due to the inadequate data set, we doubt the marine contribution of sugar alcohols (arabitol, mannitol) in Chichijima aerosols.” (Please see lines 455-467 in the revised MS).

Hence, we concentrated on local emission sources for those SCs, which showed higher concentrations during the trade wind periods (summer/autumn) over Chichijima Island. SCs are emitted from the local vegetation and metabolic activities of microbial and fungal species under favorable meteorological conditions.

2. Could you use the different wind directions, based on the backward air trajectory analysis, to categorize different sources of incoming aerosols, and investigate their influence? For example, westerly wind for aerosol transported from Asian continent; trade winds in summer/autumn for Pacific Ocean; no wind for local emission, etc.

Response: Thank you for the suggestions. We discussed a significant role of the atmospheric circulation (westerly and trade winds) in the seasonal concentrations of SCs in aerosol samples collected at Chichijima Island. We also discussed local sources for some SCs.

Westerly winds:

1. Please see lines 318-325 in the revised MS: “In winter/spring, Chichijima is influenced by strong westerly winds that deliver the air masses from the Asian continent including Mongolia, Russian Far East and North China, where vegetations are active. Consequently, declined concentrations of glucose in winter mean a depressed transport of glucose associated with continental bioaerosols from Asia despite long-range transport of Asian dusts due to strong westerly winds. The local vegetation (vascular plants) in Chichijima Island might be responsible to enhanced concentrations of glucose during growing season (spring and summer) and decaying periods of plant leaves (autumn).”

2. Please see lines 374-379 in the revised MS: “However, the possibilities of pollen transport from East Asia to Chichijima cannot be excluded because pollens can travel long distances with springtime high-speed winds by westerlies (Rousseau et al., 2006). The pollen grains emitted from flowering boreal forest in China, Mongolia, Siberia and Russian Far East, could significantly be delivered to the western North Pacific during spring, which may result in the contribution of sucrose and fructose to Chichijima aerosols.”

3. Please see lines 387-402 in the revised MS: “These observations may support that westerly winds have delivered pollen grains from the Asian.....which leads to smaller particles with longer residence time in the atmosphere.”

4. Please see lines 528-537 in the revised MS: “This study suggested the possibilities of the long-range transport of the fungal.....an increased transport of bioaerosols for the last decade may have caused a drastic increase in the concentrations of sugar alcohols during P-II and P-III compared to P-I period over the western North Pacific.”

Trade winds:

Chichijima Island is dominated by trade winds during summer/autumn. We tried to find out some oceanic sources (like marine fungi) of SCs, transported under the influence of trade winds from the central Pacific Ocean but we did not find strong supportive data set to conclude the oceanic emissions of SCs.

1. Please see lines 455-467 in the revised MS: “Several studies have described the occurrence of fungi in marine environment (Jones, 1976; Kohlmeyer and Kohlmeyer, 1991; Moss, 1986). The fungal species eject spores.....Therefore, due to the inadequate data set, we doubt the marine contribution of sugar alcohols (arabitol, mannitol) to Chichijima aerosols.”

2. Please see line no. 405-415 in the revised MS: “In China and India there are two seasons (spring and winter) for wheat crops; winter wheat is harvested from mid-May to.....suggest the transport of sucrose associated with soil particles under the influence of occasional air mass transport in summer from Southeast Asia in summer (Figure 3)”.

Local Sources:

The local sources are discussed for those SCs, which are dominant during summer/autumn.

1. Please see lines 334-337 in the revised MS: “Monthly mean concentrations of fructose show two prominent peaks in February-March and June-July, the latter peak may be due to the local vegetation in Chichijima (Figure 5g). The fructose peak in February-March may be influenced by air borne pollen grains in the spring bloom of flowering plants.”

2. Please see lines 449-454 in the revised MS: “The meteorological factors such as RH and temperature significantly affect fungal and bacterial activity (Kim and Xiao, 2005; Malik and Singh, 2004). Higher RH and temperature are crucial in increasing fungal and bacterial growth (Sharma and Razak, 2003). Their maximum growth was observed under the condition of 92-100% RH (Ibrahim et al., 2011). Higher concentrations of arabitol and mannitol in summer and autumn may be caused by the increased fungal and bacterial activities in Chichijima Island.”

3. Please see lines 471-477 in the revised MS: “The thirteen-year precipitation record over Chichijima Island shows that precipitations were lowered in July and August (Figure 2). The lower precipitation amount decreases the RH (Figure 2) and thus depresses the fungal and microbial activities. The lower precipitation also suppresses the moisture contents in the surface soil of Chichijima, which should result in a significant decline of local fungal and other microbial activities on the ground of Chichijima Island. Decreased precipitation might be a possible reason for the lower concentrations of arabitol, mannitol, and trehalose in July and August.”

4. Please see lines 507-511 in the revised MS: “PMF analysis showed that local emissions from vegetation are important contributor for primary sugars (glucose, fructose and sucrose). Therefore, a drastic increase in the concentrations of primary sugars in summer/autumn for P-II and P-III than P-I may be caused by an increased emission of primary sugars by local vegetation under the influence of meteorological conditions in the western North Pacific.”

5. Please see lines 552-556 in the revised MS: “In Chichijima aerosols, glucose and fructose are significant contributors in spring (Figure 6b), summer (Figure 6c) and autumn

(Figure 6d), Therefore, the respective factors in Figure 6 are termed as a vegetation source for both sugar species. This is reasonable because plants started growing in spring and summer seasons. In autumn, leaf senescence and decay result in the emission of glucose and fructose to the atmosphere.”

Detailed Comments:

Line 73- the sentence starting with "Previous studies..." needs revision in term of grammar.
Response: Corrected. “Sucrose is dominant sugar component in airborne pollen grains and plays a significant role in plant blossoming activity (Bieleski, 1995; Fu et al., 2012; Pacini, 2000)”. (Please see lines 80-82 in the revised MS).

Line 77- the word "form" seems to be "from".
Response: Corrected. Please see line 84 in the revised MS.

Line 82- the sentence starting with "Thus, ..." can be removed, and the narrative in this and the next paragraph can be largely refined.
Response: The sentence was removed and the next paragraph has been modified. Please see lines 89-100.

Line 183- this section about PMF can be considerably shortened by removing unrelated content and using reference.
Response: According to referee’s suggestions, “Section 2.4 Positive matrix factorization (PMF) analysis” was significantly shortened. We added a new sentence: “The detailed discussions of the determination and application of the PMF are reported in Norris et al. (2008), Paatero et al. (2002) and Zhou et al. (2004).” (Please see lines 206-208 in the revised MS).

Line 510- "20004a" seems to be "2004a"?
Response: Corrected. (Please see line 495 in the revised MS).

Line 514- be consistent about using hyphenation, for example, “BB tracers” or “BB-tracers”.
Response: We decided to use “BB-tracers” throughout the manuscript. (Please see the revised MS).

1 **Thirteen years of observations on primary sugars and sugar alcohols over remote**
2 **Chichijima Island in the western North Pacific**

3

4

5 Santosh Kumar Verma^{1,2}, Kimitaka Kawamura^{1,3*}, Jing Chen^{1,4}, and Pingqing Fu^{1,5}

6

7

8 ¹ Institute of Low Temperature Science, Hokkaido University, N19, W8, Kita-ku, Sapporo 060-
9 0819, Japan

10 ² Permanent address: State Forensic Science Laboratory, Home (Police) Department,
11 Government of Chhattisgarh, Raipur 49001, India

12 ³ Now at Chubu Institute for Advanced Studies, Chubu University, Kasugai 487-8501, Japan

13 ⁴ Now at Institute of Geographic Sciences and Source Research, Chinese Academy of Sciences,
14 Beijing 100101, China

15 ⁵ Now at LAPC, Institute of Atmospheric Physics, Chinese Academy of Sciences, Beijing
16 100029, China

17

18

19 *Corresponding author: kkawamura@isc.chubu.ac.jp

20

21

22 **Abstract.** In order to understand the atmospheric transport of bioaerosols, we conducted long-
23 term observations of primary sugars and sugar alcohols over remote Chichijima Island in the
24 western North Pacific from 2001 to 2013. Our results showed that concentrations of total sugar
25 compounds for 13 years ranged from 1.2 to 310 ng m⁻³ (average, 46±49 ng m⁻³). We found that
26 atmospheric circulations significantly affect the seasonal variations of bioaerosol distributions
27 over the western North Pacific. The primary sugars (glucose and fructose) maximized in
28 summer, possibly due to an increased emission of the vegetation products from local vascular
29 plants in Chichijima. We also found higher concentrations of sugar components (arabitol,
30 mannitol and trehalose) in more recent years during summer/autumn, suggesting an enhanced
31 emission of fungal and microbial species over the island. Sucrose peaked in late winter to early
32 spring, indicating a springtime pollen contribution by long-range atmospheric transport, while
33 elevated concentrations of sucrose in early summer could be explained by long-range transport
34 of soil dust from Southeast Asia to Chichijima. Sucrose and trehalose were found to present
35 increasing trends from 2001 to 2013, while total sugar components did not show any clear
36 trends during the thirteen years periods. Positive matrix factorization analyses suggested the
37 locally emitted sugar compounds as well as long-range transported air borne pollen grains,
38 microbes and fungal spores are the major contributors to total sugar compounds in the
39 Chichijima aerosols. Backward air mass trajectories support the atmospheric transport of
40 continental aerosols from the Asian continent during winter/spring over Chichijima.

41
42 **Keywords:** Sugar compounds, fungal and microbial tracer, pollen tracer, bioaerosols, the
43 western North Pacific.

44

45 **1 Introduction**

46 East Asia has experienced rapid economic developments and population growth since last
47 several decades (Elliot et al., 1997; Jaffe et al., 1999, 2003), whose activities emit organic and
48 bioaerosols into the atmosphere (Xu et al., 2011). The atmospheric particles are transported to
49 downwind region in the Pacific, associated with Asian desert dust from the Taklamakan and
50 Gobi Deserts, and Loess plateau (Duce et al., 1980; Iwasaka et al., 1983; Jaffe et al., 1997;
51 Prospero and Savoie, 1989; Talbot et al., 1997). The transported dust contains bacterial cells,
52 fungal spores, and microbial cells, which fall out over the Pacific and remote islands in the
53 Pacific Ocean (Lacey and West, 2006; Mims and Mims, 2003). The microbes associated with
54 bioaerosols significantly affect the natural environment of marine and land ecosystem in
55 downwind regions (Graffin et al., 2003, 2007; Prospero et al., 2005). Long-range atmospheric
56 transport plays a key role for the global distribution of microbes from source regions to
57 receptor site (Graffin et al., 2001). Fungi and bacteria are often attached to dust particles,
58 which can propagate diseases to human and plants (Brown and Hovmoller, 2002). Therefore,
59 the transported organic- and bio-aerosols have been the focus of extensive studies for the past
60 years (Yamaguchi et al., 2012).

61 Organic aerosols are composed of a complex mixture of different types of molecules, in
62 which water-soluble organic compounds (WSOCs) are enriched (Graham et al., 2002). WSOCs
63 play an important role in climate change and global radiative forcing by scattering or absorbing
64 light directly or indirectly (Fuzzi et al., 2007). They can act as cloud condensation nuclei
65 (CCN) (Kanakidou et al., 2005; Martin et al., 2010). Sugar compounds (SCs) contribute 13–
66 26% and 63% of total WSOCs identified in continental and marine aerosol samples,
67 respectively (Simoneit et al., 2004a; 2004b). Yttri et al. (2007) reported that sugars (fructose,
68 glucose, sucrose, trehalose) accounted for 0.6-3.1% of the WSOC at urban and suburban sites
69 in Norway. Tominaga et al. (2011) analyzed aerosol samples collected from urban and forest
70 suburban site from Japan and reported that the sugars (arabinose, fructose, galactose, glucose,

71 mannose, rhamnose, and xylose) accounted for 2.1% and 4.5% of the WSOC in the fine and
72 coarse mode ranges at Yokohama, respectively, and for 3.0% and 7.2% at Mt. Oyama,
73 respectively. SCs are directly emitted from biological sources such as fungi, algae, pollen,
74 spores and bacteria (Carvalho et al., 2003; Wang et al., 2009) and transported long distances in
75 the atmosphere (Wang et al., 2011). They are also derived from suspended soil particles and
76 associated biota (Rogge et al., 2006; Simoneit et al., 2004b; Wang et al., 2009), and biomass
77 burning (Schmidl et al., 2008; Simoneit et al., 2002).

78 Primary sugars are emitted from biological sources (Medeiros et al., 2006). Glucose and
79 fructose are emitted from terrestrial plant fruits, pollen, and detritus of vascular plants (Cowie
80 and Hedges, 1984; Speranza et al., 1997). Sucrose is dominant sugar component in airborne
81 pollen grains and plays a significant role in plant blossoming activity (Bielecki, 1995; Fu et al.,
82 2012; Pacini, 2000). Trehalose is emitted from fungal metabolic activities and resuspension of
83 soil particles and unpaved road dust (Rogge et al., 2007; Simoneit et al., 2004). Sugar alcohols
84 are also emitted from biological sources like fungi and microbes via metabolic activities (Bauer
85 et al., 2008). Sugar alcohols, i.e., arabitol and mannitol, are tracers for fungal spores (Jia and
86 Fraser; 2011; Yang et al., 2012). Di Filippo et al. (2013) reported that arabitol and mannitol
87 are key sugar components in fungal spores.

88 Chichijima Island is located in the western North Pacific: an outflow region of Asian
89 dust (Mochida et al., 2003a). It is one of the best remote islands to study a long-range transport
90 of Asian aerosols, because local pollutants in Chichijima are insignificant due to low
91 population density and no major sources from industrial or anthropogenic activities (Chen et al.,
92 2013). Kawamura et al. (2003) reported that concentrations of lower molecular weight fatty
93 acids (C₁₂-C₁₉) derived from marine organisms became higher in summer, while those of
94 higher molecular weight fatty acids (C₂₁-C₃₄), n-alkanes (C₂₅-C₃₅), n-alcohols (C₂₀-C₃₄) and
95 dicarboxylic acids (C₂₀-C₂₈) derived from terrestrial higher plants and soil organic matter
96 maximized in winter to spring. Seasonal variations of low molecular weight (C₂-C₁₀)

97 dicarboxylic acids and levoglucosan (biomass burning tracer) have also been reported in
98 Chichijima aerosols by Mochida et al. (2003a) and Mochida et al. (2010), respectively.
99 Although seasonal variation of saccharides was reported in Chen et al. (2013), the observation
100 period is rather short.

101 Here, we report thirteen year data set of SCs in remote Chichijima Island. The goal of
102 this study is to characterize seasonal and annual variations of SCs and specify their possible
103 source regions. We will also discuss a potential role of Asian dust to control the distributions
104 of bioaerosols over the western North Pacific. The outcomes of this study will improve our
105 understanding about a possible influence of long-range transport of bioaerosols from the
106 continent to the clean oceanic environment. We will compare the data set of SCs for the
107 periods 1990-1993, 2001-2003 and 2010-2013, which may provide imperative information
108 about decadal changes in the atmospheric conditions over Chichijima. Seasonal source
109 identifications by positive matrix factorization (PMF) analysis will also be discussed for the
110 measured SCs.

111

112 **2 Materials and methods**

113 **2.1 Sampling site and meteorological conditions**

114 The detailed information on the sampling site was reported in Kawamura et al. (2003) and
115 Chen et al. (2013). Briefly, Chichijima Island is located in the western North Pacific (27°04'N;
116 142°13'E), 1000 km south of Tokyo, Japan, and 2000 km east of the Asian continent (Figure 1).
117 Total area of the island is 24 km² with a population of 2000 (Verma et al., 2015). The climate
118 of Chichijima is classified as subtropical; it is warm to hot (temperature, 7.8-34.1 °C) and
119 humid (relative humidity, 66-88%) all year round.

120 Figure 2 shows monthly averaged variations in the meteorological parameters of
121 Chichijima during 2001-2013. It receives more precipitation in between April and July,
122 September and October during the thirteen year period. The sampling site is less influenced by

123 the East Asian monsoon to receive heavy rainfall compared to Northeast Asia. The climate
124 over Chichijima is strongly influenced by the seasonal changes in wind system. In
125 winter/spring, the westerly winds are dominant with the air masses being enriched with Asian
126 dust, industrial pollutants, biomass burning products, organic compounds and black carbon as
127 well as bioaerosols emitted from East Asia and Eurasia (Figure 3) (Seinfeld et al., 2004;
128 Simoneit et al., 2004b; Wang et al., 2009). Trade winds are dominant in summer/autumn,
129 which transport clean and pristine marine air masses from the central Pacific to Chichijima
130 (Kawamura et al., 2003, Mochida et al., 2010).

131 **2.2 Aerosol sampling and chemical analysis**

132 The details on aerosol sampling and chemical analysis are reported elsewhere (Chen et al.,
133 2013; Mochida et al., 2010). Briefly, total suspended particle (TSP) samples were collected at
134 the Ogasawara Downrange Station of the Japan Aerospace Exploration Agency (JAXA) in
135 Chichijima Island (254 m, above sea level, asl). The samples were collected on weekly basis
136 (January 2001 to November 2013) using a high-volume air sampler (Kimoto AS-810A) at a
137 flow rate of $1.0 \text{ m}^3 \text{ min}^{-1}$ and pre-combusted (450°C for 6 h) quartz fiber filters (20 x 25 cm,
138 Pallflex). Filter sample was placed in a pre-combusted glass jar with a Teflon-lined screw cap
139 and stored in a dark freezer room at -20°C prior to analysis in order to inhibit fungal growth.
140 Due to the maintenance of the JAXA facility at sampling site, TSP samples were not collected
141 for November–December 2004 and March–August 2005.

142 Total 590 aerosol samples were analyzed to determine primary sugars (xylose, fructose,
143 glucose, sucrose and trehalose) and sugar alcohols (erythritol, arabitol, mannitol and inositol)
144 during 2001 to 2013. An aliquot (21 cm^2) of the filters were extracted three times with
145 dichloromethane/methanol (2:1, v/v) mixture using ultrasonic agitation for 10 minutes. A
146 Pasteur pipette packed with quartz wool was used to remove particles and filter debris in the
147 extracts. Filtrates were then concentrated using a rotary evaporator under vacuum and blown
148 down with a stream of pure nitrogen gas. The total extracts were derivatized using 60 μl of

149 N,O-bis-(trimethylsilyl)trifluoroacetamide (BSTFA) with 1% trimethylsilyl chloride in the
150 presence of 10 μl of pyridine in a sealed vial at 70 $^{\circ}\text{C}$ for 3 hours to convert hydroxyl groups to
151 corresponding trimethylsilyl (TMS) ethers. The derivatized fractions were diluted with n-
152 hexane containing internal standard of C_{13} n-alkane ($1.43 \text{ ng } \mu\text{l}^{-1}$), prior to injection to gas
153 chromatography-mass spectrometer (GC-MS).

154 Identification of SCs have been confirmed by the comparison of GC retention times and
155 mass spectra with those of authentic standards as well as literature and library data. SCs were
156 characterized by their common base peak at m/z 217 and 204 with specific fragment ions for
157 individual sugars, i.e., $m/z = 307$ (arabitol), 205 and 319 (mannitol), 205 (erythritol), 305 and
158 318 (inositol), 361 (sucrose and trehalose), 191 (glucose), and 437 (fructose). The selected ion
159 peak area and relative response factors determined by injection of authentic standards have
160 been used for the quantification of sugar compounds. Field blank filters were analyzed as a real
161 sample, but no target compounds were detected in the field blanks. The recoveries of the target
162 compounds were better than 90%. Therefore, the data reported here were not corrected for
163 recoveries. Analytical errors of SCs were generally $<15\%$ based on duplicate analysis. The
164 detection limits of primary sugars and sugar alcohols were $105\text{-}557 \text{ pg } \mu\text{l}^{-1}$, which corresponds
165 to ambient concentrations of $0.0015\text{-}0.0081 \text{ ng m}^{-3}$ under a typical sampling volume of 9000
166 m^3 (Zhu et al., 2015).

167 The derivatized fractions were introduced into GC-MS using an Agilent model 7890 GC
168 coupled to an Agilent model 5975 mass selective detector (MSD) operated in an electron
169 impact mode at 70 eV and scanned from 40 to 650 Dalton. The GC separation was carried out
170 on a DB-5MS fused silica capillary column (30 m long, 0.25 mm i.d., 0.25 μm film thickness),
171 with a temperature program of 50 $^{\circ}\text{C}$ for 2 min at a rate of $15 \text{ }^{\circ}\text{C min}^{-1}$ from 50 to 120 $^{\circ}\text{C}$, then
172 from 120 to 305 $^{\circ}\text{C}$ at a rate of $5 \text{ }^{\circ}\text{C min}^{-1}$ with a final isotherm hold at 305 $^{\circ}\text{C}$ for 15 min. The
173 sample was injected on a splitless mode at an injector temperature of 280 $^{\circ}\text{C}$. GC-MS data
174 were acquired and processed with the Agilent GC/MSD ChemStation software.

175 **2.3 Backward air mass trajectory analysis**

176 In order to identify the source regions of sugar compounds in Chichijima aerosols, ten-day
177 backward trajectories were calculated at 00:00 UTC of each sampling period for thirteen years
178 using the NOAA Hybrid Single-Particle Lagrangian Integrated Trajectory
179 (<http://ready.arl.noaa.gov/HYSPLIT.php>) (Figure 3). The starting height of the trajectories
180 presented in this study is 500 m asl. We plotted thirteen year trajectories for each sampling day
181 but there are no significant year-to-year changes in the atmospheric circulations. Therefore, we
182 presented seasonal trajectories for recent year (December, 2011 to November, 2012) in Figure
183 3 to understand the seasonal aerosol mass transport from the source regions to Chichijima
184 Island. Backward trajectories significantly supported a long-range transport of air mass under
185 the influence of existing meteorological parameters (Figure 3). The trajectories clearly show
186 the influences of continental air masses during mid-autumn to mid-spring and of marine air
187 masses during mid-spring to mid-autumn.

188 **2.4 Positive matrix factorization (PMF) analysis**

189 Positive matrix factorization (PMF 3.0, Environmental Protection Agency, USA) has
190 been used as a powerful statistical tool that may resolve potential sources contributing to
191 atmospheric levels of particle (as presented by %) when appropriate source profiles are not
192 available (Paatero and Tapper, 1994). At the beginning PMF has been used in precipitation
193 study (Juntto and Paatero, 1994) as well as air pollution and source apportionment studies
194 (Polissar et al., 1999). Recently, it is widely used for the air quality and source apportionment
195 (Xie and Berkowitz, 2006). In addition, PMF has been applied to the wastewater
196 (Soonthornnonda and Christensen, 2008), lake sediments (Bzdusek et al., 2006) and soils (Lu
197 et al., 2008). One of the main features of PMF results is their quantitative nature; it is possible
198 to obtain the composition of the sources determined by the model.

199 PMF analysis was performed for quantitative estimation of sources for the collected
200 samples using tracer compounds for primary sugars, sugar alcohols, and anhydrosugars. Based

201 on given understanding of sugar sources, 4-7 factors were examined and total five interpretable
202 factors were characterized by the enrichment of each tracer compound, which reproduced more
203 than 94% of SCs. Minimal robust and true Q values of the base run were 3001 and 3413,
204 respectively. Concentrations and percentage of tracers in each factor of bootstrap run were
205 close of those of base run results. The Q values and factor profiles of F_{peak} rotation runs showed
206 no significant changes compared with base run, indicating stable PMF results. **The detailed**
207 **discussions of the determination and application of the PMF are reported in Norris et al. (2008),**
208 **Paatero et al. (2002) and Zhou et al. (2004).**

209 In winter/spring, Chichijima Island receives air masses enriched with anthropogenic
210 aerosols from the Asian continent by strong westerly winds, whereas during summer/autumn it
211 receives clean air masses from the Pacific Ocean under the influences of trade winds. The
212 seasonal changes in the atmospheric circulation over Chichijima may have a significant
213 influence on the seasonal distributions of SCs. **Therefore,** we performed the seasonal PMF
214 analysis on the thirteen year sugar data set to better understand the seasonal source profile of
215 individual sugar component. For seasonal PMF analysis, 3-5 factors were examined and 4
216 factors were determined for each season. We included the data set of anhydrosugars from
217 Verma et al. (2015) for PMF analysis.

218

219 **3 Results and discussion**

220 **3.1 Ambient concentrations of sugar compounds**

221 Temporal variations of primary sugars and sugar alcohols are shown in Figure 4. Nine sugar
222 compounds (SCs) including five primary sugars and four sugar alcohols were detected in the
223 aerosol samples collected from Chichijima Island. The concentrations of total SCs varied from
224 1.23 to 339 ng m^{-3} (average, $46.7 \pm 49.5 \text{ ng m}^{-3}$) during 2001 to 2013 (Table 1). Concentrations
225 of primary sugars and sugar alcohols were in the range of 0.28 to 176 ng m^{-3} ($23.3 \pm 25.7 \text{ ng m}^{-3}$)
226 3) and 0.37 to 231 ng m^{-3} ($23.4 \pm 30.8 \text{ ng m}^{-3}$), respectively. Average concentration of primary

227 sugars in Chichijima aerosols is several times lower than that of primary sugars (62.0 ± 54.9 ng
228 m^{-3}) reported from Cape Hedo, Okinawa, Japan (Zhu et al., 2015) while that of sugar alcohols
229 is equivalent to or little lower than that from Cape Hedo (29.5 ± 35.5 ng m^{-3}).

230 Interestingly, primary sugars (49.9%) and sugar alcohols (50.1%) were found to
231 contribute almost equal to total SCs during the entire study period. Mannitol (26.7%) and
232 arabitol (21.4%) were the main contributors to total SCs followed by glucose (16.7%), sucrose
233 (13.6%), fructose (10.2%), and trehalose (9.2%). Erythritol (1.6%), inositol (0.3%), and xylose
234 (0.3%) were also present in the aerosols at lower concentration levels. Temporal plots of
235 individual sugars clearly indicate a large variation of SCs (Figure 4). This large variation in the
236 concentrations of SCs might be involved with seasonal changes in the atmospheric circulations
237 over the western North Pacific (Kawamura et al., 2003).

238 3.1.1 Concentrations of primary sugars in total SCs

239 Glucose is the dominant sugar species among primary sugars with the concentration range of
240 0.05 to 64.3 ng m^{-3} (average, 7.79 ± 8.80 ng m^{-3}). Similarly, a wide concentration range of
241 fructose (0.03 - 115 ng m^{-3} ; 4.69 ± 8.04 ng m^{-3}) was also observed in Chichijima aerosols.
242 Thirteen year mean concentrations of glucose and fructose were observed to be lower than
243 those (27.2 ng m^{-3} and 16.4 ng m^{-3} , respectively) reported for the aerosol samples (TSP) from
244 Cape Hedo, Okinawa, Japan (Zhu et al., 2015). Glucose and fructose significantly contribute to
245 total primary sugars (33.5% and 20.17%, respectively) in Chichijima aerosols. Primary sugars
246 are abundant in the fragments of vascular plants in vegetated and forest areas (Medeiros et al.,
247 2006). Pacini et al. (2000) reported that primary sugars are synthesized in leaves during
248 photosynthesis and stored in root, stem, flower, pollen and fruit of growing plants. The nectars
249 and fruits of tropical and subtropical plants also contain glucose and fructose abundantly
250 (Backer et al., 1998). Graham et al. (2002) reported significant amounts of glucose and
251 fructose in pollen, fern spores, and insects in aerosol samples collected from the Amazon forest.
252 Chichijima Island is covered with endemic and vascular plants, which may emit glucose and

253 fructose. Moreover, different sources such as soil dust (Rogge et al., 2007; Simoneit et al.,
254 2004), lichens (Dahlman et al., 2003) and biomass burning (Medeiros et al., 2006; Nolte et al.,
255 2001) have also been reported as dominate sources for glucose and fructose.

256 Among all the SCs detected in the Chichijima aerosols, sucrose is the second most
257 abundant sugar species ($0.002\text{-}100\text{ ng m}^{-3}$; $6.43\pm 12.9\text{ ng m}^{-3}$), accounting for 27.3% of total
258 primary sugars. The average sucrose concentration observed in Chichijima is twice lower than
259 that (13.2 ng m^{-3}) from Cape Hedo, Okinawa, Japan (Zhu et al., 2015). Sucrose is synthesized
260 in plant leaves and circulated by phloem to different plant sections, which is accumulated in
261 root cells as well as developing flower buds (Bieleski, 1995; Jia et al., 2010). Sucrose is a
262 dominant component in airborne pollen grains derived from flowering plants (Bieleski, 1995;
263 Pacini, 2000). Simoneit et al. (2004a and 2004b) reported the presence of sucrose in surface
264 soil and paved road dust. Sucrose was also observed in dry plant materials during harvesting
265 period (Ma et al., 2009).

266 Thirteen year mean concentration of trehalose ranged from 0.01 to 70.2 ng m^{-3}
267 ($4.30\pm 7.28\text{ ng m}^{-3}$), whose average concentration accounts for 18.4% of total primary sugars
268 detected in Chichijima aerosols for 13 years. Microbes (bacterial cell), fungal spores, yeast,
269 algae, invertebrates, suspended soil dust, as well as plant species, contribute significantly to
270 trehalose in the atmosphere (Elbein, 1974; Graham et al., 2003; Medeiros et al., 2006; Rogge et
271 al., 2007; Simoneit et al., 2004; Wiemken, 1990). Xylose is a less abundant primary sugar,
272 accounting for 0.60% of total primary sugars observed in Chichijima aerosols. The
273 concentration range of xylose was $0.001\text{-}1.35\text{ ng m}^{-3}$ ($0.14\pm 0.18\text{ ng m}^{-3}$) during sampling
274 period of thirteen years. Biomass burning activities emit xylose to the atmosphere. Cowie and
275 Hedges (1984) reported that xylose is produced by angiosperm and gymnosperm plants,
276 phytoplankton, and groups of microorganisms. Simoneit et al. (2004a) have reported xylose in
277 soil dust from various locations in the United States and Japan. Wan and Yu (2007) also
278 observed xylose in soils and associated microbiota.

279 3.1.2 Concentrations of sugar alcohols in total SCs

280 Thirteen year mean concentrations of arabitol and mannitol were found to be $9.99 \pm 13.6 \text{ ng m}^{-3}$
281 and $12.5 \pm 17.5 \text{ ng m}^{-3}$, which contribute to 42.7% and 53.3% of total sugar alcohols,
282 respectively. The concentration ranges of arabitol ($0.04\text{--}106 \text{ ng m}^{-3}$) and mannitol ($0.10\text{--}118$
283 ng m^{-3}) are comparable to those from the Mediterranean region, Israel (arabitol, $1.85\text{--}58.3 \text{ ng}$
284 m^{-3} and mannitol, $5.57\text{--}138 \text{ ng m}^{-3}$) (Burshtein et al., 2011). Yttri et al. (2007) also reported
285 that arabitol and mannitol were main contributors of sugar alcohols in aerosol samples
286 collected from the different background sites in Norway. Sugar alcohols (arabitol, mannitol)
287 can be used as tracers for various fungal and algal species (Bauer et al., 2008a,b; Pashanska et
288 al., 2002; Zhang et al., 2010). Loos et al. (1994) discussed arabitol and mannitol as potential
289 sources of bacteria and other microbes. High levels of detritus from the spring bloom and
290 autumn decomposition have been reported as significant sources for arabitol and mannitol in
291 the vegetated region (Burshtein et al., 2011; Pashynska et al., 2002). Good positive correlations
292 of arabitol ($r = 0.63$) and mannitol ($r = 0.72$) with glucose indicate a vegetation contribution to
293 both sugar alcohols in Chichijima aerosols. Erythritol and inositol are less abundant sugar
294 species, accounting for 3.29% and 0.66% of total sugar alcohols. Their concentration ranges
295 are $0.01\text{--}8.32 \text{ ng m}^{-3}$ and $0.01\text{--}1.81 \text{ ng m}^{-3}$, respectively. Significant positive correlations of
296 both sugar species with arabitol and mannitol indicate similar sources for these SCs in
297 Chichijima aerosols (Table 2).

298 3.2 Seasonal variations of total sugar compounds

299 Seasonal concentration range, mean and median values of individual sugars during the study
300 periods of thirteen years are presented in Table 1. The concentrations of individual sugars were
301 extensively fluctuated from season to season in aerosol samples collected at Chichijima
302 (Figures 4 and 5a). The seasonally averaged concentrations of total SCs are higher in summer
303 ($71.5 \pm 70.9 \text{ ng m}^{-3}$) and autumn ($57.0 \pm 64.2 \text{ ng m}^{-3}$) than spring ($39.8 \pm 67.6 \text{ ng m}^{-3}$) and winter
304 ($18.2 \pm 34.0 \text{ ng m}^{-3}$) over Chichijima Island. Zhu et al. (2015) measured sugar components in

305 aerosol samples collected from Cape Hedo, Okinawa, Japan and reported 2 to 3 times higher
306 concentrations in summer (136 ng m^{-3}) and spring (133 ng m^{-3}) than autumn (86 ng m^{-3}) and
307 winter (40 ng m^{-3}), whose seasonal trends are similar to Chichijima. Wan and Wu (2007)
308 reported different seasonal variations with the highest concentration in autumn (375 ng m^{-3}),
309 followed by winter (292 ng m^{-3}) and spring (84 ng m^{-3}) for the continental urban aerosols
310 collected from Hong Kong. These concentrations in Hong Kong are 16 and 6 times higher than
311 those of the remote Chichijima samples for winter and autumn, respectively. Interestingly, the
312 different seasonal trends between the continental urban sites and two islands in the western
313 North Pacific may be associated with different sources and transport pathways between the
314 urban and marine environments.

315 **3.2.1 Seasonal variations of primary sugars**

316 Glucose maximized in summer ($11.0 \pm 9.02 \text{ ng m}^{-3}$) followed by autumn ($9.25 \pm 8.63 \text{ ng m}^{-3}$),
317 spring ($7.68 \pm 10.3 \text{ ng m}^{-3}$) and winter ($3.11 \pm 3.53 \text{ ng m}^{-3}$) (Table 1 and Figure 5h). Glucose is
318 the most abundant primary sugar in Chichijima aerosols. In winter/spring, Chichijima is
319 influenced by strong westerly winds that deliver the air masses from the Asian continent
320 including Mongolia, Russian Far East and North China, where vegetation is active.
321 Consequently, declined concentrations of glucose in winter mean a depressed transport of
322 glucose associated with continental bioaerosols from Asia despite long-range transport of
323 Asian dusts due to strong westerly winds. The local vegetation (vascular plants) in Chichijima
324 Island might be responsible to the enhanced concentration of glucose during growing season
325 (spring and summer) and decaying periods of plant leaves (autumn). Seasonal PMF analysis
326 also supports dominant sources of vegetation for glucose among four factors, which
327 contributed >75% for mixed factor in summer (Figure 6c), >80% for fungal and vegetation
328 factor in autumn (Figure 6d), and >75% for vegetation factor in spring (Figure 6b).

329 Fructose shows the highest concentrations in summer ($7.25 \pm 7.63 \text{ ng m}^{-3}$) followed by
330 spring ($4.51 \pm 9.21 \text{ ng m}^{-3}$), autumn ($3.70 \pm 2.68 \text{ ng m}^{-3}$) and winter ($3.36 \pm 10.2 \text{ ng m}^{-3}$). As

331 shown in Table 2, a significant correlation ($r=0.57$) was obtained between glucose and fructose.
332 Burshtein et al. (2011) reported similar correlations for both sugar species, suggesting a
333 **contribution** of glucose and fructose from the local vegetation in summer (Baker et al., 1998;
334 Pacini, 2000). Monthly mean concentrations of fructose show two prominent peaks in
335 February-March and June-July, the latter peak may be due to the local vegetation in Chichijima
336 (Figure 5g). The fructose peak in February-March may be influenced by air borne pollen grains
337 in the spring bloom of flowering plants. High concentration of fructose was observed in spring
338 followed by summer, indicating an input of this sugar compound from pollen grains (Fu et al.,
339 2012). The positive correlation of fructose with sucrose (pollen tracer) supports the similar
340 sources. Seasonal PMF analysis further supports the identical source for fructose and sucrose;
341 that is, among four factors, fructose contributes $>70\%$ and $>60\%$ for pollen factor in spring
342 (Figure 6b) and winter (Figure 6a), respectively.

343 Seasonal mean concentrations of trehalose showed a maximum in summer (7.06 ± 8.49 ng
344 m^{-3}) followed by autumn (6.09 ± 8.81 ng m^{-3}), spring (2.93 ± 6.08 ng m^{-3}) and winter (1.03 ± 1.26
345 ng m^{-3}) (Table 1). Monthly mean concentrations of SCs for 13 years show that concentrations
346 of trehalose are higher during June to October (Figure 5j). Ma et al. (2009) reported higher
347 concentrations of trehalose at the urban site of Guangzhou, China during summer and autumn.
348 Similarly, Wan and Wu (2007) reported a similar autumn maximum in Hong Kong. On the
349 other hand, different seasonal trends of trehalose were reported for the aerosol samples (TSP)
350 collected in USA (Medeiros et al., 2006), China (Wang et al., 2011), Australia (Hackl et al.,
351 2000), and Gosan, Jeju Island in the western North Pacific Rim (Fu et al., 2012). In the above
352 studies, highest concentrations of trehalose were reported in early spring due to the re-
353 suspension of soil particles during agricultural practice. Hackl et al. (2000) also obtained
354 abundant trehalose in spring, and proposed that trehalose can be used as a tracer of soil dust
355 emission to the atmosphere. However, we did not detect a spring peak of trehalose in
356 Chichijima aerosols, suggesting that soil dust contribution of trehalose over Chichijima is

357 insignificant via long-range atmospheric transport. Seasonal PMF analysis for autumn showed
358 that more than 85% of trehalose was contributed by microbial factor among four factors
359 (Figure 6d). An indirect contribution of trehalose from soil dust will be discussed later.

360 The seasonal mean concentrations of sucrose are almost equal during spring (8.80 ± 18.0
361 ng m^{-3}), summer ($7.31 \pm 11.5 \text{ ng m}^{-3}$) and winter ($6.60 \pm 13.1 \text{ ng m}^{-3}$), except for autumn
362 ($2.76 \pm 4.35 \text{ ng m}^{-3}$) (Table 1). The similar seasonal distributions suggest multiple sources of
363 sucrose in Chichijima aerosols. Monthly mean concentrations of sucrose show two peaks
364 during February-March and June-July (Figure 5i). The March peak of sucrose was reported in
365 the forest area of Sapporo, Japan to be 3 to 7 times more abundant than other months due to
366 springtime pollen emissions (Miyazaki et al., 2012). Fu et al. (2012) analyzed pollen samples
367 from different plant species (white birch, Chinese willow, Peking willow) for SCs and found
368 the highest concentrations of sucrose followed by fructose and glucose in pollen. The pollen
369 emissions from developing buds of plants may be the reason for the increased concentration of
370 sucrose and fructose in February and March over Chichijima Island in the western North
371 Pacific. Seasonal PMF analysis shows that sucrose contributed 100% in spring and >80% in
372 winter, suggesting a significant pollen contribution for sucrose in those seasons (Figures 6b
373 and 6a).

374 However, the possibilities of pollen transport from East Asia to Chichijima cannot be
375 excluded because pollens can travel long distances with springtime high-speed winds by
376 westerlies (Rousseau et al., 2006). The pollen grains emitted from flowering boreal forest in
377 China, Mongolia, Siberia and Russian Far East, could significantly be delivered to the western
378 North Pacific during spring, which may result in the contribution of sucrose and fructose to
379 Chichijima aerosols. Recent studies have discussed a long-range transport of airborne pollen
380 from North America to Greenland in spring (Rousseau et al., 2008). Lorenzo et al. (2006)
381 reported the long-range transport of airborne allergenic pollen to central Italy. Makra et al.
382 (2010) reported a long-range transport of airborne pollen in three European cities by applying

383 three-dimensional clustering of backward trajectories. Several studies also discussed a long-
384 range transport of pollen to the remote arctic region (Andrews et al., 1980; Bourgeois et al.,
385 2001; Campbell et al., 1999; Hicks et al., 2001; Hjelmroos and Franzen, 1994; Rousseau et al.,
386 2004).

387 These observations may support that westerly winds have delivered pollen grains from
388 the Asian continent including Mongolia, Siberia, and Russian Far East to Chichijima Island in
389 spring. Using a box model and typical settling velocity of pollens (3 cm/sec) with the grain size
390 of 30 μm in diameter (Sosnoskie et al., 2009), we estimated lifetime of pollen grains to be 9.3
391 hours in the atmospheric marine boundary layer (height of 1 km above the ocean surface). The
392 settling velocity of the pollens is ca. 20 times larger than that of typical marine aerosols (Slinn
393 and Slinn, 1980). Because pollen grain sizes range from 10 μm to 100 μm in diameter, the
394 lifetime of pollens may have a large uncertainty. If pollens could be largely transported in the
395 free troposphere (e.g., 5 km high) to the North Pacific from the Asian continent, then lifetime
396 of typical pollen grains would increase upto 2 days. These calculations for the lifetime of
397 pollen grains further support their long-range atmospheric transport from the Asian continent
398 over the western North Pacific. Based on backward air mass trajectories (Figure 3), we can
399 roughly estimate the transport time from East Asia to Chichijima site to be 2-4 days in winter
400 and spring. It is also of interest to note that pollens can rapture under condition of high relative
401 humidity (RH) (Hader et al., 2014; Miguel et al., 2006; Wright et al., 2014), which leads to
402 smaller particles with longer residence time in the atmosphere.

403 In addition, tilling process after wheat crop harvesting in farmland causes an enhanced
404 exposure of wheat root associated with soil particles into the atmosphere. China, India, and
405 USA are three largest countries for wheat production in the world. In China and India there are
406 two seasons (spring and winter) for wheat crops; winter wheat is harvested from mid-May to
407 mid-July. During those periods (early summer), Chichijima Island is highly influenced by trade
408 winds (Figure 3). However, **air mass** trajectories clearly show the occasional **atmospheric**

409 transport from Southeast Asia to Chichijima **in summer** (Pavuluri et al., 2010). PMF results of
410 sucrose for summer (Figure 6c) and autumn (Figure 6d) account for >85% and >90%,
411 respectively, for soil dust factor among four source factors, suggesting an additional source of
412 soil dust for sucrose in Chichijima **aerosols** (Simoneit et al., 2004). The elevated sucrose
413 concentrations in June and July (summer; non-flowering season) suggest the **long-range**
414 transport of sucrose associated with soil particles under the influence of occasional air mass
415 transport from Southeast Asia **in summer** (Figure 3).

416 Xylose was found as the least abundant sugar compound in the aerosol samples. The
417 maximum concentration of xylose (1.35 ng m^{-3}) was found in summer whereas minimum
418 (0.001 ng m^{-3}) in spring (Table 1). Summer mean concentration ($0.18 \pm 0.26 \text{ ng m}^{-3}$) was highest
419 (Table 1). The PMF analyses showed that xylose contributed >75% for BB factor in winter
420 (Figure 6a) and >70% in autumn (Figure 6d) for microbial factor. These results suggest
421 different sources and seasons for xylose; i.e., biomass burning in winter (Sullivan et al., 2011)
422 and groups of microorganisms in summer (Cowie and Hedges, 1984).

423 **3.2.2 Seasonal variations of sugar alcohols**

424 The seasonal mean concentrations of arabitol and mannitol are higher in summer/autumn than
425 spring/winter (Table 1). The concentrations of arabitol are equally distributed between summer
426 ($15.1 \pm 12.9 \text{ ng m}^{-3}$) and autumn ($15.8 \pm 18.3 \text{ ng m}^{-3}$) with lower levels in spring ($7.13 \pm 9.50 \text{ ng}$
427 m^{-3}) and winter ($1.73 \pm 2.60 \text{ ng m}^{-3}$). Mannitol maximized in summer ($21.7 \pm 19.7 \text{ ng m}^{-3}$)
428 followed by autumn ($18.2 \pm 19.9 \text{ ng m}^{-3}$), spring ($7.95 \pm 13.8 \text{ ng m}^{-3}$) and winter ($1.89 \pm 2.81 \text{ ng}$
429 m^{-3}). Arabitol and mannitol strongly co-varied throughout the study period. As depicted in
430 thirteen year monthly mean concentrations of total SCs (Figure 5b,c), we found elevated
431 concentrations of sugar alcohols from May to October. Similar seasonal trends were reported
432 for the aerosol samples collected from Gosan, Jeju Island in the western North Pacific Rim (Fu
433 et al. 2012) and urban aerosol samples from Ghent, Belgium (Pashynska et al., 2002). In above
434 studies, higher relative abundances of arabitol and mannitol in total sugar alcohols were

435 reported during late summer to autumn. The higher concentration of arabitol in autumn was
436 also reported for aerosol samples from the Mediterranean region in Israel (Burshtein et al.,
437 2011). Erythritol and inositol showed the similar seasonal trend, but their concentrations are
438 lower than the former two sugar species.

439 Sugar alcohols are emitted to the atmosphere from a variety of bacteria, few green algal
440 lichens and fungi (Dahlman et al., 2003; Filippo et al., 2013). Arabitol and mannitol are
441 abundant in fungal spores (Lewis and Smith, 1967; Yttri et al., 2007). Arabitol ($r = 0.73$) and
442 mannitol ($r = 0.80$) showed a strong co-variance with trehalose, suggesting identical sources of
443 sugar species in Chichijima. The PMF analysis showed that fungal factor and mixed factor
444 (fungal, vegetation, and microbial) accounted for 25% and 54.2% of total SCs observed in
445 summer, respectively (Figure 6c). In autumn, fungal and vegetation factor contributed 71% of
446 total SCs detected in Chichijima aerosols (Figure 6c). In winter (Figure 6a) and spring (Figure
447 6b) fungal and vegetation factor and mixed factor account for 31.2% and 37.2% of total SCs,
448 respectively. This is reasonable because fungal and microbial activities are lower during
449 winter/spring as compared to summer/autumn. The meteorological factors such as RH and
450 temperature significantly affect fungal and bacterial activity (Kim and Xiao, 2005; Malik and
451 Singh, 2004). Higher RH and temperature are crucial in increasing fungal and bacterial growth
452 (Sharma and Razak, 2003). Their maximum growth was observed under the condition of 92-
453 100% RH (Ibrahim et al., 2011). Higher concentrations of arabitol and mannitol in summer and
454 autumn may be caused by the increased fungal and bacterial activities in Chichijima Island.

455 Several studies have described the occurrence of fungi in marine environment (Jones,
456 1976; Kohlmeyer and Kohlmeyer, 1991; Moss, 1986). The fungal species eject spores from
457 hard materials like coral and sand grains. Some fungi also eject spores from woods associated
458 with sand in summer/autumn when higher ambient temperature and RH are available (Jones
459 and Mitchell, 1996). Marine fungal growths are observed on several mediums of substrates
460 such as wood, sediments, muds, soil, sand, algae, corals, decaying leaves of mangroves and

461 living animals in marine environment (Bremer, 1995; Kohlmeyer and Kohlmeyer, 1979;
462 Nagakiri et al., 1996). Although the above-mentioned studies have claimed the occurrence and
463 growth of marine fungi on several mediums of substrates, the knowledge of the role of marine
464 fungi in sediments and decaying dead animals are still insufficient due to a lack of appropriate
465 data set. It is still unclear if these fungi are active in sediments (Hyde et al., 1998). Therefore,
466 due to the inadequate data set, we doubt the marine contribution of sugar alcohols (arabitol,
467 mannitol) to Chichijima aerosols.

468 Thirteen year monthly mean concentrations of SCs clearly show slightly decreased
469 concentrations of arabitol, mannitol, and erythritol in July and August; a similar trend was
470 observed for trehalose (Figure 5b,c,e,j). These sugar compounds are derived from the microbial
471 activities in source regions. The thirteen year precipitation record over Chichijima Island
472 shows that precipitations were lowered in July and August (Figure 2). The lower precipitation
473 amount decreases the RH (Figure 2) and thus depresses the fungal and microbial activities. The
474 lower precipitation also suppresses the moisture contents in the surface soil of Chichijima,
475 which should result in a significant decline of local fungal and other microbial activities on the
476 ground of Chichijima Island. Decreased precipitation might be a possible reason for the lower
477 concentrations of arabitol, mannitol, and trehalose in July and August.

478 3.3 Annual variations and decadal comparisons of SCs

479 The annual variations in the concentrations of primary sugars and sugar alcohols are shown in
480 Figure 7a. The annual mean concentrations of total SCs varied randomly during 2001 to 2013.
481 As shown in Figure 7 (i, j, f and d), concentrations of sucrose, trehalose, xylose, and inositol
482 increase from 2001 to 2013 in Chichijima aerosols. Similarly, arabitol (Figure 7b), glucose
483 (Figure 7h), and fructose (Figure 7g) show clear increasing trends from 2006 to 2013 whereas
484 mannitol (Figure 7c) and erythritol (Figure 7e) show a random trend. Here, we compare the
485 data set of SCs for the periods of 1990-1993 (Period, P-I) with the current observations from
486 2001 to 2003 (P-II) and 2010 to 2013 (P-III) (Table 3) (Figures 8a, b, c).

487 The comparison for three periods indicates that concentrations of anhydrosugars are
488 highest in winter followed by autumn. Their concentrations significantly increased from P-I to
489 P-II/P-III (Figure 8a). The detailed discussions on anhydrosugars were reported in Verma et al.
490 (2015). Here, we refer the data set of anhydrosugars for the decadal comparison with SCs.
491 Interestingly, biomass-burning tracers (BB-tracers; levoglucosan, mannosan and galactosan)
492 showed a significant difference in the decadal trends among three periods (i.e., P-I, P-II, and P-
493 III) during winter and autumn. In winter, BB-tracers showed an increasing trend from P-I to P-
494 III (Figure 8a). Biomass burning is common in winter for house heating (Simoneit et al.,
495 2004a), thus it is obvious that the lower ambient temperature is the more biomass burning
496 activities are. Westerly winds abundantly transport biomass-burning products over Chichijima
497 Island in the western North Pacific with air masses derived from East Asia, Siberia, Mongolia,
498 and Russian Far East during winter (Bendle et al., 2007; Simoneit and Elias, 2010; Verma et al.,
499 2015). In contrast, BB-tracers in autumn show an opposite trend (i.e., higher concentrations in
500 P-I followed by P-II and P-III) compared to those in winter (Figure 8a).

501 The difference in the concentrations of anhydrosugars in winter and autumn during P-I is
502 insignificant while the concentrations are 3 and 5 times higher in winter than autumn for P-II
503 and P-III, respectively. This seasonal shifting in the concentrations of anhydrosugars may be
504 attributable to the changes in the strength of both westerly and trade wind systems from mid
505 autumn to early winter among three periods (Chen et al., 2013). In contrast, the concentrations
506 of primary sugars were 2 to 7 times higher during P-II and P-III than P-I period in summer and
507 autumn (Figure 8b). PMF analysis showed that local emissions from vegetation are important
508 contributor for primary sugars (glucose, fructose and sucrose). Therefore, a drastic increase in
509 the concentrations of primary sugars in summer/autumn for P-II and P-III than P-I may be
510 caused by an increased emission of primary sugars by local vegetation under the influence of
511 meteorological conditions in the western North Pacific. However, a possible soil dust

512 contribution of primary sugar (sucrose) associated with an occasional air mass transport from
513 Southeast Asia cannot be excluded.

514 Similar to primary sugars, a drastic increase in the concentrations of sugar alcohols was
515 observed for P-II and P-III compared to P-I period (Figure 8c). The concentrations of sugar
516 alcohols in P-II and P-III are 6 to 19 times higher than those of P-I in summer/autumn (Table
517 3; Figure 8c). Arabitol and mannitol are key sugar alcohols and are reported as fungal and
518 microbial tracers, which contribute significantly to total SCs (Bauer et al., 2008b; Lewis and
519 Smith, 1967; Zhang et al., 2010). Microbes such as fungi and bacteria are significantly
520 increasing in the Asian and European countries (Yamaguchi et al., 2012). They are largely
521 transported towards downwind regions in the Pacific Ocean from the Asian continent in
522 winter/spring under the influence of strong westerlies (Griffin et al., 2001, 2003, 2007; Hua et
523 al., 2007; Uno et al., 2009) and settled down by wet and dry deposition in the western North
524 Pacific according to air mass trajectories (Figure 3).

525 Hirst et al. (1967) studied the lifetime of fungal spores and long-range transport of spores.
526 They reported that the fungal spores are in between 2 and 200 μm in diameter (mostly 5 to 30
527 μm), whose settling velocity is between 0.05 and 3.0 cm/sec with a mode of less than 1 cm/sec,
528 and that different fungal spores vary significantly in shape and ornamentation. This study
529 suggested the possibilities of long-range transport of fungal spores. Jeon et al. (2011) and
530 Yamaguchi et al. (2012) have analyzed aerosol samples collected during the Asian dust event
531 over the Sea of Japan, and identified similar groups of microbes (bacterial cells) transported
532 from the source regions of Asian dust. Consequently, bacteria and fungi associated with
533 bioaerosols grow extensively during summer/autumn, when the climate conditions are
534 favorable (i.e., higher RH and temperature) for their metabolic activities (Morris et al., 2004).
535 Accordingly, an increased transport of bioaerosols for the last decade may have caused a
536 drastic increase in the concentrations of sugar alcohols during P-II and P-III compared to P-I
537 period over the western North Pacific.

538 3.4 Source apportionment of SCs

539 To investigate the source apportionment of sugar components, the data sets of Chichijima
540 aerosols were subjected to positive matrix factorization (PMF) analysis. Based on the PMF
541 analysis, total five factors were determined to be significant to classify the sources of sugar
542 compounds (SCs). Five factors successfully explored the source profile for the individual sugar
543 component. Factor profiles resolved by PMF analysis are shown in Figures 6 and 9-11, where
544 percentages of each component summed for factors 1 to 5 are calculated to be 100%.

545 Vegetation factor (Figure 9) was dominated by xylose (75%), glucose (48%), and
546 fructose (36%). Xylose is significantly produced by gymnosperm and angiosperm (Cowie and
547 Hegdes, 1984; Sjoström, 1981). Fructose and glucose are highly water-soluble sugar species,
548 and present in the bark and leaves of plants (Fu et al., 2012). Glucose is the second most
549 abundant sugar that contributed to this factor. Cowie and Hegdes (1984) reported higher
550 concentration of glucose in vascular plants and phytoplankton in the marine environment. The
551 SCs emitted by the vegetation during growing season significantly contribute to vegetation
552 factor. In Chichijima aerosols, glucose and fructose are significant contributors in spring
553 (Figure 6b), summer (Figure 6c) and autumn (Figure 6d), Therefore, the respective factors in
554 Figure 6 are termed as a vegetation source for both sugar species. This is reasonable because
555 plants started growing in spring and summer seasons. In autumn, leaf senescence and decay
556 result in the emission of glucose and fructose to the atmosphere.

557 Fungal and microbial factor (Figure 9) was characterized by trehalose (88%), mannitol
558 (64%), and arabitol (54%). These sugars that contribute to fungal and microbial factor are
559 associated with fungal spores, bacteria and yeast (Bauer et al., 2008a; Medeiros et al., 2006;
560 Wiemken, 1990). The three sugars are good tracers of fungal spores and microbes (Loos et al.,
561 1994; Rogge et al., 2007). Arabitol and mannitol are produced by a large variety of fungal
562 species (Ion et al., 2005; Medeiros et al., 2006), and considered as a suitable tracer for fungal
563 and bacterial metabolic activities (Bauer et al., 2008b; Elbein et al., 1974; Rogge et al., 2007).

564 Arabitol is strongly correlated with mannitol ($r=0.88$), suggesting similar sources for both
565 species (Table 2) (Elbert et al., 2007). Fungi, bacteria, and other microbes in soils are the main
566 sources for trehalose (Graham et al., 2003; Rogge et al., 2007; Simoneit et al., 2004). An
567 excellent correlation of trehalose with arabitol and mannitol suggested similar sources in the
568 marine environment (Table 2) (Lewis and Smith, 1967).

569 Sugar alcohols have been proposed as tracers for microbes and fungal spores (Bauer et
570 al., 2008b; Ion et al., 2005; Medeiros et al., 2006; Rogge et al., 2007). The fungal and
571 microbial activities are considered higher during summer and autumn due to higher
572 temperature and RH. The above discussions for the sources of arabitol, mannitol and trehalose
573 are well supported by the seasonal PMF analysis. Arabitol and mannitol are well contributed in
574 summer (Figure 6c, fungal factor and mixed factor) and autumn (Figure 6d, fungal and
575 vegetation factor). Correspondingly, trehalose also contributed in summer (Figure 6c, mixed
576 factor) and autumn (Figure 6d, microbial factor). Therefore, significant contributions of
577 arabitol, mannitol and trehalose are observed during the respective seasons in Chichijima
578 aerosols.

579 Mixed factor (Figure 9) is associated with erythritol (94%), arabitol (44%), mannitol
580 (34%), inositol (32%), glucose (24%), and fructose (31%). Due to the highly miscellany
581 characteristics of fungi, other microbes, and plant debris, it is quite difficult to specify the
582 particular source for individual sugar species (Percival, 1970). Arabitol and mannitol are also
583 attributed to the vegetation (photosynthesized by mature leaves) (Burshtein et al., 2011;
584 Pashynska et al., 2002). The contributions of arabitol and mannitol in winter (Figure 6a, mixed
585 factor) and spring (Figure 6b, mixed factor) indicate other sources than fungal spores in the
586 Chichijima aerosols. Because the fungal and microbial growth is less important in winter and
587 spring compared to summer/autumn. Therefore, this factor (Figure 9) should be associated with
588 mixed sources from microbial and vegetational activities. Contemplating of mixed sources is

589 very likely because sugar species that are highly apportioned to vegetation factor contribute, to
590 some extent, to the same sources that are responsible to fungal and microbial factor (Figure 9).

591 **Biomass burning (BB)** factor (Figure 9) is loaded significantly with levoglucosan (94%),
592 mannosan (91%), and galactosan (90%), and moderately with xylose (20%). These species are
593 associated with biomass burning (Fraser and Lakshmanan, 2000; Graham et al., 2002;
594 Simoneit, 2002). Nolte et al. (2001) and Medeiros et al. (2006) also reported that biomass
595 burning-influenced aerosols are enriched with levoglucosan, mannosan, and galactosan.
596 Kawamura et al. (2003) and Mochida et al. (2010) reported that biomass-burning products are
597 abundantly transported to Chichijima under the influence of westerly winds during
598 winter/spring (Figure 3). These results are well supported by the fact that BB factor is
599 associated with SCs, which are derived from biomass burning in East Asia. The seasonal PMF
600 analysis (Figure 6) also supports the above explanation; BB products are contributed highest in
601 winter/spring (18.8% and 6.3%) under the influence of westerly winds, followed by
602 summer/autumn (3.8% and 4.1%) for total sugars in the aerosol samples collected from
603 Chichijima Island.

604 Pollen factor (Figure 9) is characterized by high loading of sucrose (91%), fructose
605 (35%), and inositol (33%); these sugar species are associated with airborne pollen sources.
606 Previous studies have also reported that sucrose is an excellent tracer for airborne pollen grains
607 of flowering plants (Pacini, 2000; Graham et al., 2003; Wang et al., 2008; Medeiros et al.,
608 2006). Fructose is well correlated with inositol ($r=0.57$), indicating a similar origin for both
609 sugar species (Table 2). Two prominent peaks of sucrose, fructose, and inositol, which
610 appeared in late winter to early spring or summer, also indicate a similar source of those sugars.
611 Contributions of sucrose (pollen factor) in winter (39.5%; Figure 6a) and spring (36%; Figure
612 6b) are supported the sources of airborne pollen for sucrose in Chichijima Island. The sucrose
613 contribution (soil dust factor) in non-flowering seasons, i.e., summer (17.1%; Figure 6c) and
614 autumn (8.4%; Figure 6d), indicates different sources for sucrose in Chichijima aerosols.

615 According to the seasonal PMF analysis (Figure 6), we termed additional sources of sucrose in
616 Chichijima Island as soil dust factor as well as pollen factor.

617 As seen in Figure 6a, b, c and d, PMF analysis for seasonal source identification
618 indicated variable contributions of individual SCs in different seasons according to their
619 seasonal source origin. In winter (Figure 6a), airborne pollen (39.5%) contributed highest
620 followed by vegetation, microbial, fungal (mixed) (31.2%), biomass burning (18.8%), and
621 microbial (10.5%) sources. However, in spring, vegetation, microbial, and fungal (mixed)
622 sources (37.2%) contributed almost equal to the airborne pollen (36.0%) followed by
623 vegetation (21.2%) and biomass burning (6.3%) sources. The vegetation, microbial, fungal
624 (mixed) (54.2%), fungal (25.2%), soil particles (17.1%), and biomass burning (3.8%) sources
625 are characterized to maximize in summer. Similar to summer, vegetation and fungal (mixed)
626 sources (71.2%) are also leading to the contribution of total SCs followed by microbial
627 (16.4%), soil particles (8.4%), and biomass burning (4.1%) in total SCs observed in autumn for
628 Chichijima aerosols.

629 Overall, average contributions of each factor to measured SCs as resolved by the PMF
630 analyses are shown in Figure 10. Fungal and microbial factor accounts for 41% of total SCs
631 measured. The emission from microbes including fungal spores was found as a dominant
632 contributor to total SCs. Mixed factor (27%) indicates a common involvement of fungal,
633 microbial, and vegetation sources. Figure 11 shows annual trends in % contributions of five
634 source factors to SCs in Chichijima aerosols. Fungal and microbial factor, and mixed factor
635 contributed higher than other sources for SCs during 2001 to 2013. However, no clear trends in
636 annual % contributions were observed for both source factors during thirteen-year study period.
637 The sugar species assigned as pollen tracers were found to contribute 18% in pollen factor.
638 Vegetation accounts for 11% of total SCs, indicating less emission from vegetation as
639 compared to fungi and microbes. As indicated by BB factor, biomass-burning source
640 contributes only 3% of total SCs.

641 Interestingly, we found an increasing trend in % contribution of vegetation, pollen, and
642 BB factors to SCs in 2006 to 2013 (see Figure 11). Sugar components, which are contributed
643 from pollen (sucrose, fructose, and inositol), vegetation (glucose and fructose), and biomass
644 burning sources (levoglucosan, galactosan, and mannosan), also show a similar increasing
645 trend for the period of 2006 to 2013 (Figure 7). The increased annual trends of BB and pollen
646 factors might be due to an enhanced long-range transport of airborne pollens and biomass
647 burning products from the Asian continent to the western North Pacific under the influence of
648 strong westerly winds.

649

650 **4 Summary and conclusions**

651 We reported thirteen years of temporal, seasonal and decadal trends of sugar compounds (SCs)
652 measured in the aerosol samples collected at Chichijima Island in the western North Pacific, an
653 outflow region of Asian aerosols. The high abundances of total SCs and primary sugars are
654 found during summer, whereas sugar alcohols are almost equally distributed during summer
655 and autumn. The seasonal distributions of arabitol, mannitol, and trehalose are strongly
656 influenced by long-range atmospheric transport of bioaerosols associated with microbes such
657 as fungi and bacteria and their metabolic activities under the influences of westerly winds and
658 favorable meteorological conditions, including high RH and temperature in summer/autumn.
659 Seasonal variation of sucrose is controlled by both locally emitted and long-range transported
660 pollen from East Asia to the western North Pacific during spring bloom periods. On the other
661 hand, the increased concentrations of sucrose and fructose during summer may be caused by
662 the local activity of vegetation and possibly by long-range transport of plant root-associated
663 soil dust particles from East and Southeast Asia. PMF analysis indicated specific sources for
664 individual SCs during different seasons. The results clearly separated biogenic emissions into
665 two parts as vegetation and microbes, including fungal species. The emissions from vegetation,

666 pollen, and microbial activities contributed about 97% of total measured SCs, with the
667 remaining fraction being derived from biomass burning activities.

668 The concentration and seasonal variations of SCs at Chichijima Island are well
669 regulated by atmospheric circulations, i.e., the westerly winds passing from the Asian continent
670 during winter/spring and trade winds originated from the central Pacific Ocean during
671 summer/autumn dominate over Chichijima Island in the western North Pacific. The
672 meteorological parameters also significantly affect the concentrations and seasonal variations
673 of SCs over Chichijima. Based on a decadal observation at Chichijima, we conclude that
674 drastic increases in the concentrations of sugar alcohols and primary sugars during 2001-2003
675 and 2010-2013 can be caused by an enhanced atmospheric transport of bioaerosols from East
676 Asia to the western North Pacific.

677 Sugar components (SCs) are important compositions of organic aerosols worldwide and
678 it is recognized as a significant factor affecting air quality and possibly climate. The outcomes
679 of the thirteen-year study of SCs at Chichijima Island have an implication for global radiative
680 forcing by scattering or absorbing light and also the activity of cloud condensation nuclei
681 (CCN) in the western North Pacific that have a high sensitivity to global climate change due to
682 an outflow region of the Asian dust and bioaerosols. The NASA Global Climate Change
683 (<http://climate.nasa.gov/vital-signs/global-temperature/>) has reported a continuous increase in
684 the global land/ocean temperature. The increasing annual trends in % contribution of
685 vegetation factor to SCs suggested a significantly increased activity of local vegetation in
686 Chichijima Island from 2006 to 2013, which could be involved with a recent global warming
687 especially in the western North Pacific region.

688

689 **Acknowledgements**

690 We acknowledge the financial support by Japan Society for the Promotion of Science (JSPS)
691 through grant-in-aid Nos. 19204055 and 24221001. The authors gratefully appreciate the

692 NOAA Air Resources Laboratory (ARL) for the provision of the HYSPLIT transport and
693 dispersion model (<http://www.ready.noaa.gov>). The data for this paper are available upon
694 request to the corresponding author (Kimitaka Kawamura, kkawamura@isc.chubu.ac.jp).
695

696 **References**

- 697 Andrews, J. T., W. N. Mode, and P. T. Davis (1980), Holocene climate based on pollen transfer-
698 functions, *Eastern Canadian Arctic, Arct. Alp. Res.*, 12, 41-64.
- 699 Baker, H. G., I. Baker, S. A. Hodges (1998), Sugar composition of nectars and fruits consumed
700 by birds and bats in the tropics and subtropics, *Biotropica*. 30, 559-586.
- 701 Bauer, H., E. Schueller, G. Weinke, A. Berger, R. Hitzenberger, I. L. Marr, and H. Puxbaum
702 (2008a), Significant contributions of fungal spores to the organic carbon and to the
703 aerosol mass balance of the urban atmospheric aerosol, *Atmos. Environ.*, 42(22), 5542-
704 5549.
- 705 Bauer, H., M. Claeys, R. Vermeylen, E. Schueller, G. Weinke, A. Berger, and H. Puxbaum
706 (2008b), Arabitol and mannitol as tracers for the quantification of airborne fungal spores,
707 *Atmos. Environ.*, 42(3), 588-593.
- 708 Bendle, J., K. Kawamura, K. Yamazaki, and T. Niwai (2007), Latitudinal distribution of
709 terrestrial lipid biomarkers and n-alkane compound-specific stable carbon isotope ratios
710 in the atmosphere over the western Pacific and Southern Ocean, *Geochim. Cosmochim.*
711 *Acta*, 71(24), 5934-5955.
- 712 Bielecki, R. L. (1995), Onset of phloem export from senescent petals of Daylily, *Plant Physiol.*,
713 109(2), 557-565.
- 714 Bourgeois, J. C., K. Gajewski, and R. M. Koerner (2001), Spatial patterns of pollen deposition
715 in arctic snow, *J. Geophys. Res.*, 106, 5255-5265.
- 716 Bremer, G. B. (1995), Lower marine fungi (Labyrinthulomycetes) and the decay of mangrove
717 leaf-litter, *Hydrobiologia*, 304(3), 243-245.
- 718 Brown, J. K. M., and M. S. Hovmoller (2002), Epidemiology—Aerial dispersal of pathogens on
719 the global and continental scales and its impact on plant disease, *Science*, 297(5581),
720 537-541.
- 721 Burshtein, N., N. Lang-Yona, and Y. Rudich (2011), Ergosterol, arabitol and mannitol as tracers
722 for biogenic aerosols in the eastern Mediterranean, *Atmos. Chem. Phys.*, 11(2), 829-839.
- 723 Bzdusek P.A., E. R. Christensen, C. M. Lee, U. Pakadeesusuk, D. C. Freedman, (2006). PCB
724 congeners and dechlorination in sediments of Lake Hartwell, South Carolina, determined
725 from cores collected in 1987 and 1988. *Environ. Sci. Technol.*, 40, 109-119.
- 726 Campbell, I. D., K. McDonald, M. Flannigan, and J. Kringayark (1999), Long-distance transport
727 of pollen into the Arctic, *Nature*, 399, 29- 30.
- 728 Carvalho, A., C. Pio, and C. Santos (2003), Water-soluble hydroxylated organic compounds in
729 German and Finnish aerosols, *Atmos. Environ.*, 37(13), 1775-1783.
- 730 Chen, J., K. Kawamura, C. Q. Liu, and P. Q. Fu (2013), Long-term observations of saccharides
731 in remote marine aerosols from the western North Pacific: A comparison between 1990-
732 1993 and 2006-2009 periods, *Atmos. Environ.*, 67, 448-458.
- 733 Cowie, G. L., and J. I. Hedges (1984), Carbohydrate sources in a coastal marine-environment,
734 *Geochim. Cosmochim. Acta*, 48(10), 2075-2087.
- 735 Dahlman, L., J. Persson, T. Nasholm, and K. Palmqvist (2003), Carbon and nitrogen distribution
736 in the green algal lichens *Hypogymnia physodes* and *Platismatia glauca* in relation to
737 nutrient supply, *Planta*, 217(1), 41-48.
- 738 Duce, R. A., C. K. Unni, B. J. Ray, J.M. Prospero, J.T. Merrill (1980), Long-range atmospheric

- 739 transport of soil dust from Asia to the tropical North Pacific: temporal variability,
740 Science, 209, 1522–1524.
- 741 Elbein, A. D. (1974), The metabolism of α , α -trehalose, *Adv. Carboh. Chem. and Biochem.*, 30,
742 227–256.
- 743 Elbert, W., P. E. Taylor, M. O. Andreae, and U. Pöschl (2007), Contribution of fungi to primary
744 biogenic aerosols in the atmosphere: wet and dry discharged spores, carbohydrates, and
745 inorganic ions, *Atmos. Chem. Phys.*, 7(17), 4569-4588.
- 746 Elliot S., Blake D.R., Duce R.A., Lai C.A., McCreary I., McNair L.A., Rowland F.S., Russell
747 A.G., Streit G.E., and Turco R.P. Motorization of China implies changes in pacific air
748 chemistry and primary production. *Geophys. Res. Letts.* 24, 2671 -2674, 1997.
- 749 Filippo, P. D., D. Pomata, C. Riccardi, F. Buiarelli, and C. Perrino (2013), Fungal contribution
750 to size-segregated aerosol measured through biomarkers, *Atmos. Environ.*, 64,132-140.
- 751 Fraser, M. P., and K. Lakshmanan (2000), Using levoglucosan as a molecular marker for the
752 long-range transport of biomass combustion aerosols, *Environ. Sci. Technol.*, 34(21),
753 4560-4564.
- 754 Fu, P. Q., K. Kawamura, M. Kobayashi, and B. R. T. Simoneit (2012), Seasonal variations of
755 sugars in atmospheric particulate matter from Gosan, Jeju Island: Significant
756 contributions of airborne pollen and Asian dust in spring, *Atmos. Environ.*, 55, 234-239.
- 757 Fuzzi, S., et al. (2007), Overview of the inorganic and organic composition of size-segregated
758 aerosol in Rondonia, Brazil, from the biomass-burning period to the onset of the wet
759 season, *J. Geophys. Res.-Atmos.*, 112(D1), doi:10.1029/2005jd006741.
- 760 Graham, B., O. L. Mayol-Bracero, P. Guyon, G. C. Roberts, S. Decesari, M. C. Facchini, P.
761 Artaxo, W. Maenhaut, P. Koll, and M. O. Andreae (2002), Water-soluble organic
762 compounds in biomass burning aerosols over Amazonia - 1. Characterization by NMR
763 and GC-MS, *J. Geophys. Res.-Atmos.*, 107(D20), doi:10.1029/2001jd000336.
- 764 Graham, B., P. Guyon, P. E. Taylor, P. Artaxo, W. Maenhaut, M. M. Glovsky, R. C. Flagan, and
765 M. O. Andreae (2003), Organic compounds present in the natural Amazonian aerosol:
766 Characterization by gas chromatography-mass spectrometry, *J. Geophys. Res.-Atmos.*,
767 108(D24), doi:10.1029/2003jd003990.
- 768 Griffin, D. W., et al. (2001), African desert dust in the Caribbean atmosphere: Microbiology and
769 public health, *Aerobiologia*, 17, 203–213.
- 770 Griffin, D. W., et al. (2003), Atmospheric microbiology in the northern Caribbean during
771 African dust events, *Aerobiologia*, 19, 143–157.
- 772 Griffin, D. W., (2007), Atmospheric movement of microorganisms in clouds of desert dust and
773 implications for human health, *Clin. Microbiol. Rev.* 20, 459–477.
- 774 Hackl, E., G. Bachmann, and S. Zechmeister-Boltenstern (2000), Soil microbial biomass and
775 rhizosphere effects in natural forest stands, *Phyton.-Ann. Rei. Bot. A*, 40(4), 83-90.
- 776 Hader, J. D., T. P. Wright, and M. D. Petters (2014), Contribution of pollen to atmospheric ice
777 nuclei concentrations, *Atmos. Chem. Phys.*, 14, 5433-5449.
- 778 Hicks, S., H. Tinsley, A. Huusko, C. Jensen, M. Hättestrand, A. Gerasimides, and E. Kvavadze
779 (2001), Some comments on spatial variation in arboreal pollen deposition: First records
780 from the Pollen Monitoring Programme (PMP), *Rev. Palaeobot. Palynol.*, 117, 183-194.
- 781 Hirst J. M., O. J. Stedman, and W. H. Hogg (1967), Long-distance spore transport: Methods of
782 measurement, vertical spores profile and the detection of immigrant spores, *J. Gen.*

- 783 Microbiol. 48, 329-355.
- 784 Hjelmroos, M., and L. G. Franzen (1994), Implications or recent long distance pollen transport
785 events for the interpretation of fossil pollen records in Fennoscandia, *Rev. Palaeobot.*
786 *Palynol.*, 82, 175-189.
- 787 Hua, N.-P. et al. (2007), Detailed identification of desert-originated bacteria carried by Asian
788 dust storms to Japan, *Aerobiologia*, 23, 291-298.
- 789 Hyde, K. D., E. B. G. Jones, E. Leano, S. B. Pointing, A. D. Poonyth, and L. L. P. Vrijmoed
790 (1998), Role of fungi in marine ecosystems, *Biodivers. Conserv.*, 7(9), 1147-1161.
- 791 Ibrahim, M., A. B. Rabah, B. Liman, and N. T. Ibrahim (2011), Effect of temperature and
792 relative humidity on the growth of *Helminthosporium fulvum*, *Nigerian J. Basic Appli.*
793 *Sci.*, 19(1) 127- 129.
- 794 Ion, A. C., R. Vermeylen, I. Kourtchev, J. Cafmeyer, X. Chi, A. Gelencser, W. Maenhaut, and
795 M. Claeys (2005), Polar organic compounds in rural PM(2.5) aerosols from K-pusztá,
796 Hungary, during a 2003 summer field campaign: Sources and diel variations, *Atmos.*
797 *Chem. Phys.*, 5, 1805-1814.
- 798 Iwasaka, Y., et al. (1983), The transport and spacial scale of Asian dust-storm clouds: a case
799 study of the dust-storm event of April 1979, *Tellus*, 35B, 189–196.
- 800 Jaffe, D., et al. (1999), Transport of Asian air pollution to North America, *Geophys. Res. Lett.*,
801 26(6), 711–714.
- 802 Jaffe, D., I. McKendry, T. Anderson, and H. Price (2003), Six “new” episodes of trans-Pacific
803 transport of air pollutants, *Atmos. Environ.*, 37(3), 391–404.
- 804 Jaffe, D.A., Mahura A., Kelley, J., Atkins J., Novelli P.C. and Merrill J. Impact of Asian
805 Emissions on the Remote North Pacific Atmosphere: Interpretation of CO Data from
806 Shemya, Guam, Midway and Mauna Loa. *J. Geophys. Res.* 23, 28,627-28,636, 1997.
- 807 Jeon, E.-M. et al. (2011), Impact of Asian dust events on airborne bacterial community assessed
808 by molecular analyses, *Atmos. Environ.* 45, 4313–4321.
- 809 Jia, Y. L., A. L. Clements, and M. P. Fraser (2010), Saccharide composition in atmospheric
810 particulate matter in the southwest US and estimates of source contributions, *J. Aerosol*
811 *Sci.*, 41(1), 62-73.
- 812 Jones, E.B.G. (ed.) (1976), *Recent Advances in Aquatic Mycology*, London: Elek Science.
- 813 Jones, E.B.G., and J. I. Mitchell (1996), Biodiversity of marine fungi, In *Biodiversity:*
814 *International Biodiversity Seminar* (A. Cimerman and N. Gunde-Cimerman, eds), pp. 31-
815 42. Ljubljana: National Inst. Chemistry and Slovenia National Commission for UNESCO.
- 816 Juntto S and P. Paatero (1994), Analysis of daily precipitation data by positive matrix
817 factorization. *Environmetrics*, 5, 127–144.
- 818 Kanakidou, M., et al. (2005), Organic aerosol and global climate modelling: a review, *Atmos.*
819 *Chem. Phys.*, 5, 1053-1123.
- 820 Kawamura, K., Y. Ishimura, and K. Yamazaki (2003), Four years' observations of terrestrial
821 lipid class compounds in marine aerosols from the western North Pacific, *Global*
822 *Biogeochem. Cy.*, 17(1), doi:10.1029/2001gb001810.
- 823 Kim, V. K., and C.L. Xiao (2005), Influence of culture media and environmental factors on
824 mycelial growth and picnidial production of *Sphaeropsis pyripitrescens*, *Mycologia*,
825 97(1): 25-32.

- 826 Kohlmeyer, J., and B. Volkmann-Kohlmeyer (1991), Illustrated key to the filamentous fungi,
827 Bot. Mar. 34, 1-61.
- 828 Kohlmeyer, J., and E. Kohlmeyer (1979), Marine Mycology: The Higher Fungi, London:
829 Academic Press.
- 830 Lacey, M. E., and J. S. West (2006), The Air Spora: a Manual for Catching and Identifying
831 Airborne Biological Particles, pp. 32-33.
- 832 Lewis, D. H., and D. C. Smith (1967), Sugar alcohols (polyols) in fungi and green plants. I.
833 Distribution physiology and metabolism, New Phytologist, 66(2), 143.
- 834 Loos, H., R. Kramer, H. Sahn, and G. A. Sprenger (1994), Sorbitol promotes growth of
835 zymomonas mobilis in environments with high concentrations of sugar: evidence for a
836 physiological function of glucose-fructose oxidoreductase in osmoprotection, J.
837 Bacteriol., 176(24), 7688-7693.
- 838 Lu J., P. Jiang, et al. (2008), Assessing soil quality data by positive matrix factorization.
839 Geoderma, 145, 259-266.
- 840 Ma, S. X., Z. Z. Wang, X. H. Bi, et al. (2009), Composition and source of saccharides in
841 aerosols in Guangzhou, China, Chinese Sci. Bull., 54, 4500-4506.
- 842 Makra, L., T. Santa, I. Matyasovszky, A. Damialis, K. Karatzas, K. C. Bergmann, and D. Vokou
843 (2010), Airborne pollen in three European cities: Detection of atmospheric circulation
844 pathways by applying three-dimensional clustering of backward trajectories, J. Geophys.
845 Res.-Atmos., 115, doi:10.1029/2010jd014743.
- 846 Malik, V. K., and S. Singh (2004), Effect of temperature and relative humidity on teliospore
847 germination in *Ustilago hordei*, J. Mycol. Plant Pathol., 34, 410-411.
- 848 Martin, S. T., et al. (2010), Sources and Properties of Amazonian Aerosol Particles, Rev.
849 Geophys., 48, doi:10.1029/2008rg000280.
- 850 Medeiros, P. M., M. H. Conte, J. C. Weber, and B. R. T. Simoneit (2006), Sugars as source
851 indicators of biogenic organic carbon in aerosols collected above the Howland
852 Experimental Forest, Maine, Atmos. Environ., 40(9), 1694-1705.
- 853 Medeiros, P. M., and B. R. T. Simoneit, (2008), Source profiles of organic compounds emitted
854 upon combustion of green vegetation from temperate climate forests, Environ. Sci.
855 Technol., 42, 8310-8316.
- 856 Miguel, A. G., Taylor P. E., House J., Glovsky M. M., and Flagan R. C. (2006), Meteorological
857 influences on respirable fragment release from Chinese elm pollen, Aerosol Sci. Technol.
858 40, 690-696.
- 859 Mims, S. A., and F. M. Mims (2003), Fungal spores are transported long distances in smoke
860 from biomass fires, Atmos. Environ., 38, 651-655.
- 861 Miyazaki, Y., J. Jung, P. Fu, Y. Mizoguchi, K. Yamanoi, and K. Kawamura (2012), Evidence of
862 formation of submicrometer water-soluble organic aerosols at a deciduous forest site in
863 northern Japan in summer, J. Geophys. Res., 117, D19213, doi:10.1029/2012JD018250.
- 864 Mochida, M., A. Kawabata, K. Kawamura, H. Hatsushika, and K. Yamazaki (2003a), Seasonal
865 variation and origins of dicarboxylic acids in the marine atmosphere over the western
866 North Pacific, J. Geophys. Res.-Atmos., 108(D6), doi:10.1029/2002jd002355.
- 867 Mochida, M., K. Kawamura, N. Umemoto, M. Kobayashi, S. Matsunaga, H. J. Lim, B. J. Turpin,
868 T. S. Bates, and B. R. T. Simoneit (2003b), Spatial distributions of oxygenated organic
869 compounds (dicarboxylic acids, fatty acids, and levoglucosan) in marine aerosols over

- 870 the western Pacific and off the coast of East Asia: Continental outflow of organic
871 aerosols during the ACE-Asia campaign, *J. Geophys. Res.-Atmos.*, 108(D23),
872 doi:10.1029/2002jd003249.
- 873 Mochida, M., K. Kawamura, P. Q. Fu, and T. Takemura (2010), Seasonal variation of
874 levoglucosan in aerosols over the western North Pacific and its assessment as a biomass-
875 burning tracer, *Atmos. Environ.*, 44(29), 3511-3518.
- 876 Morris, C. E., D. G. Georgakopoulos, and D. C. Sands (2004), Ice nucleation active bacteria and
877 their potential role in precipitation, *J. Phys. Iv.*, 121, 87-103.
- 878 Moss, S.T. (ed.) (1986), *The Biology of Marine Fungi*, Cambridge: Cambridge University Press.
- 879 Nakagiri, A., S. Y. Newell, T. Ito, and T. K. Tan (1996), Biodiversity and ecology of the
880 oomycetous fungus, *Halophytophthora*, In *Biodiversity and the Dynamics of Ecosystems*.
881 (DIWPA series. Vol.1) (I.M. Turner, C.H. Diong, S.S.L. Lim, and P.K.L. Ng, eds), pp.
882 273-80. Tokyo: International Network for DIVERSITAS in Western Pacific and Asia.
- 883 Nolte, C. G., J. J. Schauer, G. R. Cass, and B. R. T. Simoneit (2001), Highly polar organic
884 compounds present in wood smoke and in the ambient atmosphere, *Environ. Sci.*
885 *Technol.*, 35(10), 1912-1919.
- 886 Paatero, P., and U. Tapper (1994), Positive Matrix Factorization - a Nonnegative Factor Model
887 with Optimal Utilization of Error-Estimates of Data Values, *Environmetrics*, 5(2), 111-
888 126.
- 889 Paatero, P. et al. (2002), Understanding and controlling rotations in factor analytic models,
890 *Chemom. Intell. Lab. Syst.*, 60, 253-264.
- 891 Pacini, E. (2000), From anther and pollen ripening to pollen presentation, *Plant Syst. Evol.* 222,
892 19-43.
- 893 Pashynska, V., R. Vermeylen, G. Vas, W. Maenhaut, and M. Claeys (2002), Development of a
894 gas chromatographic/ion trap mass spectrometric method for the determination of
895 levoglucosan and saccharidic compounds in atmospheric aerosols. Application to urban
896 aerosols, *J. Mass Spectrom.*, 37(12), 1249-1257.
- 897 Pavuluri, C. M., K. Kawamura, E. Tachibana, and T. Swaminathan (2010), Elevated nitrogen
898 isotope ratios of tropical Indian aerosols from Chennai: Implication for the origins of
899 aerosol nitrogen in South and Southeast Asia, *Atmos. Environ.*, 44(29), 3597-3604.
- 900 Polissar A.V., P. K. Hopke, P. Paatero, Y. J. Kaufmann, D. K. Hall, B.A. Bodhaine, E.G. Dutton,
901 J. M. Harris, (1999). The aerosol at Barrow, Alaska: long-term trends and source
902 locations. *Atmos. Environ.*, 33, 2441-2458.
- 903 Prospero, J.M. and D.L. Savoie (1989), Effect of continental sources of nitrate concentrations
904 over the Pacific Ocean, *Nature*, 339, 687- 689.
- 905 Prospero, J. M. et al. (2005), Interhemispheric transport of viable fungi and bacteria from Africa
906 to the Caribbean with soil dust, *Aerobiologia*, 21, 1-19.
- 907 Rogge, W. F., P. M. Medeiros, and B. R. T. Simoneit (2006), Organic marker compounds for
908 surface soil and fugitive dust from open lot dairies and cattle feedlots, *Atmos. Environ.*,
909 40(1), 27-49.
- 910 Rogge, W. F., P. M. Medeiros, and B. R. T. Simoneit (2007), Organic marker compounds in
911 surface soils of crop fields from the San Joaquin Valley fugitive dust characterization
912 study, *Atmos. Environ.*, 41, 8183-8204.
- 913 Rousseau, D.-D., P. Schevin, J. Ferrier, D. Jolly, T. Andreasen, S. E. Ascanius, S. E. Hendriksen,

- 914 and U. Poulsen (2008), Long-distance pollen transport from North America to Greenland
915 in spring, *J. Geophys. Res.-Biogeo.*, 113(G2), doi:10.1029/2007jg000456.
- 916 Rousseau, D.-D., D. Duzer, J.-L. Etienne, G. Cambon, D. Jolly, J. Ferrier, and P. Schevin (2004),
917 Pollen record of rapidly changing air trajectories to the North Pole, *J. Geophys. Res.*, 109,
918 D06116, doi:10.1029/2003JD003985.
- 919 Rousseau, D.-D., P. Schevin, D. Duzer, G. Cambon, J. Ferrier, D. Jolly, and U. Poulsen (2006),
920 New evidence of long distance pollen transport to southern Greenland in late Spring, *Rev.*
921 *Palaeobot. Palynol.*, 141, 277–286.
- 922 Schmidl, C., H. Bauer, A. Dattler, R. Hitzenberger, G. Weissenboeck, I. L. Marr, and H.
923 Puxbaum (2008), Chemical characterisation of particle emissions from burning leaves,
924 *Atmos. Environ.*, 42(40), 9070-9079.
- 925 Seinfeld, J. H., et al., (2004), ACE-ASIA - Regional climatic and atmospheric chemical effects
926 of Asian dust and pollution, *Bull. Amer. Meteorol. Soc.*, 85(3), 367-380.
- 927 Sharma, R. and R. C. Rajak (2003), Keratinophilic fungi: Nature's keratin degrading machines,
928 *Resonance*, 8, 28-30.
- 929 Simoneit, B. R. T. (2002), Biomass burning - A review of organic tracers for smoke from
930 incomplete combustion, *Appl. Geochem.*, 17(3), 129-162.
- 931 Simoneit, B. R. T. (2004), Biomarkers (molecular fossils) as geochemical indicators of life, *Adv.*
932 *Space. Res.*, 33(8), 1255-1261.
- 933 Simoneit, B. R. T., and V. O. Elias (2000), Organic tracers from biomass burning in
934 atmospheric particulate matter over the ocean, *Mar. Chem.*, 69(3-4), 301-312.
- 935 Simoneit, B. R. T., V. O. Elias, M. Kobayashi, K. Kawamura, A. I. Rushdi, P. M. Medeiros, W.
936 F. Rogge, and B. M. Didyk (2004a), Sugars-dominant water-soluble organic compounds
937 in soils and characterization as tracers in atmospheric particulate matter, *Environ. Sci.*
938 *Technol.*, 38(22), 5939-5949.
- 939 Simoneit, B. R. T., M. Kobayashi, M. Mochida, K. Kawamura, M. Lee, H. J. Lim, B. J. Turpin,
940 and Y. Komazaki (2004b), Composition and major sources of organic compounds of
941 aerosol particulate matter sampled during the ACE-Asia campaign, *J. Geophys. Res.-*
942 *Atmos.*, 109(D19), doi:10.1029/2004jd004598.
- 943 Slinn, S. A., and W. G. N. Slinn (1980), Predictions for particle deposition on natural waters,
944 *Atmos. Environ.*, 14, 1013-1016.
- 945 Soonthornnonda P. and E. R. Christensen (2008). Source apportionment of pollutants and flows
946 of combined sewer wastewater. *Water Res.*, 42, 1989–1998.
- 947 Sosnoskie, L. M., T. M. Webster, D. Dales, G. C. Rains, T. L. Grey, and A. S. Culpepper (2009),
948 Pollen grain size, density, and settling velocity for Palmer amaranth (*Amaranthus*
949 *palmeri*), *Weed Science*, 57, 404-409. <http://dx.doi.org/10.1614/WS-08-157.1>.
- 950 Speranza, A., G. L. Calzoni, and E. Pacini (1997), Occurrence of mono- or disaccharides and
951 polysaccharide reserves in mature pollen grains, *Sex. Plant Reprod.*, 10(2), 110–115.
- 952 Sullivan, A. P., N. Frank, D. M. Kenski, Jr., and J. L. Collett (2011), Application of high-
953 performance anion-exchange chromatography-pulsed amperometric detection for
954 measuring carbohydrates in routine daily filter samples collected by a national network: 2.
955 Examination of sugar alcohols/polyols, sugars, and anhydrosugars in the upper Midwest,
956 *J. Geophys. Res.* 116. <http://dx.doi.org/10.1029/2010JD014169>.
- 957 Talbot R.W. et al. (1997), Chemical characteristics of continental outflow from Asia to the

- 958 troposphere over the western Pacific Ocean during February-March 1994: Results from
959 PEM -West B. *J. Geophys. Res.*, 102, 28,255-28,274, 1997.
- 960 Tominaga, S., K. Matsumoto, N. Kaneyasu, A. Shigihara, K. Katono, N. Igawa (2011),
961 Measurements of particulate sugars at urban and forested suburban sites, *Atmos. Environ.*
962 45, 2335-2339.
- 963 Uno, I., K. Eguchi, K. Yumimoto, T. Takemura, A. Shimizu, M. Uematsu, Z. Y. Liu, Z. F.
964 Wang, Y. Hara, and N. Sugimoto (2009), Asian dust transported one full circuit around
965 the globe, *Nat. Geosci.*, 2(8), 557-560.
- 966 Verma, S. K., K. Kawamura, J. Chen, P. Q. Fu and C. Zhu (2015), Thirteen years observation of
967 biomass-burning organic tracers over Chichijima Island in the western North Pacific: An
968 outflow region of Asian aerosols, *J. Geophys. Res. Atmos.*, 120,
969 doi:10.1002/2014JD022224. .
- 970 Wan, E. C. H., and J. Z. Yu (2007), Analysis of sugars and sugar polyols in atmospheric
971 aerosols by chloride attachment in liquid chromatography/negative ion electrospray mass
972 spectrometry, *Environ. Sci. Technol.*, 41(7), 2459-2466.
- 973 Wang, G. H., C. L. Chen, J. J. Li, B. H. Zhou, M. J. Xie, S. Y. Hu, K. Kawamura, and Y. Chen
974 (2011), Molecular composition and size distribution of sugars, sugar-alcohols and
975 carboxylic acids in airborne particles during a severe urban haze event caused by wheat
976 straw burning, *Atmos. Environ.*, 45(15), 2473-2479,
- 977 Wang, G. H., K. Kawamura, and M. Lee (2009), Comparison of organic compositions in dust
978 storm and normal aerosol samples collected at Gosan, Jeju Island, during spring 2005,
979 *Atmos. Environ.*, 43(2), 219-227.
- 980 Wiemken, A. (1990), Trehalose in Yeast, Stress Protectant Rather Than Reserve Carbohydrate,
981 *Anton. Leeuw. Int. J. G.*, 58(3), 209-217.
- 982 Wright, T. P., J. D. Hader, G. R. McMeeking, and M. D. Peters (2014), High relative humidity
983 as a trigger for widespread release of ice Nuclei, *Aerosol Sci. Technol.*, 48, 11, i-v, DOI:
984 10.1080/02786826.2014.968244.
- 985 Xie Y. and C. M. Berkowitz (2006), The use of positive matrix factorization with conditional
986 probability functions in air quality studies: an application to hydrocarbon emissions in
987 Houston, Texas. *Atmos. Environ.*, 40, 3070– 3091.
- 988 Xu, W. Y., et al. (2011), Characteristics of pollutants and their correlation to meteorological
989 conditions at a suburban site in the North China Plain, *Atmos. Chem. Phys.*, 11(9), 4353-
990 4369.
- 991 Yamaguchi, N., T. Ichijo, A. Sakotani, T. Baba, and M. Nasu (2012), Global dispersion of
992 bacterial cells on Asian dust, *Sci. Rep.*, 2, DOI: 10.1038/srep00525.
- 993 Yang, Y. H., C. Y. Chan, J. Tao, M. Lin, G. Engling, Z. S. Zhang, T. Zhang, and L. Su (2012),
994 Observation of elevated fungal tracers due to biomass burning in the Sichuan Basin at
995 Chengdu City, China, *Sci. Total Environ.*, 431, 68–77.
- 996 Yttri, K. E., C. Dye, and G. Kiss (2007), Ambient aerosol concentrations of sugars and sugar-
997 alcohols at four different sites in Norway, *Atmos. Chem. Phys.*, 7(16), 4267-4279.
- 998 Zhang, T., Engling, G., Chan, C.-Y., Zhang, Y.-N., Zhang, Z.-S., Lin, M., Sang, X.-F., Li, Y.D.,
999 Li, Y.-S., 2010. Contribution of fungal spores to particulate matter in a tropical rainforest.
1000 *Environ. Res. Lett.* 5. doi:10.1088/1748-9326/1085/1082/ 024010.
- 1001 Zhou, L., E. Kim et al., (2004), Advanced factor analysis on Pittsburgh particle size distribution
1002 data, *Aerosol Sci. Technol.*, 38, 118-132.

1003 Zhu, C., K. Kawamura, and B. Kunwar (2015), Organic tracers of primary biological aerosol
1004 particles at subtropical Okinawa Island in the western North Pacific Rim, *J. Geophys.*
1005 *Res. Atmos.*, 120, 5504–5523.

1006
1007
1008
1009
1010
1011
1012
1013
1014
1015
1016
1017
1018
1019
1020
1021
1022
1023
1024
1025
1026
1027
1028
1029
1030
1031
1032
1033
1034
1035
1036
1037
1038

1039 **Figure Captions**

1040 **Figure 1.** Geographical location of Chichijima Island (27°04'N; 142°13'E; 254 m, asl) in the
1041 western North Pacific.

1042 **Figure 2.** The monthly variation of the meteorological parameters over Chichijima Island
1043 during 2001-2013 (The error bars denote the standard deviations).

1044 **Figure 3.** The seasonal ten-day air mass backward trajectories over Chichijima for 2012
1045 (winter: Dec-Feb, spring: Mar-May, summer: Jun-Aug, autumn: Sep-Nov). The trajectory
1046 calculations were performed everyday starting at Chichijima Island.

1047 **Figure 4.** Temporal plots for the concentrations (ng m^{-3}) of sugar compounds in Chichijima
1048 aerosol samples collected for 2001-2013 in the western North Pacific.

1049 **Figure 5.** Monthly mean concentrations (ng m^{-3}) of sugar compounds in aerosol samples from
1050 Chichijima Island in the western North Pacific during 2001-2013.

1051 **Figure 6.** Seasonal source contributions to sugar compounds from various sources based on
1052 PMF analyses. (BB – biomass-burning; Mixed – vegetation, fungal and microbial sources).

1053 **Figure 7.** Annual mean concentrations (ng m^{-3}) of sugar compounds in aerosol samples
1054 collected from Chichijima Island in the western North Pacific during 2001-2013.

1055 **Figure 8.** The seasonal concentrations of anhydrosugars (biomass burning tracers), primary
1056 sugars and sugar alcohols measured in Chichijima aerosols during three periods, i.e., P-I
1057 (1990-1993), P-II (2001-2003) and P-III (2010-2013).

1058 **Figure 9.** PMF analyses of sugar compounds in Chichijima aerosols based on the 2001-2013
1059 data set. (BB – biomass-burning; Mixed – vegetation, fungal and microbial sources).

1060 **Figure 10.** Source contributions to sugar compounds from various sources based on PMF
1061 analyses. (BB – biomass-burning; Mixed – vegetation, fungal and microbial sources).

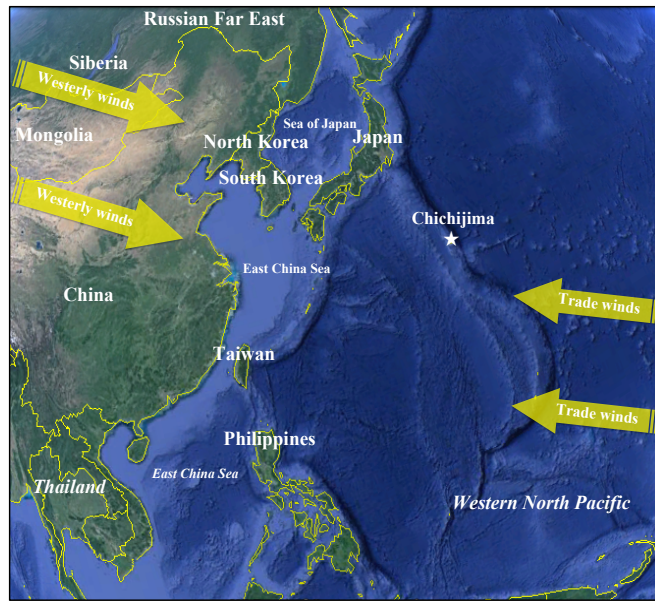
1062 **Figure 11.** Annual trends in % contributions of five source factors: (a) vegetation, (b) fungal
1063 and microbial, (c) mixed, (d) biomass burning (BB), and (e) pollen factors to SCs in
1064 Chichijima aerosols. The data of 2005 are not plotted due to limited data points.

1065

1066

1067
1068
1069
1070
1071
1072
1073
1074
1075

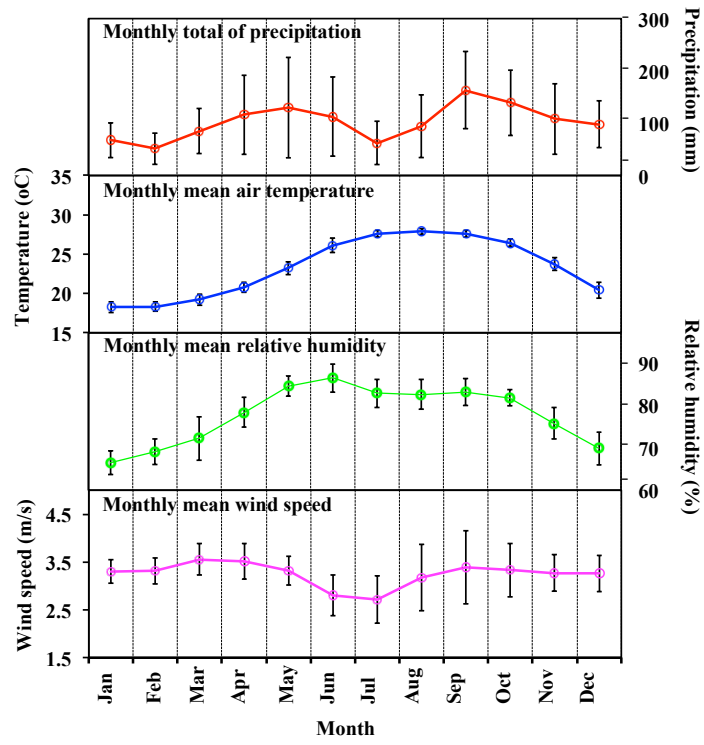
Figure 1.



1087
1088
1089
1090
1091
1092
1093
1094
1095
1096
1097
1098
1099
1100
1101

1102
1103
1104
1105
1106
1107
1108

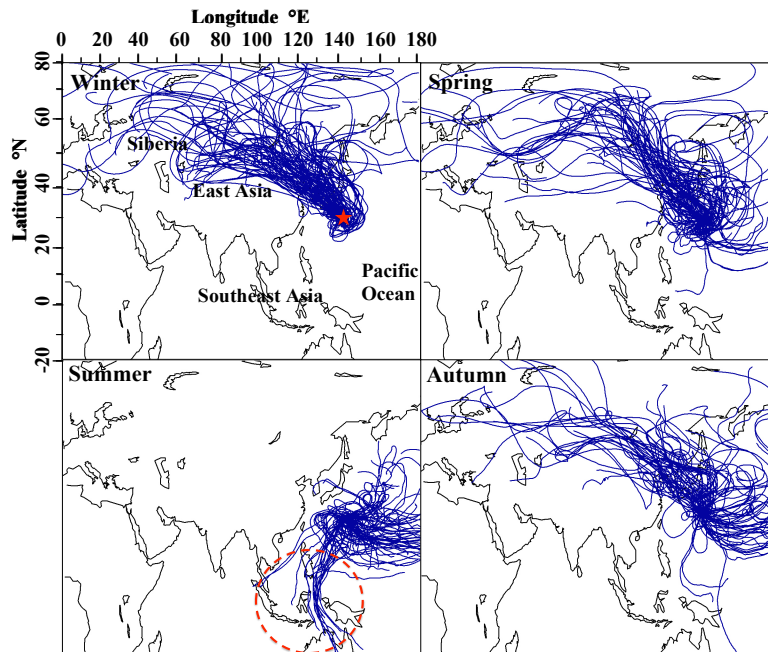
Figure 2.



1109
1110
1111
1112
1113
1114
1115

1116
1117
1118
1119
1120
1121
1122
1123
1124
1125
1126

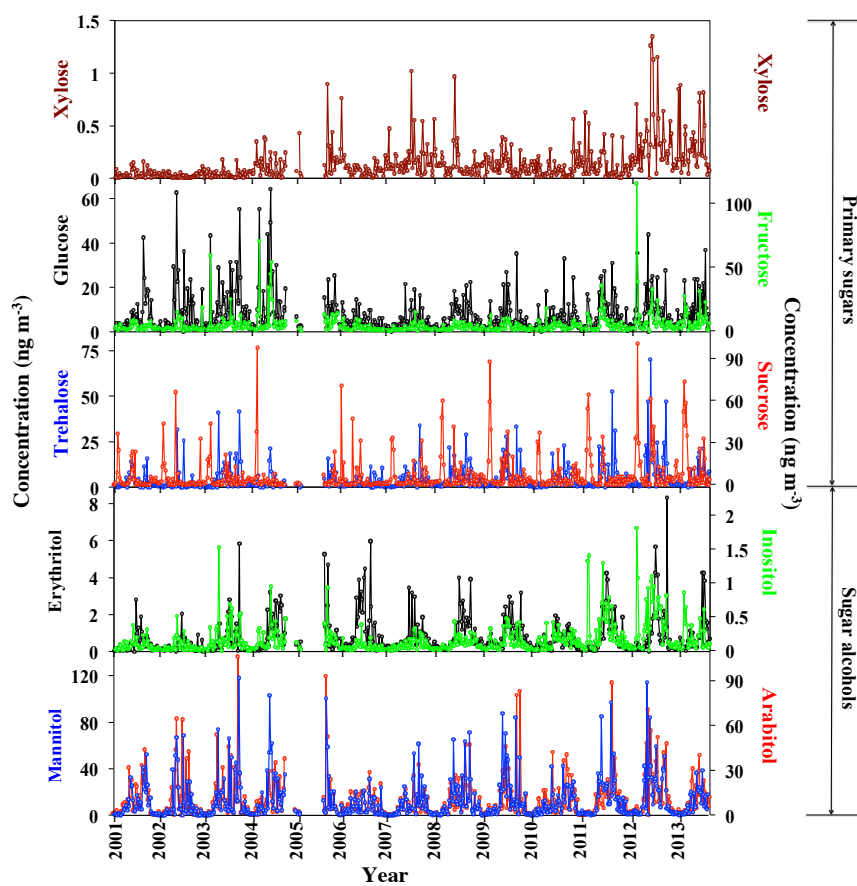
Figure 3.



1127
1128
1129
1130
1131
1132
1133
1134
1135
1136
1137
1138
1139

1140
1141
1142
1143
1144
1145
1146

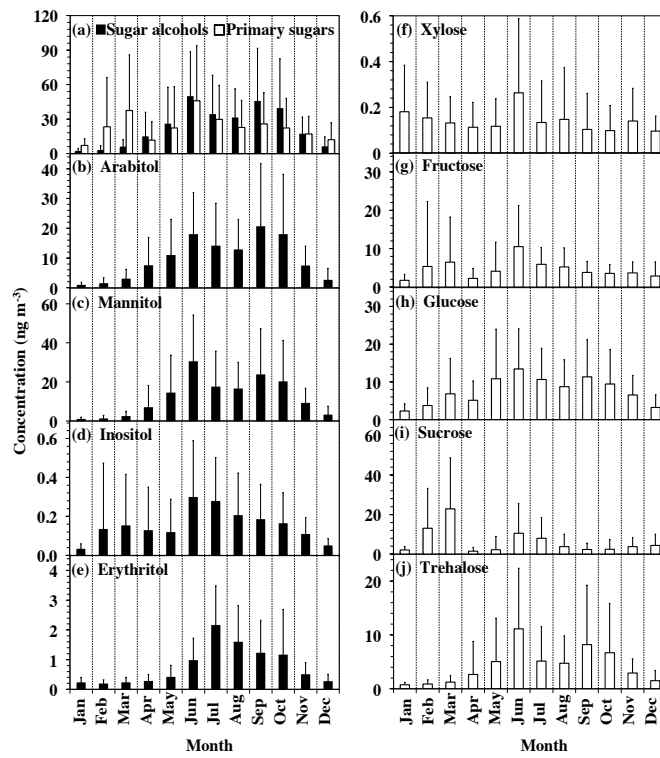
Figure 4.



1147
1148
1149

1150
1151
1152
1153
1154
1155
1156
1157
1158
1159
1160

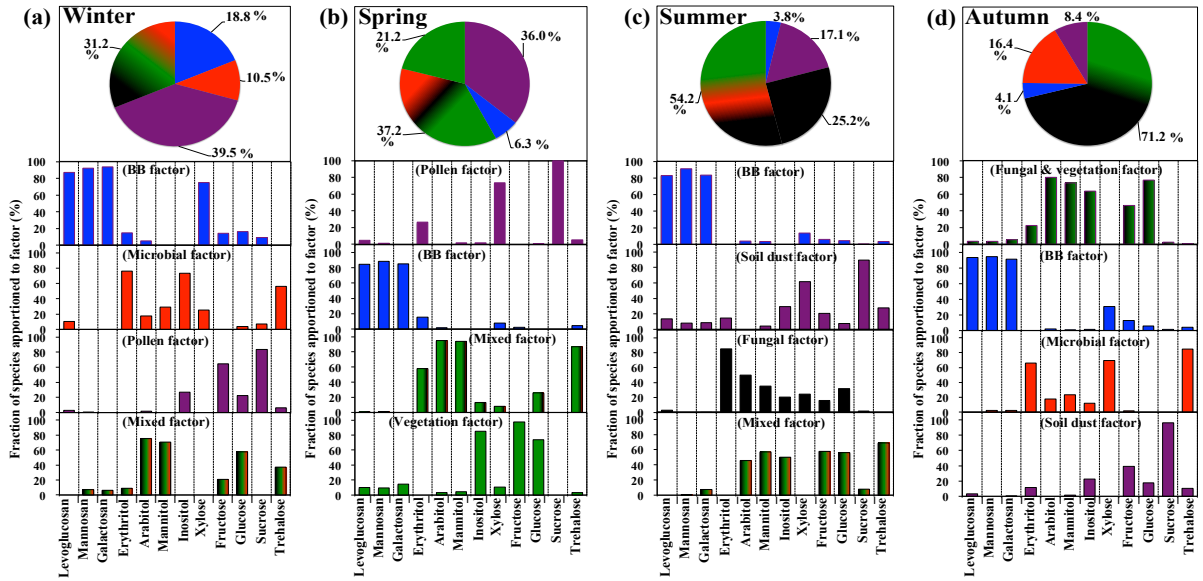
Figure 5.



1161
1162
1163
1164
1165
1166
1167
1168
1169
1170
1171

1172
 1173
 1174
 1175
 1176
 1177
 1178
 1179

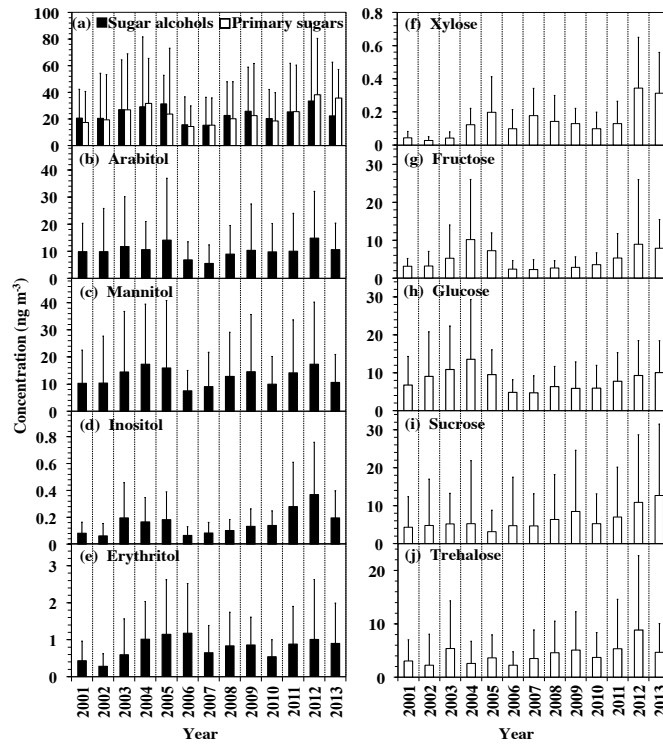
Figure 6.



1180
 1181
 1182
 1183
 1184
 1185
 1186
 1187
 1188
 1189
 1190
 1191
 1192
 1193
 1194
 1195
 1196

1197
1198
1199
1200
1201
1202
1203
1204

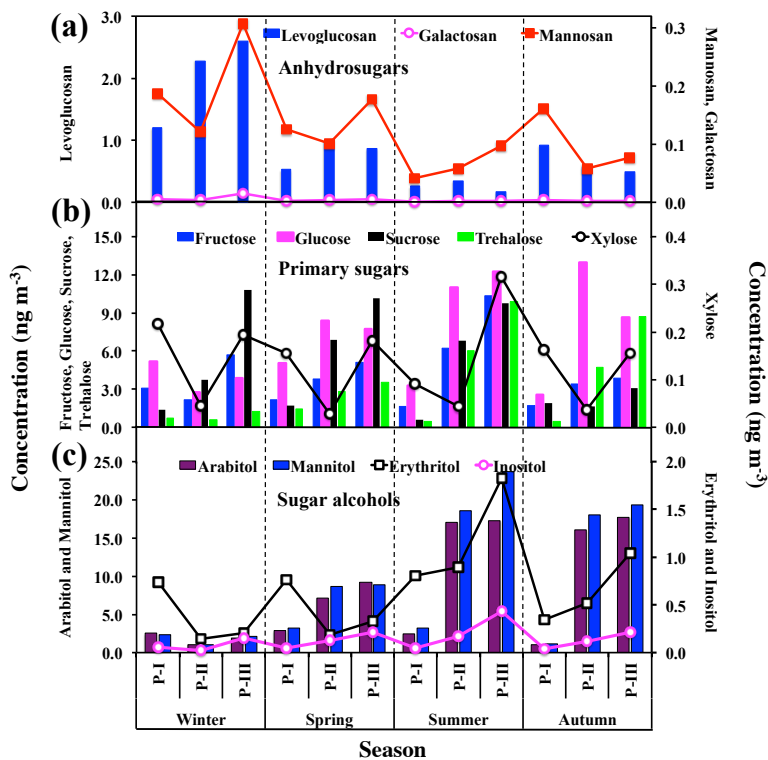
Figure 7.



1205
1206
1207
1208
1209
1210
1211
1212
1213
1214
1215
1216
1217
1218

1219
 1220
 1221
 1222
 1223
 1224
 1225
 1226

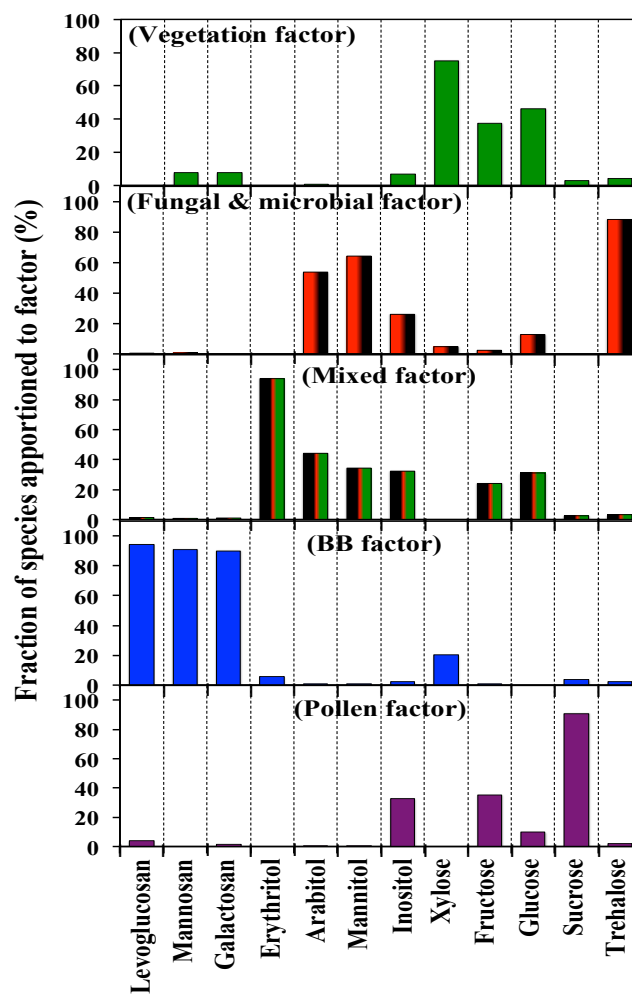
Figure 8.



1227
 1228
 1229
 1230
 1231
 1232
 1233
 1234
 1235
 1236
 1237
 1238
 1239
 1240

1241
 1242
 1243
 1244
 1245
 1246
 1247
 1248
 1249
 1250

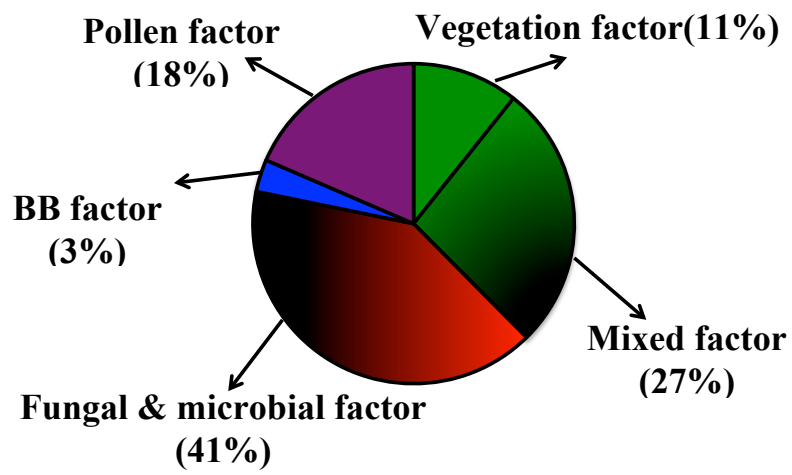
Figure 9.



1251
 1252
 1253
 1254
 1255
 1256
 1257

1258
1259
1260
1261
1262
1263
1264
1265
1266
1267
1268
1269
1270
1271

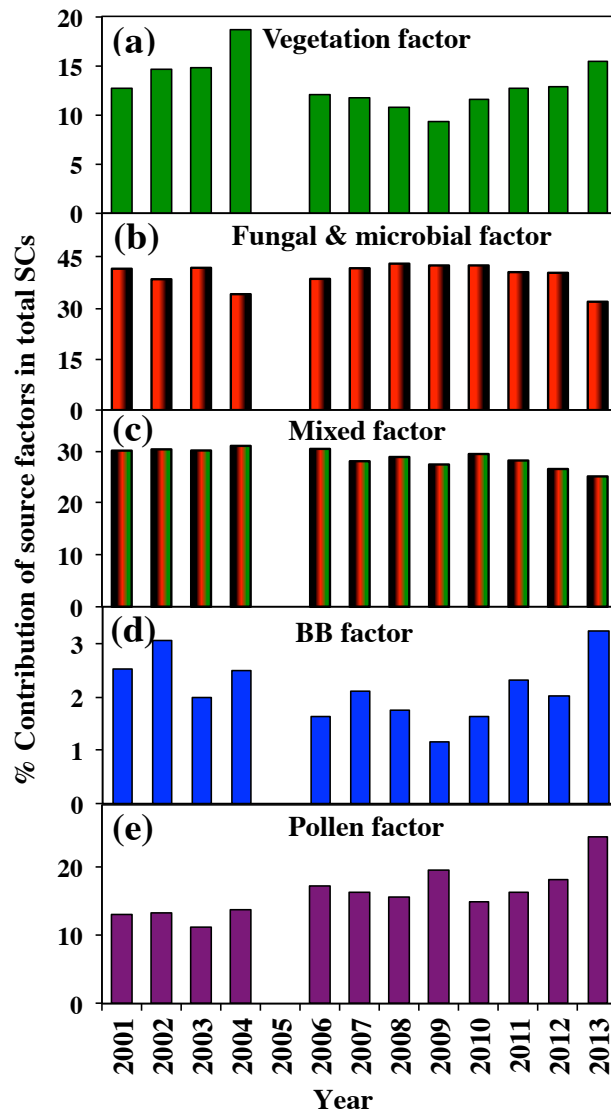
Figure 10.



1272
1273
1274
1275

1276
1277
1278
1279
1280
1281
1282
1283
1284

Figure 11.



1285

1286
1287
1288

Table 1. Seasonal concentrations (ng m⁻³) of sugar compounds (SCs) in the aerosol samples collected at ChichiJima Island in the western North Pacific during 2001-2013.

Sugars	Winter ^a (n=139)			Spring ^b (n=155)			Summer ^c (n=146)			Autumn ^d (n=150)			2001-2013 (n=590)		
	Range	Mean±SD	Med ^e	Range	Mean±SD	Med ^e	Range	Mean±SD	Med ^e	Range	Mean±SD	Med ^e	Range	Mean±SD	Med ^e
Primary sugars															
Xylose	0.05-0.89	0.14±0.15	0.12	0.001-0.56	0.12±0.11	0.08	0.002-1.35	0.18±0.26	0.08	0.004-0.89	0.11±0.14	0.07	0.001-1.35	0.14±0.18	0.08
Fructose	0.17-115	3.36±10.2	1.38	0.03-70.4	4.51±9.21	1.90	0.24-54.0	7.25±7.63	5.23	0.22-13.0	3.70±2.68	2.82	0.03-115	4.69±8.04	2.53
Glucose	0.27-23.3	3.11±3.53	2.13	0.05-62.6	7.68±10.3	4.36	0.23-64.3	11.0±9.02	7.99	0.72-55.5	9.25±8.63	5.77	0.05-64.3	7.79±8.80	4.73
Sucrose	0.02-73.4	6.60±13.1	2.08	0.005-100	8.80±18.0	1.30	0.003-66.0	7.31±11.5	2.58	0.002-31.1	2.76±4.35	1.14	0.002-100	6.43±12.9	1.71
Trehalose	0.03-10.5	1.03±1.26	0.72	0.006-47.2	2.93±6.08	1.23	0.03-70.2	7.06±8.49	4.46	0.04-52.5	6.09±8.81	2.46	0.01-70.2	4.30±7.28	1.61
Σ Primary sugars	0.49-223	14.2±28.2		0.09±281	24.2±43.8		0.51-256	32.8±36.9		0.99-153	22.0±24.6		0.28-176	23.3±25.7	
Sugar alcohols															
Erythritol	0.03-1.17	0.23±0.18	0.18	0.008-2.25	0.31±0.28	0.23	0.07-5.70	1.55±1.21	1.18	0.05-8.32	0.99±1.16	0.56	0.01-8.32	0.77±1.01	0.37
Arabitol	0.12-21.2	1.73±2.60	0.96	0.04-70.8	7.13±9.50	4.44	0.24-64.5	15.1±12.9	12.0	0.57-106	15.8±18.3	8.83	0.04-106	9.99±13.6	4.97
Mannitol	0.10-23.9	1.89±2.81	1.11	0.16-114	7.95±13.8	4.07	0.25-104	21.7±19.7	16.9	0.59-118	18.2±19.9	9.15	0.10-118	12.5±17.5	5.54
Inositol	0.01-1.81	0.07±0.20	0.03	0.008-1.51	0.13±0.22	0.06	0.01-1.29	0.26±0.25	0.17	0.01-0.93	0.16±0.15	0.10	0.01-1.81	0.16±0.22	0.08
Σ Sugar alcohols	0.26-48.2	3.93±5.79		0.22-188	15.5±23.8		0.56-175	38.6±34.1		1.22-234	35.1±39.5		0.37-231	23.4±30.8	
Σ SCs	0.75-272	18.2±34.0	10.6	0.31-469	39.8±67.6	18.2	1.07-431	71.5±70.9	50.7	2.21-387	57.0±64.2	31.1	1.23-339	46.7±49.5	30.9

^a Winter (December-February), ^b Spring (March-May), ^c Summer (June-August) and ^d Autumn (September-November); Med.=Median

1289
1290
1291
1292
1293
1294
1295

1297
1298
1299

Table 2. Pearson correlation coefficients (r) for the dataset of sugars in Chichijima aerosols during 2001-2013 (n = 590).

	Levoglucozan ^a	Mannosan ^a	Galactosan ^a	Erythritol	Arabitol	Mannitol	Inositol	Xylose	Fructose	Glucose	Sucrose	Trehalose
Levoglucozan	1.00											
Mannosan	0.79	1.00										
Galactosan	0.55	0.58	1.00									
Erythritol	-0.18	-0.12	-0.16	1.00								
Arabitol	-0.16	-0.09	-0.16	0.48	1.00							
Mannitol	-0.18	-0.12	-0.17	0.49	0.88	1.00						
Inositol	-0.06	-0.02	0.10	0.35	0.49	0.58	1.00					
Xylose	0.20	0.32	0.34	0.15	0.18	0.23	0.42	1.00				
Fructose	0.02	0.08	0.26	0.17	0.16	0.28	0.57	0.31	1.00			
Glucose	-0.11	0.00	-0.06	0.32	0.63	0.72	0.53	0.19	0.57	1.00		
Sucrose	0.05	0.07	0.18	-0.02	-0.06	0.01	0.40	0.26	0.30	0.14	1.00	
Trehalose	-0.10	-0.05	-0.10	0.33	0.73	0.80	0.55	0.33	0.22	0.54	0.13	1.00

^a data from Verma et al. (2015)

1300
1301
1302
1303
1304
1305
1306
1307
1308
1309
1310
1311
1312
1313

1314
 1315
 1316
 1317
 1318

Table 3. Comparisons of seasonal concentrations (ng m^{-3}) of primary sugars and relative contributions (%) of sugar compounds (SCs) in total SCs in Chichijima aerosols among 1990-1993^a, 2001-2003 and 2010-2013.

Season	Anhydrosugars			Sugar alcohols			Primary sugars			Total Sugars		
	1990-93 ^a	2001-03 ^b	2010-13 ^b	1990-93 ^a	2001-03	2010-13	1990-93 ^a	2001-03	2010-13	1990-93 ^a	2001-03	2010-13
Winter	1.44	2.44	3.05	5.64	2.27	4.37	10.6	9.34	21.8	17.7	14.0	29.2
%	11.4	23.7	16.6	31.6	16.5	19.8	57.0	59.9	63.6			
Spring	0.68	1.12	1.10	6.91	16.2	18.6	10.5	21.3	26.7	18.1	38.6	46.4
%	5.08	5.17	4.04	39.2	39.7	42.0	55.7	55.1	53.9			
Summer	0.31	0.43	0.29	6.50	36.7	43.1	6.07	30.0	42.5	12.9	67.1	86.0
%	2.92	0.91	0.43	55.6	51.3	51.1	41.4	47.8	48.5			
Autumn	1.12	0.65	0.60	2.57	34.7	38.3	6.79	22.6	24.5	10.5	57.9	63.4
%	12.9	3.45	1.73	29.2	50.4	57.2	57.9	46.2	41.1			

^a data from Chen et al. (2013), ^b data from Verma et al. (2015).

1319
 1320