

Anonymous Referee #1

Comments on “Measurement report: Diurnal and temporal variations of sugar compounds in suburban aerosols from the northern vicinity of Beijing, China: An influence of biogenic and anthropogenic sources” by Verma et al., October 2020.

General Comments:

In this manuscript the authors report observations of sugar compounds (SCs) in air at a rural site about 40 km north of Beijing from Aug 15- Oct 5, 2007. Diurnal variability is examined, and meteorological parameters are considered as explanatory variables. The SC time series were analyzed with positive matrix factorization to identify sugar aerosol types and the relative contributions of the sugars and aerosol types to organic aerosol mass are reported. Overall this is a useful contribution of measurements to a topic that is still not well-understood. The differentiation between daytime and nighttime samples is rightly recognized as important by the authors. However, the manuscript often reads like a laundry list of observations interspersed with comparisons to previous observations, and insufficient analysis to support some claims.

Response: Authors are thankful to the reviewer for his valuable comments and suggestions, which help to upgrade the quality of the manuscript. We make significant changes in the manuscript especially in section 2. Materials and methods and 3.1. - 3.2. Results and discussion. We deleted several phrases of comparisons with previous observations.

Some inferences made about the observations are provided as speculations with little explanation or supporting analysis. In fact there are a few claims made in the manuscript (noted below) that don't even seem to clearly follow from the evidence presented. The logic behind the claims needs to be clarified or the claims need to be changed or removed. A major driver of this issue is that day-night differences were also differences in temperature and humidity, and differences in air mass origin. How can these effects be disentangled to draw inferences? For example, sucrose is interpreted to be controlled by local emission related to temperature and radiation on the basis of correlation with those variables, and transport is not considered. But arabitol and mannitol correlate very closely with local RH- why in the Abstract is it claimed that these are related to transport from Beijing? Perhaps this would be clearer if a more comprehensive table of correlation coefficients were shown for the relevant quantities, for daytime, nighttime, and overall, but some more textual clarification would also help. Also, would be really nice to have some proxy of transport there too, e.g. average magnitude of the wind in the direction between Beijing and the site.

Response: We modified section 3.1.3 in the results and discussions added new lines in the revised MS. Following the reviewer's suggestions we added new correlation table (Table 2) in the revised MS. We have added a new figure (Figure 2b) showing the day and night time wind direction in the Mangshan site. Please see the revised MS.

After deeply digging the dataset we found some interesting facts about the mannitol contribution in the Mangshan aerosol samples. We added several lines in the revised manuscript. Please see lines 358 – 372 in the revised MS.

A major limitation is the lack of any air mass trajectory analysis. The authors state that there was a typical diurnal pattern to the wind direction, with daytime winds from the south and the large cities, and nighttime winds from more rural areas to the north. It would help the reader to see how consistent this pattern was in order to evaluate some of the claims made. I suggest at least a time series plot of wind speed and wind direction, perhaps as a sub-plot to Figure 2, or a wind rose diagram. A couple of representative air mass back trajectories could be instructive as well.

Response: Following the reviewers suggestions we added new figure (Figure 2b). Figure 2 shows the wind direction and magnitude of wind directions in the Mangshan site. Please see the revised MS.

The results section should have more description of the present data set, with more of the previous observations moved to the Introduction. In particular, the description of the PMF results could be expanded, perhaps with a figure showing the time series of the PMF factors.

Response: We significantly modified the result and discussion sections by adding new phrases and relocating the discussions on previous studies. Please see lines 211–214, 242–250, 261–264, 294–297, 306–317, 328–334, 345–347, 353–363, 368–375, 378–384.

We added information about the PMF analysis in the revised MS. Please see lines 187 – 201, 389 – 407, 418 – 429, 442 – 445, 453 – 457, 460 – 462 in the revised MS.

We added Figure S-1 and S-2 as supporting information in the revised MS.

Figure S-1. The Scatter plots between observed (input data) and predicted (modeled data) concentrations show statistical parameter (coefficient of determination (r), Intercept, and Slope) with linear equation of individual sugar compounds. A blue 1:1 line is provided on this plot for reference (a perfect fit would line up exactly on this line), and the regression line is shown as a dotted red line.

Figure S-2. The time series plots between observed (input data) and predicted (modeled data) concentrations of individual sugar compounds. Blue and red lines show observed (input data) and predicted (modeled data) concentrations, respectively.

Throughout the manuscript, the authors should to be more careful in their descriptions of how the SCs get into the atmosphere. They frequently state that a process or an organism “emits” a sugar compound, which reads somewhat ambiguously (i.e. are gas phase compounds being released?). I think it’s OK to use this language, but there first needs to be a clear statement in the Introduction about how SCs get into the atmosphere, or at least the state of the science on that question. How are SCs are released into the air, as fragments of organisms, as whole fungal spores, as individual molecules, etc?

Response: We added new paragraph in the introduction section, including a clear statement on the sources of individual sugar species in the atmosphere. Please see lines 55 – 61, 67 – 75, 78 – 100.

We modified the sentences where we used “emits” in the revised MS.

A further issue that needs to be resolved before this is publishable is the extremely sparse description of the methods. Specific questions are raised below.

At this point this manuscript is essentially a descriptive account of measurements made at a particular location, with some interpretive claims that seem a bit ambiguous. The measurements themselves are of value, but I think for publication, it needs 1) a much more thorough method description section and 2) either a) a scale-back of the claims made, or b) additional analysis in support of the claims.

Response: We significantly modified the sections of materials and methods by including new information and new section in the revised MS. Please see lines 138 - 144, 147 – 149, 161 – 201.

Specific comments:

There are several grammatical issues of subject-verb agreement and lack of pluralization throughout the manuscript. They don't usually impede understanding, but the manuscript would benefit from a thorough grammar check.

Response: We significantly corrected grammatical issues in the manuscript. Please see the revised MS.

It would be very helpful to view the data in Figure 2 directly as a part of Figure 3.

Response: We added new figure in the (Figure 2b) in the revised MS. We also separated figure 3 in three parts (Fig 4a-c, 5a-d and 6a-c) according to the groups of sugar compounds. The new arrangement is very easy to understand and clearer than the earlier version. Please see the revised MS.

Line 66: “SCs are emitted from algae, microbes, pollen, suspended soil particle[s], and associated biota into the atmosphere” This statement reads to me a bit like sugar compounds might be released into the air as individual gas phase molecules, which I don't think is the case (?). Maybe it could be phrased “SCs are emitted as part of aerosols formed from algae...”? Same thing at line 72: I don't think mannitol and arabitol are mostly emitted as individual gases, but are a part of fungal material that gets into the atmosphere. What is physically meant here needs to be a little clearer. Are these SCs usually part of biological fragments, or is this unknown?

Response: Sentence rephrased please see lines 78 – 81 and 85 - 87 in the revised MS.

Section 2.2:

Please provide more details of the sampling apparatus and methodology. Were these samples collected with a high-volume sampler? Where was it installed specifically, and at what height? What aerosol sizes were collected? Were there any measures to avoid the sampling of gas phase components? Did the 3-hr and 9-hr samples overlap

in time? What times were the samples started? Perhaps a table of sample collection times would be helpful.

Response: We added Table S1 that contains the detailed description of the sampling procedures and collections. Other information's are added in the text. Please see section "2.1. Site description and aerosol sample collection" in the revised MS.

The twelve (n = 12) aerosol samples on 15 to 18, September and 28 September to 5 October were collected for 9-h from 9 to 18 h. While twenty six (n = 26) aerosol samples of 19 to 27 september were collected from 9 to 12, 12 to 15, 15 to 18 h.

We did not use a denuder system to remove gaseous components in high volume air sampler.

Furthermore, what were the methods for determining the WSOC and total OC? How was Ca(2+) concentration determined? Was the filter cut into sections for each analysis?

Response: Information's are added in the revised manuscript. Please see lines 185 – 186 in the revised MS.

Line 132: What is meant by C13 n-alkane? Is this an isotope standard of one n-alkane? Which one? Or do you mean C13H28, n-tridecane?

Response: C₁₃ n-alkane is n-tridecane (C₁₃H₂₈).

Line 157: "Hence, it is evident that increased BB activities at nighttime are associated with cool temperature (Fig. 2)." Is this saying that because it's cool at night, it makes sense that there's more BB aerosol at night? Isn't it equally likely that the different air mass origins in the day and at night are the reason?

Response: The sentence is rephrased. Please see lines 242 - 243 in the revised MS. The day-night time difference of the air masses can also influence the concentrations of BB tracers. We added related text please see lines 242 - 250 in the revised MS.

Line 209: "the meteorological conditions". Is this referring to the strong daytime winds and convective activity? It would be clearer to state that directly.

Response: Modified. Please see lines 272 - 274 in the revised MS.

Line 254: "northeasterly (99.5%)". Does this mean 99.5% of the nighttime hours the wind was northeasterly? Please clarify in the text.

Response: Yes, sentence rephrased (Please see lines 308-310 in revised MS).

Line 255: What would cause sugar emissions to decrease with lower temperature? Is there supporting literature for this?

Response: The daytime ambient temperature and solar radiations significantly affect plant activities and, subsequently, emissions of sugar enriched plant fragments. Therefore, the contribution of primary sugars at night was lower than in daytime. Miyazaki et al. (2012) reported the emissions of sugar compounds associated with

light and ambient temperature at forest site. Please see lines 311 – 317 in the revised MS.

Line 268: Trehalose paragraph. Trehalose didn't show a strong diurnal cycle, but the authors point out a correlation between trehalose and mannitol and arabitol at night, and between trehalose and Ca (2+) in the day. It would be helpful to at least report the corresponding correlation coefficients for the day and night, respectively, for comparison, and possibly to include the corresponding figures in Figure 5.

Response: We added correlation coefficient values in the text. We also added a new Table (Table 2) for correlation of sugar species and meteorological parameters in the manuscript. Please see lines 332 – 334, 336 - 337 in the revised MS.

Line 272: Why would nighttime low RH and temp cause microorganisms to emit more trehalose? Please cite a reference. Again, the use of "emit" here can be confusing. Is it the release of spores that prefers these conditions?

Response: Several studies have reported that the meteorological conditions, i.e., high RH and low temperature, are favorable for the microbes and fungi to discharge spores. The high RH and low temperature was recorded at night time therefore the microbes emit spores at night. We added information in the revised manuscript. Please see lines 328-339 in the revised MS. References are also added in the text and references section.

Line 310: Aren't the Mt. Tai measurements higher than Mangshan, not lower?

Response: We deleted this comparison. Please see in the revised MS.

Lines 315-319: I don't understand the reasoning here. How does RH relate to transport from megacities as an explanation for fungal aerosol?

Response: Rephrased. Please see lines 368 – 375, 378-385 in the revised MS.

Line 350: Separate the Factor 3 and Factor 4 descriptions into separate paragraphs.

Response: Factor 3 and Factor 4 are described into separate paragraphs (Please see line 440 – 457 in revised MS).

Line 352: "The PMF results are very well supported by the fact that anhydrosugars are associated with BB in the Mangshan site." Is this referring to results from a previous study? Please cite it.

Response: We modified the phases. Please see lines 448-449 in the revised MS.

Line 410: "Our results also denote that secondary production of OC and WSOC from BB-derived organic precursors was crucial during nighttime at the Mangshan site." What evidence shows this? And do you mean that organic compounds went through chemical changes to form aerosol OC and WSOC, or simply that organics produced during biomass burning were incorporated into aerosol after the burning? In either case, I don't see how we know that.

Response: We deleted the sentence.

Anonymous Referee #2

Verma et al. discuss observations of sugars found in the aerosol-phase collected for 1 month in a forested site north of Beijing. The aerosol were collected onto filters and analyzed for the sugars. The authors then describe the pattern of the various sugars throughout the study period and speculate the sources via differences in day- and night-time mass concentration, wind patterns, and PMF. They discuss 5 potential sources, including biomass burning, vegetation, microbial and soil dust, pollen, and fungal.

The results presented here may be of interest to the audience and its scope generally fits within a measurement report. However, along with the concerns discussed by Reviewer #1, the authors need to address the comments and concerns presented below prior to consideration for publication in ACP.

Major: (1) Statistics: Throughout the text, the authors state that the results are statistically different. However, conducting the t-test with the mean and standard deviation values listed in the table, majority of the observations are statistically similar at the 95% confidence interval and not statistically different. The lack of statistical difference in the observations makes many of the statements the authors use to differentiate day/night and thus sources less substantiated. Further, the correlations shown by the authors in Fig. 5 have very low R values (as stated throughout the text) and suggest that many of the correlations only explain 50% or less of the mass concentration.

Response: We found a positive correlation between sucrose and ambient temperature ($r = 0.52$), sucrose and solar radiation ($r = 0.55$), mannitol and RH ($r = 0.57$), trehalose and arabinol ($r=0.58$), trehalose and mannitol ($r=0.58$), with significance levels <0.05 . Therefore we mentioned those positive linear correlation values in the text and discussed accordingly.

(2) Contextualization of results: I agree with Reviewer #1 that the listing of numbers from prior results makes it difficult to understand the conclusions in each section and the whole paper. Further, as highlighted with point (1) above, the data not being statistically different makes sections 3.1.1 thru 3.1.3 very long and repetitive. Also, the listing of numbers from prior studies to ascribe sources for the sugars makes the source apportionment very uncertain. This is also relevant for Section 3.5, where they found no differences in the levoglucosan/mannosan ratio and spend 1.5 pages on this. If this is important, it could be summarized in one paragraph at most.

Response: We deleted repeated sentences, several comparisons with previous studies and significantly modified section 3.1.1. to 3.1.3. We also shorten section 3.5. Please see section 3.1.1 – 3.1.3 and 3.5 in the revised MS.

(3) Methods: Reviewer #1 highlighted many of the methods that should be discussed in more detail. Further, PMF needs to be described in more detail to understand how the 5 results were determined (e.g., how many solutions were there allowed to be, how did the time series look, were the results compared against and investigated against external variables, etc.). Also, agree with Reviewer #1 in how were WSOC, OC, Ca²⁺, etc determined.

Response: Authors thank the reviewer's valuable comments and suggestions. We make significant changes in the manuscript especially in the section 2 (Materials and methods), and section 3 (Results and discussions). Please see the revised MS.

We added information about the PMF analysis in the revised MS. Please see lines 187 – 201, 389 – 407, 418 – 429, 442 – 445, 453 – 457, 460 – 462 in the revised MS.

We added Figure S-1 and S-2 as supporting information in the revised MS.

Figure S-1. The scatter plots between observed (input data) and predicted (modeled data) concentrations show statistical parameter (coefficient of determination (r), Intercept, and Slope) with linear equation of individual sugar compounds. (A blue 1:1 line is provided on this plot for reference (a perfect fit would line up exactly on this line), and the regression line is shown as a dotted red line).

Figure S-2. The time series plots between observed (input data) and predicted (modeled data) concentrations of individual sugar compounds. Blue and red lines are shown for observed (input data) and predicted (modeled data) concentrations, respectively.

(4) PMF: I think this is the more interesting and compelling part of the paper. I highly recommend the authors spend more time expanding on this section while reducing the discussion in the other sections. As highlighted above, there are statistical concerns, thus shortening them while increasing the discussion about PMF, which had lower statistical concern.

Response: We added more sentences in the PMF analysis section. Please see line 386 – 468 in the revised MS.

(5) Figures: The x-axis/date is very hard to read in all figures. It is unclear what the values are shown in different colors in Fig. 3.

Response: We separated Figure 3 as Figures 4a-c, 5a-d and 6a-c. We also modified x-axis in the manuscript and changes are added in the figure captions.

Minor: Please review the grammar throughout the paper, as highlighted by Reviewer #1.

Response: According to the reviewer's suggestion, whole manuscript is properly checked for the grammatical mistakes.

Fig. 2.

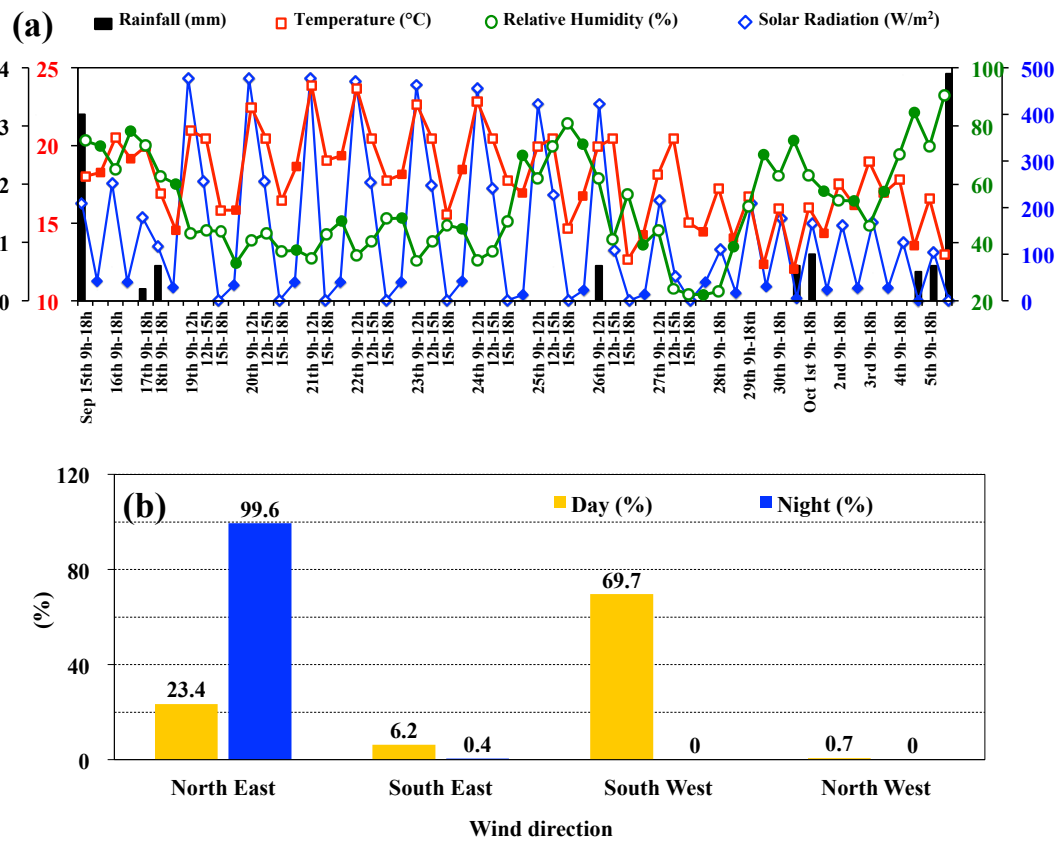


Fig. 4.

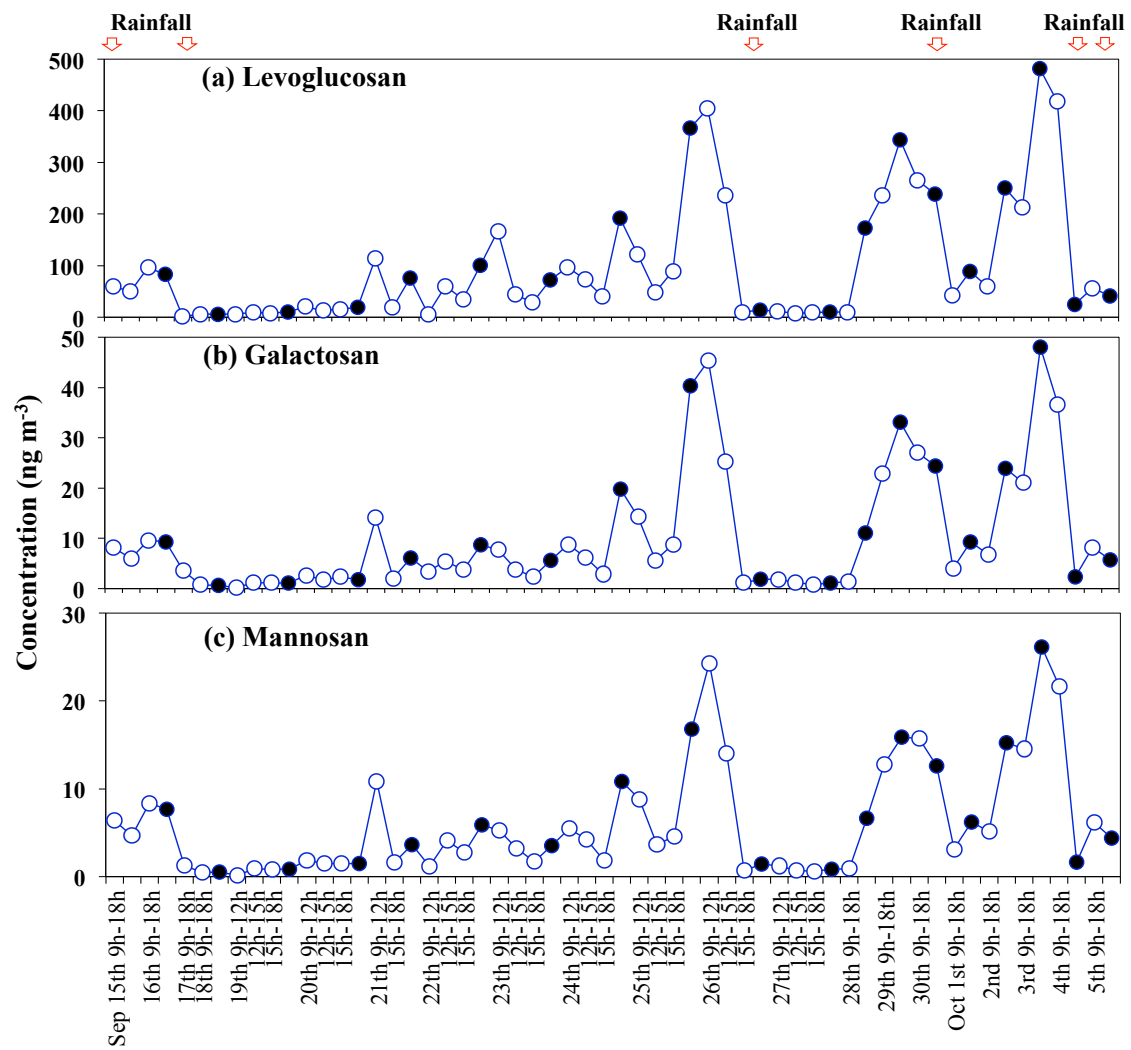


Fig. 5.

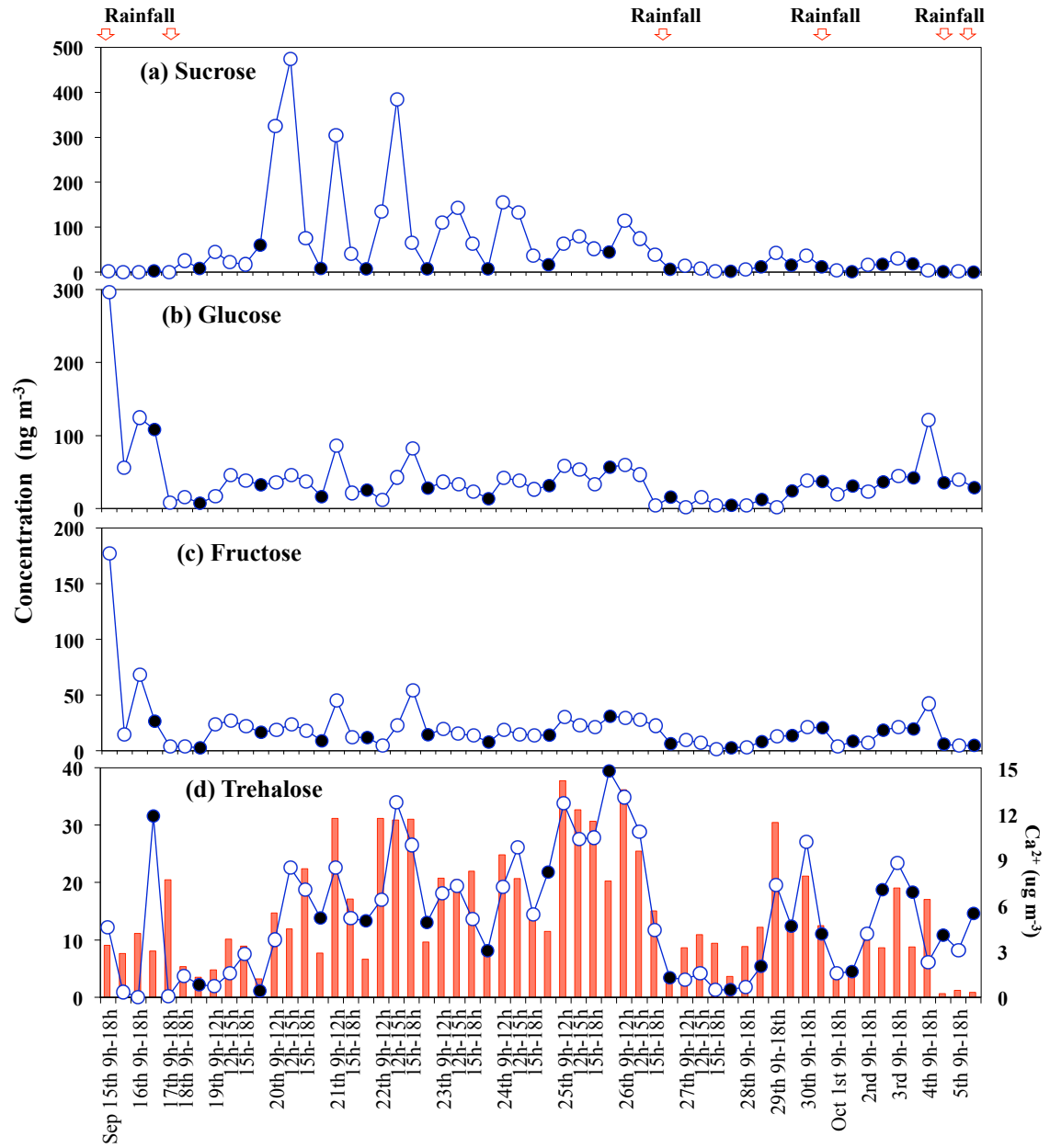


Fig. 6.

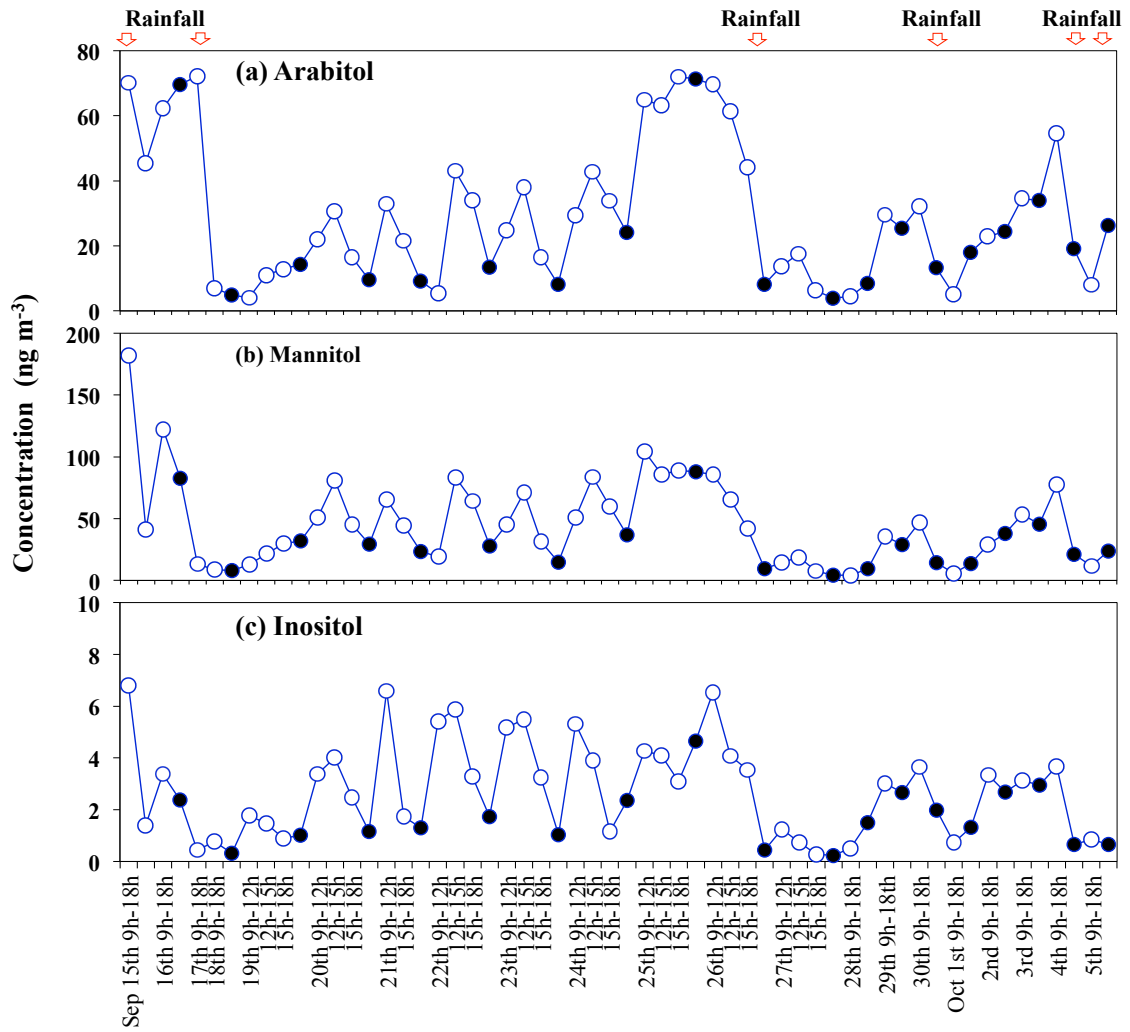


Table 2. Statistical summary of correlations among the chemical species and meteorological variables in aerosol samples collected at a forest site in northern Japan.

Linear regression	Correlation coefficient	p value	Significance of correlation at P value < 0.05
Overall (n = 58)			
Levoglucon vs. Galactosan	0.98	< 0.05	Significant
Levoglucon vs. Mannosan	0.97	< 0.05	Significant
Mannosan vs. Galactosan	0.98	< 0.05	Significant
Sucrose vs. Temperature	0.52	< 0.05	Significant
Sucrose vs. Solar radiation	0.55	< 0.05	Significant
Arabitol vs. Mannitol	0.81	< 0.05	Significant
Arabitol vs. RH	0.69	< 0.05	Significant
Mannitol vs. RH	0.57	< 0.05	Significant
Glucose vs. Fructose	0.94	< 0.05	Significant
Trehalose vs. Arabitol	0.58	< 0.05	Significant
Trehalose vs. Mannitol	0.58	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.70	< 0.05	Significant
Daytime (n = 38)			
Sucrose vs. Ca ²⁺	0.32	> 0.05	Not significant
Glucose vs. Ca ²⁺	0.02	> 0.05	Not significant
Trehalose vs. Arabitol	0.49	< 0.05	Significant
Trehalose vs. Mannitol	0.51	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.81	< 0.05	Significant
Fructose vs. Mannitol	0.79	< 0.05	Significant
Levoglucon vs. OC	0.45	< 0.05	Significant
Levoglucon vs. WSOC	0.40	< 0.05	Significant
Nighttime (n = 20)			
Sucrose vs. Ca ²⁺	0.37	> 0.05	Not significant
Glucose vs. Ca ²⁺	0.27	> 0.05	Not significant
Trehalose vs. Arabitol	0.76	< 0.05	Significant
Trehalose vs. Mannitol	0.85	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.61	< 0.05	Significant
Fructose vs. Mannitol	0.86	< 0.05	Significant
Levoglucon vs. OC	0.81	< 0.05	Significant
Levoglucon vs. WSOC	0.70	< 0.05	Significant

Supporting Information

Table S-1 Sampling informations.

Sampler:High Volume Air Sampler (Kimoto-AS810A)						
Sample ID	Start Time	Finish Time	Center time	Total Time (Hours)	Total Air V (m ³)	Filter Total (cm ²)
CHN-131	2007/9/15 9:08	2007/9/15 17:57	15/09/07 13:32	8:49	533.25	405.3
CHN-132	15/09/07 18:01	16/09/07 8:25	16/09/07 1:13	14:24	873.09	405.3
CHN-133	16/09/07 8:29	16/09/07 17:53	16/09/07 13:11	9:24	563.03	405.3
CHN-134	16/09/07 17:57	17/09/07 8:56	17/09/07 1:26	14:59	907.48	405.3
CHN-135	17/09/07 9:00	2007/9/17 17:38	17/09/07 13:19	8:38	515.44	405.3
CHN-136	18/09/07 9:58	18/09/07 18:03	18/09/07 14:00	8:05	490.5	405.3
CHN-137	2007/9/18 18:06	19/09/07 8:54	19/09/07 1:30	14:48	905.69	405.3
CHN-139	19/09/07 9:04	19/09/07 12:00	19/09/07 10:32	2:56	177.5	405.3
CHN-140	19/09/07 12:04	19/09/07 15:01	19/09/07 13:32	2:57	180.14	405.3
CHN-141	19/09/07 15:05	19/09/07 18:01	19/09/07 16:33	2:56	179.44	405.3
CHN-142	19/09/07 18:04	20/09/07 9:01	20/09/07 1:32	14:57	920.18	405.3
CHN-143	20/09/07 9:05	20/09/07 12:02	20/09/07 10:33	2:57	172.93	405.3
CHN-144	2007/9/20 12:06	2007/9/20 15:01	20/09/07 13:33	2:55	177.25	405.3
CHN-145	20/09/07 15:04	20/09/07 17:59	20/09/07 16:31	2:55	178.6	405.3
CHN-146	20/09/07 18:02	21/09/07 8:57	21/09/07 1:29	14:55	919.91	405.3
CHN-147	21/09/07 9:00	21/09/07 12:02	21/09/07 10:31	3:02	180.96	405.3
CHN-149	21/09/07 15:06	21/09/07 17:59	21/09/07 16:32	2:53	177.52	405.3
CHN-150	21/09/07 18:02	22/09/07 9:00	22/09/07 1:31	14:58	915.67	405.3
CHN-151	22/09/07 9:02	22/09/07 11:59	22/09/07 10:30	2:57	178.6	405.3
CHN-152	22/09/07 12:02	22/09/07 14:59	22/09/07 13:30	2:57	177.73	405.3
CHN-153	22/09/07 15:02	22/09/07 17:58	22/09/07 16:30	2:56	180.68	405.3
CHN-154	22/09/07 18:01	23/09/07 8:56	23/09/07 1:28	14:55	911.27	405.3
CHN-155	23/09/07 9:00	23/09/07 11:57	23/09/07 10:28	2:57	177.82	405.3
CHN-156	2007/9/23 12:00	23/09/07 14:59	23/09/07 13:29	2:59	179.3	405.3
CHN-157	23/09/07 15:02	23/09/07 17:59	23/09/07 16:30	2:57	180.75	405.3
CHN-158	23/09/07 18:03	24/09/07 8:55	24/09/07 1:29	14:52	918.89	405.3
CHN-159	24/09/07 8:59	24/09/07 11:56	24/09/07 10:27	2:57	178.16	405.3
CHN-160	24/09/07 11:59	24/09/07 14:57	24/09/07 13:28	2:58	180.62	405.3
CHN-161	2007/9/24 15:00	24/09/07 17:56	24/09/07 16:28	2:56	179.84	405.3
CHN-163	24/09/07 18:11	25/09/07 9:01	25/09/07 1:36	14:50	899.43	405.3
CHN-164	25/09/07 9:04	25/09/07 12:07	25/09/07 10:35	3:03	184.92	405.3
CHN-165	25/09/07 12:09	25/09/07 14:58	25/09/07 13:33	2:49	169.63	405.3
CHN-166	25/09/07 15:01	25/09/07 17:58	25/09/07 16:29	2:57	180.45	405.3
CHN-167	25/09/07 18:01	26/09/07 9:01	26/09/07 1:31	15:00	912.34	405.3
CHN-168	2007/9/26 9:04	26/09/07 11:59	26/09/07 10:31	2:55	176.33	405.3
CHN-169	26/09/07 12:02	26/09/07 14:58	26/09/07 13:30	2:56	176.46	405.3
CHN-170	26/09/07 15:01	26/09/07 17:58	26/09/07 16:29	2:57	181.09	405.3
CHN-171	26/09/07 18:01	27/09/07 8:56	27/09/07 1:28	14:55	916.56	405.3
CHN-172	27/09/07 8:59	27/09/07 12:00	27/09/07 10:29	3:01	183.05	405.3
CHN-173	27/09/07 12:03	27/09/07 15:01	27/09/07 13:32	2:58	180.88	405.3
CHN-174	27/09/07 15:04	27/09/07 17:59	27/09/07 16:31	2:55	179.04	405.3
CHN-175	27/09/07 18:02	28/09/07 9:15	28/09/07 1:38	15:13	936.45	405.3
CHN-176	28/09/07 9:19	28/09/07 17:57	28/09/07 13:38	8:38	519.1	405.3
CHN-178	28/09/07 18:10	29/09/07 8:59	29/09/07 1:34	14:49	907.18	405.3
CHN-179	29/09/07 9:01	29/09/07 17:59	29/09/07 13:30	8:58	539.66	405.3
CHN-180	29/09/07 18:01	30/09/07 8:57	30/09/07 1:29	14:56	922.34	405.3
CHN-181	30/09/07 9:00	30/09/07 17:57	30/09/07 13:28	8:57	531.89	405.3
CHN-182	30/09/07 18:00	01/10/07 8:55	01/10/07 1:27	14:55	896.33	405.3
CHN-183	01/10/07 8:58	01/10/07 17:54	01/10/07 13:26	8:56	538.04	405.3
CHN-184	01/10/07 17:57	02/10/07 8:56	02/10/07 1:26	14:59	926.33	405.3
CHN-185	02/10/07 8:59	02/10/07 17:56	02/10/07 13:27	8:57	531.43	405.3
CHN-186	02/10/07 17:59	03/10/07 8:59	03/10/07 1:29	15:00	913.14	405.3
CHN-187	03/10/07 9:01	03/10/07 17:52	03/10/07 13:26	8:51	523.19	405.3
CHN-188	03/10/07 17:54	04/10/07 8:57	04/10/07 1:25	15:03	919.84	405.3
CHN-189	04/10/07 9:00	04/10/07 17:56	04/10/07 13:28	8:56	535.79	405.3
CHN-190	04/10/07 18:00	05/10/07 8:59	05/10/07 1:29	14:59	918.46	405.3
CHN-191	05/10/07 9:04	05/10/07 17:56	05/10/07 13:30	8:52	535.52	405.3
CHN-192	05/10/07 17:59	06/10/07 8:53	06/10/07 1:26	14:54	906.91	405.3
CHN-138	19/09/07 9:00	blank	19/09/07 9:00			405.3
CHN-148	21/09/07 13:52	blank	21/09/07 13:52			405.3
CHN-162	24/09/07 17:58	blank	24/09/07 17:58			405.3
CHN-177	28/09/07 18:00	blank	28/09/07 18:00			405.3

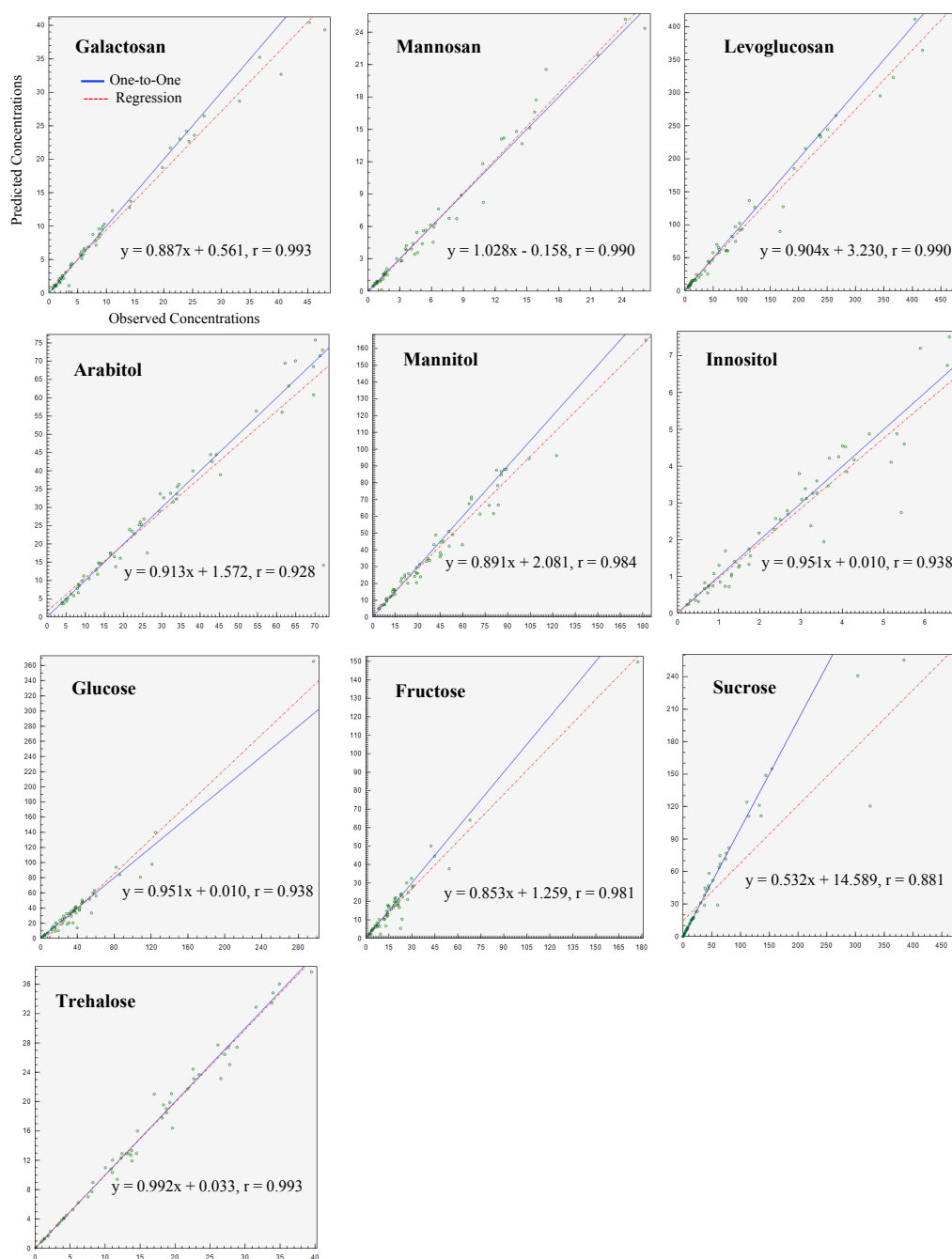


Figure S-1. The Scatter plots between observed (input data) and predicted (modeled data) concentrations show statistical parameter (coefficient of determination (r), Intercept, and Slope) with linear equation of individual sugar compounds. (A blue 1:1 line is provided on this plot for reference (a perfect fit would line up exactly on this line), and the regression line is shown as a dotted red line).

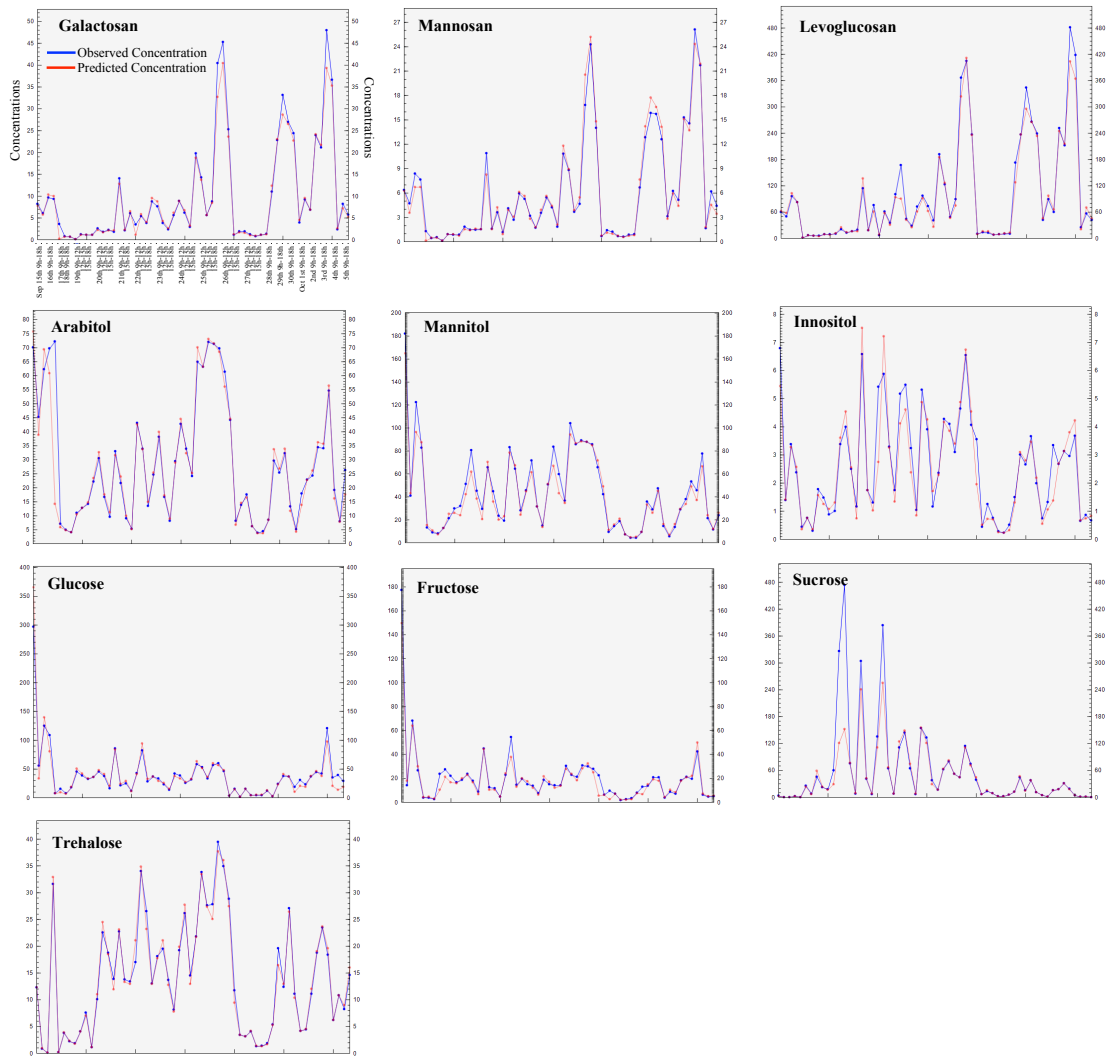


Figure S-2. The time series plots between observed (input data) and predicted (modeled data) concentrations of individual sugar compounds. (A blue line and redline shown as observed (input data) and predicted (modeled data) concentrations, respectively).

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List of all relevant changes

1. We add three key points in the revised MS.
2. We added new paragraph in the introduction section.
3. The materials and methods sections were modified by adding new information and subsection.
4. We modified the discussion section by re-organizing paragraphs.
5. We added new figure as figure 2b and moved figure 4 to Figure 3.
6. We split figure 3 into three Figures as Figure 4 (anhydrosugars), Figure 5 (primary sugars), and Figure 6 (sugar alcohols). We added the modified version of Figure 9. We changed Figure 5 to table 2.
7. We added table S-1 (sampling information), Figure S-1 (PMF results), and Figure S-2 (PMF results) as supplementary material.
8. We went through the paper and corrected the typos. We also rewrote sentences that were difficult to read.

1 **Measurement report: Diurnal and temporal variations of sugar compounds in suburban**
2 **aerosols from the northern vicinity of Beijing, China: An influence of biogenic and**
3 **anthropogenic sources**

4

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18 **Key points:**

19

20 1. Autumn time observations of sugar compounds (SCs) in the northern vicinity of Beijing, China.

21 2. Influence of natural biogenic emissions on SCs from forest area.

22 3. Influence of anthropogenic and bioaerosol on SCs from the Beijing area.

23 4. Biomass burning is a significant contributor to SCs.

24 5. Biogenic and fungal-microbial emissions are essential sources for mannitol and arabitol.

25

26

27

28 **Abstract**

29 Sugar compounds (SCs) are major water-soluble constituents in atmospheric aerosols. In
30 this study, we investigated their molecular compositions and abundances in the northern receptor
31 site (Mangshan) of Beijing, China, to better understand the contributions from biogenic and
32 anthropogenic sources using a gas chromatography–mass spectrometry technique. **The sampling
33 site receives anthropogenic air mass transported from Beijing by southerly winds, while northerly
34 winds transport relatively clean air mass from the forest areas.** Day- and nighttime variations were
35 analyzed for anhydrosugars, primary sugars, and sugar alcohols in autumn 2007. We found that
36 biomass burning (BB) tracers were more abundant in nighttime than daytime, while other SCs
37 showed different diurnal variations. Levoglucosan was found as a dominant sugar among the SCs
38 observed, indicating an intense influence of local BB for cooking and space heating at the
39 surroundings of the Mangshan site. **The high levels of arabitol and mannitol in daytime suggest a
40 significant contribution of locally emitted fungal spores and long-range transported bioaerosols
41 from the Beijing area.** The plant emissions from Mangshan forest park significantly control the
42 diurnal variations of glucose, fructose, and mannitol. **The meteorological parameters (relative
43 humidity, temperature, and rainfall) significantly affect the concentrations and diurnal variations
44 of SCs.** Sucrose (pollen tracer) showed a clear diurnal variation, peaking in the daytime due to
45 higher ambient temperature and wind speed, which influences the pollen release from the forest
46 plants. We found the contribution of trehalose from soil dust in daytime, while microbial and
47 fungal spores were responsible for nighttime. **Anhydrosugar and primary sugars are prime carbon
48 sources of the Mangshan aerosols. The high ratios of levoglucosan in organic carbon and water
49 soluble organic carbon in nighttime suggest a significant contribution of BB to organic aerosols at
50 night. Levoglucosan/mannosan ratios demonstrate that low temperature burning of hardwood is
51 dominant in Mangshan.** The positive matrix factorization analysis concluded that forest
52 vegetation, fungal species, and local BB are the significant sources of SCs.

53 **Keywords:** Anthropogenic bioaerosols, biomass burning, pollen tracer, fungal tracers, soil dust,
54 and microbial tracers

55

56 **1. Introduction**

57 Increased economic growth and massive consumption of fossil fuels from industries emit
58 anthropogenic gases, aerosols, and biomass burning (BB) products cause severe air pollution in
59 East Asian countries (Lelieveld et al., 2015; Lin et al., 2014; Kawamura et al., 2013; Li et al.,
60 2010; Sun et al., 2016). On a global scale, one-fourth of anthropogenic aerosols is contributed by
61 China, and approximately 70% of which emitted from coal burning (Streets et al., 2003). Beijing
62 is situated in the northern part of China, with a 20 million people and 5 million motor vehicles.
63 Beijing is one of the largest polluted cities in East Asia; its air quality deteriorates seriously due to
64 massive emissions of organic aerosols (OAs) from vehicles and industries (Cao et al., 2014; Qiao
65 et al., 2018; Tao et al., 2017; Wei et al., 2018; Yu et al., 2013). OAs are comprised with a complex
66 mixture of diverse molecules (Xu et al., 2011). They play essential roles in global climate changes
67 via the modification of radiative forcing and cause a serious negative impact on human health
68 (Fuzzi et al., 2007). OAs contain various water-soluble organic compounds, which can act as
69 cloud condensation nuclei (CCN) (Kanakidou et al., 2005).

70 BB is essentially a primary source of OAs, controlling the air quality levels and affecting
71 the earth's radiative forcing by scattering or absorbing incident solar radiation (Deshmukh et al.,
72 2019a; Kanakidou et al., 2005; Kanaya et al. 2013; Streets et al., 2003; Sullivan et al., 2008).
73 There are several kinds of BB, including industrial biofuel burning, open field burning (fires of the
74 forest, peatlands, and agricultural wastes), and domestic BB burning for house heating and
75 cooking, which emits BB products into the atmosphere (Akagi et al., 2011). The BB aerosols are
76 subjected to long-range atmospheric transport once they are emitted into the atmosphere (Verma et
77 al., 2015). Levoglucosan (1,6-anhydro- β -D-glucopyranose) is a pyrolysis product of cellulose and
78 hemicellulose, which is generally found as major organic constituents in the BB-influenced
79 aerosols (Simoneit et al., 1999; 2002). Levoglucosan have been reported as a specific tracer for
80 BB aerosols (Engling et al., 2009).

81 Sugar compounds (SCs) are ubiquitous in the atmosphere from different geographical
82 locations, including urban, forest, marine, and polar regions (Burshtein et al., 2011; Fu et al.,
83 2010; Wan et al., 2017). SCs are emitted from algae, microbes, pollen, suspended soil particle, and
84 associated biota into the atmosphere by various processes, and thus they are termed as primary
85 biological aerosol particles (PBAPs) (Carvalho et al., 2003; Despres et al., 2012; Elbert et al.,
86 2007). The detailed study of bio-aerosols has been emphasized in the past decades due to the
87 global impact of microbes and fungi because they can travel long distances from the source
88 regions by winds (Burshtein et al., 2011; Brown and Hovmoller, 2002; Yamaguchi et al., 2012).

89 Fungi are essential microbes in the ecosystem, which discharge spores of 8-186 Tg yr⁻¹ into the
90 atmospheric environment (Elbert et al., 2007; Heald and Spracklen, 2009). Sugar alcohols like
91 arabitol and mannitol are enriched in fungal spores; thus, they are considered as specific tracers
92 (Bauer et al., 2008).

93 Devis et al. (1988, 1990) reported that mannitol was also found in about 70 different
94 higher plant families. Loescher et al. (1992) reported that mannitol is an important photosynthetic
95 product converted by biosynthesis in plants. Keller and Matile (1989) also found the arabitol and
96 mannitol during the increased photosynthesis in growing vegetation. Pollens are the largest
97 particles that could contribute up to 65% of the PBAPs, which are the significant sources for
98 sucrose and fructose in the forest aerosols (Manninen et al., 2014; Pacini, 2000). Higher plants
99 synthesize primary sugars (glucose, fructose, and sucrose) during photosynthesis, which are
100 circulated by phloem to accumulate in root cells and to develop plant sections (Jaenicke, 2005; Jia
101 et al., 2010; Pacini, 2000). Cowie et al. (1984) also reported various sugars in terrestrial plant
102 fruits, flowers, and plant tissues. Bielecki (1995) reported that glucose, fructose, and sucrose are
103 well-known components of microbes and invertebrates. The plant debris, as well as lichens,
104 invertebrates, and soil dust, are also recognized as possible sources of primary sugars in the
105 atmosphere (Medeiros et al., 2006; Rogge et al., 2007; Simoneit et al., 2004).

106 Previous studies analyzed aerosol samples for SCs and discussed several factors to control
107 their local and global atmospheric levels. Recently, Xu et al. (2020) examined the seasonal
108 molecular distributions of primary biological aerosols and BB aerosol samples collected from
109 urban Beijing. They reported a high level of arabitol, mannitol, sucrose, glucose, and fructose in
110 the vegetation-growing season. Kang et al. (2018) also reported higher concentrations of sugars in
111 the urban aerosols from Beijing. They suggested a large contribution of coal combustion and
112 agriculture residue burning under stable meteorological conditions in winter and spring. Verma et
113 al. (2015, 2018) reported that the atmospheric circulations and long-range transport of organic-
114 /bio-aerosols from East Asia significantly control the levels and compositions of SCs over the
115 western North Pacific. The above studies discussed the several factors that affect the
116 concentrations of SCs in the aerosol samples collected from urban and remote areas.

117 In this study, we conducted analyses of SCs in the aerosol samples collected from the
118 northern vicinity of Beijing City in 2007. Here, we present comprehensive data sets of
119 anhydrosugars, primary sugars, and sugar alcohols in the suburban aerosol samples and their
120 diurnal variations to explain the source variance following the wind patterns in the day- and
121 nighttime. The positive matrix factorization (PMF) has been applied to clarify the different

122 sources of measured SCs in the aerosol. We present the influence of local meteorology of
123 sampling site and atmospheric transport from Beijing by southerly winds and Mangshan National
124 Forest Park by northerly winds on the molecular distributions of SCs. Using the mass
125 concentration ratio of levoglucosan to mannosan, we explain the relative contribution of hard and
126 softwood burning to the air quality of Mangshan. This study also discussed carbon contributions
127 of SCs and BB measured in the Mangshan aerosol samples from different sources.

128 **2. Materials and Methods**

129 **2.1. Site description and aerosol sample collection**

130 The sampling site (Mangshan: 40.28 N, 116.26 E) is located 40 km north of Beijing. A
131 detailed description of the sampling site is given in He et al. (2014, 2015). Briefly, Mangshan is
132 surrounded by urban areas in the south and forest areas with the national park in the north (Fig. 1).
133 The ambient temperature was higher in daytime (23.9°C) than nighttime (12.1°C), with an average
134 of 17.8°C during the campaign. The relative humidity (RH) varied significantly from 22.1% to
135 90.5%, with an average of 51.7% during the study period. The rainfall was observed at midnight
136 on 15th September, the morning of 17th to evening 18th September, the night of 26th September,
137 and light rain lasted from 4th October to the end of the campaign (Fig. 2a). Interestingly, the
138 sampling site is characterized by a specific wind pattern, i.e., southwest wind (69.9%) prevailed,
139 followed by northeast wind (23.4%) and southeast wind (6.2%) during the daytime (Fig. 2a). The
140 northeast wind (99.5%) was dominated at night, which is consistent with the air mass back
141 trajectories (He et al., 2014) (Fig. 2b). The daytime wind from the southwest direction passed over
142 Beijing, delivering anthropogenic air mass to the Mangshan site.

143 Detailed descriptions of the total suspended particulate (TSP) samples collected at
144 Mangshan are given in He et al. (2014, 2015). Briefly, The aerosol samples were collected near
145 the entrance of Mangshan National Forest Park. The elevation of the sampling location is 187 m
146 above sea level. TSP (size <100 μm) samples were collected on pre-combusted (450°C for 6 h)
147 quartz fiber filters (Pallflex 2500QAT-UP, 20 cm \times 25cm) from 15th September to 5th October
148 2007. A high-volume air sampler (Kimoto-AS810A) at a flow rate of 1.13-1.17 $\text{m}^3 \text{min}^{-1}$ was used
149 to collect the TSP samples. After sample collection, the individual filters were placed in pre-
150 combusted glass jars with Teflon-lined screw caps and stored in a dark, cold room at -20°C to
151 prevent microbial activity and loss of semi-volatile organic compounds from the samples. In this
152 study, a total of 58 filter samples were analyzed. We collected 3h daytime (from 9 to 12, 12 to 15,
153 15 to 18 h) (n=26), 9h daytime (from 9 to 18 h) (n=12), and 15h nighttime (from 18 to 9 h) (n=20)

154 samples together with four field blanks. Table S1 shows the details of aerosol sample collection in
155 the Mangshan site.

156 **2.2. Extraction and derivatization of samples**

157 A total of 58 aerosol samples were analyzed for anhydrosugars, primary sugars, and sugar
158 alcohols (Table 1). The sample filters (approximately 21 cm²) were extracted with a
159 dichloromethane and methanol mixture (2:1) under ultrasonication. Pasteur pipettes packed with
160 pre-combusted quartz wool were used to filter the extracts to remove filter debris. After filtration,
161 the extracts were concentrated in a rotary evaporator under vacuum and dried by nitrogen
162 blowdown. The extracts were reacted with 60 µl of N,O-bis-(trimethylsilyl)trifluoroacetamide
163 (BSTFA) with 1% trimethylsilyl (TMS) chloride in the presence of 10 µL of pyridine at 70°C for
164 three hours to derivatize hydroxyl (OH) and carboxyl (COOH) groups into corresponding
165 trimethylsilyl (TMS) ethers and esters, respectively. After the reaction, n-hexane was used for
166 dilution, and C₁₃ n-alkane was added as an internal standard before GC-MS analysis.

167 **2.3. Gas chromatography-mass spectrometry determination of sugar compounds (SCs)**

168 Details of GC-MS operation and identification of SCs are described in Verma et al. (2015,
169 2018). Briefly, GC-MS analyses were performed on Agilent model 6890 gas chromatograph (GC)
170 combined with an Agilent model 5973 mass selective detector (MSD) to determine SCs. The mass
171 spectrometer was operated in the electron ionization (EI) mode at 70 eV with a scan range of *m/z*
172 40–650. The GC separation was achieved on a DB-5MS fused silica capillary column (30 m ×
173 0.25 mm in diameter, 0.25 µm film thickness) and a split/splitless injector. The GC oven
174 temperature was programmed to maintain at 50°C for 2 min and then to increase from 50 to 120°C
175 at a rate of 15°C min⁻¹, then from 120 to 305°C at a rate of 5°C min⁻¹. The final isotherm holds at
176 305°C for 15 min. Helium was used as the carrier gas at a flow rate of 1.0 mL min⁻¹. The sample
177 was injected on a splitless mode at 280°C injector temperature. GC-MS data were acquired and
178 processed with Agilent GC/MSD ChemStation software.

179 The individual compounds (TMS derivatives) were identified by comparing the relative
180 response factors determined by the injection of authentic standards and those reported in the
181 literature and library texts (Claeys et al., 2004). Fragment ions of sugar compounds at 217 and 204
182 were used for quantifications. Total ten sugar compounds, including three anhydrosugars
183 (levoglucosan, galactosan, mannosan), four primary sugars (glucose, fructose, sucrose, trehalose
184 and xylose) and three sugar alcohols (arabitol, mannitol, and inositol), were detected in the

185 Mangshan aerosols. Field blanks were treated as a real sample and analyzed by the procedure used
186 for the real samples. Recoveries for SCs were better than 85% as obtained by the standards spiked
187 to precombusted quartz filter followed by extraction and derivatization. Based on the duplicate
188 analysis, the analytical errors in the concentrations of the detected compounds were obtained to be
189 within 10%. The detection limits of SCs corresponds to ambient concentrations of 150-620 pg μL^{-1}
190 ¹, which corresponds to ambient concentrations of 15-70 pg m^{-3} under a typical sampling volume
191 of 900 m^3 . The data of water-soluble organic carbon (WSOC), organic carbon (OC), and inorganic
192 ions (Ca^{2+}) were reported in He et al. (2015).

193 **2.4. Positive Matrix Factorization (PMF) Analysis**

194 Positive matrix factorization (PMF) is a powerful statistical tool for resolving the potential
195 sources contributing to atmospheric particles (Paatero and Tapper, 1994). The measured ambient
196 concentrations and method detection limits (MDLs) of SCs were used to calculate the
197 uncertainties. The measured concentrations of SCs below or equal to the MDLs were replaced by
198 half of the MDL, and associated uncertainties were set at 5/6 of the MDL [(5/6) \times MDL] values of
199 each sample. The geometric mean concentrations were used for missing concentrations, and the
200 uncertainty of the concentrations greater than the MDL was calculated based on the following
201 equation:

$$202 \text{ Uncertainty} = \sqrt{(\text{error fraction} \times \text{concentration})^2 + (0.5 \times \text{MDL})^2}$$

203 The error fraction is a user-provided estimation of the analytical uncertainty of the
204 measured concentration or flux. For example, Han et al. (2017) used an error fraction of 0.2-0.3
205 for organics and 0.2 for all the species. In this work, the error fraction was set to be 0.3 for all
206 species. Paatero et al. (2002) and Zhou et al. (2004) reported detailed discussions of the
207 determination and application of PMF analysis.

208 **3. Results and Discussion**

209 **3.1. Ambient concentrations and diurnal variations of SCs**

210 We detected a total of ten SCs, including three anhydrosugars, four primary sugars, and three
211 sugar alcohols in the Mangshan aerosol samples. Figure 3a-c showed the temporal variations and
212 Table 1 showed minimum, maximum, and average concentrations of anhydrosugars, primary
213 sugars, and sugar alcohols with a standard deviation. The overall concentrations of SCs varied
214 from 30.8–875 ng m^{-3} (avg. 325 ng m^{-3}), showed insignificant diurnal variations. Interestingly,
215 higher average concentrations of SCs were reported for the aerosol samples collected from at Mt.

216 Tai (daytime 640 ng m⁻³ and nighttime 799 ng m⁻³) in the North China Plain (Fu et al., 2008) than
217 the Mangshan aerosol. The diurnal concentrations of SCs may be significantly influenced by
218 vegetation and BB activities in the Mangshan site. SCs are significantly contributed by plant
219 fractions and fungus from the forest area (Zhu et al., 2016). The meteorological parameters also
220 affect the concentrations of SCs in the forest site (Miyazaki et al., 2012).

221 In addition, anthropogenic aerosols emitted from urban areas are probably transported to the
222 northern receptor site in daytime by a southerly wind (He et al., 2014; 2015). Therefore, the high
223 levels of SCs in daytime may be related to the transport of organic and bio-aerosols from urban
224 regions. The nighttime, the wind direction is shifted to northerly, delivering comparatively clean
225 air masses from the Mangshan National Forest area to the sampling site. Air mass from the forest
226 may significantly contribute to nighttime SCs in the Mangshan site. The influence of local sources
227 and long-range transported aerosols on the SCs will be discussed in sections 3.1.1 to 3.1.3.

228 3.1.1. Ambient concentrations and diurnal variations of anhydrosugars

229 The average concentrations of anhydrosugars were found 116 ng m⁻³, contributing 31.9%
230 of overall SCs in the Mangshan aerosols (Table 1). Figure 4a-c shows the temporal variations of
231 anhydrosugars. They are more abundant in nighttime (avg. 152 ng m⁻³) than daytime (avg. 97.1 ng
232 m⁻³). Levoglucosan (100 ng m⁻³) is the most abundant anhydrosugar followed by galactosan (10.1
233 ng m⁻³) and mannosan (6.05 ng m⁻³) detected in Mangshan aerosols. Kang et al. (2018) reported
234 high levels of levoglucosan (avg. 110 ng m⁻³) in autumn aerosols from Beijing, China. It is well
235 known that biofuel burning is the common energy source for cooking and house heating in China
236 in winter and autumn (Verma et al., 2015), thus the domestic BB activities in the surroundings of
237 the Mangshan site significantly contribute to the levoglucosan. BB tracers showed significant
238 positive correlations with each other (levoglucosan and galactosan, $r = 0.98$; levoglucosan and
239 mannosan, $r = 0.97$; galactosan and mannosan, $r = 0.98$), suggesting their similar sources in the
240 Mangshan aerosols (Table 2).

241 The levoglucosan concentrations showed significant diurnal variations, which was higher
242 in nighttime (avg. 132 ng m⁻³) than daytime (avg. 83.2 ng m⁻³) (Table 1). A similar diurnal pattern
243 was also found for the concentrations of galactosan and mannosan. The increased concentrations
244 of BB tracers were observed during the periods of lower ambient temperature (Figs. 2a, 4a-c). The
245 higher ambient temperature was recorded in daytime between 09h to 15h during the campaign,
246 associated with declined BB activities. In this sequence, the nighttime samples were collected
247 from 18:00h to 09:00h, including peak hours of BB for domestic purpose. Therefore, it is

248 reasonable to detect higher abundances of BB tracers in the nighttime than daytime. Hence, it is
249 evident that BB activities were increased at night because of cooking and house heating at cool
250 night in autumn. In addition, recent studies reported the widespread BB aerosols in the North
251 China Plain, including megacities such as Beijing, Nanjing, Hebei, and Tianjin (Lelieveld et al.,
252 2015; Kawamura et al., 2013; Li et al., 2010; Sun et al., 2016). Therefore, the atmospheric
253 transport of BB aerosols from the urban area to the Mangshan site by southerly winds cannot be
254 excluded. The diurnal variations of levoglucosan may be significantly influenced by the local BB
255 activities and transported BB aerosols from urban areas, where BB products are generated by
256 brown coal combustion (Yan et al., 2018).

257 3.1.2. Ambient concentrations and diurnal variations of primary sugars

258 The fragment of vascular plants contains primary sugars, including glucose, fructose,
259 sucrose, and trehalose (Medeiros et al., 2006). Primary sugars were found as the most abundant
260 sugars (avg. 133 ng m⁻³), contributing to 41.8% of the total SCs in Mangshan aerosols (Table 1).
261 They showed apparent diurnal variations with daytime high (avg. 166 ng m⁻³) and nighttime low
262 values (avg. 69.4 ng m⁻³) (Figs. 3a-c, 5a-d). Graham et al. (2003) also reported similar diurnal
263 variations of primary sugars for the Amazon forest aerosols. Sucrose was found as dominant
264 primary sugars (avg. 58.5 ng m⁻³), accounting for 44% of measured primary sugars in Mangshan
265 aerosols (Table 1). Pollen was reported as a primary source for sucrose in aerosols collected from
266 a Texas rural site (Jia et al., 2010). Fu et al. (2012) found high sucrose concentrations up to 1390
267 ng m⁻³ in the aerosols from Jeju Island, South Korea. Therefore, the plant materials, including
268 pollen spores from the local vegetation of Mangshan National Forest Park, are likely the primary
269 source of sucrose in the aerosols. Miyazaki et al. (2012) also reported higher sucrose
270 concentrations in the aerosol samples collected from the Hokkaido deciduous forest.

271 We found a significant diurnal variation of sucrose with higher daytime (82.9 ng m⁻³) than
272 nighttime (12.3 ng m⁻³). Meteorological parameters such as temperature, rainfall, wind speed, and
273 solar radiation significantly influence pollen activities and, subsequently, sucrose concentrations
274 (Verma et al., 2018). Interestingly, an elevated peak of sucrose was observed from 12h to 15h with
275 higher ambient temperature. In contrast, lower sucrose concentrations were observed from 15h to
276 9h with lower ambient temperature (Fig. 5a). Daytime increased concentrations of sucrose might
277 be related to the higher daytime ambient temperature, low RH, and high solar radiation (Miyazaki
278 et al., 2012). Taylor et al. (2002) reported the influence of the meteorological conditions, i.e.,
279 strong daytime winds and convective activity, which can result in catapulting of pollen, opening of
280 pollen-laden flower anthers, and causing enhance entrainment and dispersal of the particles into

281 the air. Pacini (2000) reported that higher levels of sucrose in daytime coincide with higher counts
282 of pollen, fern spore, and insect. The positive linear correlations of sucrose with ambient
283 temperature ($r = 0.52$) and solar radiation ($r = 0.55$) further supported the influence of
284 meteorological parameters in the sucrose concentration (Table 2).

285 Five rain events were recorded during the campaign, i.e., 15th, 17th, 18th, and 26th
286 September, and 1st and 5th October (Fig. 2a). Pollens are significantly settled down by wet
287 scavenging during rain events because their sizes are large. A low concentration of sucrose was
288 found from the beginning of sampling to the morning of 20th September and from the afternoon of
289 26th September to the end of the sampling campaign (Fig. 5a). In addition, the increased
290 concentrations of sucrose were found in the aerosol samples collected from 20th to 22nd September,
291 and moderate concentrations were observed after 23rd to the evening of 25th September during
292 non-precipitation events. Consequently, the pollens were significantly scavenged during wet
293 precipitation and washout effect from the atmosphere, resulting in lower sucrose concentrations at
294 the earlier periods, than later periods. In addition, Rogge et al. (2007) reported that surface soil
295 dust and unpaved road dust also contribute sucrose in the atmospheric aerosols. However,
296 insignificant correlations between sucrose and Ca^{2+} (daytime, $r = 0.32$; night time, $r = 0.37$) do not
297 supports soil dust contributions to sucrose in the Mangshan aerosols (Table 2).

298 Glucose was the second dominant primary sugar in the Mangshan aerosols. The average
299 concentrations of glucose and fructose were observed to be 40.0 ng m^{-3} and 20.1 ng m^{-3} ,
300 respectively (Table 1, Fig. 5b). The sampling site is characterized by the dense vegetation in the
301 Mangshan National Forest Park. Therefore, the nectars and fruits of vegetation (Baker et al.,
302 1998), plant debris (Medeiros et al., 2006) and pollens (Fu et al., 2012) in the forest significantly
303 contribute to glucose and fructose. The glucose levels are equivalent to that (50.1 ng m^{-3}) reported
304 from the Howland Experimental Forest site in the USA (Medeiros et al., 2006). Glucose and
305 fructose showed significant diurnal variations, whose concentrations were higher in daytime (44.2
306 ng m^{-3} and 23.9 ng m^{-3} , respectively) than nighttime (32.0 ng m^{-3} and 12.8 ng m^{-3} , respectively)
307 in Mangshan aerosols (Table 1, Figs. 3b, c; 5b, c). This diurnal variation could be involved with
308 emissions of pollens, fern spores, and other giant particles by strong winds (Graham et al., 2003;
309 Pacini, 2000). Similar trends of glucose and fructose were reported in the Amazon forest, being
310 coincided with plant fragments and insects (Graham et al., 2003). The autumn decay of vascular
311 plant leaves in the Mangshan forest may have contributed to the levels of glucose and fructose.

312 Although, the daytime southerly winds deliver anthropogenic air masses from megacities
313 to the sampling site. The daytime winds from the northeast direction (23.4%) also carry air masses

314 from the forest region, transporting primary sugars to the Mangshan site. However, 99.5% of the
315 nighttime hours, the wind is shifted to northeasterly, i.e., in forest region (He et al., 2015), but the
316 emissions of primary sugars at night in the form of plant fragments are lower than in daytime.
317 Because the daytime ambient temperature and solar radiations significantly induce the emissions
318 of sugar compounds in the forest site (Miyazaki et al., 2012). Therefore, low glucose and fructose
319 levels were found at nighttime than daytime aerosols at the Mangshan site (Table 1, Fig. 3).
320 Previous studies have reported lichens (Dahlman et al., 2003) and soil dust (Nolte et al., 2001;
321 Rogge et al., 2007) as significant sources of both primary sugars. The concentration of glucose
322 was insignificantly correlated with soil tracer (Ca^{2+}) in day ($r = 0.02$) and nighttime ($r = 0.27$),
323 denying their soil dust contributions in Mangshan aerosol samples.

324 Trehalose in the environment is significantly controlled by the activities of bacteria, fungi,
325 yeast, algae, invertebrates, and plant species, as well as suspended soil particles (Medeiros et al.,
326 2006, Rogge et al., 2007). The average concentration of trehalose was found 14.3 ng m^{-3} (Table 1,
327 Fig. 5d). Yttri et al. (2007) reported higher trehalose concentrations in the aerosol samples
328 collected from urban (29 ng m^{-3}) and suburban (27 ng m^{-3}) than rural (3.8 ng m^{-3}) areas in
329 Norway. The above results emphasize that fungi and microbes associated with anthropogenic and
330 bioaerosols, emitted in the urban and suburban areas, might be responsible for the trehalose
331 concentration in aerosol samples (Verma et al., 2018). Trehalose showed insignificant diurnal
332 variation, whose day and night concentrations were observed 15.3 ng m^{-3} and 12.3 ng m^{-3} ,
333 respectively, indicating its different emission sources in day and night for Mangshan aerosols (Fig.
334 3b, c; 5d).

335 The southerly winds might transport fungi and microbes associated with bioaerosols, eject
336 spores under favorable meteorological conditions (high RH and low temperature) (Jones and
337 Mitchell et al., 1996). Several microbes and fungi discharge spores at nighttime due to high RH
338 conditions (Ibrahim et al., 2011; Kim and Xiao, 2005; Malik and Singh, 2004; Sharma and Razak,
339 2003). Interestingly, trehalose is more significantly correlated with arabitol and mannitol ($r = 0.76$
340 and 0.85 , respectively) in nighttime than daytime ($r = 0.49$ and 0.51 , respectively) (Table 2),
341 suggesting that fungal and microbial spores contributed to high levels of trehalose in nighttime.
342 Hackl et al. (2000) found trehalose as dominant sugar in spring aerosols and proposed it as a tracer
343 for soil dust particles. Trehalose concentration was more significantly correlated with Ca^{2+} ($r =$
344 0.82) in daytime than nighttime ($r = 0.61$), indicating soil dust contribution (Table 2). Therefore,
345 we hypothesized that winds transported soil particles from the urban area in daytime due to the

346 active building constructions (He et al., 2015), contributing to the high levels of trehalose in
347 daytime.

348 3.1.3. Ambient concentrations and diurnal variations of sugar alcohols

349 The average concentrations of sugar alcohols were found 75.8 ng m^{-3} , contributing 26.4%
350 of total SCs measured in Mangshan aerosols (Table 1). Sugar alcohols showed clear diurnal
351 variations in daytime high (avg. 87.4 ng m^{-3}) and nighttime low (avg. 53.7 ng m^{-3}) (Table 1).
352 Mannitol was found as the dominant sugar alcohol (avg. 44.1 ng m^{-3}), followed by arabitol (avg.
353 29.1 ng m^{-3}) and inositol (avg. 2.62 ng m^{-3}) (Table 1; Fig. 6a-c). Mannitol and arabitol are
354 common polyols detected in green algae, lichens, and fungal spores (Bieleski, 1995, Dahlman et
355 al., 2003; Filippo et al., 2013; Lewis and Smith, 1967; Yttri et al., 2007). Previous studies have
356 reported that arabitol and mannitol are key components of fungal spores, and thus they are
357 considered as fungal tracers (Bieleski, 1995; Lewis and Smith, 1967). Several fungal and microbial
358 species released spores during biological activities into the atmosphere (Dahlman et al., 2003;
359 Bauer et al., 2008; Filippo et al., 2013). Therefore, the autumn time fungal and microbial species
360 significantly contribute to arabitol and mannitol in the Mangshan aerosol samples.

361 However, mannitol and arabitol showed a strong positive linear correlation ($r = 0.81$),
362 which suggested common origins as reported in earlier studies (Fu et al., 2012) (Table 2). In
363 contrast, the higher concentration of mannitol than arabitol suggested its addition sources than
364 fungal spores in the Mangshan forest site. In this sequence, several previous studies have
365 confirmed the significance of mannitol in plant photosynthesis (Loescher et al., 1992; Keller and
366 Matile, 1989; Rumpfo et al., 1983). Pashynska et al. (2002) reported that detritus of mature leaves
367 can emit mannitol into the atmosphere by wind action. Heald and Spracklen (2009) also found a
368 correlation between the atmospheric water vapor with mannitol concentrations and leaf area index.
369 They suggested that the activities of the terrestrial biosphere widely affect mannitol concentrations
370 in the air. Our PMF results also indicated the substantial contribution of mannitol for vegetation
371 factor (24.8%), which supports that mannitol is attributed by vegetation from the forest area
372 (section 3.2).

373 In addition, the meteorological parameters, including high RH and temperature affect the
374 fungal and bacterial activities (Kim and Xiao, 2005; Sharma and Razak, 2003). The maximum
375 growth of fungi and bacteria was observed at 92–100% RH (Ibrahim et al., 2011). Interestingly,
376 the concentrations of arabitol and mannitol gradually increased after the end of precipitation,
377 following the increases in ambient temperature and RH (Figs. 2a, 6a, b). Miyazaki et al. (2012)

378 also discussed the increased contributions of arabitol and mannitol with daytime ambient
379 temperature and solar radiation in the aerosol samples collected from the forest area. Similar
380 temporal trends and positive linear correlations were observed between arabitol ($r = 0.69$) and
381 mannitol ($r = 0.57$) with RH, which supports the above phenomenon for Mangshan aerosols
382 (Table 2). Therefore, we propose that a favorable meteorological condition in autumn increases
383 the emissions of fungal spores and fragments of forest vegetation, which may be responsible for
384 arabitol and mannitol contributions in the Mangshan aerosols.

385 The diurnal variation of mannitol and arabitol were characterized by higher in the daytime
386 (51.7 ng m^{-3} and 32.5 ng m^{-3} , respectively) than nighttime (29.6 ng m^{-3} and 22.5 ng m^{-3} ,
387 respectively) (Fig. 3b, c). Yamaguchi et al. (2012) reported that fungal spores and bacterial cells
388 associated with bioaerosols could be transported long distances. The Mangshan site receives
389 significant anthropogenic and bioaerosols from Beijing City by southerly winds. Therefore, the
390 daytime plant activities, influenced by solar radiation and ambient temperature and the long-range
391 transport of fungal spores from megacities (Beijing) by southwest winds govern the diurnal
392 variation of sugar alcohols in the Mangshan atmosphere. On the other hand, lower concentrations
393 in nighttime can be explained by the clean air mass transport by mountain breeze from the
394 Mangshan National Forest area.

395 **3.2. Source apportionment of SCs**

396 To investigate the source apportionment of SCs, positive matrix factorization (PMF)
397 analysis was performed for the measured aerosol samples using tracer compounds for
398 anhydrosugars, primary sugars, and sugar alcohols. It is essential to select a suitable number of
399 factor solutions in the PMF analysis. Based on the possible sources of SCs, four to six factor
400 solutions were run in PMF model. In the four-factor solutions, the SCs, including arabitol,
401 mannitol, and trehalose, were merged in a single factor; this might underestimate the soil dust
402 sources. The SCs, including glucose, fructose, trehalose, arabitol, and mannitol, were distributed in
403 more than four factors; it might be overestimated the number of factor solutions according to
404 possible sources of SCs. Therefore, a total of five interpretable factor solutions were characterized
405 by the enrichment of each tracer compound to be significant to categorize the origins of individual
406 sugars, which reproduced more than 95% of SCs.

407 These five-factor solutions were preferred based on minimum robust and true Q values
408 (goodness of fit parameters) of the base runs, which observed 3103 and 3505, respectively. In each
409 bootstrap run, the concentrations and percentages of tracers were close to those of base-run results.

410 The PMF results of SCs indicate a stability because no significant changes were found between Q
411 values and factor profiles of F_{peak} rotation runs compared with the base run. PMF results show a
412 good correlation between the values of observed and predicted (modeled) concentrations in scatter
413 plot, indicating that the model very well fits the individual sugar species. These results support the
414 perfect rationality of the source apportionment (Figure S-1). The time series plot of observed and
415 predicted concentration (modeled) also shown that the model well fits the observed data set
416 (Figure S-2). Figures 6 and 7 show the factor profile resolved by PMF analysis of the Mangshan
417 aerosol samples. The percentages of each component are summed for factors 1 to 5 to be
418 calculated as 100%.

419 Factor 1 is characterized by the high contribution of glucose (80.2%) followed by fructose
420 (69.6%), mannitol (24.8%), and inositol (15.1%) (Fig. 7a). Glucose and fructose are highly water-
421 soluble SCs present in the leaves and bark of plants (Graham et al., 2003). High concentrations of
422 glucose and fructose have been reported in vascular plants and phytoplankton by Cowie and
423 Hegdes (1984). The dominant glucose and fructose in the Mangshan aerosol samples collected in
424 autumn are rational as leaf senescence and decay results in both primary sugars being released into
425 the atmosphere during the fall season. We found an excellent correlation between glucose and
426 fructose ($r = 0.94$) in the Mangshan aerosols (Table 2), indicating the similar vegetation sources
427 for both sugar species in autumn (Baker et al., 1998; Burshtein et al., 2011; Pacini, 2000). Higher
428 concentrations of glucose and fructose in the aerosol samples collected during the autumn season
429 are reasonable because leaf senescence and decay result in an increased emission of primary
430 sugars into the atmosphere.

431 Several studies have reported that plant species significantly contribute to mannitol in the
432 atmosphere (Burshtein et al., 2011; Devis et al., 1988; 1990). Miyazaki et al. (2014) also found a
433 significant amount of trehalose, mannitol, and arabitol in the aerosol samples collected from the
434 forest and concluded their origin from the terrestrial plants within the forest. Significant positive
435 linear correlations of mannitol with fructose in daytime ($r = 0.79$) and nighttime ($r = 0.86$) further
436 denote that abundance of mannitol is due to the decay of plant leaves in autumn (Table 2).
437 Therefore, we conclude that the contributions of mannitol is from both vegetation and fungal
438 spores in the Mangshan aerosol samples. Hence mannitol showed the presence in factor 1.
439 Vegetations contribute to SCs during the campaign. Therefore, factor 1 can be termed as a
440 vegetation factor due to the high abundances of glucose, fructose, and mannitol.

441 Factor 2 is dominated by high loading of trehalose (80.2%), followed by mannitol (29.7%),
442 glucose (19.8%), and arabitol (18.2%) (Fig. 7b). The contribution of trehalose to soil dust has been

443 reported in several studies from different locations around the world, suggesting trehalose as a
444 tracer for the surface soil (Jia et al., 2010; Medeiros et al., 2006). In addition, previous studies
445 reported that bacteria and other microbes in the soil are also an essential source of trehalose
446 (Rogge et al., 2007). Trehalose is significantly correlated with arabitol ($r=0.58$) and mannitol
447 ($r=0.58$), and Ca^{2+} ($r=0.70$), demonstrating its microbial and soil dust origin. Therefore, factor 2
448 can be termed as microbial and soil dust factor.

449 Factor 3 is characterized by levoglucosan (82.2%), galactosan (77%), and mannosan
450 (73.6%) (Fig. 7c). Previous studies have reported that these SCs are associated with BB aerosols
451 (Fraser and Lakshmanan, 2000; Graham et al., 2002; Simoneit, 2002). Simoneit et al. (1999)
452 reported that the pyrolysis of cellulose and hemicellulose emitted levoglucosan, galactosan and
453 mannosan. These sugar species are major organic components emits in the atmosphere by BB
454 activities (Simoneit et al., 2002). The BB influenced aerosols are enriched with levoglucosan,
455 mannosan, and galactosan (Nolte et al., 2001; Medeiros et al., 2006). The domestic BB for
456 cooking and house heating due to low ambient temperature and field burning of agricultural
457 residues occur in East Asia (Verma et al., 2015). The PMF results are very well supported by the
458 fact that anhydrosugars are associated with BB (Simoneit et al., 1999). Therefore, factor 3 can be
459 termed as a BB factor due to the high abundance of BB products.

460 Factor 4 is dominated by high loading of sucrose (90%), followed by inositol (36.9%) and
461 fructose (11.7%) (Fig. 7d). Sucrose plays a crucial role in the plant blossoming process as the
462 dominant sugar compound of pollen grains (Pacini, 2000). Several studies also reported that
463 sucrose is abundant sugar species found in airborne pollen grains and flowering plants (Fu et al.,
464 2012; Graham et al., 2003; Medeiros et al., 2006; Pacini, 2000). Therefore, sucrose is reported as
465 an excellent tracer for airborne pollen spores (Pacini, 2000). Thus factor 4 is termed as pollen
466 factor due to the high loading of sucrose.

467 Factor 5 is characterized by a higher contribution of arabitol (61.5%) followed by mannitol
468 (39.3%) and inositol (15.3%) (Fig. 7e). Sugar species contributing to factor 5 are associated with
469 fungal spores (Bauer et al., 2008). Various fungi and microbes emit spores, which are tracers for
470 the arabitol and mannitol; therefore, both sugars are considered as specific tracers of fungal
471 activities (Medeiros et al., 2006; Rogge et al., 2007). Thus, factor 5 is termed as a fungal factor
472 due to the high loading of arabitol and mannitol. Overall, the average contributions of each factor
473 to measured SCs were estimated by PMF analyses (Fig. 8), in which BB was found to account for
474 27% of measured SCs. The vegetation and microbial and soil dust sources equally contribute
475 (21%) to total SCs. The fungal spores and pollen spores contribute 16% and 15% of total SCs,

476 respectively. Finally, biomass burning emissions from the local areas and megacities via long-
477 range atmospheric transport were identified as an important source for the Mangshan aerosols.

478 **3.3. Contributions of sugar compounds to WSOC and OC**

479 The contribution of carbon content of measured SCs varied from 14.1-371 ng m⁻³ (av.
480 145 ng m⁻³) in daytime and 12.8-322 ng m⁻³ (av. 117 ng m⁻³) in nighttime, accounting for 0.83%
481 and 0.91% of OC, respectively (Fig. 9a, b). The mean carbon contents of anhydrosugars showed
482 clear diurnal variation with higher nighttime values (67.1 ng m⁻³) than daytime (42.7 ng m⁻³),
483 accounting for 0.43 % and 0.22 % of OC, respectively. These results suggest that BB significantly
484 contributed to Mangshan aerosols. However, the carbon contents of primary sugars showed
485 opposite diurnal variations; higher (68.5 ng m⁻³) in daytime than nighttime (28.3 ng m⁻³),
486 accounting for 0.41 % and 0.28 % of OC, respectively (Fig. 9a, b). This study suggests that the
487 daytime emissions of primary sugars from local vegetation and the decay of plant leaf in forest
488 significantly contribute to OC. The carbon concentration contributed by sugar alcohols showed
489 insignificant diurnal variations i.e. 34.6 ng m⁻³ in daytime and 21.3 ng m⁻³ in nighttime,
490 accounting for 0.20 % and 0.19 % of OC, respectively. This result indicates multiple carbon
491 sources of sugar alcohols in day and night. In addition, contributions of anhydrosugars, primary
492 sugars, and sugar alcohols to WSOC were similar to those of OC in Mangshan aerosols.

493 Based on the PMF analysis, we found five sources for SCs measured in Mangshan
494 aerosols. The different tracer compounds were used to calculate carbon contents: biomass burning-
495 C (i.e., levoglucosan, galactosan, mannosan), vegetation-C (glucose, fructose), fungal-C (arabitol,
496 mannitol), pollen-C (sucrose), and microbial-soil-C (trehalose) (Fig. 9c, d). Among the five
497 sources, biomass burning-C was found as the largest carbon contributor to Mangshan aerosols
498 (36.7%), followed by fungal-C (23.7%), vegetation-C (19.7%), pollen-C (14.2%), and microbial-
499 soil-C (4.84%). Biomass burning-C accounted for 1.38% and 0.43% at night, while 0.57% and
500 0.22% in daytime for WSOC and OC, respectively. The BB for cooking and space heating in
501 winter and autumn seasons are common in central China (Akagi et al., 2011), which should
502 increase the nighttime levels of Biomass burning-C at the Mangshan site. However, the carbon
503 contribution by vegetation and fungal sources are similar during day and nighttime for the
504 Mangshan aerosols. Pollen-C accounted for 0.20% and 0.07% of OC in daytime and nighttime,
505 respectively. Higher pollen activities are key sources for the high daytime levels of pollen-C in the
506 forest site (Taylor et al., 2002).

507 **3.4. Contribution of levoglucosan to OC and WSOC**

508 We calculated the mass concentration ratios of levoglucosan to OC (Lev/OC) and WSOC
509 (Lev/WSOC) to evaluate the contributions of BB and anthropogenic emissions to Mangshan
510 aerosols (Fig. 9a-c). Fossil fuel combustion and BB emit WSOC and OC. They are also
511 secondarily produced by photochemical oxidation of volatile organic compounds in the
512 atmosphere (Wang et al., 2005; Deshmukh et al., 2019b). Coal combustion and vehicle exhaust
513 can contribute to the high levels of OC and WSOC in aerosols (Xu et al., 2020). Levoglucosan, a
514 dominant constituent of BB products, has been considered as an excellent tracer of BB (Simoneit,
515 2002; Kuo et al., 2011).

516 Average Lev/OC ratio (5.69×10^{-3}) was lower than that of Lev/WSOC (1.66×10^{-2}) in
517 Mangshan samples (Fig. 10a). Yan et al. (2018) reported similar ratios of Lev/OC (4.0×10^{-3}) and
518 Lev/WSOC (1.6×10^{-2}) for coal combustion, suggesting a significant carbon contribution to
519 Mangshan aerosols from coal combustions in the industrial areas via long range transport.
520 Interestingly, we found a substantial diurnal variation of Lev/OC and Lev/WSOC ratios. The
521 average Lev/OC and Lev/WSOC ratios are several times higher in nighttime (8.48×10^{-3} and
522 2.70×10^{-2} , respectively) than daytime (4.21×10^{-3} and 1.11×10^{-2} , respectively) (Fig. 10b, c). These
523 results indicate that BB contributed substantially to the Mangshan organic aerosols in nighttime.
524 Moreover, the correlations of levoglucosan with OC and WSOC are stronger in nighttime ($r =$
525 0.81 and 0.70 , respectively) than daytime ($r = 0.45$ and 0.40 , respectively), demonstrating the
526 dominance of BB-derived aerosols in the nighttime Mangshan samples (Table 2).

527 In addition, WSOCs are derived from various emission sources. We propose that
528 secondary organic aerosols constitute a significant fraction of WSOC and OC in daytime
529 Mangshan aerosols. The photochemical oxidation of organic precursors emitted from fossil fuel
530 combustion in industries and vehicular exhausts also contributes to secondary production of
531 WSOC and OC in daytime (He et al., 2015), suggesting that emissions from the urban Beijing area
532 may significantly influence the daytime levels of Mangshan aerosols. He et al. (2015) proposed a
533 possible contribution of photochemical formation of secondary organic aerosols to atmospheric
534 WSOC and OC in north China. Nevertheless, the photochemical degradation of levoglucosan by
535 OH radicals under ultraviolet radiations and high temperatures (Hennigan et al., 2010) may play a
536 key role in lowering the ratios of Lev/OC and Lev/WSOC in daytime Mangshan aerosols.

537 **3.5. Mass concentration ratios of levoglucosan/mannosan**

538 The mass concentration ratios of levoglucosan and mannosan (Lev/Man) were calculated
539 to better characterize the emissions sources of BB tracers (softwood vs. hardwood) in the
540 Mangshan site. Figure 10d represents the variations of Lev/Man ratios for overall, day- and

541 nighttime periods. The Lev/Man ratios have been used to distinguish the hardwood (angiosperm)
542 and softwood (gymnosperm) burning in the ice core record from the Russian Far East (Kawamura
543 et al., 2012). Hardwood contains 55–65% cellulose and 20–30% hemicellulose (Klemm et al.,
544 2005). Levoglucosan and mannosan are derived from the thermal decomposition of cellulose and
545 hemicelluloses, respectively (Simoneit, 2002). Levoglucosan is thermally more stable than
546 mannosan and galactosan (Kuo et al., 2011). Hence, a lower Lev/Man ratio is associated with
547 softwood burning, whereas a higher ratio is associated with hardwood burning (Engling et al.,
548 2006, 2009). However, we found insignificant diurnal variations of Lev/Man ratios between night
549 (9.33-25.9, avg. 15.8) and daytime aerosols (0.90-23.3, avg. 13.6). Likewise, comparable Lev/Man
550 ratios (9-13 for PM₁₀ and 10-13 for PM_{2.5}) were reported for aerosol samples from Tanzania,
551 where wood and charcoal are primary fuels used for domestic cooking and heating (Mkoma and
552 Kawamura, 2013). Interestingly, wheat straws and lignite are used in China for domestic cooking
553 and house heating, which may also contribute to levoglucosan and mannosan in the Mangshan
554 aerosols.

555 Different Lev/Man ratios were reported in the chamber and controlled field experiments,
556 e.g., 4-22 for conifer and savanna grass burning (Iinuma et al., 2007), and 41.6 for rice straw and
557 and 55.7 cereal straw burning (Engling et al., 2009; Zhang et al., 2007). Kuo et al. (2011) reported
558 higher emissions of levoglucosan during high-temperature flaming (27.5-52.3) compared to low-
559 temperature smoldering (2.43-3.08). Hence, it is not easy to differentiate hardwood and softwood
560 burning based on Lev/Man ratios alone. Several studies reported a high Lev/Man ratio for both
561 softwood and hardwood burning. Thus, there may exist some other factors that significantly
562 control the Lev/Man ratios. Yan et al. (2018) found a significant contribution of levoglucosan in
563 coal combustion with Lev/Man ratio of 7.2. The variations of Lev/Man ratios in Mangshan may be
564 significantly influenced by several factors, i.e., flaming vs. smoldering, duration of biomass
565 burning, coal combustion, and hardwood vs. softwood burning. The moderate Lev/Man ratios in
566 autumn aerosols from Mangshan suggest that low temperature smoldering processes of hardwood
567 contribute to levels of levoglucosan and mannosan. However, the contribution of coal
568 combustions for house heating could not be excluded.

569 **4. Summary and Conclusions**

570 Anhydrosugars, primary sugars, and sugar alcohols were detected with distinct diurnal
571 variations in suburban aerosol samples collected at the Mangshan site in the northern vicinity of
572 Beijing. The wind patterns indicate that daytime air masses were transported from urban Beijing to
573 Mangshan, while clean air masses were delivered in nighttime from the Mangshan National Forest

574 Park. Daytime air masses from urban Beijing significantly influence the air quality of the northern
575 forest region. We observed the highest abundance of primary sugars, followed by anhydrosugars
576 and sugar alcohols. Local emissions from the forest plants and fungal species are the main
577 contributors to the primary sugars and sugar alcohols in the Mangshan aerosols. The
578 meteorological parameter significantly influenced the levels of SCs in the Mangshan samples. We
579 observed a significant influence of enhanced ambient temperature and solar radiation on the pollen
580 rupture and increased RH on fungal and microbial growth. This study suggested the source
581 variation for trehalose, i.e., local microbes at night and soil dust particles transported from Beijing
582 areas by southerly wind in daytime. We found that vegetation and fungal spores are not a specific
583 source of glucose and mannitol, respectively. Both sugars may have multiple sources in the forest
584 aerosols.

585 PMF results concluded the contributions of 36% from vegetation (21% vegetation factor
586 and 15% pollen factor) and 37% from microbial and fungal species (21% microbial soil dust and
587 16% fungal factor). The BB activities for domestic cooking and space heating in north China
588 contributed higher organic carbon at nighttime (0.43%) than in daytime (0.22%). Therefore, local
589 BB seriously affected the air quality of the Mangshan site. Lev/Man ratio suggested that low
590 temperature smoldering burning of hardwood is the main source for BB aerosols. SCs were
591 recognized as a significant aerosol component at Mangshan, northern suburbs of Beijing. SCs can
592 influence the air quality and thus climate because they are essential components of organic
593 aerosols on a global scale. This study of SCs at Mangshan demonstrates that ambient levels of SCs
594 are highly sensitive to the emissions of anthropogenic and biogenic aerosols. Higher contribution
595 of levoglucosan to SCs demonstrated a significant BB activity around the Mangshan site in north
596 China.

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598 **Data availability.** Raw data are available on request by contacting the corresponding author.

599 **Author contributions.** This research was designed YK, KK and ZW. Laboratory measurements
600 were performed by FY with a support of PF. The paper was prepared by SKV and KK.

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

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900 **Figure Captions**

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902 Figure 1. Geographical location of Mangshan, China. The map was downloaded from © Google
903 Maps 2019.

904 Figure 2. (a) The meteorological parameters at Mangshan during sampling periods, (b) Fractions
905 of local wind directions at Mangshan site, north of Beijing, China.

906 Figure 3. Concentrations (ng m^{-3}) of sugar compound (a) overall, (b) daytime and (c) nighttime in
907 aerosol samples from Mangshan during September-October 2007 (The error bars denote the
908 standard deviation).

909 Figure. 4. Temporal variations in the concentrations (ng m^{-3}) of anhydrosugars in the Mangshan
910 aerosol samples collected for September-October 2007. (Solid circle represents nighttime
911 samples collected from 18:00 to 09:00 hours. Hollow circle represents daytime samples).

912 Figure. 5. Temporal variations in the concentrations (ng m^{-3}) of primary sugars in the Mangshan
913 aerosol samples collected for September-October 2007. (Solid circle represents nighttime
914 samples collected from 18:00 to 09:00 hours. Hollow circle represents daytime samples).

915 Figure. 6. Temporal variations in the concentrations (ng m^{-3}) of sugar alcohols in the Mangshan
916 aerosol samples collected for September-October 2007. (Solid circle represents nighttime
917 samples collected from 18:00 to 09:00 hours. Hollow circle represents daytime samples).

918 Figure 7. PMF analyses of sugar compounds in Mangshan aerosols based on the autumn 2007 data
919 set.

920 Figure 8. Source contributions to sugar compounds from various sources based on PMF analyses.

921 Figure 9. The concentrations and relative contributions of the carbon content of anhydrosugars,
922 primary sugars and sugar alcohols to the carbon concentrations of measured sugar
923 compounds, water-soluble organic carbon (WSOC) and organic carbon (OC) fraction of
924 Mangshan aerosols (a = daytime and b = nighttime). The concentrations and relative
925 contribution of the carbon content of five sources of sugar compounds to total sugar
926 compounds measured, WSOC and OC fraction of Mangshan aerosols (c = daytime and d =
927 nighttime).

928 Figure 10. Mass concentrations ratio of carbon contents of (a) levoglucosan (Lev) to organic
929 carbon (OC) and water soluble organic carbon (WSOC), (b) levoglucosan (Lev) to organic
930 carbon (OC) daytime and night time, (c) levoglucosan (Lev) to water soluble organic carbon
931 (WSOC) daytime and night time, (d) average levoglucosan to mannosan ratios (Lev/Man) in
932 the Mangshan aerosol samples for autumn 2007.

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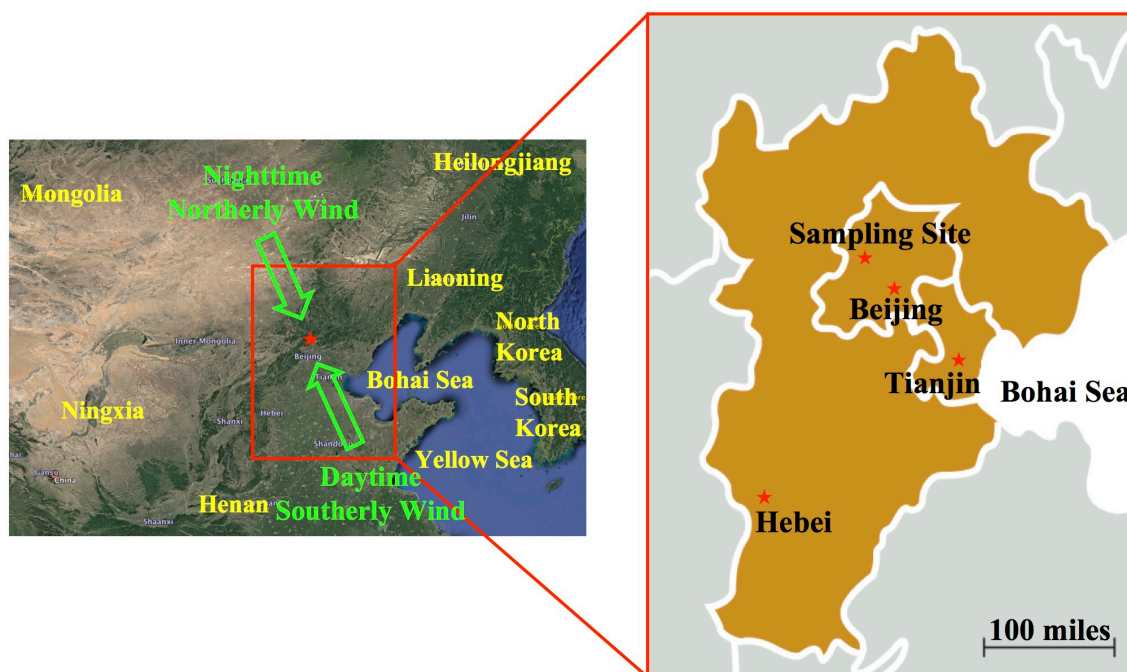
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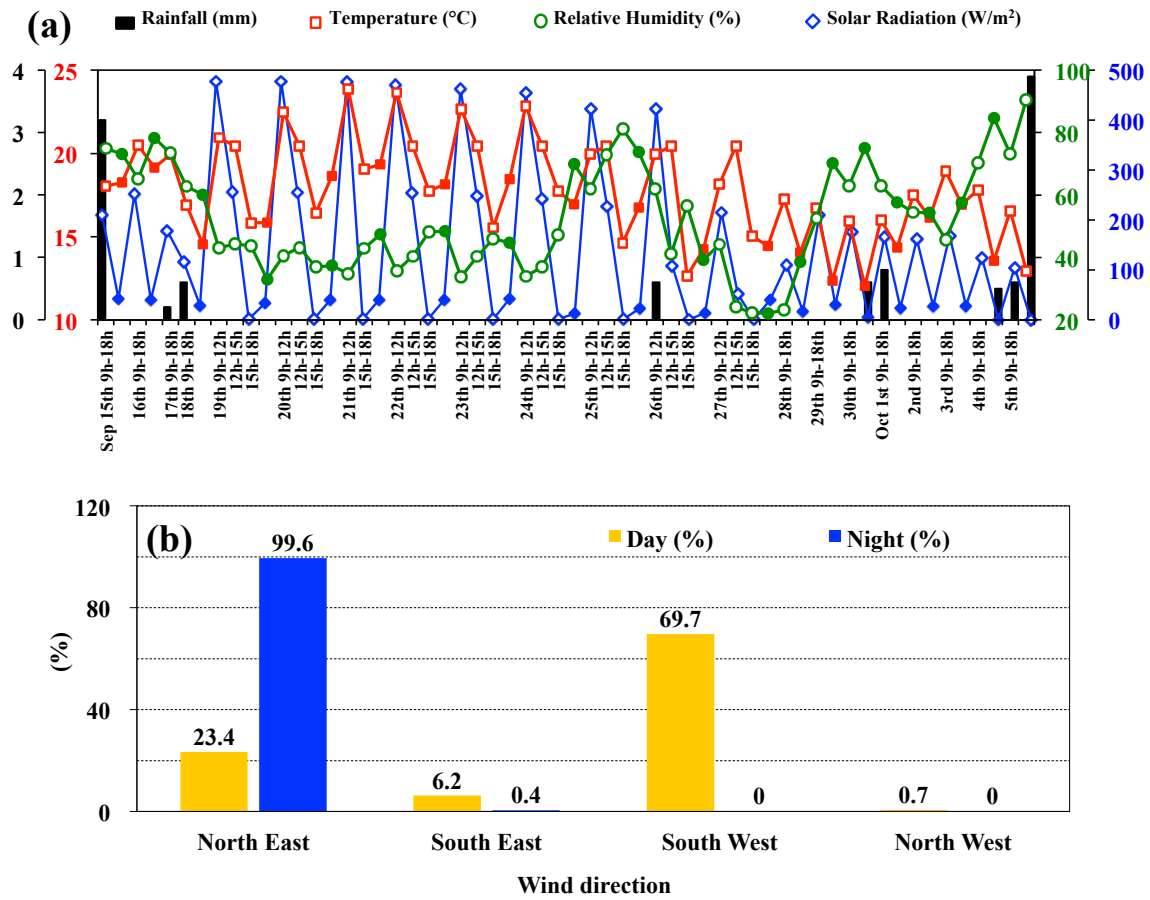
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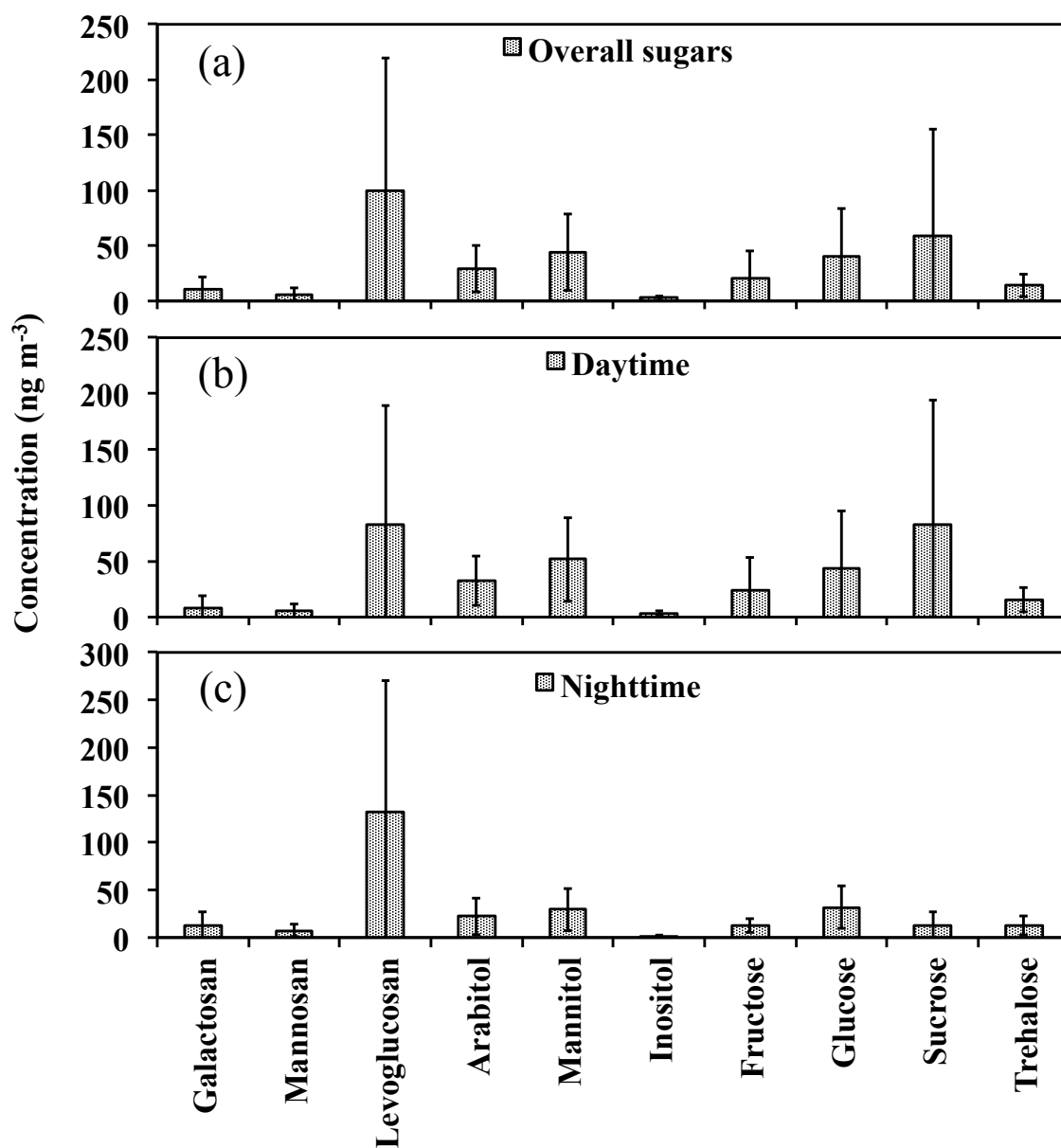
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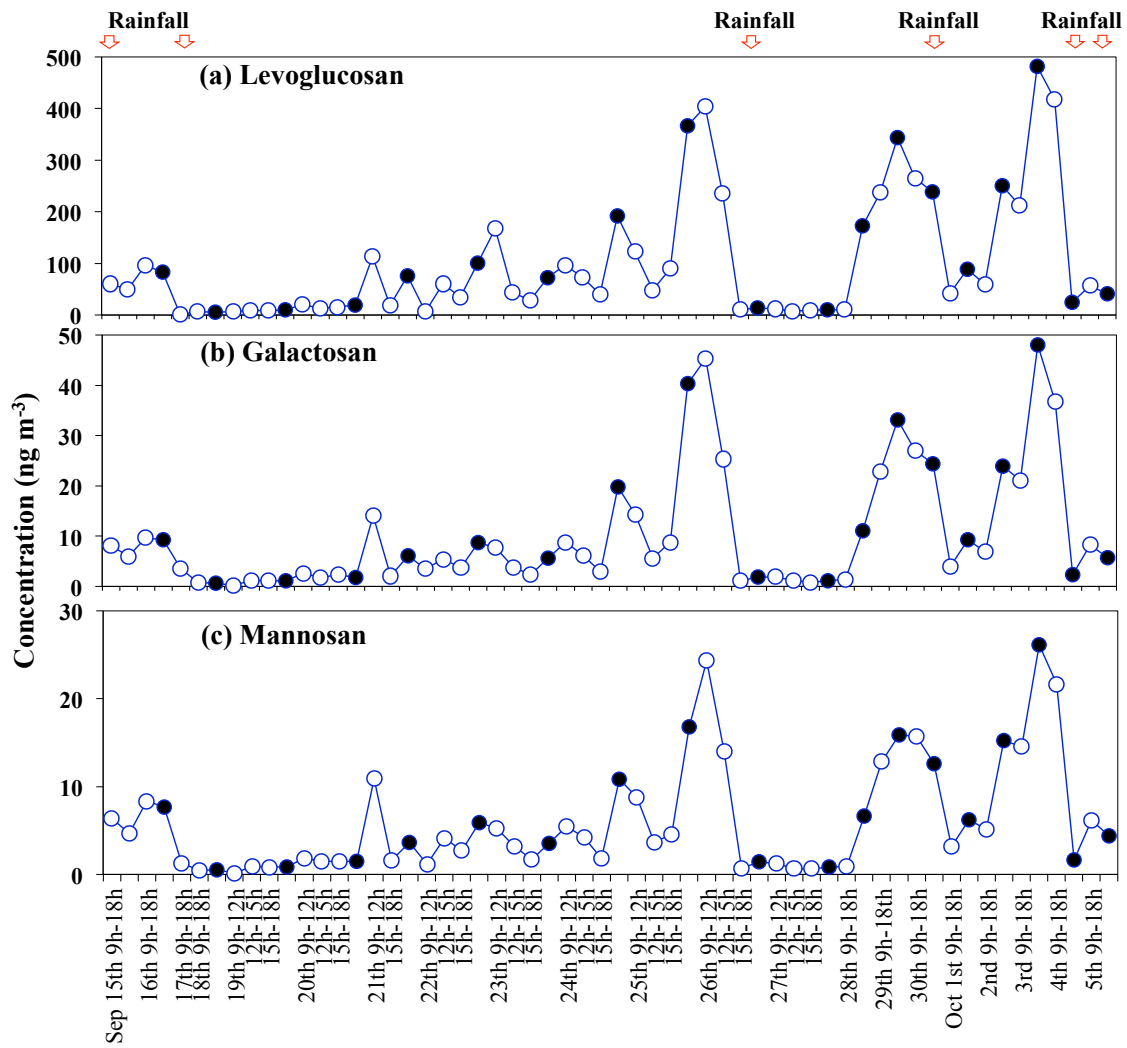
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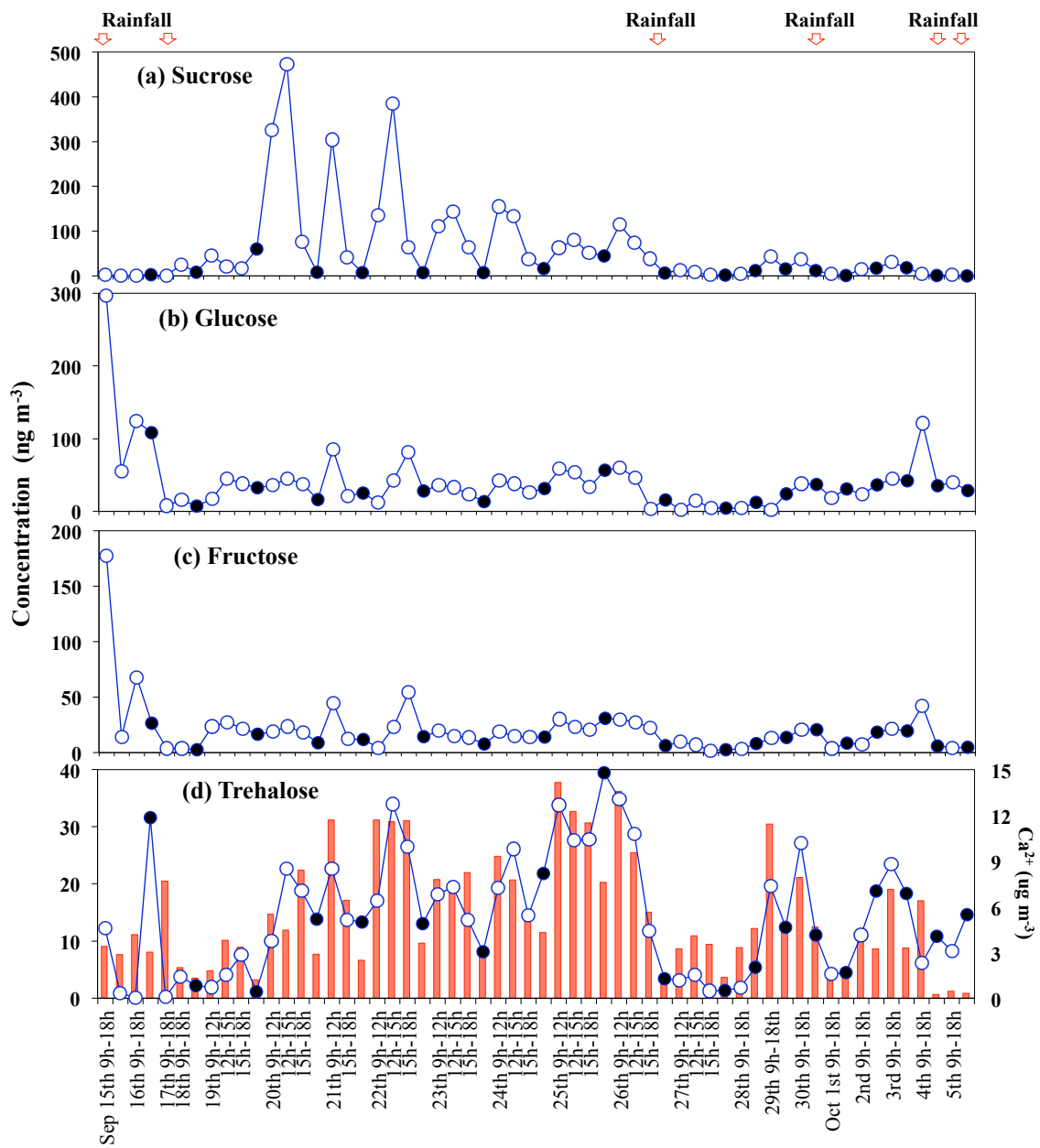
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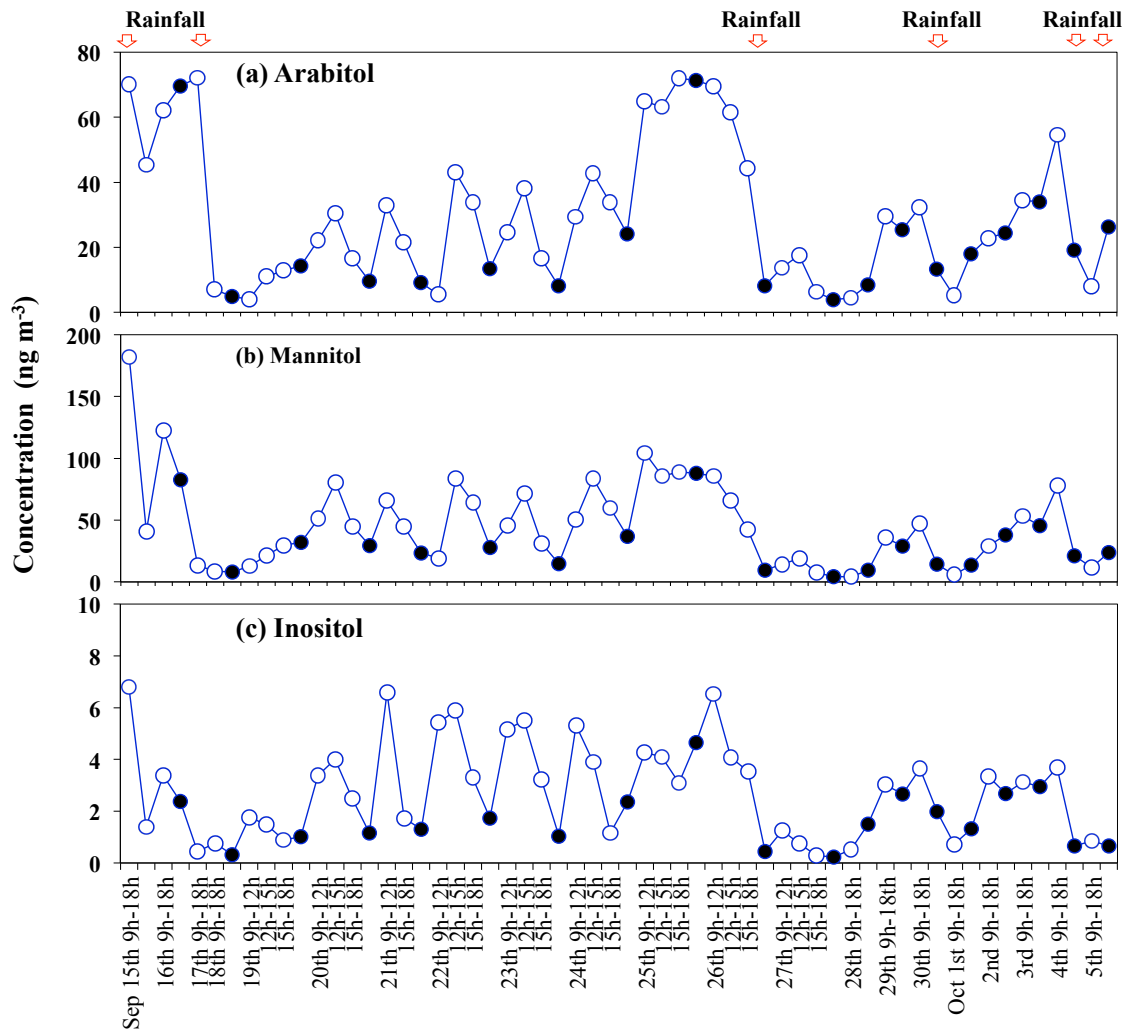
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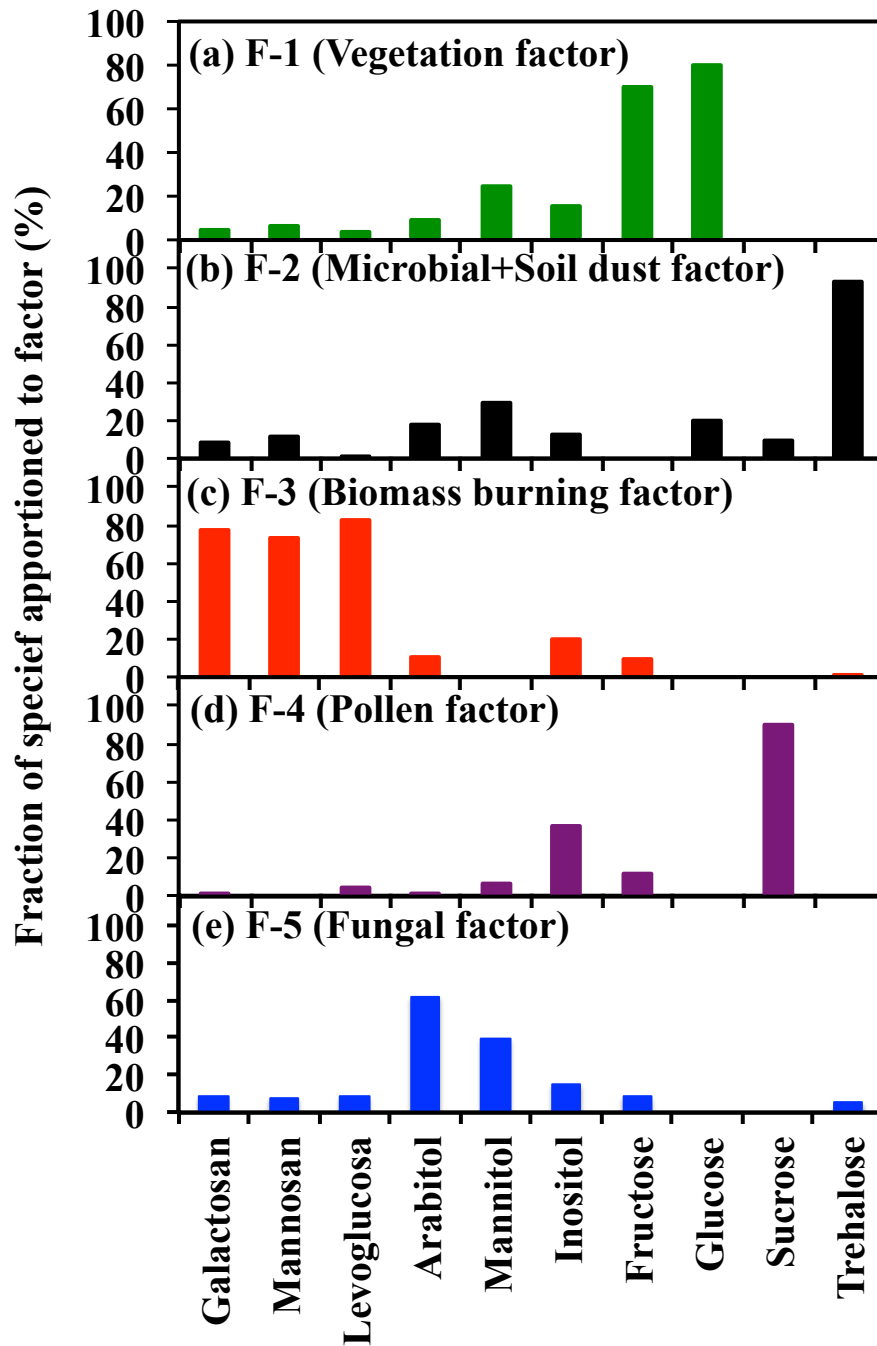
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Fig. 6.



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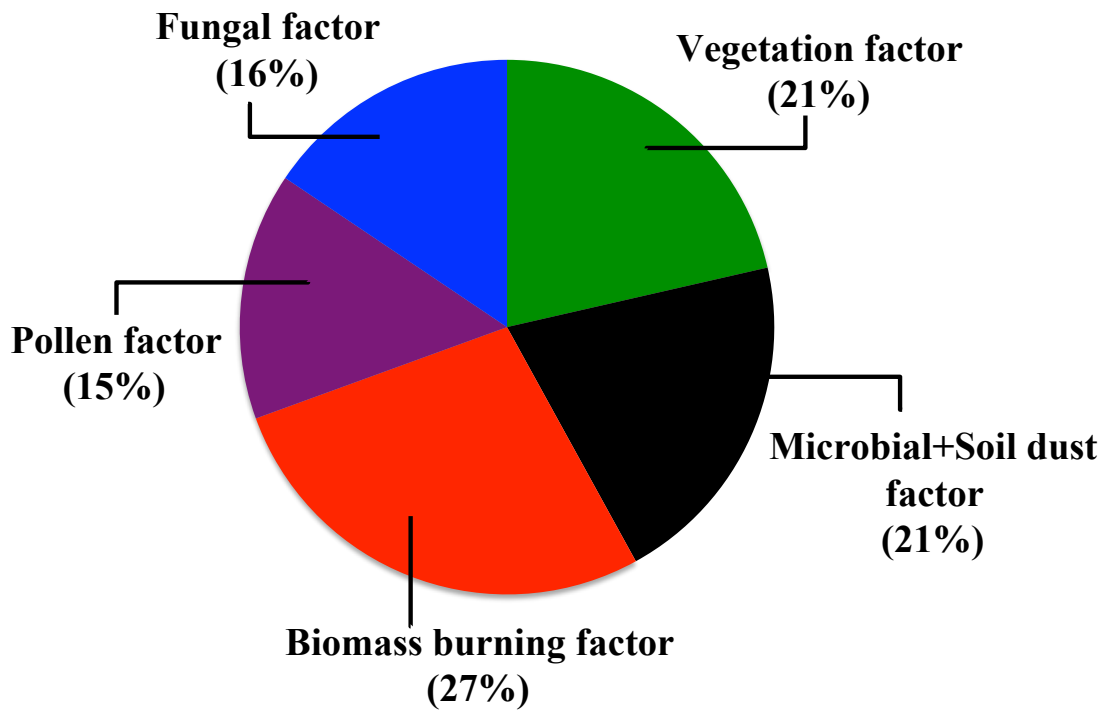
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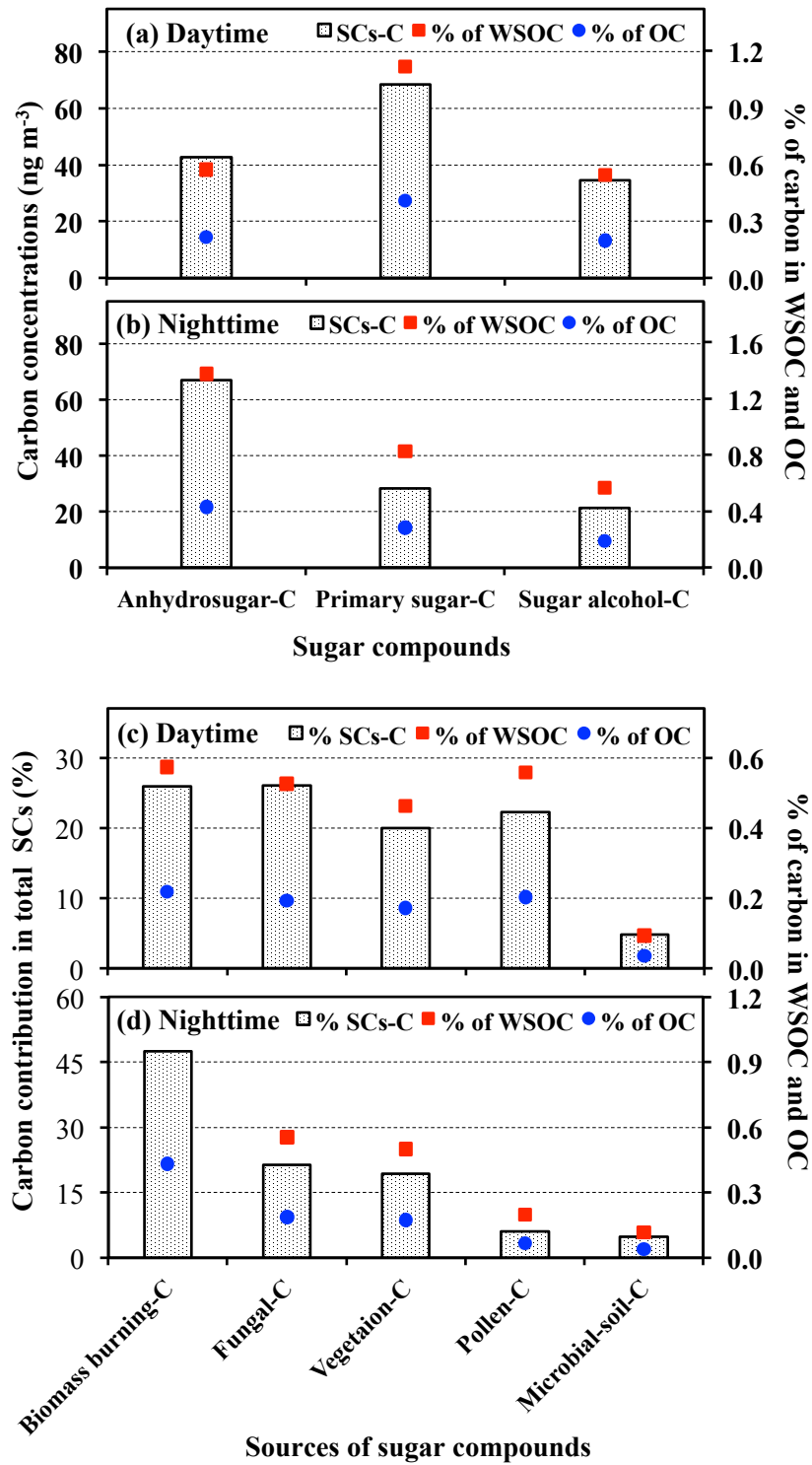
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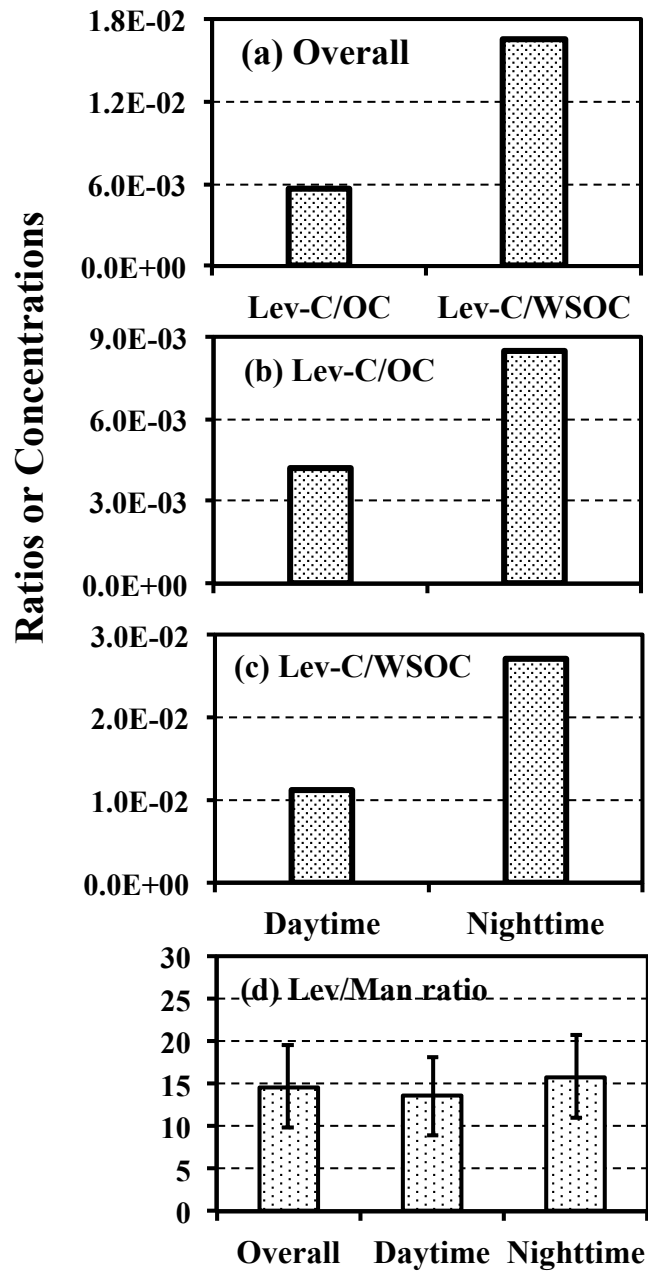
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Fig. 9.



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Table 1. Minimum, maximum, average and standard deviations of concentrations of sugar compounds in aerosol samples (TSP) from Mangshan, China.

Sugar Compounds	Overall				Daytime (n = 38)				Nighttime (n = 20)			
	Min	Max	Avg.	S.D.	Min	Max	Avg.	S.D.	Min	Max	Avg.	S.D.
Anhydrosugars												
Galactosan	0.14	48.0	10.1	11.9	0.14	45.3	8.53	10.5	0.69	48.0	13.0	14.0
Mannosan	0.13	26.1	6.05	6.33	0.13	24.3	5.37	6.01	0.53	26.1	7.35	6.87
Levoglucoosan	1.17	482	100	119	1.17	418	83.2	106	5.66	482	132	138
Sugar alcohols												
Arabitol	3.89	72.2	29.1	21.5	3.99	72.2	32.5	22.0	3.89	71.3	22.5	19.4
Mannitol	4.19	182	44.1	34.5	4.19	182	51.7	37.5	4.40	87.7	29.6	22.3
Inositol	0.23	6.8	2.62	1.81	0.27	6.80	3.14	1.90	0.23	4.65	1.62	1.09
Primary sugars												
Fructose	1.72	177	20.1	24.6	1.72	177	23.9	29.3	2.64	30.9	12.8	7.67
Glucose	1.86	297	40.0	43.4	1.86	297	44.2	50.8	4.52	108	32.0	22.8
Sucrose	0.02	474	58.5	96.5	0.02	474	82.9	112	0.04	60.1	12.3	15.1
Trehalose	0.06	39.5	14.3	10.5	0.06	34.9	15.3	10.6	0.87	39.5	12.3	10.2
Anhydrosugars	6.01	556	116	137	6.01	476	97.1	122	6.88	556	152	159
Primary sugars	9.41	565	133	125	9.41	565	166	141	10.5	172	69.4	43.0
Sugar alcohols	8.53	259	75.8	54.7	9.09	259	87.4	57.5	8.53	164	53.7	41.9
Total Sugars	30.8	875	325	232	34.1	875	351	240	30.8	759	276	212
Anhydrosugars (%)			31.9				24.6				45.7	
Primary sugars (%)			41.8				47.3				31.3	
Sugar alcohols (%)			26.4				28.1				23.0	

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Table 2. Statistical summary of correlations among the chemical species and meteorological variables in aerosol samples collected at a forest site in northern Japan

Linear regression	Correlation coefficient	p value	Significance of correlation at P value < 0.05
Overall (n = 58)			
Levogluconan vs. Galactosan	0.98	< 0.05	Significant
Levogluconan vs. Mannosan	0.97	< 0.05	Significant
Mannosan vs. Galactosan	0.98	< 0.05	Significant
Sucrose vs. Temperature	0.52	< 0.05	Significant
Sucrose vs. Solar radiation	0.55	< 0.05	Significant
Arabitol vs. Mannitol	0.81	< 0.05	Significant
Arabitol vs. RH	0.69	< 0.05	Significant
Mannitol vs. RH	0.57	< 0.05	Significant
Glucose vs. Fructose	0.94	< 0.05	Significant
Trehalose vs. Arabitol	0.58	< 0.05	Significant
Trehalose vs. Mannitol	0.58	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.70	< 0.05	Significant
Daytime (n = 38)			
Sucrose vs. Ca ²⁺	0.32	> 0.05	Not significant
Glucose vs. Ca ²⁺	0.02	> 0.05	Not significant
Trehalose vs. Arabitol	0.49	< 0.05	Significant
Trehalose vs. Mannitol	0.51	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.81	< 0.05	Significant
Fructose vs. Mannitol	0.79	< 0.05	Significant
Levogluconan vs. OC	0.45	< 0.05	Significant
Levogluconan vs. WSOC	0.40	< 0.05	Significant
Nighttime (n = 20)			
Sucrose vs. Ca ²⁺	0.37	> 0.05	Not significant
Glucose vs. Ca ²⁺	0.27	> 0.05	Not significant
Trehalose vs. Arabitol	0.76	< 0.05	Significant
Trehalose vs. Mannitol	0.85	< 0.05	Significant
Trehalose vs. Ca ²⁺	0.61	< 0.05	Significant
Fructose vs. Mannitol	0.86	< 0.05	Significant
Levogluconan vs. OC	0.81	< 0.05	Significant
Levogluconan vs. WSOC	0.70	< 0.05	Significant

The data of Ca²⁺, OC and WSOC are adapted from He et al. (2015).

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