Measurement report: Chemical characteristics of PM2.5 during typical biomass

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burning season at an agricultural site of the North China Plain

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12 Abstract:

Biomass burning activities are ubiquitous in China, especially in North China, where there is an 13 enormous rural population and winter heating custom. Biomass burning tracers (i.e., levoglucosan, 14 15 mannosan -(MN)-and potassium (K⁺)), as well as other chemical components were quantified at a 16 rural site (Gucheng, GC) in North China from 15 October to 30 November, during a transition 17 heating season, when the field burning of agricultural residues was becoming intense. The measured 18 daily average concentrations of LGlevoglucosan, mannosanMN and K⁺ in PM_{2.5} during this study were $0.79 \pm 0.75 \ \mu g \ m^{-3}$, $0.03 \pm 0.03 \ \mu g \ m^{-3}$ and $1.52 \pm 0.62 \ \mu g \ m^{-3}$, respectively. Carbonaceous 19 20 components and biomass burning tracers showed higher levels at nighttime than daytime, while 21 secondary inorganic ions were enhanced during daytime. An episode with high levels of biomass 22 burning tracers was encountered at the end of October, 2016, with high LG levoglucosan at 4.37 µg 23 m⁻³. Based on the comparison of chemical components during different biomass burning pollution 24 periods, it appeared that biomass combustion can obviously elevate carbonaceous components 25 levels, whereas no essentially effect on secondary inorganic aerosols in the ambient air. Moreover, 26 the levoglucosanLG/mannosanMN ratios during different biomass burning pollution periods 27 remained at high values (in the range of 18.3 - 24.9), however, the levoglucosan LG/K^+ ratio was 28 significantly elevated during the intensive biomass burning pollution period (1.67) when air temperatures decreasing, substantially higher than in other biomass burning periods (averaged at0.47).

31 *Keywords*: Biomass burning; Organic tracers; Levoglucosan; Mannosan; Potassium

32 **1. Introduction**

Particulate air pollution is attracting more and more concerns in China because of their obvious 33 34 adverse impact on visibility reduction, as well as health implication and regional or global climate 35 change (Kanakidou et al., 2009; Pope and Dockery, 2006; Cheng et al., 2016). Carbonaceous species, i.e., organic carbon (OC) and elemental carbon (EC), and water-soluble inorganic ions, e.g., SO_4^{2-} , 36 37 NO_3^- and NH_{4^+} are the major components of ambient aerosols (Liang et al., 2017; Du et al., 2014; 38 Zheng et al., 2015; Tan et al., 2016). Biomass burning (BB) emissions constitute a large source of 39 ambient particulate pollution, especially for carbonaceous components, i.e., primary organic carbon 40 (POC) and black carbon (BC) on global scale (Bond et al., 2004; Tang et al., 2018; Titos et al., 2017). 41 As an important aerosol component, black carbon from industrial and combustion emissions 42 contributes to the enhanced $PM_{2.5}$ (particulate matter with aerodynamic diameters less than 2.5 μ m) 43 mass concentrations and influences regional radiative forcing (Chen et al., 2017). Fresh biomass burning aerosol was found to be mainly comprised of carbonaceous species which typically 44 45 constitutes 50-60% of the total particle mass (Hallquist et al., 2009). Yao et al. (2016) identified 46 approximately half of carbonaceous aerosols being contributed by biomass burning at Yucheng, a 47 rural site in the North China Plain.

48 Biomass burning emissions also represent a potentially large source of secondary organic aerosol (SOA). The precursors and formation pathways of SOA from biomass burning emissions 49 50 were investigated by extensive field observations (e.g., Zhu et al., 2015; 2017; Adler et al., 2011; 51 Zhang et al., 2010; 2015). Based on morphological particle analysis, Yao et al. (2016) investigated 52 the smoke emitted from biomass burning impacting SOA production. Sun et al. (2010) found that phenolic compounds, which were emitted in large amounts from wood combustion, can form SOA 53 54 at high yields in aqueous-phase reactions. In addition, smoke from biomass burning can be 55 transported thousands of kilometers downwind from the source areas. Biomass burning aerosol from 56 Southeast Asia can be transported to China, Singapore and even further to North America (Liang et al., 2017; Hertwig et al., 2015; Peltier et al., 2008). Based on molecular tracer measurements,
synoptic data as well as air mass back trajectory analysis, a fire episode was captured at a
background site of East China with smoke advected from Southeast Asia (Liang et al., 2017).

60 The North China Plain (NCP) is one of the most polluted regions in China. Severe haze-fog of 61 longer duration and more extensive coverage has occurred frequently in the NCP area, especially 62 during the seasons of autumn and winter. NCP covers one quarter of China's cultivated land and 63 yields 35% of the agricultural products in China (Boreddy et al., 2017). The rural population in NCP 64 is also large and dense, and biomass burning activities are common in this region in form of cooking 65 and heating. Intense fire activity typically occurs in October after the corn harvest. Abundant smoke is emitted from agricultural burning, i.e., residential biofuel combustion, open field burns, etc. 66 67 Various field observations have investigated different aspects of biomass burning, e.g., seasonal 68 variations, chemical and physical properties of smoke particles, spatial distribution, sources, 69 transport, etc., in the NCP region (Cheng et al., 2013; Shen et al., 2018; Sun et al., 2013; 2016; 70 Boreddy et al., 2017; Xu et al., 2019). However, these field investigations of the contribution of 71 biomass burning to ambient aerosols in the NCP region were concentrated on the city of Beijing 72 (Cheng et al., 2013; Zheng et al., 2015; Duan et al., 2004). Little field research about biomass 73 burning was reported for rural areas in the NCP. In fact, biomass burning activities are common in 74 the rural areas of the NCP region, and the resulting smoke aerosol can be transported to urban areas, 75 e.g., the city of Beijing, resulting in haze episodic events.

76 The objective of this study is gain insights about the abundance of smoke during the typical 77 biomass burning season, i.e., autumn-winter transition season, following the corn harvest. In this 78 paper, we focus on quantifying multiple biomass burning tracers, i.e., levoglucosanLG, 79 mannosanMN and K⁺ as well as other chemical species in PM2.5 in the rural areas of the NCP region 80 during the typical biomass burning season. In order to characterize the biomass burning pollution 81 status within rural areas of the NCP region, multiple biomass burning tracers, i.e., levoglucosan, 82 mannosan and K^{\pm} in PM_{2.5} sampled at a heavily polluted rural site in Hebei province were quantified 83 during a typical biomass burning season, i.e., autumn-winter transition season, following the corn 84 harvest. Combined with the analysis of other chemical components, it reveals different levels of 85 biomass combustion pollution impacting on different types of chemical components in ambient 86 aerosol. Meanwhile, based on the multi-analysis of biomass burning molecular tracers, back
87 trajectory analysis, fire activity data and synoptic conditions, the results of this study demonstrate
88 the biomass burning pollution status, as well as the formation process of severe biomass burning
89 pollution episode in the rural atmosphere of North China. These results can provide valuable
90 information about the biomass burning activities in all of Northern China.
91 The results of this study demonstrated the biomass burning pollution status, as well as

92 chemical properties of ambient aerosols under different biomass burning pollution levels in the rural
 93 atmosphere of North China.

94 **2. Site description and experimental Methods**

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2.1 Site description and sampling

96 Samples were collected at a rural site, Gucheng (GC, 39°09'N, 115°44'E; 15.2 m a.s.l), located 97 on a platform at the China Meteorological Administration farm in the town of Gucheng (GC site), 98 approximately 110 km southwest of Beijing and 35 km north of the city of Baoding (population of 99 about 5 million) in Hebei province, as shown in Figure S1. The station is surrounded by agricultural 100 fields, with major crop species being corn and wheat. The dominant wind direction at GC is southwest and northeast during the study period. This site is upwind of Beijing, when the wind 101 102 blows from the south or southwest, where heavily polluted cities and regions of Hebei province, i.e., 103 Baoding, Shijiazhuang, Xingtai, Handan, are located. Thus, it is an appropriate station for representing the air pollution situation in the NCP region (Sheng et al., 2018; Chi et al., 2018; Xu 104 105 et al., 2019; 2020; Kuang et al., 2020).

106 Daytime and nighttime PM_{2.5} samples were collected from 15 October, 2016 to 23 November, 107 2016, by using PM_{2.5} High-volume (Hi-Vol) sampler (GUV-15HBL1, Thermo Fisher Scientific CO., LTD), at the nominal flow rate of 1.13 m³ min⁻¹. The daytime samples were collected from 07:00 to 108 19:00, while nighttime samples were collected from 19:00 to 07:00 local time of the next day. All 109 $PM_{2.5}$ samples were collected on quartz fiber filters, prebaked at 850 °C for at least 5 h to remove 110 carbonaceous material. A total of 33 couples of daytime/nighttime samples and 6 whole-day samples 111 112 as well as 4 field blank samples were collected during the sampling period. The filters were stored 113 at -20 °C after sample collection.

114 **2.2 Experimental Methods**

115 2.2.1 Anhydrosugar and water-soluble inorganic ion analysis

116 The quartz filter samples were analyzed for biomass burning anhydrosugar tracers, i.e., 117 levoglucosan and mannosanLG and MN, using an improved high-performance anion-exchange 118 chromatography (HPAEC) method with pulsed amperometric detection (PAD) on a Dionex ICS-119 5000+ system. LevoglucosanLG and mannosanMN were separated by a Dionex Carbopac MA1 120 analytical column and guard column with an aqueous sodium hydroxide (NaOH, 480 mM) eluent 121 at a flow rate of 0.4 mL min⁻¹. The detection limit of levoglucosanLG and mannosanMN was 0.002 mg L^{-1} and 0.005 mg L^{-1} , respectively. More details about the HPAEC-PAD method can be found 122 123 elsewhere (Iinuma et al., 2009).

124 The quartz filter samples were also analyzed for water-soluble inorganic ions by a Dionex ICS-5000+ ion chromatograph, including SO₄²⁻, NO₃⁻, NH₄⁺, Cl⁻, Ca²⁺, Na⁺, K⁺ and Mg²⁺, and the 125 method detection limits for the individual ionic species were 0.18 μ g L⁻¹, 0.15 μ g L⁻¹, 0.03 μ g L⁻¹, 126 $0.048 \ \mu g \ L^{-1}$, $0.08 \ \mu g \ L^{-1}$, $0.01 \ \mu g \ L^{-1}$, $0.01 \ \mu g \ L^{-1}$, $0.008 \ \mu g \ L^{-1}$, respectively. The cations were 127 128 separated on an Ionpac CS12 analytical column and CG12 guard column with a 20 mM 129 methanesulfouic acid as eluent at a flow rate of 1.0 mL min⁻¹, while the anions were separated on 130 an Ionpac AS11-HC column and AG11-HC guard column with 21.5 mM KOH eluent at a flow rate 131 of 1.0 mL min⁻¹. The water-soluble inorganic ion data were corrected by field blanks.

132 2.2.2 Organic carbon/elemental carbon analysis

OC and EC were measured on a punch (0.526 cm^2) of each quartz sample by a thermal/optical carbon analyzer (DRI Model 2001, Desert Research Institute, USA), using the Interagency Monitoring of Protected Visual Environments (IMPROVE) thermal evolution protocol with reflectance charring correction. The analytical error of OC was within 10%, and one sample of every 10 samples was selected at random for duplicate analysis. The detection limit of OC was 0.82 µgC cm⁻² (Liang et al., 2017).

139 2.2.3 Gas online monitoring (i.e., NO, NO₂, SO₂, O₃, CO and NH₃)

140During this campaign, commercial instruments from Thermo Fisher Scientific Co., LTD were

- 141 used to measure O_3 (TE 49C), NO/NO₂/Nox (Model 42CTL), CO (TE 48CTL), and SO₂
- 142 (TE43CTL), while NH₃ was measured by an ammonia analyzer (DLT-100, Los Gatos Research,

143 USA) at GC station. All measurement data quality was controlled according to standard gases (Xu

144 et al., 2019; Lin et al., 2011; Meng et al., 2018; Ge et al., 2018).

145 2.2.4 Meteorological parameters

The meteorological parameters, including air temperature, relative humidity (RH) and wind speed at a 24-h resolution at the GC site are presented in Figure 1. During this campaign, the daily average RH value was observed at $77 \pm 13\%$, with a range from 48% to 99%, while the daily wind speed was observed with an average value of 1.07 ± 1.14 m s⁻¹, exhibiting moist and stable synoptic conditions at this rural site during the autumn-winter transition season. Moreover, there was rare precipitation during the sampling period at the GC site, except for two days, i.e., 20 and 27 October, 2016 (Figure 1).

153 2.2.5 Back trajectory and fire spot analysis

To characterize the transport pathways of the aerosol at the Gucheng site, back-trajectories were calculated with the NOAA Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model via NOAA ARL READY Website (http://ready.arl.noaa.gov/HYSPLIT.php).

To investigate the influence of biomass burning activities in surrounding areas, fire hot spot counts were obtained from the Fire Information for Resource Management System (FIRMS) (available at <u>https://firms.modaps.eosdis.nasa.gov/download/</u>).

160 <u>2.2.6 –Statistical analysis</u>

Statistical analysis of data, i.e., the correlation analysis between the concentrations of
 levoglucosan, mannosan and K[±] at the Gucheng site during the sampling period were conducted
 with the linear fitting method.

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165 **3. Results and discussion**

166 **3.1 Characteristics of chemical components in PM2.5**

167 In this study, the mass concentration of $PM_{2.5-cal}$ was reconstituted by the sum of carbonaceous 168 components (1.6×OC + EC) and inorganic ions ($SO_4^{2-} + NH_4^+ + NO_3^- + CI^- + Ca^{2+} + Na^+ + K^+ +$ 169 Mg²⁺). Figure 1 describes the time-series variation obtained for daily PM_{2.5-cal}, and its major 170 components (OC, EC, SO_4^{2-} , NO_3^- and NH_4^+), biomass burning tracers (<u>levoglucosanLG</u>,

mannosanMN and K⁺), ratios of levoglucosan/OC and meteorological factors (temperature (T), 171 172 relative humidity (RH), wind speed (WS) and planetary boundary layer (PBL) height-rainfall) 173 during the sampling period. The average daily PM_{2.5-cal} mass concentration in the autumn-winter 174 transition season at GC reached 137 \pm 72.4 µg m⁻³, ranging from 23.3 µg m⁻³ to 319 µg m⁻³ (Table 175 1, Figure 1a), which is higher than during the severe winter haze in January, 2013 at an urban site 176 in Beijing (121 µg m⁻³) (Zheng et al., 2015). The mass concentrations of these chemical species during the day are distributed as follows (from highest to lowest): $OC > EC > NO_3^- > SO_4^{2-} >$ 177 $NH_4^+ > Cl^- > Ca^{2+} > K^+ > Na^+ > Mg^{2+}$. Organic matter (OM), <u>calculated by multiplying OC values</u> 178 179 with a coefficient of 1.6, was the most abundant PM component, the daily average value of which 180 was 70.4 \pm 49.6 μ g m⁻³, accounting for nearly half (46.7%) of PM_{2.5-cal} mass, indicating obvious 181 organic pollution at the rural site in the North China Plain during the sampling season.

182 The measured daily average concentrations of biomass burning tracers, i.e., levoglucosan-LG, 183 mannosanMN and K⁺ in PM_{2.5} during our study were $0.79 \pm 0.75 \ \mu g \ m^{-3}$, $0.03 \pm 0.03 \ \mu g \ m^{-3}$ and 184 $1.52 \pm 0.62 \ \mu g \text{ m}^{-3}$, respectively (Table 1). The anhydrosugar ambient concentrations of 185 levoglucosan levels (levoglucosanLG and mannosanMN) in this study were higher than those observed in the city of Beijing during the summer (averaged at $0.23 \pm 0.37 \ \mu g \ m^3$, in the range of 186 187 <u>0.06 to 2.30 μ g m⁻³) and winter (averaged at 0.59 \pm 0.42 μ g m⁻³, in the range of 0.06 to 1.94 μ g m⁻³</u> 188 $\frac{3}{2}$) of 2010-2011 (Cheng et al., 2013). The biomass burning tracer levels and ratios observed in this 189 study and other field studies are summarized in Table S1. The highest concentrations of 190 levoglucosan LG-in GC were observed on 31 October, 2016 with 4.37 μ g m⁻³, which is a sharp increase (over 30 times) of the minimum concentration (0.14 μ g m⁻³) during that period (Figure 1c). 191 192 Accordingly, the PM2.5-cal concentration during that period was also elevated (as high as 236 µg m⁻ ³) (Figure 1a). Secondary inorganic aerosol (sulfate, SO_4^{2-} ; nitrate, NO_3^{-} and ammonium, NH_4^+ , 193 194 SNA) species, were the major water soluble ions, accounting for 82.8% of total water soluble ions, the daily average values of which were $10.5 \pm 6.87 \ \mu g \ m^{-3}$, $15.9 \pm 9.29 \ \mu g \ m^{-3}$ and $10.9 \pm 5.51 \ \mu g$ 195 196 m⁻³, respectively (Table 1). SNA species exhibited a synchronous temporal trend (Figure 1c), while the NO₃ concentrations exceeded those of SO₄²⁻ at the GC site, in contrast to the results of previous 197 198 studies, e.g., Tan et al. (2016), who found SO_4^{2-} to be the dominant species in PM_{2.5} during winter 199 in 2006 in Beijing. Similarly, Chi et al., (2018) also found NO₃⁻ concentrations exceeded those of 200 SO_4^{2-} at both Beijing and GC sites during the winter in 2016, although they observed that NH_4^+ 201 was the dominant component of SNA (the concentrations of SO_4^{2-} , NO_3^- and NH_4^+ were 14.0 µg 202 m⁻³, 14.2 µg m⁻³, and 24.2 µg m⁻³, respectively).

3.2 Day-night variations in the characteristics of PM2.5 chemical components

204 Carbonaceous components and biomass burning tracers exhibited higher levels during 205 nighttime than daytime, while secondary inorganic ions showed the opposite pattern, i.e., higher concentrations during daytime than nighttime (Figure 2 and Figure S2). Besides, the gap of 206 207 carbonaceous components and anhydrosugars between daytime and nighttime (two-fold) was more 208 significant than for secondary inorganic ions. EC and primary organic componentsPOC are not 209 subject to significant differences in chemical reactions in ambient air between daytime and 210 nighttime, and they will be mainly influenced by the variations of the planetary boundary layer 211 (PBL) height. In the night, the PBL height decreases, compressing air pollutants into a shallow layer, 212 and subsequently resulting in faster accumulation and higher concentrations of pollutants (Zheng et al., 2015; Zhong et al., 2018; 2019). The contributions of OM and EC to PM_{2.5-cal} were observed to 213 214 be higher at nighttime (53.9% and 16.6%) than daytime (43.8% and 13.7%) as well (Figure 3).

215 Besides the influence from variations of the PBL height, the chemical degradation of 216 levoglucosan may occur due to photochemical reaction in the ambient aerosols during day-time, further enlarging the gap of levoglucosan levels in between daytime and nighttime (Sang et al., 2016; 217 218 Gensch et al., 2018). Consequently, T the contribution of levoglucosan to PM_{2.5-cal} during daytime 219 (0.370.45%) was observed to be considerably lower than that during nighttime (0.64%) (Figure 3). 220 However, secondary inorganic ions have an important formation pathway, i.e., photochemical 221 processing, during daytime. Thus, the secondary inorganic species (SO_4^{2-} , NO_3^{-} and NH_4^{+}) were 222 enhanced during daytime due to photochemical formation (Sun et al., 2013; Zheng et al., 2015; Wu et al., 2018). The mass contributions of SO_4^{2-} , NO_3^{-} and NH_4^+ to $PM_{2.5-cal}$ were decreased from 223 224 daytime (9.9%, 14.5% and 10.0%) to nighttime (6.5%, 9.6% and 7.1%) (Figure 3). Such an 225 enhancement in secondary transformations during daytime is more evident in terms of the sulfur 226 and nitrogen oxidation ratios (SOR and NOR, molar ratio of sulfate or nitrate to the sum of sulfate 227 and SO_2 or nitrate and NO_2), which have been used previously as indicators of secondary 228 transformations (Sun et al., 2013; Zheng et al., 2015). Both SOR and NOR during daytime were

229 <u>higher than those during nighttime (Figure S3), further confirming the elevated secondary</u>
230 formations of sulfate and nitrate during daytime.

231 In addition, the concentrations of other water-soluble inorganic ions, i.e., K⁺ and Cl⁻ during 232 nighttime (1.78 \pm 0.95 µg m⁻³ and 6.08 \pm 4.00 µg m⁻³) were higher than those in daytime (1.43 \pm $0.54 \,\mu g \,\mathrm{m}^{-3}$ and $4.33 \pm 2.30 \,\mu g \,\mathrm{m}^{-3}$), while their contributions to PM_{2.5-cal} were reversed, due to the 233 234 significant accumulation and higher concentrations of pollutants during nighttime. As Ca2+, Mg2+ 235 and Na⁺, mainly emitted from primary natural sources, such as dust, soil resuspension and sea salt, 236 are subject to more activity during the daytime and also influenced by the airflow dynamics, the contribution of those species in nighttime were lower than those during daytime, especially for Ca^{2+} , 237 238 decreasing from 2.2% in daytime to 0.9% at nighttime (Figure 3).

3.3 Biomass burning episodes and the impacts on chemical PM2.5 characteristics

240 An episode with high biomass burning tracer levels was encountered on 31 October, 2016. 241 The concentrations of <u>levoglucosanLG</u> in PM_{2.5} during this one-day episode (4.37 μ g m⁻³) were significantly higher than those during typical transition season at the GC site $(0.69 \pm 0.47 \ \mu g \ m^{-3})$ 242 243 (Figure 1d). Meanwhile, there was significant change in the meteorological conditions, i.e., the wind 244 direction changed from southwesterly to northerly winds (Figure S4). Northerly winds advected 245 cold and dry air masses, with the lowest hourly temperature observed at -5.3 °C (Figure S5). This notable temperature decline before the commencing of the operation of the central heating systems 246 247 should have caused intense combustion activities for heating purposes at the rural site. Moreover, 248 the synoptic situation on 31 October, 2016 was under weaker turbulence with low PBL height and 249 small wind speeds (Figure 1f). These worsened meteorological conditions would further enhance 250 aerosol accumulation.

Here, we mainly distinguish four sub-periods based on daily___levoglucosanLG concentrations during the time frame from 15 October to 23 November, 2016. The four periods were separated as follows: 15-30 October (Period I: Minor biomass burning), 31 October (Period II: Intensive biomass burning), 1-14 November (Period III: Major biomass burning), 15-23 November (Period IV: Heating season). Table 2 compares the concentrations of PM_{2.5-cal} mass, chemical components and gases at the GC site during these four periods, as well as the ratios between the intensive, major BB periods and heating season to minor BB period. The level of <u>levoglucosan LG</u>

258 during the intensive BB episode II was about 12 times of that during the minor BB period I. K⁺ and 259 Cl⁻, the common biomass burning tracers utilized in many studies (Duan et al., 2004; Cheng et al., 260 2013), were also observed with increased abundance during intensive BB episode II. When entering 261 into November, the weather was becoming cold, and thus combustion activities for heating in the 262 rural areas commenced, resulting in the ambient levels of levoglucosan LG to increase to 0.92 ± 0.47 263 µg m⁻³ during period III, about 3 times of those in Period I. The central heating systems in North 264 China cities were operated during period IV, and the ambient level of levoglucosanLG was observed 265 at $0.96 \pm 0.63 \,\mu g \,\mathrm{m}^{-3}$, which was similar to that observed in period III.

266 The concentrations of OC and EC were also observed to be strongly elevated in period II (Table 267 2), and especially OC levels increased to $\frac{59.996.3 \pm 25.3 \,\mu g}{1000}$ m⁻³ during the intensive BB episode II, nearly 6 times of that those during the minor BB period $(16.2 \pm 7.52 \,\mu g \, m^{-3})$. The levoglucosan/OC 268 269 ratio was utilized to estimate the effect of biomass burning to ambient organic aerosols. Accordingly, 270 levoglucosanLG/OC ratios sharply increased to 0.045 during period II, which is-was 271 obvious noticeably higher than during other periods in this study (Figure 1e). Moreover, this level is 272 also higher than most of the published field observations, i.e., at urban sites (Zhang et al., 2008; Cheng et al., 2013; Zhang et al., 2014), rural sites (Sang et al., 2013; Ho et al., 2014; Pietrogrande 273 274 et al., 2015; Mkoma et al., 2013) and agricultural sites (Ho et al., 2014; Jung et al., 2014), yet lower 275 than at an urban site in northern Italy during winter time (in the range of 0.01 to 0.13) (Pietrogrande 276 et al., 2015). This illustrates that biomass combustion played an important role in organic aerosol 277 pollution during the intensive BB episode II. However, During the major BB (period III) and heating season (period IV), due to other emissions of OC enhanced during the major BB episode (period III) 278 279 and heating season (period IV), i.e., combustion of coal and biofuel for heating, OC increased to a higher level (55.2 \pm 17.1 µgC m⁻³ and 69.4 \pm 24.6 µgC m⁻³, respectively). Due to the abundance of 280 281 organic aerosols, the contribution from biomass burning emission was thereby reduced and the 282 <u>levoglucosanLG</u>/OC ratios during periods III and IV decreased to 0.016 ± 0.005 and 0.014 ± 0.006 , 283 respectively, even lower than those <u>observed</u> in the minor BB period I (0.025 ± 0.008). 284 Compared to the carbonaceous components, the concentrations of secondary inorganic aerosol

species (SO₄²⁻, NO₃⁻, NH₄⁺) exhibited a different pattern, i.e., showing no obvious differences between minor BB period I and other three periods. The ratios of SO₄²⁻, NO₃⁻, NH₄⁺ during periods

287 II, III and IV to period I were all around 1.0 (Table 2), with no increasing trend. Moreover, the 288 relationships between levoglucosan-LG and OC (and EC) were better than those between 289 levoglucosanLG and SNA during daytime and nighttime (Figure S3). The precursor gases of SNA, 290 i.e., SO₂, NO, NO₂ and NH₃, were observed to have an increasing trend when biomass burning was 291 prevalent during periods III and IV, with the ratios to period I arranged from 1.13 to 1.90 (Table 2). 292 The time-series variations of the gases (SO₂, NO_x, NH₃, CO and O₃) and PBL during the sampling 293 period are shown in Figure S4. The primary emission gases were exhibited negative relationships 294 with PBL, while O₃ exhibited obvious positive relationship with PBL (Figure S5). Combustion from 295 different fossil fuels (coal, gasoline, diesel, etc.) and biomasses (straws, woods, leaves, etc.) can all 296 emit CO into the atmosphere (Streets et al., 2003; Chantara et al., 2019; Merico et al., 2020). Due 297 to the more abundant combustion in the colder weather, the concentrations of CO also increased to 298 1.65 ± 0.53 ppm and 1.18 ± 0.83 ppm during the major biomass burning period III and the heating 299 season period IV, respectively.

300 The combustion of biomass, especially of agricultural residues (e.g., wheat and corn straw) is 301 very common in the rural areas in North China during the autumn-winter transition period. During 302 the autumn harvest season in North China, wheat and corn straw burning is common practice, 303 resulting in more abundant fire spots when entering into November than period I (Figure 4). The 304 intense biomass burning event on 31 October, 2016 was also supported by air mass back trajectory 305 analysis (Figure 5), performed with the TrajStat software. Based on the 48 h back trajectories at the GC site at 00:00 (UTC time) on 1 November, 2016, the air mass at the GC site was restricted in the 306 region of Bejing-Tianjing-Hebei, the polluted area where fire spots were numerous. However, on 307 308 the previous and following day of this episode, i.e., 31 October and 2 November onward, the air 309 masses arriving at GC were advected from the northwest of Mongolia, where mostly desert areas 310 are present, with less farm land and rare biomass burning activities (Figure 5).

Mean percentiles of major components in $PM_{2.5}$ with respect to different BB pollution periods at GC site during the sampling time are shown in Figure 6. With the variation of BB pollution periods, the EC fraction seems to exhibit no obvious change during periods I, II and III, but slightly increased during the heating season (period IV), while the OC fraction increased significantly from 34.0% during the minor BB period I elevated to 65.4% during the intense BB period II. The

316 contributions of sulfate, nitrate and ammonium to PM2.5-cal all decreased sharply from the minor BB period to the intense period (Figure 6). This suggests that organic aerosol species become more 317 318 important during BB pollution periods, concerning their contribution to the PM_{2.5-cal}, while EC has no such character. The OM percentage during intense BB period II was 65.4%, about double of that 319 320 during the minor biomass burning period (34.0%), indicating that there was a large fraction of OM 321 in PM_{2.5-cal} originating from BB at the GC site during intensive BB period II. Opposite to OM, 322 contributions of secondary inorganic ions to PM_{2.5-cal} significantly decreased with the BB pollution becoming more severe. The contributions of SO₄²⁻, NO₃⁻ and NH₄⁺ to PM_{2.5-cal} during the minor 323 324 BB episode (11.6%, 20.5% and 12.5%) obviously declined during the intense BB episode (1.7393%, 325 7.7367% and 4.24%).

326 **3.4 Relationships among tracers during different biomass burning pollution**

327 periods

328 In addition to pollution level information of biomass burning molecular tracers, the ratios 329 between them could also be used to identify the different biomass types or indicate the burning 330 formation processes of atmospheric aerosols. Levoglucosan and mannosan showed a good 331 relationship during the entire sampling period (Figure 7a, r = 0.97, p < 0.01). The 332 levoglucosanLG/mannosanMN ratios during minor, intense, major biomass pollution and heating 333 season periods were observed at high values, i.e., 24.9, 24.1, 24.8 and 18.3 respectively (Table 2, Figure 7). LG and MN in the four periods showed a good relationship (Figure 7a, R^2 = 334 335 0.94). Compared to the former three episodes (24.1 to 24.9, averaged at 24.6), the 336 levoglucosan/mannosan ration during the heating season period (18.3) decreased by 25.6%. Based 337 on source emission studies, the levoglucosanLG/mannosanMN ratios from crop residue burning, 338 i.e., rice straw, wheat straw and corn straw, are similar and are characterized by high values 339 (averaged at 29, in the range of 12 to 55) (Zhang et al., 2007; Engling et al., 2009; Cheng et al., 340 2013; Jung et al., 2014), yet overlapped overlapping with those from hardwood (averaged at 28, in the range of 11 to 146) (Bari et al., 2009; Jung et al., 2014) and grass burning ($18.2 \pm 10.2 \Rightarrow 10.0$) 341 342 (Sullivan et al., 2008), while softwood is characterized by relatively lower 343 levoglucosanLG/mannosanMN ratios (averaged at 4.3, in the range of 2.5 to 4.7 < 5.0) (Engling et

al., 2006; Cheng et al., 2013; Jung et al., 2014). Subsequently, this declining trend of the
levoglucosan/mannosan ratios during the heating season period was partly causedcontributed by the
higherraised proportion of softwood combustion, which is characterized by relatively lower
levoglucosan/mannosan ratios. According to the local habits, soft woods, e.g. China fir and pine are
also commonly used as biofuels for stove heating in North China, since they allow sustainedlongtime heating duration.

350 The concentrations of <u>levoglucosanLG</u> and K^+ during minor, major BB episode and heating season were correlated well (Figure 7b, r = 0.84, p < 0.01), while the red dot of period II being off 351 352 from the fitted regression line. The levoglucosan LG/K^+ ratios during periods III and IV (0.520.51) 353 and 0.53) were similar to those during a BB episode at an urban site in Beijing during winter time 354 (levoglucosanLG/K⁺ = 0.51) (Cheng et al., 2013). However, the levoglucosanLG/K⁺ ratio during 355 the intense BB period II increased to 1.67, which was significantly higher than that in typical straw 356 combustion (< 1.0). Correspondingly, there was a significant drop in temperatures at the GC site 357 during period II, with the average daily temperature sharply decreasing from 7.5 °C on 30 Oct to 0.31 °C on 31 October, 2016, and the average temperature at night of 31 October even decreased to 358 359 -3.4 °C (Figure 1g). Hence, the combustion activities were apparently intense around the sampling 360 site for heating purposes. Compared to K^+ , there is a large enrichment of levoglucosan LG in wood 361 burning emissions, based on the results from previous biomass source combustion studies (Engling 362 et al., 2006; Chantara et al., 2019). The influence of softwood and/or other materials from softwood, which are commonly used as biofuels for stove heating in North China (Cheng et al., 2013; Zhou et 363 364 al., 2017), should be larger during this low temperature period. Moreover, levoglucosanLG/K⁺ ratios 365 also can be influenced by combustion conditions, i.e., smoldering versus flaming burns. Biofuels 366 are typically subject to smoldering combustion condition in residential stoves for heating purposes 367 in the rural areas in North China, which was reflected in relatively higher levoglucosanLG/K⁺ ratios 368 than during flaming combustion (Schkolnik et al., 2005; Lee et al., 2010).

369 **4. Summary and conclusion**

Anhydrosugars, including levoglucosan and mannosan, and water-soluble potassium ion were
employed as molecular tracers to investigate the characteristics of biomass burning activities as well
as chemical properties of ambient aerosols under different biomass burning pollution levels. The

measured daily average concentrations of evoglucosanLG, mannosanMN and K⁺ in PM_{2.5} during 373 a typical biomass burning season from 15 October to 30 November, 2016 were $0.79 \pm 0.75 \ \mu g \ m^{-3}$, 374 $0.03 \pm 0.03 \ \mu g \ m^{-3}$ and $1.52 \pm 0.62 \ \mu g \ m^{-3}$, respectively. The concentrations of carbonaceous 375 376 components and biomass burning tracers were observed higher at nighttime than daytime, while the 377 patterns of secondary inorganic ions (SO₄²⁻, NO₃⁻ and NH₄⁺) were opposite, since they were 378 enhanced by photochemical formation during daytime. An episode with extreme biomass burning 379 tracer levels was encountered on 31 October, 2016, with concentrations of levoglucosanLG as high as 4.37 µg m⁻³. Comparing the chemical compositions between different biomass burning periods, 380 it was apparent that biomass burning can considerably elevate the levels of organic components, 381 382 while not showing a significant effect on the production of secondary inorganic ions. Compared to 383 the other biomass burning episodes, the levoglucosan/mannosan ratios during the heating season 384 period slightly decreased, Moreover, due to more local soft wood and smoldering combustion taking 385 place for heating under the low temperatures, the while $levoglucosanLG/K^+$ ratio during the 386 intensive BB period was unusually higher than those in the other three biomass burning periods.

- 387
- 388

389 *Data availability*. The data used in this study can be obtained from this open
390 link: <u>https://pan.baidu.com/s/11bKUZff1KJbzNVxS3VsLaA</u> code: jvqx. It is also available from
391 the corresponding author upon request (lianglinlin@cma.gov.cn).

Author contributions. LL designed conducted all observations and drafted the paper. GE revised
the paper and improved the English writing. XL drew the Figure 4 and Figure 5. CL, WX, YC, ZD,
GZ, JS and XZ interpreted the data and discussed the results. All authors approved the final version
for publication.

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

Special issue statement. This article is part of the special issue– "In-depth study of air pollution
sources and processes within Beijing and its surrounding region (APHH-Beijing) (ACP/AMT
interjournal SI)". It is not associated with a conference.

400 Acknowledgements. This research is supported by the Beijing Natural Science Foundation

401 (8192055) and CAMS Fundamental Research Funds (No. 2017Z011). The authors would like to

- 402 acknowledge Yingli Yu and Ye Kuang for their help with PM_{2.5} samples collection; Hongbing
- 403 Cheng for help with chemical analyses.
- 404 *Financial support*. This research has been supported by the Beijing Natural Science Foundation
- 405 (No. 8192055), State Environmental Protection Key Laboratory of Sources and Control of Air
- 406 Pollution Complex (No. SCAPC201701) and Chinese Academy of Meteorological Sciences
- 407 Fundamental Research Funds (No. 2017Z011).

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657 658	Table 1. Average concentrations and the range of PM _{2.5-ca1} and its chemical components, biomass burning
659	tracers ($\mu g m^{-3}$), gaseous species, ratios of OC/EC and <u>levoglucosan</u> LG/OC, as well as meteorological
660	data observed at GC site at daytime, nighttime and whole day, respectively, during the sampling period
661	from 15 Oct to 23 Nov 2016.

Species	Daytime $(N = 34)$	Nighttime ($N = 3$)	3)	Whole period (N	Whole period $(N = 37)^*$	
	<u>Average</u> <u>c</u> Concentration	Range	<u>Average</u> <u>concentration</u> Co ncentration	Range	<u>Average</u> <u>concentration</u> C oncentration	Range
PM _{2.5-cal}	117 ± 58.8	19.0 - 225	170 ± 116	21.1 - 465	137 ± 72.4	23.3 - 319
OC	26.8 ± 15.7	3.78 - 64.8	61.6 ± 49.5	2.88 - 175	44.0 ± 31.0	4.13 - 117
EC	13.4 ± 8.49	1.44 - 34.0	30.9 ± 28.5	2.21 - 129	21.7 ± 15.8	2.46 - 74.9
TC	49.3 ± 27.6	5.76 - 124	92.5 ± 73.6	5.10 - 289	65.8 ± 44.1	7.36 - 192
OC/EC	2.02 ± 1.26	1.09 - 3.31	2.25 ± 1.04	1.04 - 6.72	1.95 ± 0.60	0.83 - 3.10
SO 4 ²⁻	12.1 ± 9.31	1.65 - 39.7	9.02 ± 6.22	1.55 - 23.2	10.5 ± 6.87	1.66 - 29.5
NO ₃ -	16.9 ± 9.96	1.85 - 41.2	13.1 ± 8.52	1.56 - 38.0	15.9 ± 9.29	2.40 - 45.2
Cl-	4.33 ± 2.30	0.82 - 9.46	6.08 ± 4.00	0.62 - 16.0	4.90 ± 2.46	0.93 - 9.37
$\mathbf{NH_{4}^{+}}$	11.7 ± 6.76	1.84 - 26.0	10.0 ± 5.75	1.33 - 22.2	10.9 ± 5.51	1.99 - 25.4
K^+	1.43 ± 0.54	0.20 - 2.64	1.78 ± 0.95	0.22 - 4.19	1.52 ± 0.62	0.50 - 2.96
Mg^{2+}	0.26 ± 0.14	0.07-0.64	0.19 ± 0.09	0.06 - 0.38	0.14 ± 0.12	0.04 - 0.43
Ca ²⁺	2.24 ± 1.01	1.02-4.75	1.56 ± 0.08	0.77 - 3.56	1.54 ± 0.90	0.49 - 3.84

Na ⁺	0.44 ± 0.17	0.10 - 0.79	0.43 ± 0.24	0.10 - 1.31	0.42 ± 0.17	0.11 - 0.88
NO3 ⁻ /SO4 ²⁻	1.67 ± 0.82	0.75 - 5.52	1.54 ± 0.57	0.74 - 3.50	1.65 ± 0.62	0.78 ± 3.96
Levoglucosan	0.57 ± 0.62	0.05 - 3.74	1.10 ± 0.99	0.05 - 4.82	0.79 ± 0.75	0.14 - 4.37
Mannosan	0.024 ± 0.023	0.00 - 0.14	0.05 ± 0.04	0.00 - 0.21	0.03 ± 0.03	0.00 - 0.18
levoglucosanLG/OC	0.018 ± 0.011	0.005 - 0.067	0.020 ± 0.010	0.004 - 0.047	0.020 ± 0.009	0.006 - 0.04
NO (ppb)	23.0 ± 14.7	2.07 - 56.0	45.9 ± 29.5	1.59 - 96.9	31.8 ± 18.3	1.81 - 68.5
NO ₂ (ppb)	25.8 ± 10.4	8.18 - 51.6	29.3 ± 9.37	8.81 - 51.1	26.6 ± 8.74	8.62 - 51.4
SO ₂ (ppb)	9.78 ± 4.96	3.11 - 22.5	9.63 ± 5.67	2.91 - 28.7	8.61 ± 4.04	3.37 - 20.4
CO (ppm)	0.96 ± 0.73	0.03 - 2.49	1.29 ± 1.04	0.02 - 3.26	1.05 ± 0.76	0.12 - 2.48
O ₃ (ppb)	13.0 ± 9.10	1.42 - 41.84	5.00 ± 5.73	1.60 - 24.30	9.25 ± 5.78	1.67 - 24.0
NH ₃ (ppb)	16.4 ± 11.3	1.68 - 46.2	18.3 ± 10.7	1.03 - 42.7	17.1 ± 9.88	1.46 - 44.4
Temperature (°C)	7.71 ± 4.01	- 2.07-15.9	3.30 ± 4.69	- 6.60 - 14.5	6.95 ± 4.58	- 4.33 - 15.4
Relative Humidity (%)	68 ± 17	31 - 98	85 ± 14	34 - 100	77 ± 13	48 - 99
Wind speed (m s ⁻¹)	1.43 ± 1.17	0.09 - 5.65	0.79 ± 1.55	0.03 - 7.19	1.07 ± 1.14	0.04 - 5.02

662	* Six whole-day samples were included used-in the data analysis of the "Whole period".
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668 669	Table 2. Concentrations of chemical components in PM _{2.5} aerosols as well as their ratios and gaseous
670	species collected at the GC site, during the four biomass burning periods (i.e., Minor, Intensive, Major
671	and Heating period) from 15 Oct to 23 Nov 2016.

Species	Period I (15-30 Oct) Minor BB	0 Oct) Period II (31 Oct) Intensive BB		Period III (1 -14, Nov) Major BB		Period IV (15 -23, Nov) Heating period	
	<u>Average</u> Cconcentration	<u>Average</u> concentration C oncentration	Ratio*	<u>Average</u> <u>concentration</u> Concentration	Ratio*	Average concentration ncentration	Ratio*
PM _{2.5-cal}	81.0 ± 44.5	235	2.91	163 ± 46.7	2.01	189 ± 83.0	2.33
Levoglucosan	0.36 ± 0.14	4.37	12.1	0.90 ± 0.37	2.50	0.96 ± 0.63	2.67
Mannosan	0.015 ± 0.005	0.18	12.0	0.038 ± 0.015	2.53	0.050 ± 0.026	3.33
OC	16.2 ± 7.52	96.3	5.93	55.2 ± 17.1	3.41	69.4 ± 24.6	4.28
EC	12.2 ± 5.85	36.0	2.96	25.5 ± 10.1	2.09	36.4 ± 21.5	2.98
TC	28.4 ± 13.1	132	4.66	80.9 ± 34.6	2.85	106 ± 55.3	3.73
SO 4 ²⁻	10.3 ± 8.96	4.56	0.44	11.8 ± 6.02	1.15	9.08 ± 3.87	0.88
NO ₃ -	16.6 ± 12.9	18.1	1.09	16.5 ± 6.42	0.99	12.6 ± 5.76	0.76
\mathbf{NH}_{4}^{+}	10.1 ± 7.40	10.0	0.99	12.0 ± 4.35	1.19	10.3 ± 3.62	1.02
\mathbf{K}^+	1.16 ± 0.36	2.61	2.25	1.76 ± 0.46	1.52	1.65 ± 0.84	1.42
Cl-	3.46 ± 1.97	7.49	2.16	5.58 ± 2.16	1.61	6.27 ± 2.58	1.81
OC/EC	1.53 ± 0.35	2.67	1.75	2.31 ± 0.59	1.51	2.04 ± 0.31	1.33
NO3 ⁻ /SO4 ²⁻	1.74 ± 0.60	3.96	2.28	1.50 ± 0.35	0.86	1.42 ± 0.47	0.82

levoglucosan- LG/OC	0.025 ± 0.008	0.045	1.80	0.016 ± 0.005	0.64	0.014 ± 0.006	0.56
<u>levoglucosan</u> LG /EC	0.039 ± 0.019	0.121	3.10	0.038 ± 0.017	0.97	0.028 ± 0.013	0.72
levoglucosan-	24.9 ± 4.44	24.1	0.97	24.8 ± 6.46		18.3 ± 4.27	
<u>LG/</u> <u>mannosan</u> MN			0.97		1.00		0.73
<u>levoglucosan</u> L G/K ⁺	0.36 ± 0.081	1.67	4.64	0.51 ± 0.16	1.42	0.53 ± 0.15	1.47
NO (ppb)	21.7 ± 12.5	21.7	1.00	39.6 ± 15.4	1.82	39.3 ± 23.6	1.81
NO ₂ (ppb)	21.8 ± 4.95	26.5	1.22	32.7 ± 7.27	1.50	24.6 ± 10.2	1.13
NO _X (ppb)	43.6 ± 16.3	48.2	1.11	72.4 ± 17.8	1.66	64.0 ± 33.4	1.47
SO ₂ (ppb)	5.83 ± 2.46	8.04	1.38	11.1 ± 4.10	1.90	9.75 ± 3.31	1.67
CO (ppm)	0.44 ± 0.33	0.70	1.59	1.65 ± 0.53	3.75	1.18 ± 0.83	2.68
O ₃ (ppb)	9.79 ± 4.88	23.2	2.37	7.51 ± 3.87	0.77	9.59 ± 7.55	0.98
NH ₃ (ppb)	14.3 ± 6.12	11.1	0.78	18.6 ± 8.03	1.30	21.2 ± 14.2	1.48

672 *: indicates that the ratios of the heating period, intense BB period or major biomass burning period

673 were divided by those from the minor BB period.

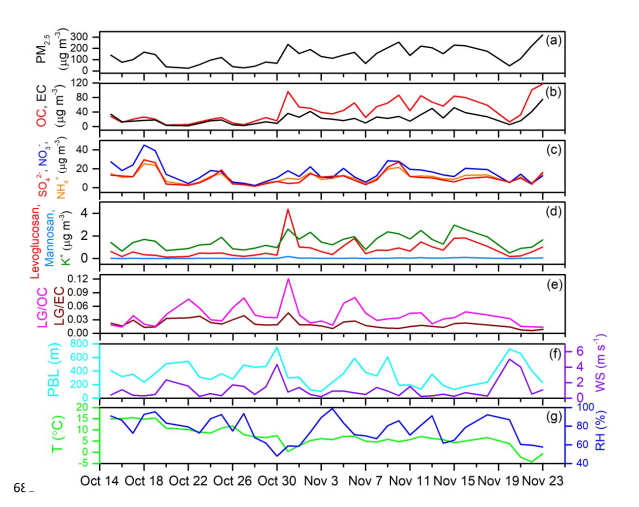
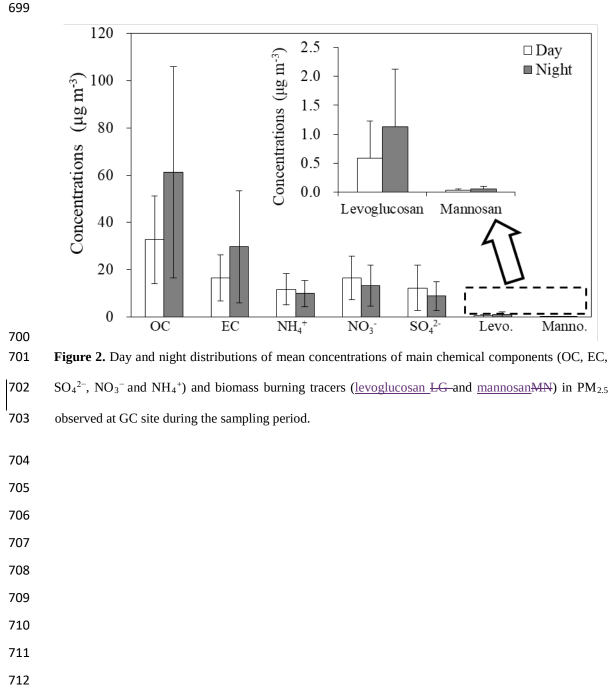
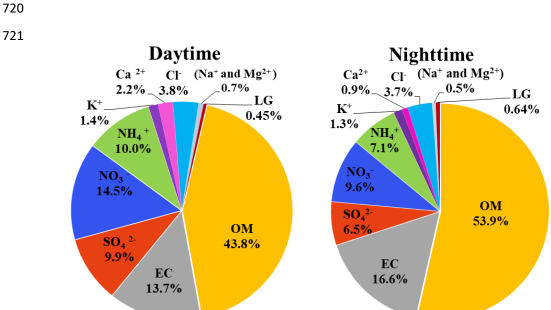


Figure 1. Time-series variation obtained for PM_{2.5-cal} and its major components, biomass burning tracers as well as meteorological factors at the GC site during the sampling period from 15 Oct to 23 Nov 2016. (a) PM_{2.5-cal_a} and its major components (b) OC,— and EC, (c) secondary inorganic aerosols, i.e., SO_4^{2-} , MO₃⁻⁻ and NH₄[±], (d) biomass burning tracers (levoglucosanLG, mannosanMN and K⁺)_a (e) ratios of levoglucosan to OC (LG/OC) and levoglucosan to EC (LG/EC), and meteorological factors (f) PBL and wind speed (WS), (g) temperature (T) and; relative humidity (RH), wind speed (WS) and rainfall) at the GC site during the sampling period from 15 Oct to 23 Nov 2016.

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- 723 Figure 3. Percent contributions of individual component mass concentrations to total estimated PM_{2.5-cal}
- mass in daytime and nighttime during the sampling period.

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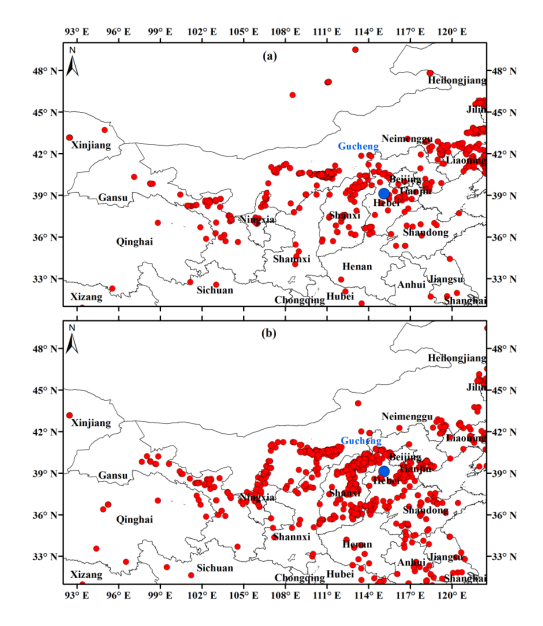




Figure 4. Fire spots at GC site and the surrounding provinces from (a) 15-30 October, 2016 and (b) 1 -



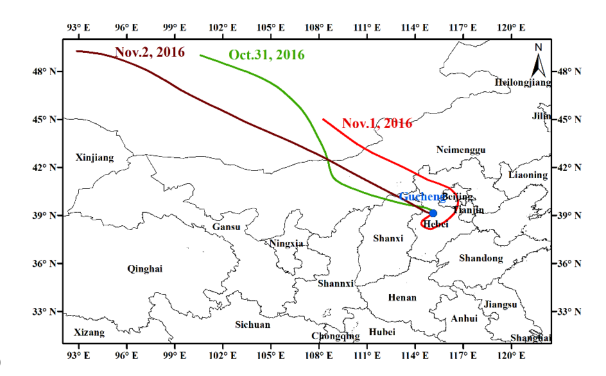




Figure 5. 48 h back trajectories at 500 m at GC site (39°09'N, 115°44'E) at 00:00 (UTC time) from 31

- 742 October to 2 November, 2016.

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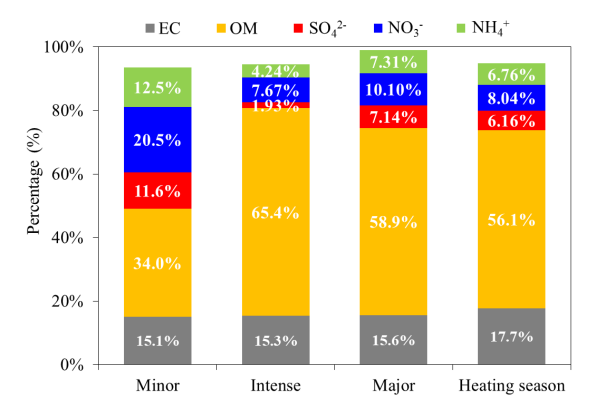


Figure 6. Mean percentiles of major components in PM_{2.5} with respect to different biomass burning
pollution periods at GC site during the sampling time.

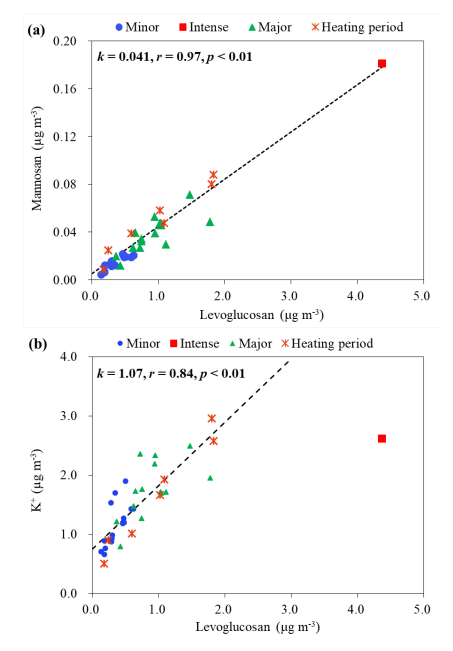






Figure 7. Scatter plots of (a) levoglucosan versus mannosan, (b) levoglucosan versus K⁺. <u>Statistical</u>

analysis of sampling data was conducted with the linear fitting method.