Title: “Chemical characteristics of PM$_{2.5}$: Impact of biomass burning at an agricultural site of the North China Plain during a season of transition”

Anonymous Referee #1

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General Comments:

The study of Liang et al. aimed to gain insights about the abundance of biomass burning smoke during the autumn-winter transition season, and explore the impact of biomass burning activities on the chemical properties of ambient aerosols. The data analysis was relatively straightforward, however, there were some issues with interpretation, especially there were some places with subjective interpretations. Although chemical characteristics of PM$_{2.5}$ composition in minor, intensive and major biomass burning periods were presented in this study, contributions by biomass burning to PM$_{2.5}$ or carbonaceous components was not quantified. There is nothing that is particularly novel in the study compared to previous work, and the environmental significance of the findings is not clear. The manuscript cannot be accepted for publication in its current form. Although the manuscript is not difficult to read, there are numerous grammar and language issues, which need to be addressed and improved. Several examples are listed below.

Our reply: We thank the referee for his/her comments. In fact, to the best of our knowledge, this study is the first one to characterize the biomass burning pollution status at a heavily polluted rural site in Hebei province during the autumn-winter transition season, following the corn harvest. The results can provide valuable source information of the biomass burning activities in the entire North China region. Moreover, we captured a unique episode with extreme biomass burning pollution, with concentrations of levoglucosan as high as 4.37 µg m$^{-3}$. Based on the multi-analysis of biomass burning molecular tracers, back trajectory analysis, fire activity data and synoptic condition, the formation process and chemical character of this severe biomass burning pollution episode were discussed in detail. In addition, combined with other chemical components analysis,
it revealed that the type of biomass burning impacts the different types of chemical components in ambient aerosol, which were have rarely been reported by previous work. Therefore, we believe that the results obtained in this study are novel and valuable for gaining additional insights into the status and impacts of biomass burning pollution in source regions of North China. Moreover, our study provides evidence to the government that it’s important to pay more attention to the residential burning activities in the North China Plain, and impose burning restrictions to reduce the air pollution contributed by biomass burning.

As for the contributions of biomass burning to carbonaceous aerosol and PM$_{2.5}$, we quantified them by the molecular tracer approach and discussed the results in a companion paper, as it would render this paper too long otherwise. Nevertheless, we have added the discussion of LG/OC ratios in the revised paper.

(1) Line 19: It's better to say “The measured daily average concentrations of LG, MN and K$^+$ during this study period were 0.79 ± 0.75 μg m$^{-3}$, 0.03±0.03 μg m$^{-3}$ and 1.52±0.62μg m$^{-3}$, respectively”

Our reply: According to the referee’s comment, we changed the sentence as follows:

“The measured daily average concentrations of LG, MN and K$^+$ during this study period were 0.79 ± 0.75 μg m$^{-3}$, 0.03 ± 0.03 μg m$^{-3}$ and 1.52 ± 0.62 μg m$^{-3}$, respectively” (See Lines 19-20)

(2) Line 53: “abundant” can be replaced by “extensive”

Our reply: According to the referee’s comment, we replaced “abundant” with “extensive” in the revised paper.

“The precursors and formation pathways of SOA from biomass burning emissions were investigated by extensive field observations.” (See Line 51)

(3) Line 84-85: “The study results demonstrate” is better to be revised to, for example, “The results of this study demonstrate”, or “The results presented in this study demonstrate”

Our reply: According to the referee’s suggestion, we changed the “The study results
“The results of this study demonstrated” the biomass burning pollution status in the rural atmosphere of North China and explore the impact of biomass burning activities on the chemical properties of ambient aerosols.” (See Lines 83-85)

(4) Line 133: “All measurement data quality was controlled according to standards...”

**Our reply:** According to the referee’s suggestion, we corrected the sentence as:

“All measurement data quality was controlled according to standard gases (Xu et al., 2019; Lin et al., 2011; Meng et al., 2018; Ge et al., 2018)” in the revised paper. (See Lines 133-134)

(5) Line 137: “on” should be replaced by “at”

**Our reply:** According to the referee’s suggestion, we changed the “on” to “at” in the revised paper.

“The meteorological parameters, including air temperature, relative humidity (RH) and wind speed at a 24-h resolution at the GC site are presented in Fig. 1” (See Lines 136-137)

(6) Line 197: repeated word “average” in “daily average wind speed averaged at: : : :” ...

**Our reply:** According to the referee’s suggestion, we rewrote the sentence in the revised paper.

“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.” (See Lines 196-197)

**Other comments:**

(7) Line 30-32: Why the finding that K⁺ did not increased as much as LG during extensive BB episodes indicated there were other sources of K⁺ in the study region?

**Our reply:** Generally, the ratio of LG to K⁺ should be consistent for the same type of biomass burning emission. Based on the published controlled biomass burning experiments, LG/K⁺ ratios of straws, i.e., wheat straw, corn straw and rice straw, were measured with low values (< 1.0),
while those for soft and hard wood were averaged at high values (>10.0) (Engling et al. 2006; Cheng et al., 2013; Sullivan et al., 2008). The LG/K⁺ ratios during periods I, III and IV (0.36, 0.52 and 0.53) in this study were similar to the observations from emission of straw burning. On the other hand, the average LG/K⁺ ratio during the intense BB period II was observed at 1.67, significantly higher than typical straw burning ratios. This indicated that besides straw burning, there should be other types of biomass burning during the intense BB period II.

The indication of other sources of K⁺ during the intense biomass burning period II was not suitable, and we corrected the description in the revised manuscript.

“Moreover, the LG/MN ratios in different BB periods were consistent, while the LG/K⁺ ratio during intensive BB period was significantly elevated. This may be due to more local soft wood and smoldering combustion taking place for heating under the low prevailing temperatures; these processes tend to show more efficient formation mechanisms of levoglucosan relative to K⁺. ” (See Lines 27-31)

(8) Lin 84: “GC” should be defined here.

Our reply: According to the referee’s suggestion, we defined the GC in the revised paper.

“In this paper, we focus on quantifying multiple biomass burning tracers, i.e., LG, MN and K⁺ as well as other chemical species in PM₂.₅ in (Gucheng, GC) during the autumn-winter transition biomass burning season.” (See Line 82)

(9) Line 103: Why quartz filters were prebaked in such a high temperature (850°C)? This is different from the temperature widely used in other studies.

Our reply: In this study, quartz fiber filters were prebaked at 850 °C to remove carbonaceous material. When analyzed the IMPROVE-A temperature protocol, the samples are heated as high at to 840 °C to detect OC and EC. In order to be consistent with the OC/EC analysis by IMPROVE-A temperature protocol, the quartz filters were prebaked at 850 °C to make sure all carbonaceous material was removed prior to sample collection.

(10) Line 128: it should be “0.82 gC/cm²”
Our reply: According to the referee’s suggestion, the detection limit of OC was corrected as 0.82 µgC cm\(^{-2}\) in the revised paper. (See Line 128)

(11) Line 163: In this study, concentrations of EC were higher than SO\(_4^{2-}\), NO\(_3^{-}\) and NH\(_4^{+}\), accounting for 13 - 17% of the PM\(_{2.5}\) mass. This is different from previous studies. Could the authors please explain why there were such high EC concentrations?

Our reply: The average concentrations of SO\(_4^{2-}\), NO\(_3^{-}\) and NH\(_4^{+}\) in this study were similar as those observed by Chi et al., (2018) at GC during winter time in 2016. Unfortunately, there is no observation of OC/EC at Gucheng during the autumn or winter season published to compare our results to. There maybe three reasons for the high EC values measured in this study. Firstly, GC is a rural site in the heavily polluted Hebei province, with the worst air quality in China, where abundant industries are located, such as power, steel, chemical industry, etc. Heavily polluted cities and regions are located throughout Hebei province, i.e., Shijiazhuang, Xingtai, Handan, Cangzhou, Hengshui, all of which are located south and southwest of the GC site. Based on the source identification of trace elements, industry is the second source in PM\(_{2.5}\) at the GC site, and it contributed ~ 23% to PM\(_{2.5}\) (Liu et al., 2020). Moreover, the GC station is also surrounded by agricultural fields. The observation time in this study was following the autumn crop harvest season, and agricultural diesel vehicles were widely used during this time, which emit abundant EC. In addition, when entering into November, coal combustion for heating in the North China started to increase. Thus, due to these abundant sources, i.e., industry, diesel vehicles, coal combustion, together with biomass burning, EC was observed at such high values at the GC site.

(12) Line 170-181: There might be other reasons for the different distributions of secondary inorganic ions in different studies. For example, the major sources might be variable with seasons and sites.

Our reply: We agree with this comment, and realize that there were many reasons, i.e., meteorological conditions, emissions of precursors, regional transport, etc., for the different distributions of secondary inorganic ions in different studies. We compared the values of SNA in our study with the results from Chi et al., (2018), which were also observed at the GC site, yet
during winter of 2016, both studies found NO\textsubscript{3}\textsuperscript{-} concentrations to exceed those of SO\textsubscript{4}\textsuperscript{2-}. However, these authors observed NH\textsubscript{4}\textsuperscript{+} to be the dominant component of SNA, while NO\textsubscript{3}\textsuperscript{-} was the most abundant ion observed in this study.

(13) Line 188-192: What about the variations of levoglucosan/OC ratio?

**Our reply:** According to the referee’s suggestion, we added the LG/OC ratios in Table 1, and also added the discussion of the variation of LG/OC ratios in the revised paper.

“Accordingly, the LG/OC ratio increased to 0.045 during period II, which is 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 et al., 2015; Mkoma et al., 2013) and agricultural sites (Ho et al., 2014; Jung et al., 2014), yet lower than at an urban site in northern Italy during winter time (Pietrogrande et al., 2015). During the major BB (period III) and heating season (period IV), due to the combustion of coal and biofuel for heating, the organic carbon component remained at a high level (55.2 ±17.1 µg m\textsuperscript{-3} and 69.4 ± 24.6 µg m\textsuperscript{-3}, respectively). These levels are more than 3 times of that during the minor BB period I. Due to the abundance of organic aerosols, the LG/OC ratios during periods III and IV decreased to 0.016 ± 0.005 and 0.014 ± 0.006, respectively, even lower than those in the minor BB period I (0.025 ± 0.008).” (See Lines 276-286)

(14) Line 216-220: The authors seemed to attribute the more significant differences of carbonaceous components during the nighttime vs. daytime compared to secondary inorganic ions to PBL. I can’t agree. Please explain/clarify how PBL cause different accumulations or influences on carbonaceous components and secondary inorganic ions.

**Our reply:** In fact, we don’t mean PBL can cause different influences on carbonaceous components and secondary inorganic ions. We meant carbonaceous components, except secondary organic components, as well as anhydrosugars, were mainly influenced by the variations of the PBL height. SNA can not only be impacted by PBL variations, but is also affected by the photochemical formation processes during daytime.

We discuss this in the revised paper as follows: “That may be due to the variations of
pollutant concentrations not only being controlled by the chemical reactions but also being subject to the influence of the planetary boundary layer (PBL) development. In the night, the PBL height decreases, compressing air pollutants into a shallow layer, and subsequently resulting in faster accumulation and higher concentrations of pollutants (Zheng et al., 2015; Zhong et al., 2018; 2019).” (See Lines 216-220)

(15) Line 220-223: I don’t agree with the authors that there are no significant differences in chemical reactions of carbonaceous components and anhydrosugars during daytime and nighttime. For example, OC include primary and secondary organics, not only the sources but also the formation mechanisms or chemical reactions could be diffident during daytime and nighttime. Besides, levoglucosan in the atmosphere is also not stable and could undergo atmospheric chemical degradations according to previous studies.

Our reply: We agree that the chemical reactions of carbonaceous components and anhydrosugars during daytime and nighttime are not the same. In order to make the description more rigorous, we rewrote this section in the revised paper.

“Elemental carbon and primary organic components are not subject to significant differences in chemical reactions in ambient air between daytime and nighttime, and they will be mainly influenced by the variations of the PBL height. The contributions of OM and EC to PM$_{2.5}$ were observed to be higher at nighttime (53.9% and 16.6%) than daytime (43.8% and 13.7%) as well. Moreover, the chemical degradation of levoglucosan may occur due to photochemical reaction in the ambient aerosols, further lowering the levoglucosan levels in daytime (Sang et al., 2016; Gensch et al., 2018). Correspondingly, the contribution of LG to PM$_{2.5}$ during nighttime (0.64%) was observed to be higher than that during daytime (0.37%). However, secondary inorganic ions have an important formation path, i.e., photochemical processing, during daytime. Thus, the secondary species (SO$_4^{2-}$, NO$_3^-$ and NH$_4^+$) were enhanced during daytime due to photochemical formation (Fig. 2 and Fig. 3).” (See Lines 220-231)

(16) Line 226-231: How about the difference in secondary transformations of secondary inorganic ions during the daytime and nighttime?
Our reply: The topic of this paper is about the impact of biomass burning activities on the chemical properties of ambient aerosols. The difference of secondary inorganic ion processes during the daytime and nighttime is not associated with the main focus of this paper. It is a very complex research topic and should be discussed in more detail in a separate paper. In any case, we thank the referee for this valuable comment.

(17) Line 277-279: What did the authors mean “the relationships between LG and OC, EC during daytime and nighttime were both better than those with SNA”?

Our reply: We meant to say that the relationships between LG and OC (and EC) during daytime and nighttime, were better than the relationships between LG and SNA during daytime and nighttime. In order to make the description more clear, we have rewritten this sentence in the revised manuscript.

“Moreover, the relationships between LG and OC (and EC) were better than those between LG and SNA during daytime and nighttime (Fig. S3).” (See Lines 290-292)

Figure S3. Correlations between LG and OC as well as EC during (a) daytime and (b) nighttime, and scatter plot
of LG versus SNA (i.e., SO$_4^{2-}$, NO$_3^{-}$ and NH$_4^{+}$) during (c) daytime and (d) nighttime.

(18) Line 268-269: Are there any evidences for “frequent heating activities in form of straw burning”?

Our reply: We thank the anonymous referees for this comment. In fact, because the ambient levels of K$^+$ and Cl$^-$ (which are common tracers for biomass burning sources, and emissions from straw burning in particular) strongly increased during period II, it can be deduced that heating activities occurred in form of straw burning in period II. In order to make the description more reasonable, the sentence was rewritten in the revised manuscript.

“K$^+$ and Cl$^-$, the common biomass burning tracers utilized in many studies (Duan et al., 2004; Cheng et al., 2013), were also observed with increased abundance during period II. Thus, it can be inferred that the episode on 31 October, 2016 with high PM$_{2.5}$ levels was apparently caused by intensive biomass combustion activities in the North China Plain.” (See Lines 263-266)

(19) Line 287: “in the range of 1.38 to 1.82” seems only for SO$_2$ but not for other SNA precursors.

Our reply: According to the referee’s suggestion, the sentence was rewritten in the revised paper.

“The precursor gases of SNA, i.e., SO$_2$, NO, NO$_2$ and NH$_3$, were observed to have an increasing trend when biomass burning was prevalent during periods III and IV, with the ratios to period I arranged from 1.13 to 1.90 (Table 2).” (See Lines 296-299)

(20) Line 288-290: It’s difficult to see the positive and negative relationships between gas precursors and PBL. Maybe better to change to scatter plots.

Our reply: According to the referee’s suggestion, we added the scatter plots figure of the relationships between gas precursors and PBL in the revised paper. (See Figure S5)
If so, how to explain the lower CO concentrations in period II, compared to period III?

**Our reply:** CO can be emitted from varied combustion sources, i.e., fossil fuel combustion and biomass burning. The ambient CO levels are not only impacted by different kinds of burning materials, but also influenced by meteorological conditions and regional transport. In order to make the description more rigorous, we rewrote the sentences in the revised paper.

“Combustion from different fossil fuels (coal, gasoline, diesel, etc.) and biomasses (straws, woods, leaves, etc.) can all emit CO into the atmosphere (Streets et al., 2003; Chantara et al., 2019; Merico et al., 2020). Due to the more abundant combustion in the colder weather, the concentrations of CO also increased to 1.65 ± 0.53 ppm and 1.18 ± 0.83 ppm during the major biomass burning period III and the heating season period IV, respectively.” (See Lines 302-306)

(22) Line 305-308: Seen from Fig. 4 and Fig. 5, there were also a large number of fire spots in the northwest. But why the authors stated that the air masses were “with rare biomass burning activities”?

**Our reply:** Fig. 4 reflects the total fire spots in two periods, i.e., before November (period I) and after November (period III and IV). Fig. 5 shows the back trajectories for three days (31 Oct, 1-2
Nov, 2016). In fact, we meant to say that there were rare biomass burning activities 
\textbf{northwest of Monglia, NOT northwest of the GC site.}

“the air masses arriving at GC were advected from the northwest of Mongolia, where mostly desert areas are present, with less farm land and rare biomass burning activities (Fig. 5).”  

(23) \textbf{Fig. 6: The authors presented LG together with OM. Do you mean LG is not belonging to OM?}

\textbf{Our reply:} We thank the anonymous referees for this valuable comment. LG certainly belongs to OM. In order to make the description more accurate, we deleted the LG data and re-draw the Figure 6 in the revised paper.  

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig6}
\caption{Mean percentiles of major components in PM$_{2.5}$ with respect to different biomass burning pollution periods at GC site during the sampling time.}
\end{figure}

(24) \textbf{Line 312-313: Similar to comments on Line 163, please explain why EC accounted for such a large fraction of PM$_{2.5}$ mass, and stable in all days.}

\textbf{Our reply:} With the variation of biomass burning pollution periods, the EC fraction seems to exhibit no obvious change during periods I, II and III, but slightly increased in the heating season (period IV). However, the daily mass concentrations of EC obvious variability, in the range of \(2.46 - 74.9 \ \mu g \ m^{-3}\). Compared to the minor period I, the average EC concentrations during other
polluted periods were all observed to be elevated (Table 2).

As for why EC accounted for such a large fraction of PM$_{2.5}$ mass in this study, there may be two main reasons. First, there were abundant sources of EC at the GC site during the observation period, i.e., industry, vehicle exhaust, coal combustion, together with biomass burning, as we replied to the earlier comment (11). Secondly, the mass concentration of PM$_{2.5}$ was reconstituted by the sum of carbonaceous components ($1.6 \times$OC + EC) and inorganic ions ($\text{SO}_4^{2-} + \text{NH}_4^+ + \text{NO}_3^- + \text{Cl}^- + \text{Ca}^{2+} + \text{Na}^+ + \text{K}^+ + \text{Mg}^{2+}$) in this study. Although this approach was widely used by many researchers, it may underestimated the actual value of PM$_{2.5}$, since there may exist some undetectable chemical components, besides carbonaceous species and water soluble ions. Thus, EC taking such large faction of PM$_{2.5}$ mass in this study appears to be reasonable.

(25) Line 354-359: Please revise “a time of strong process decrease in temperature”. And maybe it’s better to look into the reason why temperature dropped so quickly and significantly, and plot time series of temperature during these days. Besides, please add the sampling time for daytime and nighttime samples.

Our reply: According to the referee’s suggestion, we rewrote the sentence as “There was a significant drop in temperatures at the GC site during period II, with the average daily temperature sharply decreasing from 7.5 °C on 30 Oct to 0.31 °C on 31 Oct, 20106, while the average temperature at night of 31 Oct decreased to -3.4 °C (Fig.1).” (See Lines 390-392)

Moreover, we added the sampling time for daytime and nighttime samples section 2.1, site descriptions and sampling.

“The daytime samples were collected from 07:00 to 19:00, while nighttime samples were collected from 19:00 to 07:00 local time of the next day.” (See Lines 100-101)

(26) Line 359-361: I agree there are other sources for K$^+$. But I’m wondering if there were fireworks and fertilizers during the study period.

Our reply: We thank the anonymous referees for this valuable comment. As far as we know, there were no fireworks or obvious use of fertilizers during the observation period. We admit that the description of fireworks and fertilizers as other sources of K$^+$ was not suitable here, and we
corrected the discussion in the revised manuscript.

“Based on the results from previous biomass source combustion studies (Engling et al., 2006; Chantara et al., 2019), compared to K+, it appears that there is a large enrichment of LG during wood burning with efficient formation of LG during the flaming phase. This high LG/K+ ratio in period II is probably the most representative of the local BB aerosol, similar to the smoke aerosols from soft wood combustion (China fir and red pine) (Sang et al., 2013; 2020).” (See Lines 385-389)

(27) Line 361-363: Since K+ is widely used as a tracer for biomass burning. It's better to illustrate from the emission characteristics of K+ and levoglucosan from biomass burning, and influences of combustion conditions (flaming and smoldering) and fuel types.

Our reply: We thank the anonymous referees for this valuable comment. We extended the discussion of the reasons for high values of the LG/K+ ratio in period II, from the emission characteristics of LG and K+ of different types of biomass burning and combustion conditions (flaming and smoldering) in the revised paper.

“Based on the results from previous biomass source combustion studies (Engling et al., 2006; Chantara et al., 2019), compared to K+, it appears that there is a large enrichment of LG in wood burning with efficient formation of LG during the flaming phase. This high LG/K+ ratio in period II is probably the most representative of the local BB aerosol, similar to the smoke aerosols from wood combustion (China fir and red pine) (Sang et al., 2013; 2020). There was a significant drop in temperatures at the GC site during period II, with the average daily temperature sharply decreasing from 7.5 °C on 30 Oct to 0.31 °C on 31 Oct, 20106, while the average temperature at night of 31 Oct decreased to -3.4 °C (Fig.1). Thus, combustion activities were apparently intense around the sampling site for heating purposes, in form of burning of straws, as well as local soft woods, since these fuels are also commonly used in rural areas of North China, i.e., pine, poplar, China fir, etc.

Moreover, LG/K+ ratios also can be influenced by combustion conditions, i.e., smoldering and flaming burns. Compared to flaming combustion, smoldering burns are characterized by more efficient levoglucosan formation, whereas K+ emissions are relative lower. Consequently,
relatively high LG/K⁺ ratios were observed in smoldering combustion experiments (e.g., Schkolnik et al., 2005; Lee et al., 2010). Thus, the elevated LG/K⁺ ratios during period II can be partially explained by more abundant smoldering combustion under the low prevailing temperatures. This is consistent with the common heating custom in the rural areas in North China, where biofuels are typically subject to smoldering combustion in residential stoves for heating purposes.” (See Lines 385-402)

References:


