

Manuscript No.: EGUSPHERE-2023-830

Title: Vertical profiles of trace gas and aerosol properties over the Eastern North Atlantic: Variations with season and synoptic condition

SUMMARY:

1. In the manuscript the authors study the vertical profile of aerosol particle properties over the Southern Great Plains (SGP) atmospheric observatory. The vertical profiles were measured using an instrumented Gulfstream I aircraft during two intensive campaigns in spring and summer 2016.

The observed vertical profiles are explained by the authors using aerosol dynamics and transport processes. The results add to the understanding of aerosol properties at SGP and motivate further studies. The text is clear, well-structured and the topic is scientifically relevant. I can recommend the manuscript for publication in ACP after the below comments have been addressed.

Response: We thank the reviewer for the positive feedback. We have thoroughly revised the manuscript based on the comments provided. The point-to-point responses can be found as follows.

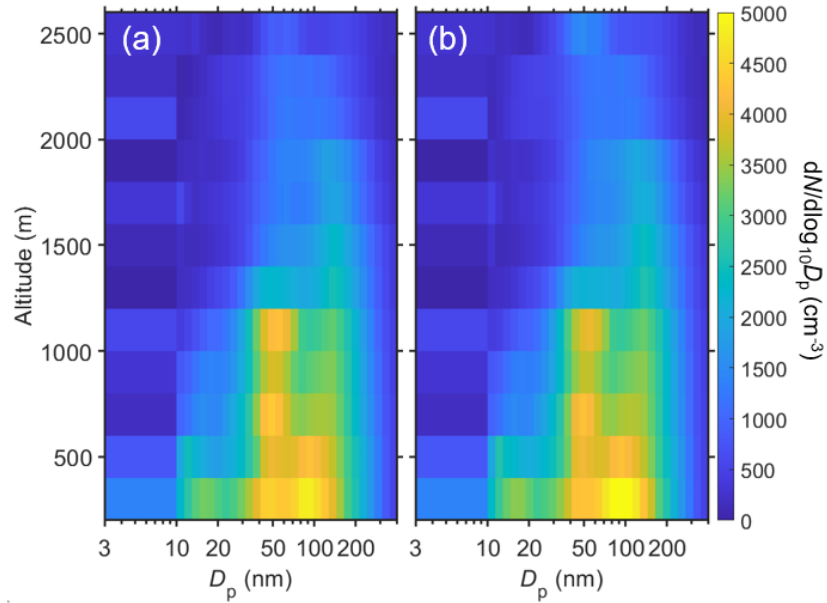
COMMENTS:

2. Fig 1: It appears that the flights were done on average about two hours later during the spring campaign than during the summer campaign. Would the time difference contribute to a possible bias when comparing the time periods? Is it possible to check that the results don't change by comparing only overlapping times of day?

Response: The U.S. daylight-saving time of the year 2016 (when HI-SCALE was conducted) starts on March 13 and ends on November 6. The campaign was conducted between April 25 and September 23, which means that there was no change in the local time due to daylight saving. The reason the flights were earlier and shorter (in general) during the summer campaign is because of the hotter temperatures, which affect flight operations. It is more difficult to lift off with a full payload with hot temperatures (when the temperature is above ~ 32 °C), and it consumes more fuel to fly. Therefore, the science team flew earlier in the morning during the summer campaign to have the coolest temperatures possible.

Regarding the impact of the flight time on temperature, the difference in time by a few hours is unlikely to cause significant bias. The boundary layer (BL) development is different for spring and summer, with BL likely developing faster during the warmer summer period. Moreover, time is one of the many factors influencing the vertical heterogeneity of aerosols, and our manuscript tries to identify these factors and analyze their influences. We also discussed the time effect in the manuscript regarding how boundary layer development affected aerosol vertical distribution in Section 3.4.1.

To address the reviewer's comment, we plotted the averaged vertical size distributions during the summer IOP excluding the two late afternoon flights on September 15, 2016 and September 20, 2016 (panel (a) of the figure below), and the results are very similar compared to the results that include all flight times (panel (b) of the figure below).



3. Fig 2 a and b: Add a scale bar indicating horizontal distance.

Response: The scale bars indicating horizontal distance have been added to **Fig. 2**. The revised figure can be found below.

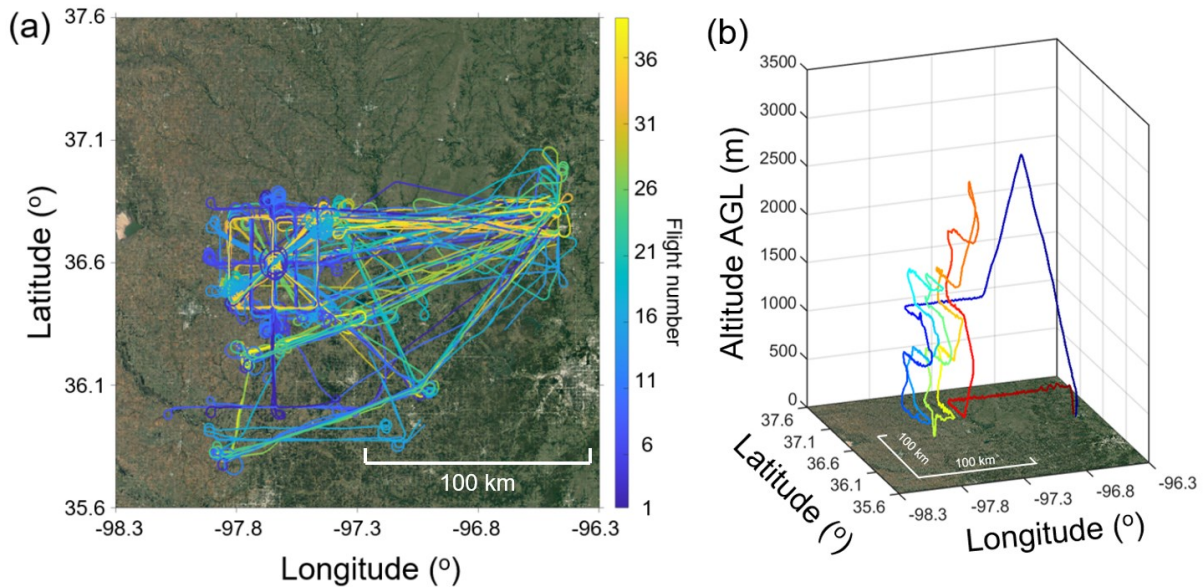


Fig. 2. (a) Flight patterns used during the HI-SCALE campaign. A total number of 38 flights were conducted over the SGP observatory. Different colors of the flight patterns represent different flight numbers. (b) An example of the flight pattern on May 3, 2016. The color represents the local time in hours. Maps are obtained from ©Google Maps.

4. Fig 3: For evaluating the source area better it would be helpful to see the distribution of back trajectories associated with each cluster.

Response: The trajectories associated with each cluster in **Fig. 3** are now added as **Figs. S3** and **S4**. The added figures can be found below.

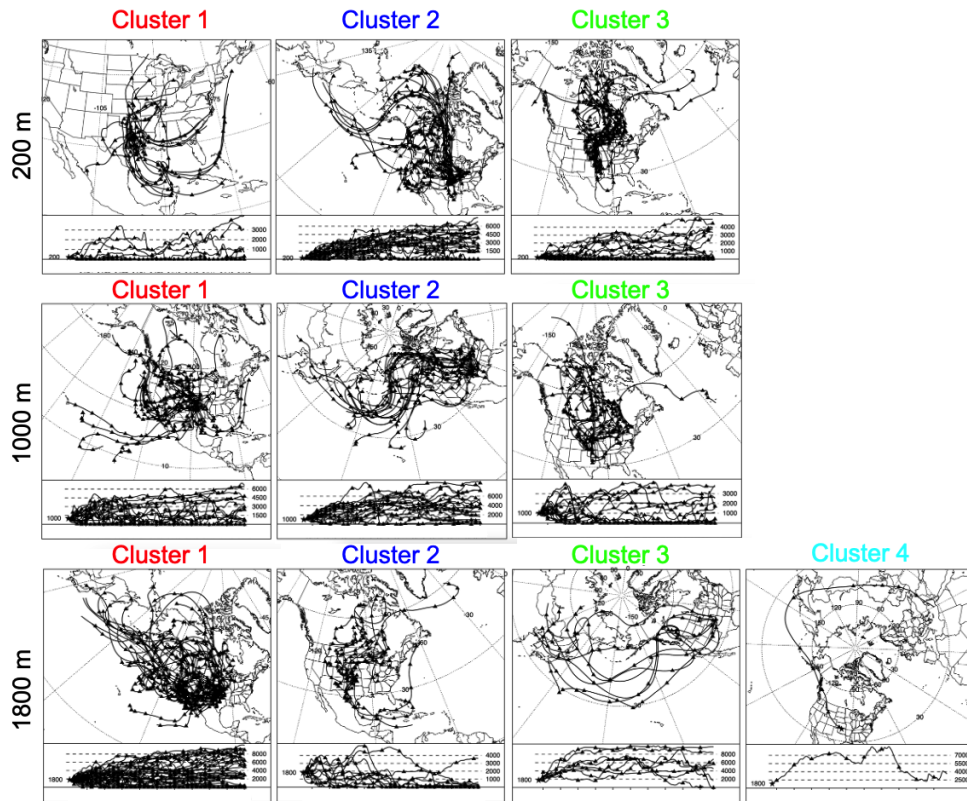


Fig. S3. Trajectories associated with each cluster during the spring campaign shown in Fig. 3.

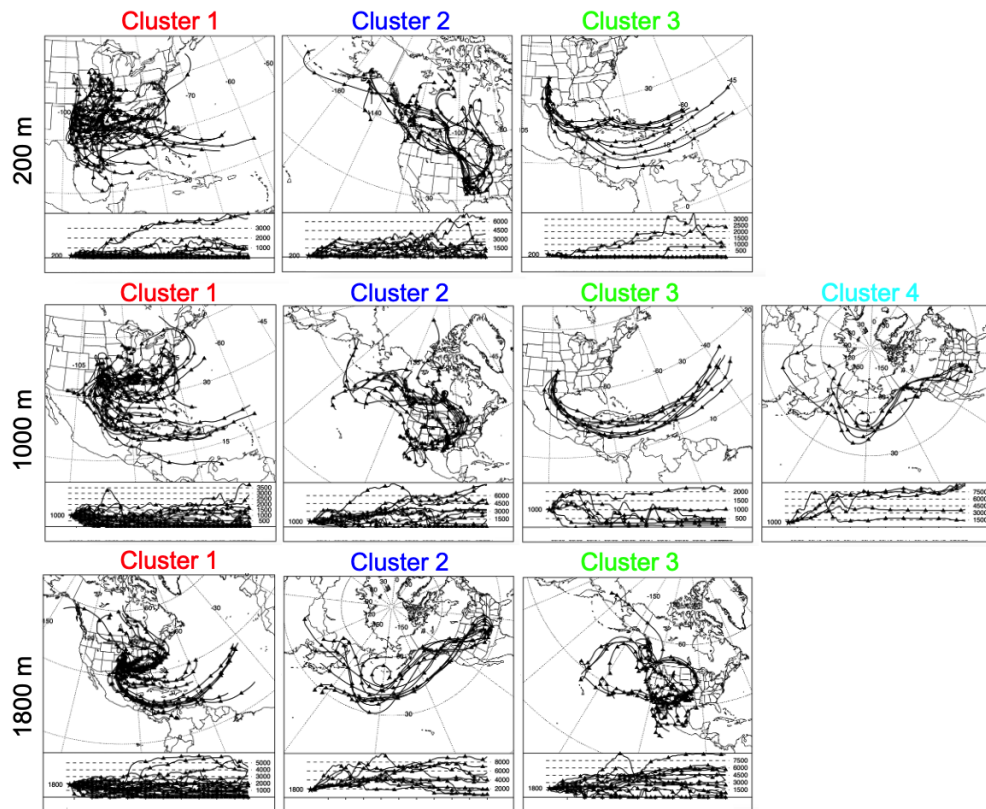


Fig. S4. Trajectories associated with each cluster during the summer campaign shown in Fig. 3.

5. Fig 4: Add a size bin for 3-10 nm size range from the two onboard CPCs.

Response: The size bin for 3 to 10 nm particles is now added in the revised Fig. 4. The figure caption is also revised to reflect the methods for calculating the size distributions. The revised figure can be found below.

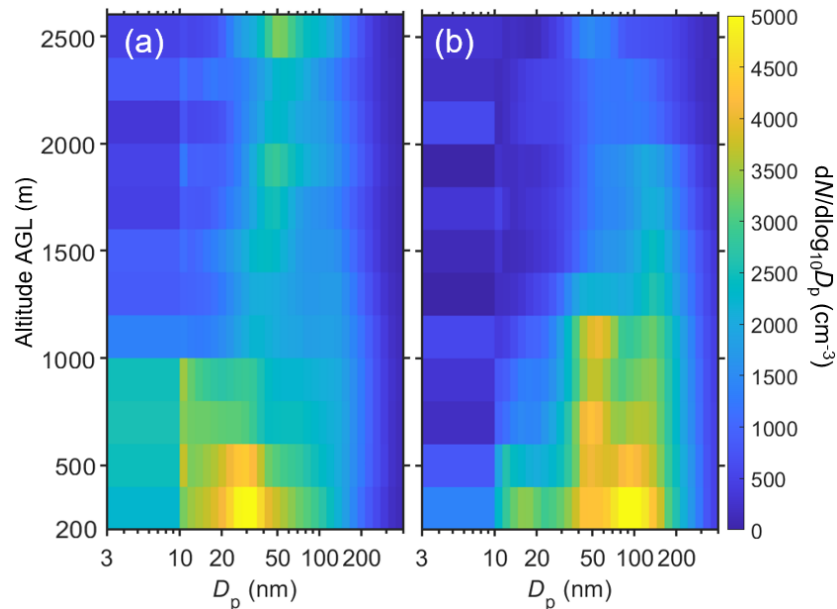


Fig. 4. Vertical profiles of the mean aerosol size distributions during the (a) spring campaign and (b) summer campaign. Note that the size distributions between 3 and 10 nm are calculated based on the difference of concentrations monitored by the two CPCs, and the size distributions above 10 nm are obtained by the FIMS. The size distributions are normalized to standard temperature and pressure (273.15 K and 101.325 kPa; STP).

6. Fig 5: Add the vertical profile of 3-10 nm number concentration calculated from the CPCs. Also add a legend showing that red is summer and blue is spring (same for Fig. 6).

Response: We have now included the analysis of N_{3-10} in the updated Fig. 5 (also shown below). N_{3-10} showed a consistent trend with N_{10-20} , meaning that these aerosols are associated with the nucleation-mode aerosols generated from NPF events. To include the contribution of N_{3-10} to total aerosol concentration, we have also changed the analysis and used $N_{>3}$ (instead of $N_{>10}$) to represent total aerosol concentration. Because of this change, the discussion regarding the vertical profiles of total aerosol concentration has changed, because now, springtime $N_{>3}$ is consistently higher than summertime $N_{>3}$ at all altitudes below 2500 m AGL due to the large contribution of nucleation mode aerosols to total aerosol number concentrations. These changes have been included in both the discussions regarding Fig. 5 and the conclusion. Legends are now added to differentiate spring and summer campaigns in Figs. 5, 6, S6 and S7.

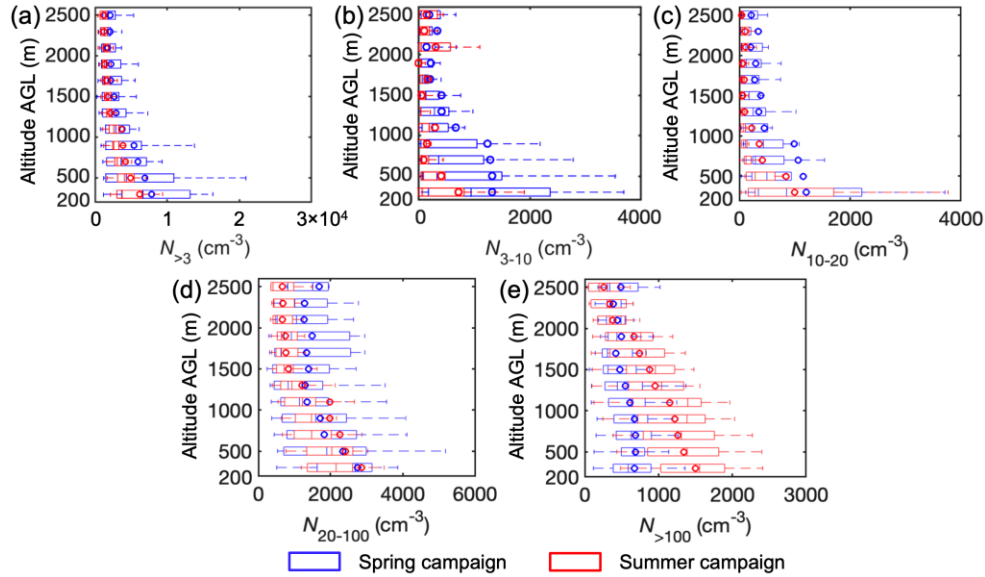


Fig. 5. Vertical profiles showing the concentrations of particles (a) larger than 3 nm ($N_{>3}$), (b) with sizes between 3 and 10 nm (N_{3-10}), (c) with sizes between 10 and 20 nm (N_{10-20}), (d) with sizes between 20 and 100 nm (N_{20-100}), (e) with sizes above 100 nm ($N_{>100}$) over the SGP site during the spring campaign (blue) and summer campaign (red). The line and circle markers represent the median and mean of the data, and the edges of the box indicate the 25th and 75th percentiles, respectively. The concentrations are normalized to standard temperature and pressure (273.15 K and 101.325 kPa; STP).

7. In addition to the vertical profiles of number concentrations and mass concentrations could you include a figure with average vertical profiles of the meteorological variables such as potential temperature as well as LWC. Alternatively add them as subplots to Fig. 5.

Response: We have now included the boxplots showing the potential temperature and LWC (figure below). However, as they are not directly related to the aerosol measurements, we have now included the figure in the supplementary materials as **Fig. S2**. The following discussion has been added to Section 3.1 in the manuscript:

“**Figure S2** shows the vertical profiles of potential temperature and LWC during the spring and summer campaigns. Note that overall, there is minimal LWC at altitudes above 1500 m (**Fig. S2b**). However, the inversion of the potential temperature may occur even when there is no LWC, meaning that the profile of LWC in **Fig. S2b** could only qualitatively show the BL height.”

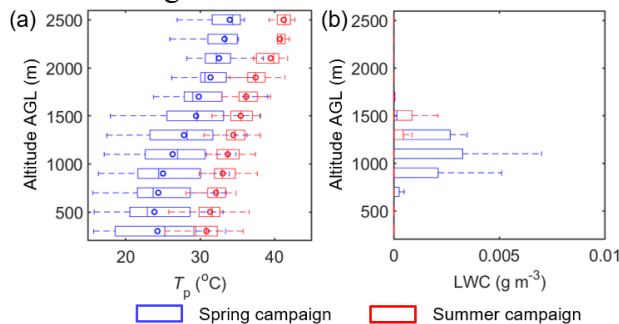


Fig. S2. Vertical profiles of (a) potential temperature (T_p) and (b) liquid water content (LWC) during the spring and summer campaigns.

8. For the NPF case studies (Figs. 7-9) it would be useful to include horizontal track of the aircraft colored by the 3-10 nm number concentration. It was mentioned that several vertical profiles were made during each flight, it would be good to see all of them. Also include the number size distribution as a function of time measured at the surface for further comparison.

Response: We have added the tracks (including both the horizontal and vertical ones) of the G-1 aircraft during these three flights, and the tracks are colored by the ratio between $N_{>3}$ and $N_{>10}$ (Figs. S9 to S11). We are showing Fig. S9 below as an example. We chose to plot the ratio ($N_{>3}/N_{>10}$) because there were periods where the difference in the concentration was small, but the aerosol population was dominated by N_{3-10} . Using $N_{>3}/N_{>10}$ could show the scale and influence of the NPF events. Together with the flight tracks, we also included the surface measurements of aerosol size distributions for comparison against the flight measurements (Figs. S9 to S11). Since the SMPS data corresponding to the case in Fig. 9 is not available, we showed the normalized size distribution data measured by the UHSAS. From the size distributions, we found that the flight measurements near the surface had a general agreement with the surface measurements. We further discussed these figures when introducing the NPF cases in Section 3.3.1. Please also see the response to comment #10.

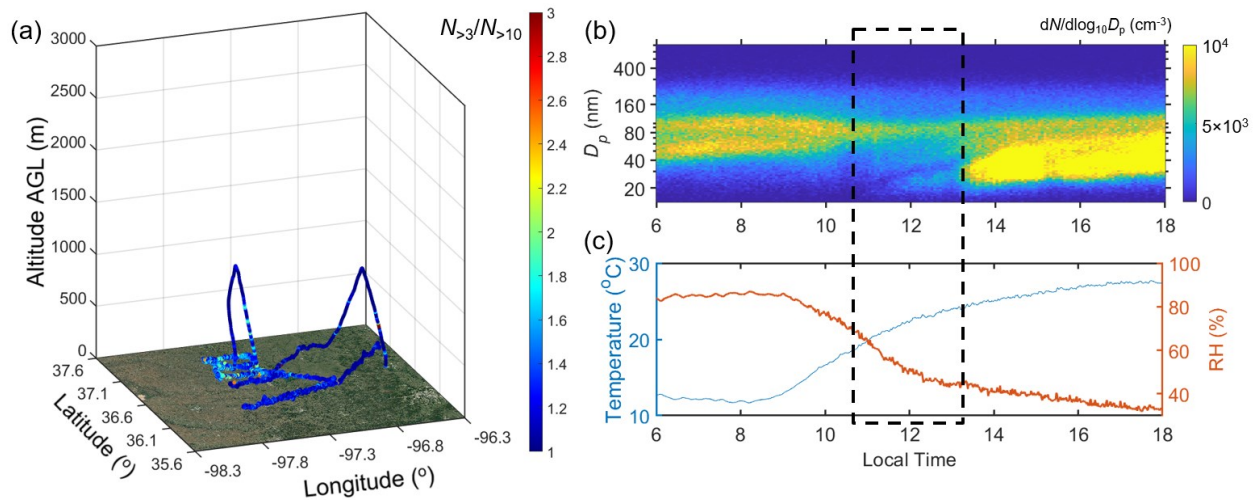


Fig. S9. (1) Track of the G-1 aircraft on September 11, 2016 colored by the ratio between $N_{>3}$ and $N_{>10}$. Map is obtained from ©Google Maps. (2) Aerosol size distribution and (3) temperature and relative humidity (RH) measured at the SGP observatory on September 11, 2016. The dashed box shows the corresponding time for the aircraft measurement.

9. Can you be sure that the observed heterogeneity in the vertical profiles is not actually due to horizontal heterogeneity? This is especially relevant in the case studies. If horizontal heterogeneity cannot be ruled out you should mention it in the text.

Response: The horizontal heterogeneity could play a role in the analysis, as indicated in Fast et al. (2022), which characterized the subgrid variability of aerosol properties using grid spacings of 3, 9, 27, and 81 km. However, as indicated by that study, the averaged aircraft-measured aerosol properties within the 81 km cell were similar to many aerosol properties measured at the SGP observatory, suggesting that averaging aircraft measurements over large spatial scales could reduce the influence of horizontal heterogeneity on aerosol properties. Since this study analyzed the averaged aerosol properties collected during the entire campaign, we believe that horizontal heterogeneity is reduced. However, we could not rule out the influence of horizontal

heterogeneity when analyzing the vertical profiles in a case-by-case manner. This discussion is now added to the Conclusion as:

“We should note that the horizontal heterogeneity could influence the analysis in this study, as indicated in Fast et al. (2022), which characterized the subgrid variability of aerosol properties using grid spacings of 3, 9, 27, and 81 km. However, as indicated by that study, the averaged aircraft-measured aerosol properties within the 81 km cell were similar to many aerosol properties measured at the SGP observatory, suggesting that averaging aircraft measurements over large spatial scales could reduce the influence of horizontal heterogeneity on aerosol properties. Since this study analyzed the averaged aerosol properties collected during the entire campaign (which covers a large special scale), we believe that horizontal heterogeneity is reduced. However, we could not rule out the influence of horizontal heterogeneity when analyzing the vertical profiles on a case-by-case manner. Nonetheless, analyzing the vertical profiles of aerosols could significantly improve our understanding of the sources and sinks of aerosols in the atmosphere and greatly compliments ground observations.”

10. Fig 9: If the particles were formed in the upper parts of the well-mixed layer you would expect them to be more vertically mixed since it takes several hours for the particles to grow to such sizes. Do you see at the surface when the layer was mixed down?

Response: We presume that the reviewer is referring to **Fig. 8**, as it is the figure describing the NPF in the upper BL. It is a good idea to consider the time needed for the growth of the newly formed particles. The NPF layer in the upper BL is indeed relatively well mixed, as indicated by the similar magnitudes of N_{3-10} between 600 and 1000 m AGL in **Fig. 9b**. The surface measurements related to the example NPF events are now included in **Fig. S10**. As shown in **Fig. S10**, the SGP observatory did not observe the formation of particles below 30 nm. Instead, particles of ~30 nm appeared at 16:00 local time (flight measurement was conducted between 12:40 and 15:30 local time), suggesting that 30 nm particles were transported vertically due to the mixing of the upper BL air to the surface. We have added this discussion in the manuscript as:

“The spatial scale of this NPF event is shown in **Fig. S10a**, where NPF occurred in the upper BL, while $N_{>3}/N_{>10}$ near the surface is relatively low. This is indicated by the surface measurements of aerosol size distributions, which did not observe the formation of particles below 30 nm. Instead, particles of ~30 nm appeared at 16:00 local time (flight measurement was conducted between 12:40 and 15:30 local time), suggesting that 30 nm particles were transported vertically due to the mixing of the upper BL air to the surface.”

11. Lines 339-441: Another explanation might be that the particles were formed in the residual layer and mixed down (Lampilahti et al., 2021). Precursor gases could be present in the residual layer as well (Beck et al., 2022).

REFERENCES:

Lampilahti et al. (2021) <https://doi.org/10.5194/acp-21-7901-2021>

Beck et al. (2022) <https://doi.org/10.5194/acp-22-8547-2022>

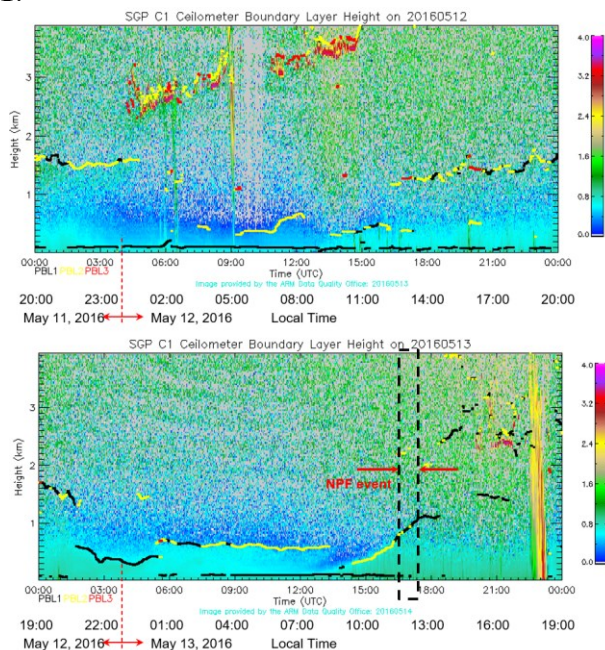
Response: We thank the reviewer for providing these references. It is indeed probable that NPF occurred in the residual layer, and the newly formed particles were further mixed down to the BL. However, due to the narrow vertical range of the residual layer above the BL, the current

aircraft cannot segregate the aerosol properties inside of this layer from the BL. We have included this discussion as:

“It may be because of the following reasons: (1) the low S_{tot} plays a dominating role in promoting NPF, and a SO_2 mixing ratio of 0.5 ppb is already sufficient for initiating NPF; (2) NPF events occurred in the residual layer above the BL and the newly formed particles were further mixed down to the upper BL (Lampilahti et al., 2021; Beck et al., 2022), where the SO_2 in the upper BL is not related to the NPF events; (3) SO_2 was depleted in the upper BL due to its oxidation to sulfate that promoted the NPF event in the upper BL; (4) different nucleating precursors, such as amines and extremely low volatility organic compounds (ELVOCs), were involved in this NPF event.”

12. Fig 10: S_{tot} and number size distribution show discontinuities between 1300-1500 m and between 2000-2200 m. Gradient in potential temperature also slightly increases around or before these altitudes indicating weak inversions. Is it possible that the BL reached 2 km on a previous day and the sub-10 nm particles are in fact inside the residual layer? Cases in Figs. 9-10 might both be residual layer NPF events observed before (Fig 10) and after (Fig 9) entrainment into the well-mixed layer.

Response: Based on the description, we presume that the reviewer is discussing the event in Fig. 9 instead of Fig. 10. This is an interesting point. To address this comment, we examined the BL height on the previous day and during the day of the event shown in Fig. 9 (NPF in FT). The figure below shows the BL height determined by the ceilometer. As can be seen from the figure, the BL height for May 12, 2016 (the day before the NPF event shown in Fig. 9) reached around 1700 m, and a nocturnal boundary layer was developed overnight. On May 13, 2016, the NPF event was observed on G-1 between 12:00 and 13:00 local time at the altitudes between 1800 and 2200 m. Therefore, this NPF event is more likely an event in the lower FT and not influenced by the residual layer. Moreover, if the newly formed particles are entrained from the residual layer to the upper BL, these newly formed particles should be observed in a wider altitude range in Fig. S11.



However, we cannot completely rule out the impact from the residual layer NPF as the BL height (1700 m) on May 12, 2016 is close to the NPF region (1800 to 2200 m) on May 13, 2016. This discussion has been added to the manuscript as follows:

“We should note that it is also possible that these newly formed particles may be generated from NPF in the residual layer developed overnight (a mechanism similar to the discussion regarding NPF in the upper BL), as the BL height reached 1700 m on the previous day before the development of the nocturnal boundary layer. If so, the NPF cases shown in **Figs. 8** and **9** may be both occurring in the residual layer but observed before (**Fig. 9**) and after (**Fig. 8**) entrained into the BL. Given these NPF events occur in the higher altitudes and there are limited observations regarding NPF in these locations, the mechanisms associated with NPF events require more investigation.”

12. Lines 429-430: Show the ground-based number-size distribution measurements. Does it support the analysis of what happens between the morning and the afternoon flights in the well-mixed part of the BL.

Response: We have now included the size distributions measured at the SGP observatory as **Fig. S12** (also shown below). The ground observation agrees with the measurements conducted in the morning and afternoon flights, where the size of the accumulation mode aerosols increased, and at the same time, aerosols below 30 nm emerged. We have added the following discussion in Section 3.4.1:

“Surface measurements conducted on the same day agreed in general with the flight observations (**Fig. S12**), showing a gradual growth in the size of the accumulation-mode aerosols during the day and the emergence of aerosols below 30 nm between 14:00 and 15:00 local time.”

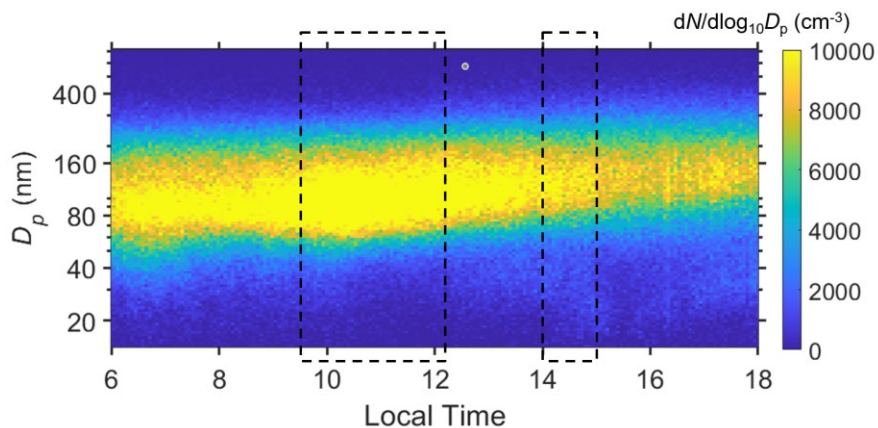


Fig. S12. Aerosol size distribution measured at the SGP observatory on September 4, 2016. The black dashed boxes correspond to morning and afternoon flight times.

13. Line 467: “FL” -> “FT”

Response: Corrected. Thank you.

14. Fig 13: Show the time evolution of the number concentrations in the relevant size ranges ($N_{<50}$ and $N_{>100}$) at the surface.

Response: We have now included the concentrations of aerosols above 100 nm ($N_{>100}$) and below 80 nm ($N_{<80}$) during this event (**Fig. 13d**, also shown below). We chose 80 nm as the size

cut because the Aitken mode had an upper size boundary of ~ 80 nm during the MCS event. We added the discussion:

“We further calculated the concentrations of aerosols above 100 nm ($N_{>100}$) and below 80 nm ($N_{<80}$) during this event. There was an increase in $N_{<80}$ between 21:10 and 23:00 (increased from ~ 600 to ~ 1000 cm^{-3} , **Fig. 13d**, while $N_{>100}$) remained relatively constant during this period. This change in aerosol properties may be explained by the horizontal transport of air mass, but at the same time, the vertical transport of FT air to the BL may also contribute to the change in aerosol properties.”

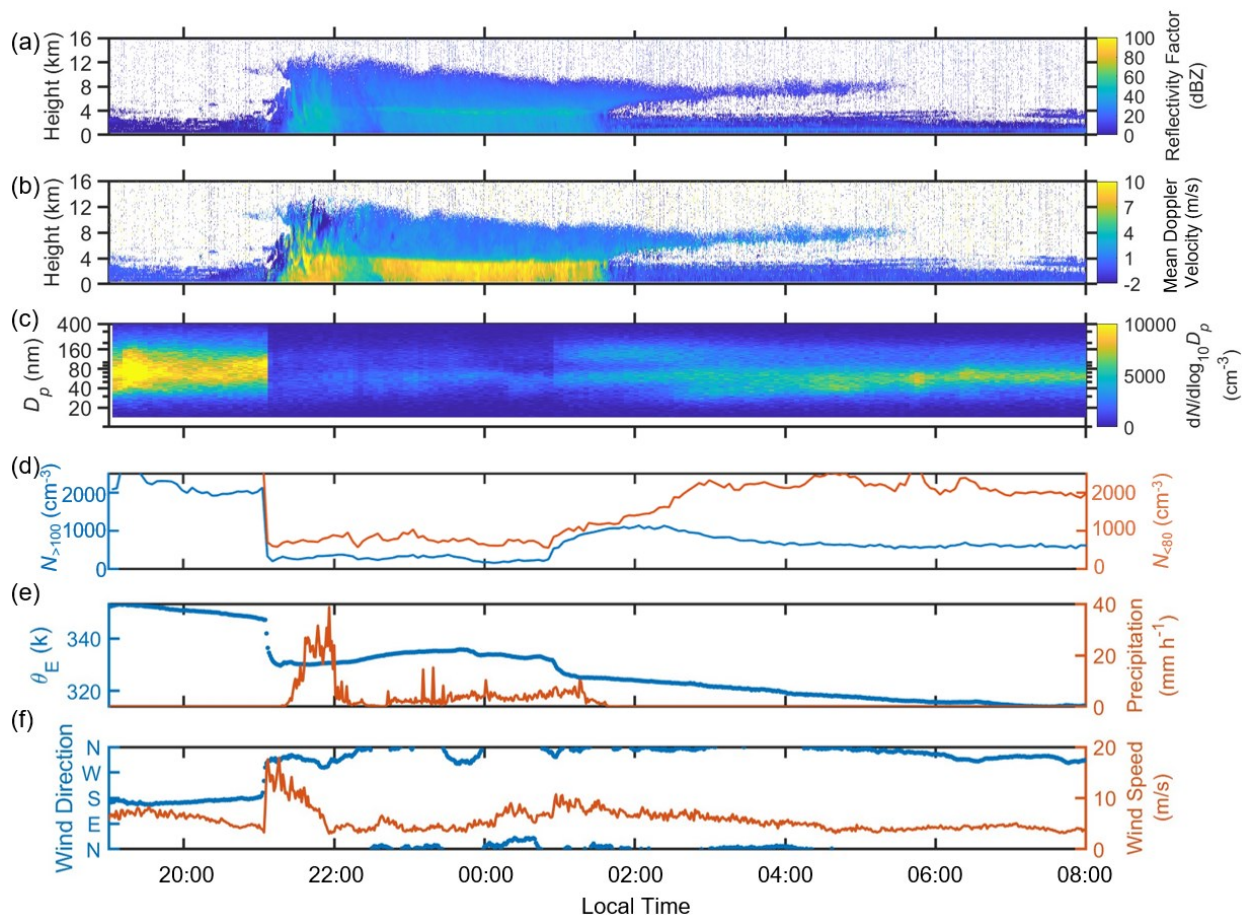


Fig. 13. Ground measurements of aerosol, cloud, and meteorological properties during the convective downward transport event from September 9 to 10, 2016. (a) Cloud reflectivity measured by RWP. (b) Mean Doppler velocities measured by the radar wind profiler (RWP). A positive value indicates downward motion. (c) Aerosol size distributions measured by a scanning mobility particle sizer (SMPS). (d) Concentration of aerosols with sizes above 100 nm ($N_{>100}$) and below 80 nm ($N_{<80}$). (e) Equivalent potential temperature and precipitation rate measured by the weather station. (f) Wind direction and wind speed measured by the weather station.

15. Line 525: Does the calculated particle exchange agree with the ground-based number concentration measurements during the convective event?

Response: Thank you for the suggestion. We calculated the increase of surface-measured aerosol concentration due to the downdraft, assuming that the BL is well mixed and there are no other sources or sinks of aerosols. This can be done via dividing the total exchange by the BL

height (assumed to be 1 km), which yields an increase in the aerosol concentration of 450 cm^{-3} . Note that this value is similar to that observed in the increase of $N_{<80}$ between 21:30 and 23:00 (from ~ 600 to $\sim 1000 \text{ cm}^{-3}$), suggesting that the increase of the Aitken-mode aerosols during MCS may be related to the convective downdraft. We have added the following discussion to Section 3.4.3:

“Assuming that the BL is well mixed and there are no other sources or sinks of aerosols, we could estimate the impact of this event to the total aerosol concentration measured at the surface. This can be done via dividing the total exchange ($4.50 \times 10^7 \text{ cm}^{-2}$) by the BL height (assumed to be 1 km), which yields an increase of the aerosol concentration of 450 cm^{-3} . Note that this value is similar to that observed in the increase of $N_{<80}$ between 21:30 and 23:00 (from ~ 600 to $\sim 1000 \text{ cm}^{-3}$), suggesting that the increase of the Aitken-mode aerosols during MCS may be related to the convective downdraft.”

We thank the reviewer for his/her comments. This has improved the quality of the manuscript and we look forward to this manuscript being accepted.

Manuscript No.: EGUSPHERE-2023-830

Title: Vertical profiles of trace gas and aerosol properties over the Eastern North Atlantic: Variations with season and synoptic condition

1. In this paper an in-depth assessment of the vertical heterogeneity of aerosols over this region is presented. The authors must be commended to present such a thorough investigation of several factors influencing the formation and growth atmospheric particulates at different altitudes. Detailed comments related to scientific, technical and language issues are indicated in the annotated PDF file. In general, the paper is well-written, especially from Section 3.3 onwards. There are, however, instances of incoherent writing and discussions as highlighted in the annotated PDF file. There are also some speculative explanations of observations, which are, however, plausible.

Response: We thank the reviewer for the positive feedback. We have thoroughly revised the manuscript based on the comments provided. The point-to-point responses can be found as follows.

2. Line 25: Do you mean in general or only at ground level?

Response: We mean in general. The sentence has been corrected.

3. Line 29: Not exactly sure what you mean here? Condensation being a sink for smaller particles growing into larger particles?

Response: The reduced condensation sink can lead to stronger NPF. The sentence has been corrected into

“..., a result of the stronger new particle formation (NPF) events due to the reduced condensation sink in spring.”

4. Line 32: Is this considered considerable?

Response: We have removed “considerable” and reworded the sentence as

“Through the vertical profiles of aerosol properties, we observed NPF events in the upper BL during 7 out of 38 research flights, where the newly formed particles continue to grow as they are mixed down to the surface.”

5. Line 46 and Line 50: Should be defined in main text.

Response: The abbreviations have been defined.

6. Line 50: Not sure whether this definition of NPF is required.

Response: Since this is an overview of the aerosol measurements during the research campaign, we feel it is necessary to include the definition of NPF to accommodate readers with different backgrounds.

7. Line 54: sulphate

Response: Changed.

8. Line 65: Please define in main text.

Response: The definition has been added.

9. Line 66: Do not like this word.

Response: The word has been replaced by “study.”

10. Line 68: What schemes? In the model or actual formation mechanisms?

Response: In the model. This has been clarified in the manuscript.

11. Line 68: Maybe give a reference here.

Response: Reference is added.

12. Line 69: Not exactly sure what is meant here.

Response: The sentence has been rephrased as

“Apart from newly formed particles, vertically compact aerosol layers, which were traced back to forest fires in East Asia, were observed at altitudes over 3000 m at the SGP.”

13. Line 70: ??

Response: The definition is now added to the main text.

14. Line 85: I do agree with these type of made up abbreviated terms.

Response: Understood.

15. Line 86: I will just say “the spring campaign”

Response: Corrected.

16. Line 89: Nothing mentioned about horizontal coverage.

Response: The horizontal coverage is now added. The sentence is now reworded as

“The Gulfstream-1 (G-1) aircraft (Schmid et al., 2014) was deployed to examine the spatial and temporal variation of aerosol properties over the SGP observatory (36°36’26” N, 97°29’16” W), with a horizontal coverage of around 100 km (Fast et al. 2022) and an altitude range of 200 to 3000 m above ground level (AGL).”

17. Line 92: Who mean diurnal. Temporal refer to seasonal and diurnal.

Response: The word “temporal” has been replaced by the word “diurnal.”

18. Line 92: ???

Response: We meant that there were four to six vertical profiles of G-1 measurements in addition to the one departing from the airport and the one arriving at the airport. The departing and arriving sections are excluded from the analysis to avoid interference from the airport and urban pollution. The sentence has been rephrased as

“Each flight consisted of four to six vertical profiles (in addition to those leaving and arriving at the airport), providing the aerosol and corresponding meteorological properties as a function of altitude.”

19. Line 94: ???

Response: The sentence has been rephrased as

“The flight tracks used during the HI-SCALE campaign and the flight track of an example research flight are shown in **Fig. 2.**”

20. *Line 111: Not sure what is meant is this term?*

Response: The term “non-refractory” is assigned to species that evaporate rapidly at 600 °C under vacuum conditions. It is used for discussing aerosol composition measured by the aerosol mass spectrometers. This definition is now added to the manuscript.

21. *Line 112: ??*

Response: Please see the response to the previous comment.

22. *Line 119: In general, this section can be written more coherently.*

Response: Thanks. We have reorganized the structure of this section. Please see our revised manuscript.

23. *Line 120: Are these averages?*

Response: Yes. This information is added to the manuscript.

24. *Line 124: Maybe consider to put in Experimental/Methods section.*

Response: The description of the back trajectory calculation has been shortened.

25. *Line 128: A map indicating vegetation and agricultural coverage, as well as burned areas (e.g. MODIS satellite imagery) would be useful and support your interpretation.*

Response: A satellite imagery showing the terrain and vegetation surrounding the SGP observatory is now included as **Fig. S1**. However, it is challenging to show the burned area in the imagery.

26. *Line 138: Air masses does not originate from a region, it passes over regions. This is very important.*

Response: Thanks. The sentences have been revised.

27. *Line 139: How and why?*

Response: Further discussion has been added as

“Also, the short trajectory from the east reflects slower wind speeds from the forests during the summertime. The longer residence time of air mass and the more abundant fresh volatile organic compounds in the forest region can lead to a higher mass of aerosols and less oxidized organic aerosols during the summer (Liu et al., 2021).”

28. *Line 146: Why?*

Response: The sentence has been rephrased as

“The wildfire-generated biomass burning aerosols in the north and west of SGP may be injected into the FT, and further transported and aged before they arrive at the SGP.”

29. *Line 149: Reference*

Response: The reference has been added.

30. *Line 149: Do you mean in relation to the impact on ground observations?*

Response: Yes. The sentence has been rephrased as

“Long-range transport of aerosols may impact ground observations under such situations (Wang et al., 2006), but the influence of this air mass on ground observations is also largely subject to the coupling between BL and FT.”

31. Line 153: Very long sentence. Can you show the seasonality of fire frequencies? It is important to support your interpretation here.

Response: The seasonality of the fire frequencies can be found in *Fire Season Climatology in by NWCG: Fire Behavior Field Reference Guide, PMS 437, 2021* (<https://www.nwcg.gov/publications/pms437/weather/fire-season-climatology>). The report shows that overall, regions to the north and west of SGP have higher fire frequencies in spring and summer, while the southeast region has a lower fire frequency. The sentence has been rephrased as

“Fire season climatology shows that, in general, there are stronger fire activities in late spring and summer to the north and west of SGP (NWCG, 2021). The stronger fire activities coincide with the warmest and driest conditions, together with wind events and dry lightning potential. There is a much weaker fire activity to the southeast of SGP in late summer, especially from August to September.”

32. Line 158: Not potentially, it does.

Response: The word has been removed.

33. Line 162: What is meant here? Dependency of SO₂ levels or dependency of NPF?

Response: The sentence has been clarified as

“SO₂ can contribute to the occurrence of NPF. At the same time, NPF depends on the concentration of pre-existing aerosols and other meteorological parameters (Section 3.3.1).”

34. Line 171: I think you mentioned that you did not consider measurements in clouds.

Response: Yes, but aqueous chemistry in the cloud droplets can contribute to the size increase of the accumulation mode aerosols.

35. Line 174: Still not sure what is meant here.

Response: The sentence has been rephrased as

“The larger size of the aerosols in the summer also leads to larger condensation and coagulation sinks for nucleating vapors and newly formed particles, suppressing the occurrence of NPF in the summer. As a result, the concentration of aerosols below 30 nm in the BL (e.g., below 1000 m) during spring is higher than that during summer.”

36. Line 175: Maybe include reference.

Response: The reference has been included.

37. Line 177: Term previously defined, i.e. CCN.

Response: The term is previously defined in the abstract only. The abbreviation is introduced in the main text here again.

38. Line 180: “due to” or “through”?

Response: The word is replaced.

39. Line 193: Evidence to support this?

Response: This can be supported by the fact that the mean values of N_{10-20} are outside the 25th to 75th percentile box in the altitude range of 500 and 1000 m. In the spring campaign, there were three NPF events where N_{10-20} values were above 2000 cm^{-3} in the altitude range of 500 and 1000 m, significantly higher than the median values between 400 and 600 cm^{-3} . More discussion has been added as

“This can also be indicated by the mean values of N_{10-20} being outside the 25th to 75th percentile box in the altitude range between 500 and 1000 m (**Fig. 5b**). In the spring campaign, there were three NPF events where N_{10-20} values were above 2000 cm^{-3} in the altitude range of 500 and 1000 m, significantly higher than the median values between 400 and 600 cm^{-3} .”

40. Line 202: A better site description/map will greatly assist in understanding/supporting interpretations.

Response: To assist in supporting the interpretations, we included a satellite image showing the terrain and vegetation surrounding the SGP observatory as **Fig. S1**.

41. Line 216: in the BL?

Response: Yes. The sentence has been rephrased as

“A breakdown of the total aerosol concentration into different size modes shows that, in the BL, N_{10-20} is higher during the spring (**Fig. 5b**), while $N_{>100}$ is higher during the summer (**Fig. 5d**).”

42. Line 216: Evidence?

Response: Please see the response to comment 39.

43. Line 243: Why?

Response: The sentence has been rephrased as

“Since nitrate aerosols are more stable under low temperatures, the higher nitrate concentrations during spring are likely driven by the lower temperature in the spring (Liu et al., 2021; Parworth et al., 2015).”

44. Line 247: Refer to figure.

Response: Figures 3 and S1 are now referred to in the discussion.

45. Line 247: Not sure how this is relevant to your observation? Sulphate is usually important for formation of particulate and organic species for particle growth. Maybe just explain it better.

Response: As Xu et al. (2015) pointed out, isoprene-derived SOA is directly mediated by the abundance of sulfate through chemical reactions among sulfate, organics, and water. These reactions allow the increase in both the sulfate and organics in the summer. The discussion has been added as

“Moreover, the previous study by Xu et al. (2015) showed that the formation of isoprene SOA is mediated by the abundance of sulfate through chemical reactions among sulfate,

organics, and water, which allows the increase of both organics and sulfates in the summer.”

46. Line 251: Usually indicative of fresher plumes.

Response: These organics are more likely caused by fresh biogenic emissions instead of biomass-burning plumes. The study by Liu et al. (2021) analyzing the aerosol mass spectrometer data showed that the biomass-burning organic aerosol (BBOA) factor was not identified during the summer season. This result is also consistent with the low concentrations of BBOA observed in summer at SGP in a previous study (Parworth et al., 2015). More discussion has been added as “These summertime organics are not generated by biomass-burning plumes as the study by Liu et al. (2021) analyzing the aerosol mass spectrometer data showed that the biomass-burning organic aerosol (BBOA) factor was not identified during the summer season. This result is also consistent with the low concentrations of BBOA observed in summer at SGP in a previous study (Parworth et al., 2015).”

47. Line 253: This means that the aerosol might not even reach SGP.

Response: The sentence has been rephrased. The springtime aerosols are more aged because they have a longer residence time in the atmosphere and different oxidant concentrations, or a combination of both effects. The discussion has been revised as “According to Liu et al. (2021), the different oxidation levels in the spring and summer are likely due to (1) aerosols in the spring are more aged due to a longer residence time in the atmosphere, potentially different oxidant concentrations, or a combination of both effects.”

48. Line 254: Not sure whether I agree.

Response: More discussion has been added as “(2) different VOC species could contribute to aerosol formation in spring and summer (e.g., higher monoterpene and isoprene concentrations were observed in the summer), and (3) the more abundant biogenic VOCs in the summer were not transformed into a higher-oxygenated form in the aerosol phase, either due to radical chemistry, oxidants, or their residence time in the atmosphere.”

49. Line 320: I would expect this.

Response: Agreed.

50. Line 338: Due to oxidation to SO₄?

Response: Agreed. More discussion has been added as “(2) SO₂ was already depleted in the upper BL due to its oxidation to sulfate that promoted the NPF event.”

51. Line 375: This seems to be repeating information.

Response: The term “volume-controlled processes” was discussed in the earlier section of the manuscript. However, “coagulation” and “condensation” were not explained. The sentence has been reworded as

“The particles can grow through different dynamic approaches, including coagulation, condensation, and volume-controlled processes.”

52. Line 381: Also, please look at the following paper and cite: Vakkari et al, 2015, Reevaluating the contribution of sulfuric acid and the origin of organic compounds in atmospheric nanoparticle growth, *Geophysical Research Letters*, 42, 10,486–10,493, doi:10.1002/2015GL066459

Response: Thank you. The discussion of this reference is now added to the manuscript as “The aerosol measurement in a South African grazed savannah grassland environment showed that depending on the gaseous precursors and size of the newly formed particles, the aerosol growth can be dominated by either sulfuric acid accompanied by ammonium or organic compounds originating in either biogenic emissions or savannah fires. The contribution of sulfuric acid was larger during the early phases of the growth, but in clean conditions, organic compounds dominated the growth from 1.5 nm up to climatically relevant sizes.”

53. Line 386: *Is abbreviated term defined?*

Response: Thanks for pointing this out. The term “extremely low volatility organic compounds (ELVOCs)” is now defined in the earlier section of the manuscript.

54. Line 517: *average?*

Response: Yes, it is the mean concentration. The word has been added.

55. Line 564: *You did it only for two seasons. Not full seasonal pattern.*

Response: Thanks. The sentence has been revised as “In this study, we present aerosol properties and meteorological parameters characterized onboard the G-1 aircraft during the spring and summer HI-SCALE campaigns. The key processes that drive the aerosol population in the BL are investigated by examining the vertical variation of aerosol properties.”

56. Line 588: *Strong word. “Significantly improved understanding” maybe?*

Response: The sentence has been revised. Thank you.

We thank the reviewer for his/her comments. This has improved the quality of the manuscript and we look forward to this manuscript being accepted.