

## **RE: A point-to-point response to reviewers' comments**

“A proxy for atmospheric daytime gaseous sulfuric acid concentration in urban Beijing” (acp-2018-1132) by Yiqun Lu, Chao Yan, Yueyun Fu, Yan Chen, Yiliang Liu, Gan Yang, Yuwei Wang, Federico Bianchi, Biwu Chu, Ying Zhou, Rujing Yin, Rima Baalbaki, Olga Garmash, Chenjuan Deng, Weigang Wang, Yongchun Liu, Tuukka Petäjä, Veli-Matti Kerminen, Jingkun Jiang, Markku Kulmala, Lin Wang

We are grateful to the helpful comments from the reviewers, and have carefully revised our manuscript accordingly. A point-to-point response to the comments, which are repeated in italic, is given below.

In addition to the reviewers' comments, we have noticed and corrected a key typo from our previous version of manuscript. “The  $[NO_2]$  concentration” in our manuscript is in fact “the  $[NO_x]$  concentration”. Correction of this term does not lead to changes in our conclusions.

### Reviewer #1's comments:

*This study examines the relationship of [SO<sub>2</sub>] to [H<sub>2</sub>SO<sub>4</sub>] as a function of light intensity, particle concentration, and other gas phase reactants in Beijing. As the authors point out, this relationship is likely different in Beijing than in European and US cities. Overall, this study is straight forward and generally useful for research conducted in megacities. However, several issues should be address before this study can be considered for publication in ACP.*

Reply: We are very grateful to the positive viewing of our manuscript by Reviewer #1, and have now revised our manuscript accordingly.

### Major comments:

1. *P5 147: Sulfuric acid concentration was measured using nitrate LToF-CIMS. It would be useful for the reader to know more details on how the sulfuric acid concentration was determined from the signals of the instrument. Does this measurement include sulfuric acid in molecular clusters (i.e. is fragmentation contributing to the sulfuric acid signal?) What are the estimated uncertainties of the sulfuric acid measurement? How do these uncertainties compare to the model predicted amounts?*

Reply: We measured signals of charged sulfuric acid as HSO<sub>4</sub><sup>-</sup>, HSO<sub>4</sub><sup>-</sup> HNO<sub>3</sub>, and HSO<sub>4</sub><sup>-</sup> (HNO<sub>3</sub>)<sub>2</sub>, and charged clusters of HSO<sub>4</sub><sup>-</sup> H<sub>2</sub>SO<sub>4</sub>. The clusters of HSO<sub>4</sub><sup>-</sup> H<sub>2</sub>SO<sub>4</sub> come from ion-induced clustering of neutral sulfuric acid and bisulfate ions within the LTOF-CIMS ion reaction zone, and also from the evaporation of dimethylamine (DMA) and the replacement of one molecule of H<sub>2</sub>SO<sub>4</sub> with one bisulfate ion, HSO<sub>4</sub><sup>-</sup>, during the NO<sub>3</sub><sup>-</sup> reagent ion charging of a stabilized neutral sulfuric acid dimer in the real atmosphere in presence of DMA or a molecule that works in the same way as DMA. Although the total signals of HSO<sub>4</sub><sup>-</sup>, HSO<sub>4</sub><sup>-</sup> HNO<sub>3</sub>, and HSO<sub>4</sub><sup>-</sup> (HNO<sub>3</sub>)<sub>2</sub> were normally orders higher than that of HSO<sub>4</sub><sup>-</sup> H<sub>2</sub>SO<sub>4</sub>, the sum of HSO<sub>4</sub><sup>-</sup>, HSO<sub>4</sub><sup>-</sup> HNO<sub>3</sub>, HSO<sub>4</sub><sup>-</sup> (HNO<sub>3</sub>)<sub>2</sub> and HSO<sub>4</sub><sup>-</sup> H<sub>2</sub>SO<sub>4</sub> has been used to derive the gaseous sulfuric acid concentration.

The uncertainty in measured concentrations of sulfuric acid is estimated to vary between 21% and 51%, which is comparable with that in Kürten *et al.* (2012).

The uncertainties of proxies came from both those of CIMS and those of proxy methods. We have elaborated how the sulfuric acid concentration was determined in our revised manuscript, which reads (L164-L183), “For example, the atmospheric H<sub>2</sub>SO<sub>4</sub> molecules would be charged by nitrate reagent ion NO<sub>3</sub><sup>-</sup>(HNO<sub>3</sub>)<sub>0-2</sub> and mainly produce HSO<sub>4</sub><sup>-</sup> ions (m/z = 96.9601 Th), HSO<sub>4</sub><sup>-</sup> HNO<sub>3</sub> ions (m/z = 159.9557 Th), and HSO<sub>4</sub><sup>-</sup> (HNO<sub>3</sub>)<sub>2</sub> ions (m/z = 222.9514 Th). In addition, HSO<sub>4</sub><sup>-</sup> H<sub>2</sub>SO<sub>4</sub> ions (m/z = 194.9275 Th) were formed from ion-induced clustering of neutral sulfuric acid and bisulfate ions within the LToF-CIMS ion reaction zone, and also from the evaporation of dimethylamine (DMA) and the replacement of one molecule of H<sub>2</sub>SO<sub>4</sub> with one bisulfate ion, HSO<sub>4</sub><sup>-</sup>, during the NO<sub>3</sub><sup>-</sup> reagent ion charging of a stabilized neutral sulfuric acid dimer in the real atmosphere in presence of DMA or a molecule that works in the same way as DMA. During the campaign, the sample flow rate was kept at 8.8 slpm, since mass flow controllers fixed the sheath flow rate and the excess flow rate, and the flow into the mass spectrometer (around 0.8 slpm) was fixed by the size of a pinhole between the ionization source and the mass spectrometer. The

concentration of gaseous sulfuric acid was then determined by Eq. (2).

$$[H_2SO_4] = \frac{HSO_4^- \cdot (HNO_3)_{0-2} + HSO_4^- \cdot H_2SO_4}{NO_3^-(HNO_3)_{0-2}} \cdot C \quad (2)$$

where C is the calibration coefficient, and  $NO_3^-(HNO_3)_{0-2}$ ,  $HSO_4^-(HNO_3)_{0-2}$  and  $HSO_4^- H_2SO_4$  represent the signals of corresponding ions and are in units of counts per second (cps). The unit of resulting  $[H_2SO_4]$  is molecule  $cm^{-3} \dots$ ".

2. *P6 162: Along these same lines, the authors comment that the calibration coefficient takes into account diffusion losses in the sampling line. Was this loss measured? It is a bit surprising that the calibration coefficient that Kurten et al. (2012) determined was  $1.1 \times 10^{10} cm^{-3}$  is the same in this study. I would have thought differences in instrument and sampling line losses (1.6 m is quite long) would have impacted this number. The authors should more clearly lay out how the sulfuric acid concentration was determined since it is an integral measurement for this paper.*

Reply: The loss rate was calculated assuming a diffusion loss of sulfuric acid in a circular tube under a laminar flow condition (Gormley and Kennedy, 1949). The identical values for the effective calibration coefficient are just by coincidence. Our calibration coefficient is  $3.79 \times 10^9$  molecule  $cm^{-3}$  in absence of the diffusion loss correction. We have elaborated our discussion on the calibration coefficient, which reads (L186-L190), "We obtain a calibration coefficient of  $3.79 \times 10^9$  molecule  $cm^{-3}$  for our instrument and use  $1.1 \times 10^{10}$  molecule  $cm^{-3}$  as the effective calibration coefficient, after taking into account the diffusion losses in the stainless-steel tube and the nitrate chemical ionization source".

3. *P6 L 176: The Fuchs-Sutugin transition kernel is used. There is associated error with using this kernel in the transition regime where sulfuric acid condenses on preexisting particles. Can the authors comment on this error? How sensitive is the fitting parameters to changes in the collision kernel? It may be helpful for the authors to use the empirically-derived collision kernel for the full regime from (Gopalakrishnan and Hogan Jr., 2011) to help limit the uncertainties from this parameter.*

Reply: We have calculated  $H(Kn_D)$  from the nondimensionalized form of Fuchs-Sutugin and the First passage regression (Gopalakrishnan and Hogan Jr., 2011) in our  $Kn_D$  range. The difference between the calculated  $H(Kn_D)$  from two methods is within 8 %. Hence, we don't think calculation of the collision kernels in this study will lead to a significant uncertainty. Nevertheless, we have added this citation to give a more comprehensive discussion on the transition-regime correction factor, which reads (L211-L213), " $\beta_m$  represents a transition-regime correction factor dependent on the Knudsen number (Fuchs and Sutugin, 1971; Gopalakrishnan and Hogan Jr., 2011)".

4. *P5 134: The authors state that two months of measurements were conducted. It is not clear from the paper if all two months of measurements were used to determine the proxy relationship. Have the authors examined how the proxy relationship changes from day to day? Or week to week? The authors correctly state that the proxy relationship likely depends on location but does it also depend on time? It is possible that other processes that affect sulfuric acid*

*concentrations (like Criegee intermediates) are not captured in the proxy relationship may play a larger role during some times of day than others.*

Reply: The intensive campaign was carried out from 9 February to 14 March, 2018 (L172), which lasts a bit more than a month. The performance of proxies could be different more or less on different days as the proxies are derived from a statistical analysis of the entire data set in this study. We have expanded the discussion about the applicability of the proxies, which reads (L491-L496) “Furthermore, the proxies might be site-specific and season-specific. Since we derived the proxies in winter in urban Beijing, the exponents of factors in the proxy for other cities or other seasons could have different values. Thus, the proxies in this study should be further tested before their application to other Chinese megacities or other seasons”.

Criegee intermediates are not intended to be included in the proxy because Criegee intermediates seem not to play an important role in the daytime formation of sulfuric acid (L135-L139) (Boy et al., 2013; Mauldin et al., 2012).

5. *P13 line 389: If two months of measurements were taken, why was only one day used to compare measured to predicted sulfuric acid concentrations? How does the comparison for the other days look? It's not necessary to add graphs of these comparison, but a few lines stating the comparison for other days is necessary for the reader to determine how useful this proxy relationship is.*

Reply: The measurements lasted a bit more than a month as mentioned above. Figure 3 (now updated to a new version) presents a statistical comparison between measured and predicted sulfuric acid concentrations in all the measurement days. Relevant discussions can be found in section 4.4.

6. *P13 line 396: Authors state that the proxy relationship developed for the boreal forest and applied to Beijing is a factor of 20 too low due to differences in CS. Why didn't the authors use the Beijing CS values when applying Petäjä et al.'s proxy relationship? Would the differences between measured and predicted from Petäjä then be smaller?*

Reply: We actually used atmospheric variables including CS values from our measurements when applying Petäjä et al.'s proxy relationship. The reason for the poor performance of Petäjä et al.'s proxy relationship on Beijing data could be the much higher CS values in Beijing with a much more complex atmosphere.

7. *Figure 4: It would be useful for the reader to see timelines of all the measured concentrations that go into the proxy relationships as well.*

Reply: Here we present results from a comprehensive campaign with participation from multiple universities and institutions. As a result, this manuscript will only focus on the development of the statistical analysis of the sulfuric acid proxy, and other manuscripts in preparation will discuss the variations of atmospheric variables.

### **Minor comments:**

1. *P1 Line 28: desirable for the atmospheric. . .*

Reply: We have revised our manuscript accordingly, which reads (L28) “...highly desirable for the atmospheric chemistry community”.

2. *P1 36-27 change one of the “using”*

Reply: We have revised our manuscript accordingly, which reads (L37-L39) “A proxy for atmospheric daytime gaseous sulfuric acid concentration was derived through a statistical analysis method by using the UVB intensity, [SO<sub>2</sub>], condensation sink (CS), [O<sub>3</sub>], and [HONO] (or [NO<sub>x</sub>]) as the predictor variables”.

3. *P3 Line 57: sulfuric acid DMA system. The citation for Petäjä et al. (2011) might not be the best. Several studies have pointed out potential experimental issues with this study (Jen et al., 2014; Kürten et al., 2014).*

Reply: We have updated the citations, which reads (L59-L61) “...H<sub>2</sub>SO<sub>4</sub>-DMA-H<sub>2</sub>O ternary nucleation (Almeida et al., 2013; Jen et al., 2014; Kürten et al., 2014; Petäjä et al., 2011; Yao et al., 2018)”.

4. *P3 line 57: demand participation is a strange phrase. Maybe necessitates participations?*

Reply: We have revised our manuscript, which reads (L61) “...involve the participation of gaseous sulfuric acid molecules”.

5. *P3 line 59: Would be worth reading and citing (Kuang et al., 2012) for sulfuric acid growth rates.*

Reply: We have added this citation, which reads (L61-L63) “In addition, the condensation of gaseous sulfuric acid onto newly-formed particles contributes to their initial growth (Kuang et al., 2012; Kulmala et al., 2013)”.

6. *P3 line 62: Knowing sulfuric acid concentrations prior to a nucleation event is also important.*

Reply: We have revised our manuscript, which reads (L63-L67) “Quantitative assessments of the contribution of gaseous sulfuric acid to both the new particle formation rates and the particle growth rates require real-time measurements of gaseous sulfuric acid concentrations prior to and during the NPF events (Nieminen et al., 2010; Paasonen et al., 2010) ”.

7. *P3 Line 68: NO<sub>3</sub>- and ligands.*

Reply: We have revised our manuscript, which reads (L72) “...with NO<sub>3</sub><sup>-</sup> and its ligands as reagent ions”.

8. *P3 line 68: CIMS is actually a pretty broad class of instruments. The low detection limit for sulfuric acid is because the instrument ionizes and samples at atmospheric pressure, which is different than the traditional CIMS.*

Reply: We have revised the sentence accordingly, which reads (L72-L73) “because nitrate CIMS with an atmospheric pressure interface (API) has a low detection limit ...”.

9. *P3 line 80: (Chen et al., 2012) shows a nice figure of sulfuric acid concentrations measured at numerous locations around the world. Not critical to add the citation but worth taking a look at.*

Reply: We appreciate that this reviewer points out a very important paper presenting the measurements of sulfuric acid in different locations. We would like to add the citation, which reads (L84-L88) “Thereafter, measurements of sulfuric acid using CIMS have been performed around the world (e.g., Berresheim et al., 2000; Bianchi et al., 2016; Chen et al., 2012; Jokinen et al., 2012; Kuang et al., 2008; Kürten et al., 2014; Kurtén et al., 2011; Petäjä et al., 2009; Weber et al., 1997; Zheng et al., 2011) ...”.

10. *P3 line 83: has been proven*

Reply: We have revised our manuscript accordingly, which reads (L88) “CIMS has been proven to be a robust tool ...”.

11. *P4 line 105: After reading this, the reader will naturally wonder why is there a positive correlation between CS and sulfuric acid concentration?*

Reply: We have revised our manuscript accordingly, which reads (L108-L111) “In several proxies developed by Mikkonen et al. (2011), the correlation between the gaseous sulfuric acid concentration and CS is positive, which is against what one would expect because a larger CS normally leads to a faster loss for gaseous sulfuric acid”. In addition, we put the detailed discussion of this issue in section 4.2 and 4.3.

12. *P4 line 108: locations that characterize with an... one or two of those words are not correct.*

Reply: We have revised our manuscript, which reads (L114-L115) “...in locations with atmospheric environments different from those in the six sites of that study”.

13. *P4 line 110: Please state the range of CS in addition to how much higher it is compared to other locations.*

Reply: We have stated the range of CS, which reads (L116-L117) “Beijing is a location with typical values of CS (e.g., 0.01-0.24 s<sup>-1</sup> in the 5-95% percentiles in this study) being 10-100 times higher ...”.

14. *P4 line 113: For north America: how do these numbers compare to Mexico City?*

Reply: The level of SO<sub>2</sub> in Beijing has decreased significantly in recent years as we have presented

in L488-L489. Nevertheless, the SO<sub>2</sub> concentration in Mexico City in 2003 is comparable with our measured SO<sub>2</sub> in Beijing. We have added one citation regarding the SO<sub>2</sub> measurements and two citations regarding the sulfuric acid concentration in Mexico City, which reads (L118-L123) “...typical SO<sub>2</sub> concentrations being 1-10 times higher (Wang et al., 2011a; Wu et al., 2017) than those in Europe and North America (Dunn et al., 2004; Mikkonen et al., 2011), yet measured gaseous sulfuric acid concentrations are relatively similar in these environments (Chen et al., 2012; Smith et al., 2008; Wang et al., 2011b; Zheng et al., 2011).”

*15. P5 119: OH radicals*

Reply: We have revised our manuscript, which reads (L126) “... a potentially important source of OH radicals in the atmosphere”.

*16. P5 119: remove the not only and but also. It is harder to read with them there.*

Reply: We have removed the two expressions.

*17. P5 line 128: Criegee should be capitalized*

Reply: We have capitalized “C”.

*18. P6 line 153: was guided through. . . strange phrasing*

Reply: We have revised our manuscript, which reads (L160) “...was introduced into a PhotoIonizer ...”.

*19. P6 line 154: Is this a custom-built inlet? If so, could the authors provide a diagram and write in the dimensions?*

Reply: The inlet is a commercial product from Aerodyne Research, Inc.

*20. P6 line 160: CIMS was calibrated. How? It would be useful to describe this procedure in brief.*

Reply: We have added a brief introduction of the calibration process, which reads (L183-L186) “The CIMS was calibrated during the campaign with a home-made calibration box that can produce adjustable concentrations of gaseous sulfuric acid from SO<sub>2</sub> and OH radicals following the protocols in previous literatures (Kürtén et al., 2012; Zheng et al., 2015).”

*21. P6 line 164: should it be ToFTools?*

Reply: The code maker just named it as tofTools.

*22. P6 line 166: 1 nm. Is this mobility diameter?*

Reply: Yes, it is mobility diameter.

*23. P7 line 213: Authors should better justify pseudo-steady state assumption*

Reply: The Mikkonen et al. (2011) study has indicated that the pseudo-steady state assumption holds well for typical atmospheric conditions. Furthermore, the errors for nonlinear proxies derived from the pseudo-steady state assumption in Mikkonen et al. study are in a range of 40-42 %, whereas ours are in a range of 17.6-19.2 % when evaluating the performances of the proxies with the “error” metric in Mikkonen et al. study. Therefore, we think that the pseudo-steady state assumption can be applied to our proxies.

*24. P8 paragraph starting on line 228: This was a difficult paragraph to understand. Can the authors better phrase it to explain the differences in parameters?*

Reply: We have revised the paragraph, which reads (L272-L283) “In practice, the exponents for variables in nonlinear fitting procedures are rarely equal to 1 (Mikkonen et al., 2011), so we replaced the factors  $x_i$  with  $x_i^{w_i}$  in the proxy, where  $x_i$  can be an atmospheric variable and  $w_i$  defines  $x_i$ ’ exponent in the proxy. Since  $k$  is a temperature-dependent reaction constant and varies within a 10 % range in the atmosphere temperature range of 267.6 - 292.6 K, *i.e.*, the actual atmospheric temperature variation in this study, we approximately regard  $k$  as a constant and use a new scaling factor  $k_0$ . This methodology has been used previously in the proxies of gaseous sulfuric acid in Hyytiälä, Southern Finland (Petäjä et al., 2009). As a result, the general proxy equation can be written as Eq. (7), with the UVB intensity,  $[\text{SO}_2]$ , condensation sink (CS),  $[\text{O}_3]$ , and  $[\text{HONO}]$  (or  $[\text{NO}_x]$ ) as predictor variables:”.

*25. P8 line 242: a matlab software. A custom-made one? Or just a function in matlab?*

Reply: The nonlinear curve-fitting procedures are performed by a custom-made MATLAB software. We have revised our manuscript, which reads (L287-L289) “The nonlinear curve-fitting procedures using iterative least square estimation for the proxies of gaseous sulfuric acid concentration based on Eq. (7) were performed by a custom-made MATLAB software.”

*26. P9: 1-2 orders of magnitude. Maybe change to 10-100 times higher to be more clear.*

Reply: We have revised our manuscript, which reads (L303-L304) “which is about 10-100 times higher ...”.

*27. P9 line 261: 60% RH does not seem dry.*

Reply: The mean RH in this campaign is 28%. We have revised our manuscript, which reads (L312-L313) “In addition, Beijing is dry in winter with a mean ambient relative humidity of 28% during the campaign”.

*28. P9 272: I do not understand how the correlation coefficient numbers are consistent with accepted formation pathways? Does the formation pathways have powers that are less than 1?*

Reply: We have rephrased the sentence, which reads (L324-L326) "...which indicate that  $[\text{SO}_2]$  and UVB have important influences on the formation of atmospheric gaseous sulfuric acid".

*29. P9 line 276: Authors should explain potential reasons why sulfuric acid positively correlates with CS.*

Reply: We discussed in P26 Line 108 that "In several proxies developed by Mikkonen *et al.* (2011), the correlation between the gaseous sulfuric acid concentration and CS is positive, which is against what one would expect because a larger CS normally leads to a faster loss for gaseous sulfuric acid". In this campaign, CS correlates well with  $[\text{SO}_2]$  ( $r = 0.83$ ), which suggests that a high CS value could serve as an indicator of atmospheric particulate pollution, and be accompanied with a high concentration of  $\text{SO}_2$  that is propitious for the formation of gaseous sulfuric acid. Please also refer to our discussion of this issue in section 4.3.

*30. P10 300: molecules  $\text{cm}^{-3}$  is normally written as just  $\text{cm}^{-3}$ .*

Reply: the unit of molecule  $\text{cm}^{-3}$  has been extensively used in the literature and we decide to keep this unit.

*31. P10 line 316: Authors mention that proxy relationship is location specific. Why then did the authors use the justification for not including RH based upon conclusions drawn from a different location?*

Reply: We made a test by introducing RH into the proxy N1 (containing CS terms) and resulted in a RH-corrected CS term (CS·RH) instead of CS as what Mikkonen *et al.* have done (2011). The performance of proxy has not significantly improved (REs changed from 20.04 % to 19.83 %, see our reply to comments #6 and #14 from Reviewer #2 for REs) and the exponent of CS was still close-to-zero (from 0.03 to -0.02). We have rephrased our discussion on RH, which reads (L371-L374) "...because a test by introducing RH into the proxies do not result in a significantly better performance, which is consistent with those conclusions in the Mikkonen *et al.* study (2011)."'

*32. P11 line 324: "unlike assumed in Eq. (3)" wording seems incorrect*

Reply: We have revised the sentence, which reads (L379-L380) "...unlike the assumption in Eq. (6) ...".

*33. P11 line 324: The naming convention between the equations in table 3 and the equations in the paper is confusing. Which equation 3 does this line refer to?*

Reply: We have used the term "function" in Table 3 to avoid confusion.

*34. Page 12 line 356: "Only occasionally slightly higher" too many adverbs. Rephrase*

Reply: We have revised the sentence, which reads (L412-L413) "Occasionally, higher  $[\text{HONO}]/[\text{NO}_x]$  ratios could be seen in the morning".

35. Page 12 line 356: The authors refer to a previous study to justify linearity of NO<sub>2</sub> and HONO.

Where was the location of this study? This paragraph is general is difficult to discern results from previous studies and results from this study. Please make this more clear.

Reply: We have revised this paragraph to focus on the measurements in Beijing, which reads (L406-L420) “Although so far the proxy N5 had the best fitting quality, it is impractical to explicitly include [HONO] because HONO measurements are very challenging. As shown in Fig. 2, [HONO] and [NO<sub>x</sub>] tend to correlate linearly with each other in the daytime during this campaign, with a linearly fitted [HONO]/[NO<sub>x</sub>] ratio of around 0.03 and a relative error of 0.42. Occasionally, higher [HONO]/[NO<sub>x</sub>] ratios could be seen in the morning, which might be due to the fact that HONO concentration could have an accumulation process during the nighttime and lead to a deviation from the steady state. Therefore, due to the good correlation, the proxy N7 replaces [HONO] by [NO<sub>x</sub>], a more easily measured variable, and performs equally well with the proxy N5.”

36. Page 12 line 376: authors should specific that this cover sulfuric acid concentrations for this location. 10<sup>6</sup> cm<sup>-3</sup> does not cover sulfuric acid concentrations around the world.

Reply: We have revised the sentence, which reads (L425-L427) “...in the sulfuric acid concentration range of  $(2.2 - 10) \times 10^6$  cm<sup>-3</sup>, which covers the 5-95% percentiles of sulfuric acid concentrations in this study.”

37. Page 13 line 416: It is a bit confusing that the authors mention that proxy N5 is the most accurate when they spend most of the paper justifying the use of N7. Maybe change the wording “for the best proxy accuracy” or consider rewording this section to make it a bit less confusing/

Reply: We have revised the sentence, which reads (L463) “For a comprehensive consideration of the formation pathways of OH radicals...”.

38. Page 14 line 439: I do not understand how this work has shown the importance of heterogenous chemistry as a potential source of OH. Was this mentioned somewhere else in the main paper? The authors should better justify this point if they want to include in the summary.

Reply: We have removed this statement.

39. Figure 1-2: What day were these measurements done?

Reply: Figure 1-2 show all the measured data points during the campaign from 9 February to 14 March, 2018. We have included the duration of measurements in the revised figure caption.

40. Figure 2: Can the authors explain why there seems to be clear break up group of points during the early morning that do not follow the linear trend?

Reply: We have expanded the discussion, which reads (L412-L415) “Occasionally, higher [HONO]/[NO<sub>x</sub>] ratios could be seen in the morning, which might be due to the fact that HONO concentration could have an accumulation process during the nighttime and lead to a deviation from

the steady state.”

41. *Figure 4: As mentioned above, it would be useful to show the time lines for the other measured concentrations (CS, OH, NO<sub>2</sub>, etc.) that the proxy model uses.*

Reply: Here we present results from a comprehensive campaign with participation from multiple universities and institutions. As a result, this manuscript will only focus on the development of the statistical analysis of the sulfuric acid proxy, and other manuscripts in preparation will discuss the variations of atmospheric variables.

#### Reference

Almeida, J., Schobesberger, S., Kürten, A., Ortega, I. K., Kupiainen-Määttä, O., Praplan, A. P., Adamov, A., Amorim, A., Bianchi, F., Breitenlechner, M., David, A., Dommen, J., Donahue, N. M., Downard, A., Dunne, E., Duplissy, J., Ehrhart, S., Flagan, R. C., Franchin, A., Guida, R., Hakala, J., Hansel, A., Heinritzi, M., Henschel, H., Jokinen, T., Junninen, H., Kajos, M., Kangasluoma, J., Keskinen, H., Kupc, A., Kurtén, T., Kvashin, A. N., Laaksonen, A., Lehtipalo, K., Leiminger, M., Leppä, J., Loukonen, V., Makhmutov, V., Mathot, S., McGrath, M. J., Nieminen, T., Olenius, T., Onnela, A., Petäjä, T., Riccobono, F., Riipinen, I., Rissanen, M., Rondo, L., Ruuskanen, T., Santos, F. D., Sarnela, N., Schallhart, S., Schnitzhofer, R., Seinfeld, J. H., Simon, M., Sipilä, M., Stozhkov, Y., Stratmann, F., Tomé, A., Tröstl, J., Tsagkogeorgas, G., Vaattovaara, P., Viisanen, Y., Virtanen, A., Vrtala, A., Wagner, P. E., Weingartner, E., Wex, H., Williamson, C., Wimmer, D., Ye, P., Yli-Juuti, T., Carslaw, K. S., Kulmala, M., Curtius, J., Baltensperger, U., Worsnop, D. R., Vehkamäki, H. and Kirkby, J.: Molecular understanding of sulphuric acid-amine particle nucleation in the atmosphere, *Nature*, 502(7471), 359–363, doi:10.1038/nature12663, 2013.

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### **Reviewer #2's comments:**

*The manuscript by Lu et al. evaluates different equations for the calculation of gas-phase sulfuric acid from proxy parameters (like  $SO_2$ , UVB radiation, condensation sink, etc.). Different sets of parameters are tested and the performance of the proxy equations is evaluated against in-situ measurements of gas phase  $H_2SO_4$  (measured with a chemical ionization mass spectrometer) and the measured proxy parameters. The measurements took place in Beijing during February/March 2018. While similar proxy expressions for sulfuric acid have been derived for other locations (see Mikkonen et al., 2011, ACP, 11(21), 11319–11334) this is the first one applying such methods for the conditions of a Chinese mega city. Unlike previous studies, the present one includes also HONO as an OH precursor and not just ozone. This leads to a slightly better correlation coefficient between the proxy-derived and measured  $H_2SO_4$ . The manuscript is scientifically relevant and should be published in ACP after the authors have considered the points listed below. Besides the general comments, I have also listed a number of suggestions to improve the language.*

Reply: We are very grateful to the positive viewing of our manuscript by Reviewer #2, and have now revised our manuscript accordingly.

### **General comments:**

1. *Page 1, line 25: “remains a major challenge” is a bit exaggerated, please revise this statement*

Reply: This statement now reads (L25) “but its measurement remains a difficulty”.

2. *Page 2, line 43: How is it known that HONO is formed heterogeneously? Isn't the gas-phase reaction between OH and NO also an efficient source?*

Reply: We agree with reviewer #2 that HONO can be formed from both homogeneous and heterogeneous processes. “heterogeneously-formed” has been removed.

3. *Page 6, line 169: Is the upper size limit of 700 nm sufficient to include all relevant particles contributing to the condensation sink? During dust events, larger particles can probably contribute quite significantly to the CS.*

Reply: The CS values were actually calculated based on the particle size distributions up to 10  $\mu m$ . We have corrected our description of the upper size limit of the PSD system, which reads (L202) “...a conventional particle size distribution system (PSD,  $\sim 3$  nm - 10  $\mu m$ )”.

On the other hand, particle number size distributions up to 700 nm will allow a reasonable calculation of CS, given the fact that most particles are smaller than 700 nm and that there were not significant dust events during our measurements.

4. *Page 8, line 231 to 234: It should be explained in more detail how  $k$  is calculated and how it relates to  $k_0$ .*

Reply: An explanation on how  $k$  is calculated has been added, which reads (L247-L255)

“...where  $k$  is a temperature-dependent reaction constant given by Eq. (5) (DeMore *et al.*, 1997; Mikkonen *et al.*, 2011).

$$k = \frac{A \cdot k_3}{(A + k_3)} \cdot \exp \left\{ k_5 \cdot \left[ 1 + \log_{10} \left( \frac{A}{k_3} \right)^2 \right]^{-1} \right\} \quad \text{cm}^3(\text{molecule} \cdot \text{s})^{-1} \quad (5)$$

where  $A = k_1 \cdot [M] \cdot \left( \frac{300}{T} \right)^{k_2}$ ,  $[M]$  represents the density of the air in molecule  $\text{cm}^{-3}$  as calculated by  $0.101 \cdot (1.381 \cdot 10^{-23} \cdot T)^{-1}$ ,  $k_1 = 4 \cdot 10^{-31}$ ,  $k_2 = 3.3$ ,  $k_3 = 2 \cdot 10^{-12}$  and  $k_5 = -0.8$ .”

We have also elaborated the explanation on how  $k$  relates to  $k_0$ , which reads (L316-L320) “Since  $k$  is a temperature-dependent reaction constant and varies within a 10 % range in the atmosphere temperature range of 267.6 - 292.6 K, *i.e.*, the actual atmospheric temperature variation in this study, we approximately regard  $k$  as a constant and use a new scaling factor  $k_0$ . This methodology has been used previously in the proxies of gaseous sulfuric acid in Hyytiälä Southern Finland (Petäjä *et al.*, 2009)”.

5. *Page 9, line 281 to 284: Could the correlation between HONO and NO<sub>2</sub> also be caused by the gas phase reaction between OH and NO (because NO correlates probably strongly with NO<sub>2</sub>)?*

Reply: As stated in the very beginning, “The [NO<sub>2</sub>] concentration” in our previous manuscript is in fact “the [NO<sub>x</sub>] concentration” due to a key typo. Hence, the correlation coefficients (Spearman type) between NO<sub>x</sub> and HONO, between NO and HONO, and between NO<sub>2</sub> and HONO are 0.88, 0.74 and 0.88, respectively. Although a slightly better correlation between NO<sub>2</sub> and HONO was observed, we cannot exclude the role of the gas phase reaction between OH and NO, and the interconversion between NO and NO<sub>2</sub>. In fact, we agree with reviewer #2 that HONO can be both homogeneously and heterogeneously formed, although heterogeneous formation from NO<sub>2</sub> is likely the reason for the daytime HONO production in urban Beijing (Liu *et al.*, 2014). In the revised manuscript, we have stated (L335-L339) that “A strong correlation between [HONO] and [NO<sub>x</sub>] ( $r = 0.88$ ) in our measurement is supported by the fact that HONO can be either heterogeneously formed by reactions of NO<sub>2</sub> on various surfaces (Calvert *et al.*, 1994) or homogeneously formed by the gas phase NO + OH reaction, between which the former likely dominate for the daytime HONO production in urban Beijing (Liu *et al.*, 2014).”

6. *Page 10, line 317 to 319: The mean absolute error does not seem to be the best metric for evaluating the fitting of data that vary over more than one order of magnitude. This can lead to a bias where the high values are well represented but the relative deviation for the small values can be large. A better metric could be the average ratio (sum over all max(V<sub>i</sub>\_measured, V<sub>i</sub>\_fitted) / min(V<sub>i</sub>\_measured, V<sub>i</sub>\_fitted), where V<sub>i</sub> is a data point for the considered parameter, e.g. [H<sub>2</sub>SO<sub>4</sub>]).*

Reply: In this study, the gaseous sulfuric acid concentrations are in a range of (2.2-10.0)  $\times 10^6$  molecule cm<sup>-3</sup> in the 5-95% percentiles, whose variation is less than one order of magnitude. Nevertheless, we have now defined a metric of “relative error” (RE) to evaluate the fitting of data, which turns out to be 20.04 % (N1), 20.00 % (N2), 19.95 % (N3), 19.95 % (N4), 19.11 % (N5), 19.66 % (N6), and 19.34 % (N7), respectively. These results are consistent with our previous MAE results. The new metric is used throughout the revised manuscript and introduced as Eq. (8) (L293)

$$RE = \frac{1}{n} \cdot \sum_{i=1}^n \frac{|[H_2SO_4]_{proxy,i} - [H_2SO_4]_{meas,i}|}{[H_2SO_4]_{meas,i}} \quad (8)$$

7. *Page 11, line 346/347: The improvement relates to an increase of R from 0.85 to 0.86; this does not justify the word “significantly”; “improved slightly” instead of “improved significantly” is more appropriate.*

Reply: We have revised our manuscript, which reads (L400-L402) “When both [O<sub>3</sub>] and [HONO] were introduced as OH precursors in the proxies N5-N7, REs have noticeable improvements, and correlation coefficients improved slightly.”

8. *Page 13, line 396: Aren’t the different values of CS taken into account in the calculation, or do the authors mean that the different exponents for CS make the difference? What is the exponent for the CS in the Petäjä *et al.* study?*

Reply: We think it is the value of CS<sup>c</sup> in the proxies that makes the difference. For example, if the exponent C is very near to zero, then no matter how CS changes, the value of CS<sup>c</sup> would always be very close to 1, which means that this term would not influence the proxies at all. The CS in the Petäjä *et al.* study did not have an exponent.

9. *Page 14, line 439: How can the proxies be evaluated for past measurements? If measurements exist for both sulfuric acid and the proxy parameters, these should be included in the present study.*

Reply: We failed to obtain previous data sets that include both gaseous sulfuric acid concentrations and other proxy parameters. As far as we know, there are two studies that measured gaseous sulfuric acid concentrations in Beijing (Zheng *et al.*, 2011; Cai *et al.*, 2017), but other key inputs for the proxy are not available from the two studies.

10. *Page 14, line 440: The importance of heterogeneous chemistry cannot be concluded from the presented data; this statement should be removed or supported with further data.*

Reply: We have removed this statement.

11. *Section 5: Discussion on the expected seasonality of the proxy-H<sub>2</sub>SO<sub>4</sub> relation should be discussed. Very likely, the exponents can have different values for other meteorological*

*condition, e.g., when desert dust contributes to the particle burden. In such a case, the exponent for the CS could become negative. This possibility should be mentioned/discussed.*

Reply: We have revised the conclusion section, and expanded the discussion on the applicability of the proxies in this study, which reads (L491-L496) “Furthermore, the proxies might be site-specific and season-specific. Since the proxies were derived with atmospheric parameters in winter, in urban Beijing, the exponents for atmospheric variables in the proxy could have different values for other cities or other seasons. Thus, the proxies in this study should be further tested before their application to other Chinese megacities or other seasons”.

*12. Table 1: The values for ozone are quite low. Are these low values typical for wintertime conditions in Beijing?*

Reply: The ozone concentration in winter 2018 in Beijing is actually lower than those in the past years. In addition, as our station is not far from a traffic-heavy road, sometimes,  $O_3$  could be completely diminished by NO.

*13. Table 3: What is the definition of the scaling factor  $k_0$ ? What are its units?*

Reply: The scaling factor  $k_0$ , which scales the calculated values from the proxy variables to match the measured sulfuric acid concentrations, is derived from the ratio of measured sulfuric acid concentrations and the proxy concentrations (without  $k_0$  itself). The units of  $k_0$  in different proxies are different, but together with units of all variables would result in a unit of molecule  $cm^{-3}$ .

*14. Figure 3: The yellow areas are hardly visible; it would probably be better to use colored lines instead of filled areas for this figure. In addition, the shape of the curves suggests a pronounced bias (the high values are on average underestimated, while the low values are overestimated). This bias can also be seen in the SI Figures. Using a different metric for the fitting (ratios instead of absolute differences, see above) could solve this issue.*

Reply: We have revised Figure 3. Colored lines are now used to present the performance of Proxy N2 and Proxy N7. A new metric of “relative error” is used to evaluate the fitting quality.

### **Language comments:**

*1. Page 2, line 46: “less than” rather than “up to”?*

Reply: we have revised our manuscript, which reads (L49) “the relative errors were reduced up to 20 %”.

*2. Page 3, line 57: “involve” instead of “demand”*

Reply: We have revised our manuscript accordingly, which reads (L61) “...involve the participation

of gaseous sulfuric acid molecules”.

3. *Page 3, line 73/74: delete “using a passive CIMS”*

Reply: We have removed this expression.

4. *Page 3, line 76: delete “associated equilibrium or”*

Reply: We have removed this expression.

5. *Page 3, line 79: replace “will be” with “are”*

Reply: We have revised our manuscript accordingly, which reads (L83-L84) “...known concentrations of OH radicals that are titrated into gaseous sulfuric acid”.

6. *Page 4, line 90: replace “supposing” with “assuming”*

Reply: We have revised our manuscript accordingly, which reads (L95-L96) “assuming that gaseous sulfuric acid is formed from reactions between SO<sub>2</sub> and OH radicals”.

7. *Page 4, line 102: delete “; a proxy for condensational sink”*

We have removed this expression.

8. *Page 4, line 108: delete “that characterized”*

We have removed this expression.

9. *Page 4, line 114: replace “between” with “in”*

Reply: We have revised our manuscript accordingly, which reads (L120-L122) “...measured gaseous sulfuric acid concentrations are relatively similar in these environments...”.

10. *Page 6, line 156: “the sample flow”*

Reply: We have revised our manuscript accordingly, which reads (L163) “...the middle of the sample flow”.

11. *Page 6, line 177: “dependent on” instead of “that could be defined as a function of”*

Reply: We have revised our manuscript accordingly, which reads (L211-L212) “... a transition-regime correction factor dependent on the Knudsen number ...”.

12. *Page 8, line 230: What is meant by the symbol xi’?*

Reply: We have revised our manuscript accordingly to make it clear, which reads (L274-L275) “...

where  $x_i$  can be an atmospheric variable such as UVB and  $[\text{SO}_2]$ ".

13. *Page 9, line 255: "in the 5-95% percentile range" instead of "was in . . . range of"*

Reply: We have revised our manuscript accordingly, which reads (L306) "in the 5-95 % percentile range, relatively similar to...".

14. *Page 9, line 280: delete "an"*

Reply: We have removed this word.

15. *Page 10, line 290: "lamination" does not seem to be the right word here, maybe better to use "layering with"*

Reply: We have revised our manuscript accordingly, which reads (L345) "A good correlation layering with  $[\text{SO}_2]$  is evident...".

16. *Page 10, line 305: "has" instead of "have"*

Reply: We have revised our manuscript accordingly, which reads (L360) "...that has the UVB intensity and  $[\text{SO}_2]$  ...".

17. *Page 11, line 352: either delete "are tended to" or use "tend to"*

Reply: We have revised our manuscript accordingly, which reads (L408) "[HONO] and  $[\text{NO}_x]$  tend to correlate linearly with each other ...".

18. *Page 12, line 382: Do the authors mean "deviations" instead of "derivations"?*

Reply: Following this reviewer's comments #6 and #14, we have used a new metric of "relative error" (L293) instead of relative deviation in Figure 3.

19. *Page 13, line 390: "when averaged to a time resolution of 5 min" instead of "with a time resolution of 5 min"*

Reply: We have revised our manuscript accordingly, which reads (L437) "when averaged to a time resolution of 5 min".

20. *Page 13, line 411: Please provide the units for the parameters in this equation.*

Reply: We have added the units for the parameters, which reads (L457-L458) "The units of  $[\text{H}_2\text{SO}_4]$  and  $[\text{SO}_2]$  are molecule  $\text{cm}^{-3}$ , and the unite of UVB is  $\text{W m}^{-2}$ ".

21. *Page 14, line 418/419: Please provide the units for the parameters in this equation.*

Reply: We have added the units for the parameters, which reads (L465-L466) “The units of  $[H_2SO_4]$ ,  $[SO_2]$ ,  $[O_3]$  and  $[HONO]$  are molecule  $cm^{-3}$ , the unite of UVB is  $W\ m^{-2}$ , and the unite of CS is  $s^{-1}$ ”.

22. *Page 14, line 425/426 Please provide the units for the parameters in this equation.*

Reply: We have added the units for the parameters, which reads (L473-L474) “The unites of  $[H_2SO_4]$ ,  $[SO_2]$ ,  $[O_3]$  and  $[NO_x]$  are molecule  $cm^{-3}$ , the unite of UVB is  $W\ m^{-2}$ , and the unite of CS is  $s^{-1}$ ”.

23. *Page 14, 433/434: Please provide also the equation numbers and not just the proxy numbers (N2 and N7).*

Reply: We have now provided both the equation numbers and proxy numbers, which reads (L484-L485) “As a summary, we recommend using the simplest proxy (proxy N2 as shown in Eq. (9)) and a more accurate proxy (Proxy N7 as shown in Eq. (11)) for calculating daytime gaseous sulfuric acid …”.

## Reference

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**Santtu Mikkonen's comments:**

*It is interesting to see how the sulphuric acid concentration can be approximated in highly polluted environment, as we did not have such data when we were making our paper Mikkonen et al. (2011). Even more interesting is, that your recommended proxy N2 is quite close to our second recommendation, simple proxy L3, having SO<sub>2</sub> power to 0.5 when you have power of 0.4. In addition, I was surprised that the H<sub>2</sub>SO<sub>4</sub> concentration was not higher than shown in Table 1. We had similar average concentrations in San Pietro Capofiume and considerably higher in Atlanta, even though they are less polluted environments. Could you add a comment on that?*

Reply: We are very grateful to the positive viewing of our manuscript by Dr. Santtu Mikkonen, and have now revised our manuscript accordingly.

As Dr. Mikkonen has noticed, Beijing did not in this campaign have a higher average concentration of sulfuric acid than other cities, which can be potentially explained by the fact that, firstly, the averaged condensation sink in Beijing in this campaign is around 0.11 s<sup>-1</sup> that corresponds to a very efficient removal of gaseous sulfuric acid, and secondly, the SO<sub>2</sub> concentration has dramatically reduced in recent years in Beijing as we have mentioned in L488-L489.

*I just want to ask about Figure 4: Why only one day, and not averages over whole period such that uncertainty would also be indicated, is shown in the figure? In addition, why comparison only to Boreal forest-proxy from Petäjä et al, why not to Mikkonen et al., who had data from multiple sites?*

Reply: Figure 3 (now updated to a new version) presents a statistical comparison between measured and predicted sulfuric acid concentrations over the whole period.

We compared our results with the Petäjä et al. study instead of the Mikkonen et al. study, simply because we only measured the UVB and a correlation between UVB and the global radiation cannot be established.

*A minor comment on the use of p-value as a screening factor for correlation (in line 266): it is not recommended. See e.g. Greenland et al. (2016): DOI 10.1007/s10654-016-0149-3*

Reply: We removed p-values as a screening factor for correlations. Now all the correlations are shown in Table 2.

1   **A proxy for atmospheric daytime gaseous sulfuric acid**  
2   **concentration in urban Beijing**

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24   **Abstract.** Gaseous sulfuric acid is known as one of the key precursors for atmospheric  
25   new particle formation processes, but its measurement remains a difficulty~~but its~~  
26   measurement remains a major challenge. A proxy method that is able to derive gaseous  
27   sulfuric acid concentrations from parameters that can be measured relatively easily and  
28   accurately is therefore highly desirable for the atmospheric chemistry community~~highly~~  
29   desirable among the atmospheric chemistry community. Although such methods are  
30   available for clean atmospheric environments, a proxy that works well in a polluted  
31   atmosphere, such as those in Chinese megacities, is yet to be developed. In this study,  
32   the gaseous sulfuric acid concentration was measured in February-March, 2018, in  
33   urban Beijing by a nitrate based - Long Time-of-Flight Chemical Ionization Mass  
34   Spectrometer (LToF-CIMS). A number of atmospheric parameters were recorded  
35   concurrently including the ultraviolet radiation B (UVB) intensity, concentrations of O<sub>3</sub>,  
36   NO<sub>x</sub> (sum of NO and NO<sub>2</sub>), SO<sub>2</sub> and HONO, and aerosol particle number size

37 distributions. A proxy for atmospheric daytime gaseous sulfuric acid concentration was  
38 derived ~~using through~~ a statistical analysis method by using the UVB intensity,  $[\text{SO}_2]$ ,  
39 condensation sink (CS),  $[\text{O}_3]$ , and  $[\text{HONO}]$  (or  $[\text{NO}_x]$ ) as the predictor variables. In this  
40 proxy method, we considered the formation of gaseous sulfuric acid from reactions of  
41  $\text{SO}_2$  and OH radicals during the daytime, and loss of gaseous sulfuric acid due to its  
42 condensation onto the pre-existing particles. In addition, we explored formation of OH  
43 radicals from the conventional gas-phase photochemistry using ozone as a proxy and  
44 from the photolysis of ~~heterogeneously formed~~ HONO using HONO (and subsequently  
45  $\text{NO}_x$ ) as a proxy. Our results showed that the UVB intensity and  $[\text{SO}_2]$  are dominant  
46 factors for the production of gaseous sulfuric acid, and that the simplest proxy could be  
47 constructed with the UVB intensity and  $[\text{SO}_2]$  alone. ~~When the OH radical production~~  
48 ~~from both homogenously- and heterogeneously-formed precursors were considered, the~~  
49 ~~relative errors were reduced up to 20 %, resulting in up to 29% relative deviations~~  
50 ~~when sulfuric acid concentrations were larger than  $2.0 \times 10^6$  molecules  $\text{cm}^{-3}$ . When~~  
51 ~~the OH radical production from both homogenously- and heterogeneously formed~~  
52 ~~precursors were considered, the relative deviations were lower than 24%.~~

53 **1 Introduction**

54 Gaseous sulfuric acid ( $\text{H}_2\text{SO}_4$ ) is a key precursor for atmospheric new particle  
55 formation (NPF) processes (Kerminen, 2018; Kirkby et al., 2011; Kuang et al., 2008;  
56 Kulmala and Kerminen, 2008; Sipilä et al., 2010). A number of atmospheric nucleation  
57 mechanisms including  $\text{H}_2\text{SO}_4\text{-H}_2\text{O}$  binary nucleation (Benson et al., 2008; Duplissy et  
58 al., 2016; Kirkby et al., 2011),  $\text{H}_2\text{SO}_4\text{-NH}_3\text{-H}_2\text{O}$  ternary nucleation (Kirkby et al., 2011;  
59 Korhonen et al., 1999; Kürten et al., 2015), and  $\text{H}_2\text{SO}_4\text{-DMA-H}_2\text{O}$  ternary nucleation  
60 ([Almeida et al., 2013](#); [Jen et al., 2014](#); [Kürten et al., 2014](#); [Petäjä et al., 2011](#); [Yao et al., 2018](#)) ~~involved~~demanded the participation of gaseous sulfuric acid molecules. In addition,  
61 the condensation of gaseous sulfuric acid onto newly-formed particles contributes to  
62 their initial growth ([Kuang et al., 2012](#); [Kulmala et al., 2013](#)). Quantitative assessments  
63 of the contribution of gaseous sulfuric acid to both the new particle formation rates and  
64 the particle growth rates require real-time measurements of gaseous sulfuric acid  
65 concentrations ~~prior to and~~ during the NPF events (Nieminens et al., 2010; Paasonen et  
66 al., 2010).

67 Measurements of gaseous sulfuric acid in the lower troposphere are challenging  
68 because its ambient concentration is typically quite low ( $10^6\text{-}10^7$  molecule  $\text{cm}^{-3}$ )  
69 (Kerminen et al., 2010; Mikkonen et al., 2011). Reported real-time measurements of  
70 gaseous sulfuric acid are currently based on Chemical Ionization Mass Spectrometry  
71 with  $\text{NO}_3^-$  ~~and its ligands~~ as reagent ions (nitrate CIMS) because [nitrate CIMS with an](#)  
72 [atmospheric pressure interface \(API\)CIMS](#) has a low detection limit for the  
73 atmospheric concentration range of gaseous sulfuric acid (Jokinen et al., 2012), and a  
74 constant fraction of sulfuric acid present in the air sample will be ionized by excessive  
75 nitrate ions in CIMS under constant instrumental conditions (Kürten et al., 2012; Zheng  
76 et al., 2010), which makes the quantification of gaseous sulfuric acid feasible.

77 Arnold and Fabian (1980) measured the negative ions in the stratosphere ~~using a~~  
78 ~~passive CIMS~~ and derived the concentration of stratospheric gaseous sulfuric acid from  
79 the fractional abundances of a series of stratospheric negative ions as well as the  
80 ~~associated equilibrium or~~ rate constants. Later, real-time measurement of sulfuric acid  
81 in the lower troposphere was performed using nitrate CIMS (Eisele and Tanner, 1993),  
82 with laboratory calibrations by production of known concentrations of OH radicals that  
83 ~~will bear~~ titrated into gaseous sulfuric acid. Thereafter, measurements of sulfuric acid  
84 using CIMS have been performed around the world ([e.g., Berresheim et al., 2000](#);  
85 [Bianchi et al., 2016](#); [Chen et al., 2012](#); [Jokinen et al., 2012](#); [Kuang et al., 2008](#); [Kürten](#)

87 [et al., 2014; Kurtén et al., 2011; Petäjä et al., 2009; Weber et al., 1997; Zheng et al.,](#)  
88 [2011](#)), and CIMS has been proved to be a robust tool for gaseous sulfuric acid  
89 detection. However, sulfuric acid measurements are still rather sparse because of the  
90 high cost of the CIMS instrument and the extensive demand of specialized expertise on  
91 the instrument calibration, maintenance, and data processing, etc. Therefore, a proxy  
92 for gaseous sulfuric acid concentration is highly desirable.

93 Proxies for the estimation of atmospheric gaseous sulfuric acid concentrations  
94 were previously developed to approximate measurement results of sulfuric acid  
95 in Hyytiälä, Southern Finland (Petäjä et al., 2009), [assuming](#) that gaseous  
96 sulfuric acid is formed from reactions between  $\text{SO}_2$  and OH radicals, and lost due to its  
97 condensation onto pre-existing particles. The derived simplest proxy can be written as  
98 Eq. (1) below, and the authors recognized that the proxies might be site-specific and  
99 should be verified prior to utilization in other environments.

$$100 \quad [H_2SO_4] = k \cdot \frac{[SO_2] \cdot (UVB \text{ or Global radiation})}{CS} \quad (1)$$

102  
103 Mikkonen et al. (2011) later developed a couple of statistical proxies based on  
104 measurements of sulfuric acid in six European and North American sites, including  
105 urban, rural and forest areas. Their results showed that the radiation intensity and  $[SO_2]$   
106 are the most important factors to determine the concentration of sulfuric acid, and that  
107 the impact of condensation sink (CS), [a proxy for condensational sink](#) for gaseous  
108 sulfuric acid, is generally negligible. In several proxies developed by Mikkonen et al.  
109 (2011), the correlation between the gaseous sulfuric acid concentration and CS is  
110 positive, which is against what one would expect [because a larger CS normally leads](#)  
111 [to a faster loss for gaseous sulfuric acid](#). In addition, the performance of a proxy  
112 equation is site-specific because of varying atmospheric conditions from one site to  
113 another, which implies that the proxy suggested by Mikkonen et al. (2011) might not  
114 work well in locations [that characterized with an atmospheric environments](#) different  
115 from those in the six sites of that study.

116 Beijing is a location with typical values of CS ([e.g., 0.01-0.24 s<sup>-1</sup> in the 5-95%](#)  
117 [percentiles in this study](#)) being 10-100 times higher (Herrmann et al., 2014; Wu et al.,  
118 2007; Xiao et al., 2015; Yue et al., 2009; Zhang et al., 2011) and typical  $\text{SO}_2$   
119 concentrations being 1-10 times higher (Wang et al., 2011a; Wu et al., 2017) than those

120 in Europe and North America (Dunn et al., 2004; Mikkonen et al., 2011), yet measured  
121 gaseous sulfuric acid concentrations are relatively similar ~~between-in~~ these  
122 environments (Chen et al., 2012; Smith et al., 2008; Wang et al., 2011b; Zheng et al.,  
123 2011). Whether previous proxies developed for European and North American sites  
124 work in Beijing remains to be tested. Furthermore, in addition to the gas phase reaction  
125 between O(<sup>1</sup>D) and water molecules (Crutzen and Zimmermann, 1991; Logan et al.,  
126 1981), photolysis of HONO could be a potentially important source of OH radicals~~s~~ in  
127 the atmosphere ~~not only~~ in the early morning (Alicke et al., 2002, 2003; Elshorbany et  
128 al., 2009; Li et al., 2012) ~~and~~ ~~but also~~ during the daytime (Acker et al., 2005; Aumont  
129 et al., 2003; Kleffmann, 2007). An experimental study measuring HONO near the  
130 surface layer estimated that HONO was a main contributor to OH production in Beijing,  
131 with HONO's contribution being larger than 70 % at around 12:00-13:00, except for  
132 summer when the contribution of O<sub>3</sub> dominated (Hendrick et al., 2014). Given the  
133 distinct characteristics of these two OH radical formation pathways, they both should  
134 be included and evaluated separately when a proxy for atmospheric gaseous sulfuric  
135 acid concentration is being built. The reactions between SO<sub>2</sub> and ~~C~~riegee  
136 intermediates formed from the ozonolysis of atmospheric alkenes could be a potential  
137 source of sulfuric acid only in the absence of solar radiation (Boy et al., 2013; Mauldin  
138 et al., 2012), so these reactions are expected to provide a minor contribution to the  
139 formation of gaseous sulfuric acid during the daytime in urban Beijing.

140 In this study, gaseous sulfuric acid concentration was measured by a Long Time-  
141 of-Flight Chemical Ionization Mass Spectrometer (LToF-CIMS) in February - March,  
142 2018, in urban Beijing. A number of atmospheric parameters were recorded  
143 concurrently, including the ultraviolet radiation B (UVB) intensity, concentrations of  
144 O<sub>3</sub>, NO<sub>x</sub>, SO<sub>2</sub> and HONO, and particle number size distributions. The objective of this  
145 study is to develop a robust daytime gaseous sulfuric acid concentration proxy for  
146 Beijing, a representative Chinese megacity with urban atmospheric environments.

147 **148 2 Ambient measurements**

149 An intensive campaign was carried out from 9 February to 14 March, 2018 on the  
150 fifth floor of a teaching building in the west campus of Beijing University of Chemical  
151 Technology (39°94' N, 116°30' E). This monitoring site is 2 km to the west of the  
152 West 3<sup>rd</sup> Ring Road and surrounded by commercial properties and residential dwellings.  
153 Hence, this station can be regarded as a representative urban site.

154 The sulfuric acid concentration was measured by a LToF-CIMS (Aerodyne  
 155 Research, Inc.) equipped with a nitrate chemical ionization source. Ambient air was  
 156 drawn into the ionization source through a stainless-steel tube with a length of 1.6 m  
 157 and a diameter of 3/4 inch. A mixture of a 3 standard cubic centimeter per minute (sccm)  
 158 ultrahigh purity nitrogen flow containing nitric acid and a 20 standard liter per minute  
 159 (slpm) pure air flow supplied by a zero-air generator (Aadco 737, USA), together as a  
 160 sheath flow, was ~~introduced into guided through~~ a PhotoIonizer (Model L9491,  
 161 Hamamatsu, Japan) to produce nitrate reagent ions. This sheath flow was then  
 162 introduced into a co-axial laminar flow reactor concentric to the sample flow. Nitrate  
 163 ions were pushed to the middle of the sample flow under an electric field and  
 164 subsequently charged sample molecules. For example, the atmospheric H<sub>2</sub>SO<sub>4</sub>  
 165 molecules would be charged by nitrate reagent ion NO<sub>3</sub><sup>-</sup>(HNO<sub>3</sub>)<sub>0-2</sub> and mainly produce  
 166 HSO<sub>4</sub><sup>-</sup> ions (m/z = 96.9601 Th), HSO<sub>4</sub><sup>-</sup>·HNO<sub>3</sub> ions (m/z = 159.9557 Th), and  
 167 HSO<sub>4</sub><sup>-</sup>·(HNO<sub>3</sub>)<sub>2</sub> ions (m/z = 222.9514 Th). In addition, HSO<sub>4</sub><sup>-</sup>·H<sub>2</sub>SO<sub>4</sub> ions (m/z =  
 168 194.9275 Th) were formed from ion-induced clustering of neutral sulfuric acid and  
 169 bisulfate ions within the LToF-CIMS ion reaction zone, and also from the evaporation  
 170 of dimethylamine (DMA) and the replacement of one molecule of H<sub>2</sub>SO<sub>4</sub> with one  
 171 bisulfate ion, HSO<sub>4</sub><sup>-</sup>, during the NO<sub>3</sub><sup>-</sup> reagent ion charging of a stabilized neutral  
 172 sulfuric acid dimer in the real atmosphere in presence of DMA or a molecule that works  
 173 in the same way as DMA. During the campaign, the sample flow rate was kept at 8.8  
 174 slpm, since mass flow controllers fixed the sheath flow rate and the excess flow rate,  
 175 and the flow into the mass spectrometer (around 0.8 slpm) was fixed by the size of a  
 176 pinhole between the ionization source and the mass spectrometer. The concentration of  
 177 gaseous sulfuric acid was then determined by Eq. (2).

$$[H_2SO_4] = \frac{HSO_4^- \cdot (HNO_3)_{0-2} + HSO_4^- \cdot H_2SO_4}{NO_3^-(HNO_3)_{0-2}} \cdot C \quad (2)$$

181 where C is the calibration coefficient, and NO<sub>3</sub><sup>-</sup>(HNO<sub>3</sub>)<sub>0-2</sub>, HSO<sub>4</sub><sup>-</sup>·(HNO<sub>3</sub>)<sub>0-2</sub> and  
 182 HSO<sub>4</sub><sup>-</sup>·H<sub>2</sub>SO<sub>4</sub> represent the signals of corresponding ions and are in units of counts per  
 183 second (cps). The unit of resulting [H<sub>2</sub>SO<sub>4</sub>] is molecule cm<sup>-3</sup>. The CIMS was calibrated  
 184 during the campaign with a home-made calibration box that can produce adjustable  
 185 concentrations of gaseous sulfuric acid from SO<sub>2</sub> and OH radicals following the  
 186 protocols in previous literatures (Kürten et al., 2012; Zheng et al., 2015). We obtain a  
 187 calibration coefficient of 3.79 × 10<sup>9</sup> molecule cm<sup>-3</sup> for our instrument and use 1.1 ×

188 ~~10<sup>10</sup> molecule cm<sup>-3</sup> as the effective calibration coefficient, after taking into account~~  
189 ~~the diffusion losses in the stainless-steel tube and the nitrate chemical ionization~~  
190 ~~source. During the campaign, the sample flow rate was kept at 8.4 slpm, since mass flow~~  
191 ~~controllers fixed the sheath flow rate and the excess flow rate and the flow into the mass~~  
192 ~~spectrometer (around 0.4 slpm) was fixed by the size of a pinhole between the~~  
193 ~~ionization source and the mass spectrometer. The CIMS was calibrated twice during~~  
194 ~~the campaign following the protocols in previous literatures (Kürten et al., 2012; Zheng~~  
195 ~~et al., 2015). Here we use 1.1 × 10<sup>10</sup> molecule cm<sup>-3</sup> as the calibration coefficient, after~~  
196 ~~taking into account diffusion losses in the stainless steel tube and the nitrate chemical~~  
197 ~~ionization source. The obtained mass spectra were analyzed with a tofTools package~~  
198 ~~based on the MATLAB software (Junninen et al., 2010).~~

199       Ambient particle number size distributions down to about 1 nm were measured  
200 using a combination of a scanning mobility particle sizer spectrometer (SMPS)  
201 equipped with a diethylene glycol-based condensation particle counter (DEG-CPC, ~1-  
202 10 nm) and a conventional particle size distribution system (PSD, ~3 nm - 10  $\mu$ m<sup>3</sup> 700  
203 ~~nm~~) consisting of a pair of aerosol mobility spectrometers developed by Tsinghua  
204 University (Cai et al., 2017; Jiang et al., 2011; Liu et al., 2016). The values of CS were  
205 calculated following Eq. (3) (Dal Maso et al., 2002):

206

$$207 \quad CS = 2\pi D \int_0^{\infty} D_p \beta_m(D_p) n(D_p) dD_p = 2\pi D \sum_i \beta_i D_{pi} N_i \quad (3)$$

208

209 where  $D_{pi}$  is the geometric mean diameter of particles in the size bin  $i$  and  $N_i$  is the  
210 particle number concentration in the corresponding size bin.  $D$  is the diffusion  
211 coefficient of gaseous sulfuric acid, and  $\beta_m$  represents a transition-regime correction  
212 factor ~~dependent on that could be defined as a function of~~ the Knudsen number (Fuchs  
213 and Sutugin, 1971; Gopalakrishnan and Hogan Jr., 2011).

214       SO<sub>2</sub>, O<sub>3</sub> and NO<sub>x</sub> concentrations were measured using a SO<sub>2</sub> analyzer (Model 43i,  
215 Thermo, USA), a O<sub>3</sub> analyzer (Model 49i, Thermo, USA) and a NO<sub>x</sub> analyzer (Model  
216 42i, Thermo, USA) with the detection limits of 0.1 ppbv, 0.5 ppbv and 0.4 ppbv,  
217 respectively. The above instruments were pre-calibrated before the campaign. The UVB  
218 (280 - 315 nm) intensity (UV-S-B-T, KIPP&ZONEN, The Netherlands) was measured  
219 on the rooftop of the building. Atmospheric HONO concentrations were measured by  
220 a home-made HONO analyzer with a detection limit of 0.01 ppbv (Tong et al., 2016).

221 Particle number size distributions and concentrations of gaseous sulfuric acid,  $\text{SO}_2$ ,  
222  $\text{O}_3$ ,  $\text{NO}_{\text{x}2}$  and  $\text{HONO}$  were recorded with a time resolution of 5 min, and the UVB  
223 intensity with time resolution of 1 min. A linear interpolation method was used for  
224 deriving the variables with the same time intervals, *i.e.*, 5 min. Only data between local  
225 sunrise and sunset were used in the subsequent analysis.

### 227 3 Development of a proxy for atmospheric gaseous sulfuric acid

228 We derived the gaseous sulfuric acid concentration proxy on the basis of currently  
229 accepted formation pathways of sulfuric acid in the atmosphere (R1-R3) (Finlayson-  
230 Pitts and Pitts, 2000; Stockwell and Calvert, 1983):



235 The reaction (R1) is the rate-limiting step of this formation pathway (Finlayson-Pitts  
236 and Pitts, 2000), so our proxy will consider the two major processes that determine the  
237 abundance of gaseous sulfuric acid: the formation of gaseous sulfuric acid from  
238 reactions between  $\text{SO}_2$  and OH radicals, and the loss of gaseous sulfuric acid due to its  
239 condensation onto pre-existing particles (Dal Maso et al., 2002; Kulmala et al., 2012;  
240 Pirjola et al., 1999).

241 The rate of change of sulfuric acid concentration can be written as Eq. (4)  
242 (Mikkonen et al., 2011):

$$245 \quad \frac{d[\text{H}_2\text{SO}_4]}{dt} = k \cdot [\text{OH}] \cdot [\text{SO}_2] - [\text{H}_2\text{SO}_4] \cdot CS \quad (4)$$

246  
247 where  $k$  is a temperature-dependent reaction constant given by Eq. (5) (DeMore et al.,  
248 1997; Mikkonen et al., 2011).

$$249 \quad k = \frac{A \cdot k_3}{(A + k_3)} \cdot \exp \left\{ k_5 \cdot \left[ 1 + \log_{10} \left( \frac{A}{k_3} \right)^2 \right]^{-1} \right\} \quad \text{cm}^3(\text{molecule} \cdot \text{s})^{-1} \quad (5)$$

250  
251 where  $A = k_1 \cdot [M] \cdot \left( \frac{300}{T} \right)^{k_2}$ ,  $[M]$  represents the density of the air in molecule  $\text{cm}^{-3}$

253 as calculated by  $0.101 \cdot (1.381 \cdot 10^{-23} \cdot T)^{-1}$ ,  $k_1 = 4 \cdot 10^{-31}$ ,  $k_2 = 3.3$ ,  $k_3 = 2 \cdot$   
254  $10^{-12}$  and  $k_5 = -0.8$ , where  $k$  is a temperature-dependent reaction constant (DeMore  
255 et al., 1997).

256 To simplify the calculation, the production and loss of sulfuric acid can be assumed  
257 to be at pseudo steady-state (Mikkonen et al., 2011; Petäjä et al., 2009). Then the  
258 sulfuric acid concentration can be written as Eq. (6).

259

$$260 [H_2SO_4] = k \cdot [OH] \cdot [SO_2] \cdot CS^{-1} \quad (6)$$

261

262 Atmospheric OH radical measurements represent a major challenge as well. Since  
263 previous studies suggest that the OH radical concentration is strongly correlated with  
264 the intensity of UVB,  $[OH]$  could be replaced with UVB intensity in the proxy equation  
265 (Petäjä et al., 2009; Rohrer and Berresheim, 2006). Although photolysis of  $O_3$   
266 ( $\lambda < 320$  nm) and subsequent reactions with  $H_2O$  are considered to be the dominant  
267 source of OH radicals in the atmosphere (Logan et al., 1981), recent studies argue that  
268 photolysis of HONO ( $\lambda < 400$  nm) is a potentially important OH radical formation  
269 pathway (Hendrick et al., 2014; Kleffmann, 2007; Su et al., 2011; Villena et al., 2011).  
270 Thus, we attempt to introduce both  $O_3$  and HONO into the proxy equation and evaluate  
271 their effects on the concentration of OH radicals.

272 In practice, the ~~exponents for variables values of the exponential factors~~ in  
273 nonlinear fitting procedures are rarely equal to 1 (Mikkonen et al., 2011), so we  
274 replaced the factors  $x_i$  with  $x_i^{w_i}$  in the proxy, where  $x_i$  can be an atmospheric  
275 variable ~~such as UVB and  $[SO_2]$~~ , and  $w_i$  defines  $x_i$ 's weight in the proxy. Since  $k$  is a  
276 temperature-dependent reaction constant and varies within a 10 % range (in the  
277 atmosphere temperature range of 267.6 - 292.6 K), *i.e.*, ~~the actual atmospheric~~  
278 ~~temperature variation in this study, we approximately regard  $k$  as a constant and use a~~  
279 ~~new scaling factor  $k_0$ . This methodology has been used previously in the proxies of~~  
280 ~~gaseous sulfuric acid we further replaced  $k$  with a scaling factor  $k_0$  that is also used in~~  
281 ~~the proxy methods built~~ in Hyytiälä, Southern Finland (Petäjä et al., 2009). As a result,  
282 the general proxy equation can be written as Eq. (7), with the UVB intensity,  $[SO_2]$ ,  
283 condensation sink (CS),  $[O_3]$ , and  $[HONO]$  (or  $[NO_x]$ ) as predictor variables:

284

$$285 [H_2SO_4] = f(k_0, x_i^{w_i}), \quad x_i = UVB, [SO_2], CS, [O_3], [HONO] \dots \quad (7)$$

286

287 The nonlinear curve-fitting procedures using iterative least square estimation for  
288 the proxies of gaseous sulfuric acid concentration based on Eq. (7) were performed by  
289 a custom-made MATLAB software. In addition to the correlation coefficient (R),  
290 relative error (RE) is used to evaluate the performance of proxies in the statistical  
291 analysis and can be written as Eq. (8).

$$293 RE = \frac{1}{n} \cdot \sum_{i=1}^n \frac{|[H_2SO_4]_{proxy,i} - [H_2SO_4]_{meas,i}|}{[H_2SO_4]_{meas,i}} \quad (8)$$

## 294 4 Results and discussion

### 295 4.1 General Characteristics of daytime sulfuric acid and atmospheric parameters

296 Table 1 summarizes the mean, median and 5-95 % percentiles of gaseous sulfuric  
297 acid concentrations and other variables measured during the daytime of the campaign.  
298 The 5-95 % percentile ranges of the UVB intensity,  $[SO_2]$ ,  $[NO_{x2}]$  and  $[O_3]$  were 0-0.45  
299 W  $m^{-2}$ , 0.9-11.4 ppbv, 3.3-61.4 ppbv and 3.5-23.3 ppbv, respectively. Compared with  
300 the sites in the study by Mikkonen et al. (2011), Beijing was characterized with a factor  
301 of 1.4-13.1 higher mean  $[SO_2]$  but a factor of 3.4-5.4 lower mean  $[O_3]$ . The 5-95 %  
302 percentile range of CS in Beijing was 0.01-0.24  $s^{-1}$ , which is about 10-100 times  
303 higher+2 orders of magnitude larger than corresponding value ranges in Europe and  
304 North America. The concentration of gaseous sulfuric acid during this campaign was  
305  $(2.2 - 10.0) \times 10^6$  molecule  $cm^{-3}$  ~~was in thea~~ 5-95 % percentile range ~~of~~, relatively  
306 similar to observed elsewhere around the world. A diurnal mean concentration of 0.74  
307 ppbv for HONO was observed in this campaign, consistent with previous long-term  
308 HONO measurements of about 0.48-1.8 ppbv (averaged values) in winter in Beijing  
309 (Hendrick et al., 2014; Spataro et al., 2013; Wang et al., 2017), which is a factor of 4-  
310 10 higher than HONO concentrations measured in Europe (Alicke et al., 2002, 2003).  
311 In addition, Beijing is dry in winter with a mean ambient relative humidity of 28 %  
312 during the campaign.an ambient relative humidity generally lower than 60%.

### 315 4.2 Correlations between $[H_2SO_4]$ and atmospheric variables

316 Table 2 summarizes the correlation coefficients between  $[H_2SO_4]$  and atmospheric  
317 variables using a Spearman-type correlation analysis. ~~Note that only correlations with~~  
318 ~~p-values smaller than 0.01 were included to ensure a statistical significance.~~ Clearly,  
319 the UVB intensity is an isolated variable that is independent of all the other variables

320 but that imposes a positive influence on  $O_3$  because of photochemical formation of  
321 ozone, and a negative influence on HONO because of HONO's photochemical  
322 degradation. The sulfuric acid concentration shows positive correlations with all the  
323 other variables. The correlation coefficients between  $[H_2SO_4]$  and  $[SO_2]$  and between  
324  $[H_2SO_4]$  and UVB intensity are 0.74 and 0.46, respectively, which indicate that  $[SO_2]$   
325 and UVB have important influences on the formation of atmospheric gaseous sulfuric  
326 acid.eonsistent with the accepted formation pathway of gaseous sulfuric acid from the  
327 reaction between  $SO_2$  and OH radicals. Accordingly,  $[O_3]$  and  $[HONO]$  show positive  
328 correlations with  $[H_2SO_4]$  because both  $O_3$  and HONO could be precursors of OH  
329 radicals. Surprisingly, a high positive correlation coefficient (0.6) was found between  
330  $[H_2SO_4]$  and CS, which is in contrast to the conventional thought that CS describes the  
331 loss of gaseous sulfuric acid molecules onto pre-existing particles and thus should show  
332 a negative correlation. CS correlates well with  $[SO_2]$  ( $r = 0.83$ ) and  $[NO_{x2}]$  ( $r = 0.77$ ):  
333 a high CS value, as an indicator of an atmospheric particle pollution, is thus usually  
334 accompanied with a high concentration of both  $SO_2$  and  $NO_{x2}$  in urban China,  
335 indicating co-emissions. A strong correlation between  $[HONO]$  and  $[NO_{x2}]$  ( $r = 0.88$ )  
336 in our measurement is supported by the fact that HONO can be either heterogeneously  
337 formed by reactions of  $NO_2$  on various surfaces (Calvert et al., 1994) or homogeneously  
338 formed by the gas phase NO + OH reaction, between which the former likely dominate  
339 for the daytime HONO production in urban Beijing (Liu et al., 2014).

340 Since the UVB intensity and  $[SO_2]$  have been reported as the dominating factors  
341 for the formation of sulfuric acid (Mikkonen et al., 2011; Petäjä et al., 2009), we further  
342 explored the relationship of the measured sulfuric acid concentrations with the UVB  
343 intensity and  $[SO_2]$  using the nonlinear curve-fitting method with a single variable.  
344 Figure 1a presents a scatter plot of  $[H_2SO_4]$  against the UVB intensity, color-coded by  
345  $[SO_2]$ . A good correlation layering withwith a clear lamination by  $[SO_2]$  is evident,  
346 indicating that the UVB intensity and  $[SO_2]$  together play an important role in the  
347 formation of sulfuric acid. A similar scatter plot (Figure 1b) of  $[H_2SO_4]$  against  $[SO_2]$ ,  
348 color-coded by the UVB intensity, leads to a similar conclusion.

349

### 350 4.3 Proxy construction

351 Similar to the non-linear proxies suggested by Mikkonen et al. (2011), we tested a  
352 number of proxies for gaseous sulfuric acid, listed in Table 3 with their respective fitting  
353 parameters and performance summarized in Table 4. The scatter plots of observed

354 [H<sub>2</sub>SO<sub>4</sub>] *versus* predicted values given by proxies are presented in Fig. S1. In these  
355 proxies, the concentration of a gaseous species is in the unit of molecule cm<sup>-3</sup>, the unit  
356 of the UVB intensity is W m<sup>-2</sup>, the unit of CS is s<sup>-1</sup>, and  $k_0$  is a scaling factor.

357 The proxy N1 was built by using the UVB intensity and [SO<sub>2</sub>] as the source terms  
358 and CS as the sink term, which follows the conventional idea of the H<sub>2</sub>SO<sub>4</sub> formation  
359 and loss in the atmosphere. CS was then removed from this proxy to examine the  
360 performance of the proxy N2 that has~~ve~~ the UVB intensity and [SO<sub>2</sub>] as the only  
361 predictor variables. Since the formation of OH radicals in the atmosphere depends on  
362 precursors in addition to UVB, we further attempted to introduce the OH precursor term  
363 into the H<sub>2</sub>SO<sub>4</sub> proxy. The proxies N3 and N4 were built by introducing O<sub>3</sub> as the only  
364 OH precursor to evaluate its influence on the formation of sulfuric acid. Furthermore,  
365 we added HONO as another potential precursor for OH radicals, resulting in the proxies  
366 N5 and N6. Lastly, the proxy N7 was built by replacing [HONO] with [NO<sub>x2</sub>] because  
367 firstly, HONO is not regularly measured, and secondly, a good linear correlation  
368 between [HONO] and [NO<sub>x2</sub>] was generally observed in the daytime during this  
369 campaign, although higher [HONO]/[NO<sub>x2</sub>] ratios were observed in the morning due to  
370 the accumulation of HONO during the night (Figure 2). RH was not considered in the  
371 current study because ~~a test by introducing RH into the proxies do not result in a~~  
372 ~~significantly better performance, which is consistent with those conclusions~~  
373 ~~the introduction of RH into the proxy did not yield significantly better results~~ in the  
374 Mikkonen et al. study (2011).

375 As shown in Table 4, the correlation coefficients are in the range of 0.83-0.86 and  
376 ~~RMA~~Es are in the range of ~~19.1-20.0 % (0.94 – 1.03) × 10<sup>6</sup> molecule cm<sup>-3</sup>~~. The  
377 exponents for the UVB intensity range from 0.13 to 0.16, and those for [SO<sub>2</sub>] generally  
378 range from 0.38 to 0.41, except in case of the proxy N6 ( $b=0.33$ ). The obtained  
379 exponent  $b$  for [SO<sub>2</sub>] is significantly smaller than 1 unlike ~~the assumptioned~~ in Eq.  
380 (63), mainly because [SO<sub>2</sub>] is also an indicator of air pollution that usually influences  
381 the sinks of both OH radicals and sulfuric acid. The exponent for [SO<sub>2</sub>] ranged from  
382 0.5 to 1.04 in the previous proxy study for European and North American sites  
383 (Mikkonen et al., 2011), including values from 0.48 to 0.69 in Atlanta, GA, USA, which  
384 was probably quite a polluted site because the measurements were conducted only 9  
385 km away from a coal-fired power plant. The obtained value range of the exponent  $b$   
386 for [SO<sub>2</sub>] in our study is probably related to the urban nature of Beijing. The value of  
387 exponent  $c$  for CS in the proxy N1 is as low as 0.03, which either might be due to the

388 covariance of CS and certain  $\text{H}_2\text{SO}_4$  sources that cancels the dependence on CS, or it  
389 might indicate that CS is actually insufficient in regulating the  $\text{H}_2\text{SO}_4$  concentration, as  
390 recently suggested by Kulmala et al. (2017). By comparing the proxies N1 and N2, we  
391 can see that CS plays a minor role because the exponents of  $[\text{SO}_2]$  and UVB, the overall  
392 correlation coefficient and the RMAEs are almost identical with and without CS. We  
393 can see the negligible role of CS also when comparing the results of the proxies N3 and  
394 N4 where  $\text{O}_3$  is considered. However, the role of CS becomes evident between the  
395 proxies N5 and N6 when HONO is introduced: the exponents of  $[\text{SO}_2]$ ,  $[\text{O}_3]$ , and  
396  $[\text{HONO}]$  significantly increased when taking into account the CS, suggesting that the  
397 covariance between HONO and CS can explain, at least partially, the close-to-zero  
398 exponent of CS in the proxies N1-N4. In addition, when  $[\text{O}_3]$  is introduced as the only  
399 precursor for OH radicals, minor improvements in the correlation coefficient and  
400 RMAE were obtained, as suggested by comparing the proxies N3 and N1. When both  
401  $[\text{O}_3]$  and  $[\text{HONO}]$  were introduced as OH precursors in the proxies N5-N7, REs have  
402 noticeable improvements, and correlation coefficients improved slightly. MAE and  
403 correlation coefficient significantly improved. Altogether, these observations suggest  
404 that it is crucial to introduce HONO into the proxy, both in our study and also likely for  
405 the previous work where the exponent of CS is close-to-zero (Mikkonen et al., 2011).

406 Although so far the proxy N5 had the best fitting quality, it is impractical to  
407 explicitly include  $[\text{HONO}]$  because HONO measurements are very challenging. As  
408 shown in Fig. 2,  $[\text{HONO}]$  and  $[\text{NO}_{\text{x2}}]$  ~~are~~ tended to correlate linearly with each other in  
409 the daytime during this campaign, with a linearly fitted  $[\text{HONO}]/[\text{NO}_{\text{x2}}]$  ratio of around  
410 0.03 and a relative error of 0.42 mean absolute error (MAE) of 0.3 ppbv. Similar, strong  
411 linearity was observed in a previous study by Hao et al. (2006) who attributed this  
412 observation to the heterogeneous conversion of  $\text{NO}_2$  to HONO. Only occasionally  
413 slightly higher  $[\text{HONO}]/[\text{NO}_{\text{x2}}]$  ratios in the morning could be seen in the morning,  
414 which might be due to the fact that HONO concentration could have an accumulation  
415 process during the nighttime and lead to a deviation from the steady state. deviation  
416 from the steady state. Bernard et al. (2016) reported that  $[\text{NO}_2]$  has a similar diurnal  
417 behavior to that of  $[\text{HONO}]$  and hence the ratio of  $[\text{HONO}]/[\text{NO}_2]$  varies slightly during  
418 the diurnal cycle. Therefore, due to the good correlation, the proxy N7 replaces  $[\text{HONO}]$   
419 by  $[\text{NO}_{\text{x2}}]$ , a more easily measured variable, and performs equally well with the proxy  
420 N5.

421 Clearly, the proxy N2 provides the simplest parameterization, but the proxies N5

422 and N7 result in the best fitting quality because of the introduction of [HONO]. [Figure](#)  
423 [3 presents the RE values for the proxies N2 and N7, respectively, as a function of linear](#)  
424 [bins of measured sulfuric acid concentrations. The performance of the proxy N7 is](#)  
425 [considerably better than that of the proxy N2 in the sulfuric acid concentration range of](#)  
426 [\(2.2 – 10\) × 10<sup>6</sup> molecule cm<sup>-3</sup>, which covers the 5–95% percentiles of sulfuric acid](#)  
427 [concentration in this study. In the worst scenario, RE of proxy N2 is 1.2 times as high](#)  
428 [as that of proxy N7, e.g., REs are 16.75 % and 13.99 %, respectively, in the sulfuric](#)  
429 [acid concentration bin of \(4 – 5\) × 10<sup>6</sup> molecule cm<sup>-3</sup>, and 16.71 % and 14.42 %,](#)  
430 [respectively, in the bin of \(7 – 8\) × 10<sup>6</sup> molecule cm<sup>-3</sup>.](#)

431

#### 432 **4.4 Comparison of measured and predicted [H<sub>2</sub>SO<sub>4</sub>]**

433 A comparison between measured and predicted [H<sub>2</sub>SO<sub>4</sub>] was performed. Figure 4  
434 includes calculated results from the proxies N2 and N7 as well as from a proxy  
435 constructed according to measurement in a boreal forest site, Finland, *i.e.*, Eq (1) (Petäjä  
436 et al., 2009). The measured daytime [H<sub>2</sub>SO<sub>4</sub>] on 10 March, 2018, was above 4 × 10<sup>6</sup>  
437 molecules cm<sup>-3</sup> [when averaged towith](#) a time resolution of 5 min. The predicted [H<sub>2</sub>SO<sub>4</sub>]  
438 using the proxies N2 and N7 both track the measured [H<sub>2</sub>SO<sub>4</sub>] pretty well, even when  
439 an unexpected dip in the sulfuric acid concentration was observed at around 10:00–  
440 11:00. The performance of the proxy N7 is better than that of proxy N2 during the entire  
441 day, consistent with our results in Fig. 3. The proxy by Petäjä et al. (2009)  
442 underestimated the concentrations of sulfuric acid by a factor of 20 or so, which can be  
443 attributed to the very different values of CS between Beijing and the boreal forest. The  
444 fact that [H<sub>2</sub>SO<sub>4</sub>]<sub>Petäjä et al.</sub> does not track the measured [H<sub>2</sub>SO<sub>4</sub>] even after including  
445 a scaling factor indicates that proxies are site-specific and do not necessarily work well  
446 in locations other than where they have originally been developed for. In addition, the  
447 direct performance comparison between the proxy N2 and the proxy by Petäjä et al.  
448 (2009) indicates the importance of assigning exponential weights to variables in the  
449 nonlinear fitting procedures, which is consistent with results by Mikkonen et al. (2011).

450

#### 451 **5 Summary and conclusions**

452 Sulfuric acid is a key precursor for atmospheric new particle formation. In this  
453 study, we constructed a number of proxies for gaseous sulfuric acid concentration  
454 according to our measurements in urban Beijing during the winter. According to the  
455 obtained proxies and their performance, the UVB intensity and [SO<sub>2</sub>] were the

456 dominant influencing factors. Hence, the simplest proxy (Proxy N2) only involves  
457 UVB intensity and  $[\text{SO}_2]$  as shown by Eq. (9). The units of  $[\text{H}_2\text{SO}_4]$  and  $[\text{SO}_2]$  are  
458 molecule  $\text{cm}^{-3}$ , and the unit of UVB is  $\text{W m}^{-2}$ .

460

$$459 \quad [\text{H}_2\text{SO}_4] = 280.05 \cdot \text{UVB}^{0.14} \cdot [\text{SO}_2]^{0.40} \quad (9)$$

461

462 This proxy resulted in a relative deviation of up to 29 %.

463 For a comprehensive consideration of the formation pathways of OH radicals, For  
464 the best proxy accuracy,  $[\text{O}_3]$  and  $[\text{HONO}]$  as well as CS should be included (Proxy  
465 N5), as shown by Eq. (10). The units of  $[\text{H}_2\text{SO}_4]$ ,  $[\text{SO}_2]$ ,  $[\text{O}_3]$  and  $[\text{HONO}]$  are molecule  
466  $\text{cm}^{-3}$ , the unit of UVB is  $\text{W m}^{-2}$ , and the unit of CS is  $\text{s}^{-1}$ .

467

$$468 \quad [\text{H}_2\text{SO}_4] = 0.0072 \cdot \text{UVB}^{0.15} \cdot [\text{SO}_2]^{0.41} \cdot \text{CS}^{-0.17} \cdot ([\text{O}_3]^{0.36} \\ 469 \quad + [\text{HONO}]^{0.38}) \quad (10)$$

470

471 Since HONO measurements are not a regular practice, we can further replace  $[\text{HONO}]$   
472 with  $[\text{NO}_{\text{x}2}]$ , shown in Eq. (11), which can be justified by the strong linear correlation  
473 between  $[\text{HONO}]$  and  $[\text{NO}_{\text{x}2}]$  observed in this study. The units of  $[\text{H}_2\text{SO}_4]$ ,  $[\text{SO}_2]$ ,  $[\text{O}_3]$   
474 and  $[\text{NO}_x]$  are molecule  $\text{cm}^{-3}$ , the unit of UVB is  $\text{W m}^{-2}$ , and the unit of CS is  $\text{s}^{-1}$ .

475

$$476 \quad [\text{H}_2\text{SO}_4] = 0.0013 \cdot \text{UVB}^{0.13} \cdot [\text{SO}_2]^{0.40} \cdot \text{CS}^{-0.17} \cdot ([\text{O}_3]^{0.44} \\ 477 \quad + [\text{NO}_{\text{x}2}]^{0.41}) \quad (11)$$

478

479 We consider this last proxy more reasonable than the others due to the following reasons:  
480 first, it makes the equation physically meaningful as the CS starts to be involved as a  
481 sink term, and second, the absolute and relative fitting error RE was reduced considerably compared with the other proxies. Overall, this suggests that the photolysis  
482 of  $\text{O}_3$  and HONO are both important OH sources in urban Beijing.

483

484 As a summary, we recommend using the simplest proxy (proxy N2 as shown in  
485 Eq. (9)) and a more accurate proxy (Proxy N7 as shown in Eq. (11) for calculating  
486 daytime gaseous sulfuric acid concentrations in the urban Beijing atmosphere. It is clear  
487 that the current proxies are based on only a month-long campaign of sulfuric acid  
488 measurements in urban Beijing during winter. Given the dramatic reduction in the  
489 concentration of  $\text{SO}_2$  in recent years (Wang et al., 2018) and the strong dependence of

490 calculated  $[H_2SO_4]$  on  $[SO_2]$ , the performance of the proxies in the past and future years  
491 remain to be evaluated. Furthermore, the proxies might be site-specific and season-  
492 specific. Since the proxies were derived with atmospheric parameters in winter, in  
493 urban Beijing, the exponents for atmospheric variables in the proxy could have different  
494 values for other cities or other seasons. Thus, the proxies in this study should be further  
495 tested before their application to other Chinese megacities or other  
496 seasons. Nevertheless, our work here shows the importance of heterogeneous chemistry  
497 as a potential source of OH radicals in an urban air; however, the proxies might be site-  
498 specific and should be further tested before their application to other Chinese  
499 megacities.

500

#### 501 **Author contributions**

502 LW designed this study. YL (Yiqun Lu), CY, YF, YC, YL (Yiliang Liu), GY, YW, YZ, RY, RB  
503 and CD conducted the field campaign. YL (Yiqun Lu) analyzed data with contributions from  
504 LW and all the other co-authors. YL (Yiqun Lu) and LW wrote the manuscript with  
505 contributions from all the other co-authors.

506

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817

Table 1 Mean, median, 5-95 % percentiles of key atmospheric variables and  $[H_2SO_4]$  in the daytime.

	UVB (W m <sup>-2</sup> )	$[SO_2]$ (ppbv)	CS (s <sup>-1</sup> )	$[O_3]$ (ppbv)	$[HONO]$ (ppbv)	$[NO_{x2}]$ (ppbv)	$[H_2SO_4]$ ( $\times 10^6$ molecule cm <sup>-3</sup> )	RH (%)
mean	0.17	4.6	0.11	10.5	0.74	25.3	5.4	28
median	0.14	3.7	0.11	9.0	0.51	23.0	4.9	26
5-95% percentiles	0.00-0.45	0.9-11.4	0.01-0.24	3.5-23.3	0.09-2.65	3.3-61.4	2.2-10.0	9-59

Table 2 Correlation coefficients (Spearman type) between  $[H_2SO_4]$  and atmospheric variables in the daytime. ~~Only correlation coefficients with p-values less than 0.01 are included to ensure a statistical significance.~~

	UVB	$[SO_2]$	CS	$[O_3]$	$[HONO]$	$[NO_2]$	$[H_2SO_4]$
UVB	1	<u>0.01/</u>	<u>-0.02/</u>	0.14	-0.23	<u>-0.04/</u>	0.46
$[SO_2]$		1	0.83	0.25	0.64	0.70	0.74
CS			1	0.36	0.75	0.77	0.60
$[O_3]$				1	<u>-0.02/</u>	<u>-0.04/</u>	0.29
$[HONO]$					1	0.88	0.39
$[NO_2]$						1	0.53
$[H_2SO_4]$							1

Table 3 Proxy functions for the nonlinear fitting procedure.

Proxy	Function	Equation <sup>#</sup>
N1	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot CS^c$	
N2	$k_0 \cdot UVB^a \cdot [SO_2]^b$	
N3	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot CS^c \cdot [O_3]^d$	
N4	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot [O_3]^d$	
N5	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot CS^c \cdot ([O_3]^d + [HONO]^e)$	
N6	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot ([O_3]^d + [HONO]^e)$	
N7	$k_0 \cdot UVB^a \cdot [SO_2]^b \cdot CS^c \cdot ([O_3]^d + [NO_{x2}]^f)$	

<sup>#</sup>UVB is the intensity of ultraviolet radiation b in  $\text{W cm}^{-3}$ ;  $[\text{SO}_2]$  is the concentration of sulfur dioxide in molecule  $\text{cm}^{-3}$ ; CS is the condensation sink in  $\text{s}^{-1}$ ;  $[\text{O}_3]$  is the concentration of ozone in molecule  $\text{cm}^{-3}$ ;  $[\text{HONO}]$  is the concentration of nitrous acid in molecule  $\text{cm}^{-3}$ ;  $[\text{NO}_{x2}]$  is the concentration of nitrogen dioxide in molecule  $\text{cm}^{-3}$ ;  $k_0$  is a scaling factor.

Table 4 Results of the nonlinear fitting procedure for different proxy functions, together with correlation coefficient (R, Pearson type) and relative error (RE),mean absolute error (MAE).

Proxy	$k_0$	$a$	$b$	$c$	$d$	$e$	$f$	$R$	$RMAE$ ( $\times 10^{-4}$ )
N1	515.74	0.14	0.38	0.03				0.83	<u>20.041.03</u>
N2	280.05	0.14	0.40					0.83	<u>1.0320.00</u>
N3	9.95	0.13	0.39	-0.01	0.14			0.85	<u>1.0019.95</u>
N4	14.38	0.13	0.38		0.14			0.85	<u>1.0019.95</u>
N5	0.0072	0.15	0.41	-0.17	0.36	0.38		0.86	<u>0.9419.11</u>
N6	2.38	0.14	0.33		0.24	0.24		0.85	<u>0.9819.66</u>
N7	0.0013	0.13	0.40	-0.17	0.44		0.41	0.86	<u>0.9519.34</u>

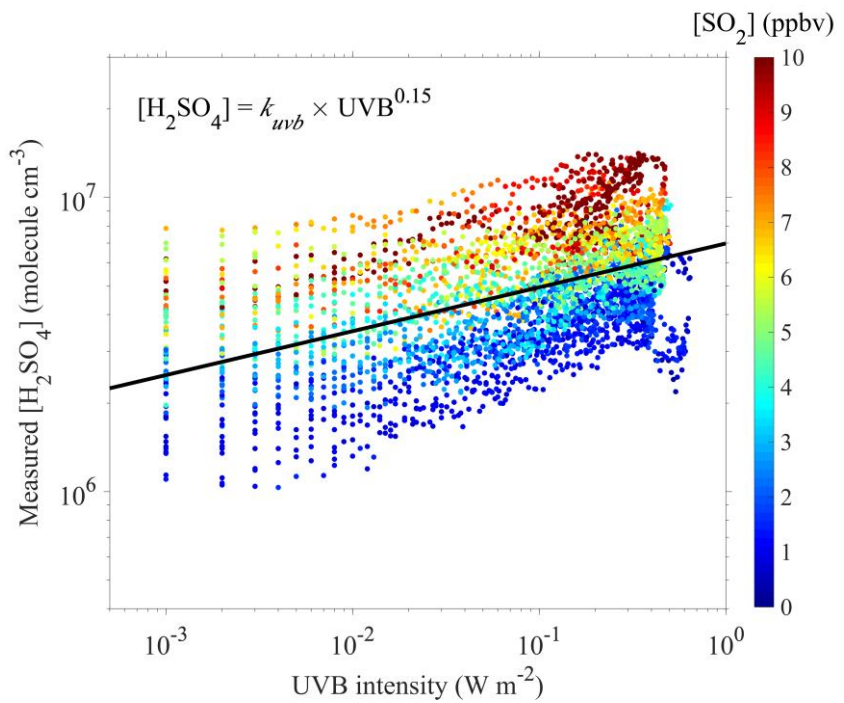
## Figure Captions

**Figure 1.** Correlations (a) between  $[\text{H}_2\text{SO}_4]$  and UVB intensity, and (b) between  $[\text{H}_2\text{SO}_4]$  and  $[\text{SO}_2]$  during the campaign from 9 February to 14 March, 2018.  $k_m$  is a constant term.

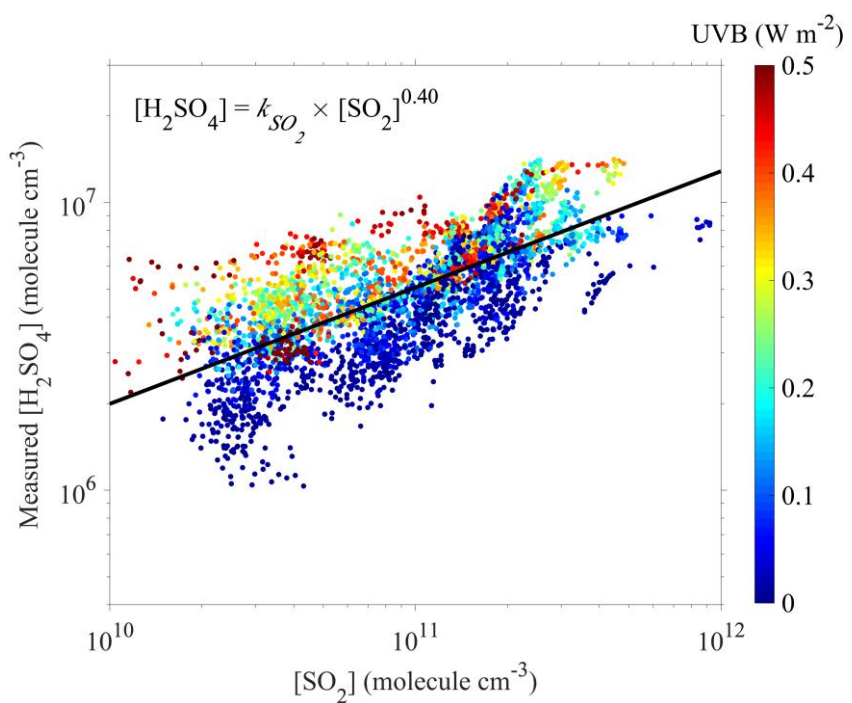
**Figure 2.** Correlation between  $[\text{HONO}]$  and  $[\text{NO}_{\text{x2}}]$  during the campaign from 9 February to 14 March, 2018. The black line represents a linear fitting with a zero intercept.

**Figure 3.** Performance assessments of proxy N2 and proxy N7. The REs are used to evaluate the performances of proxy N2 and N7, respectively as a function of linear bins of measured sulfuric acid concentrations. The averaged deviation and the relative deviation in the plots are defined by Eq. (6) and Eq. (7) and used to evaluate the performance of proxy N2 and N7, respectively. “Overlap” refers to the smaller values between proxy N2 and proxy N7, and the larger ones are indicated by the color code of proxies N2 and N7.

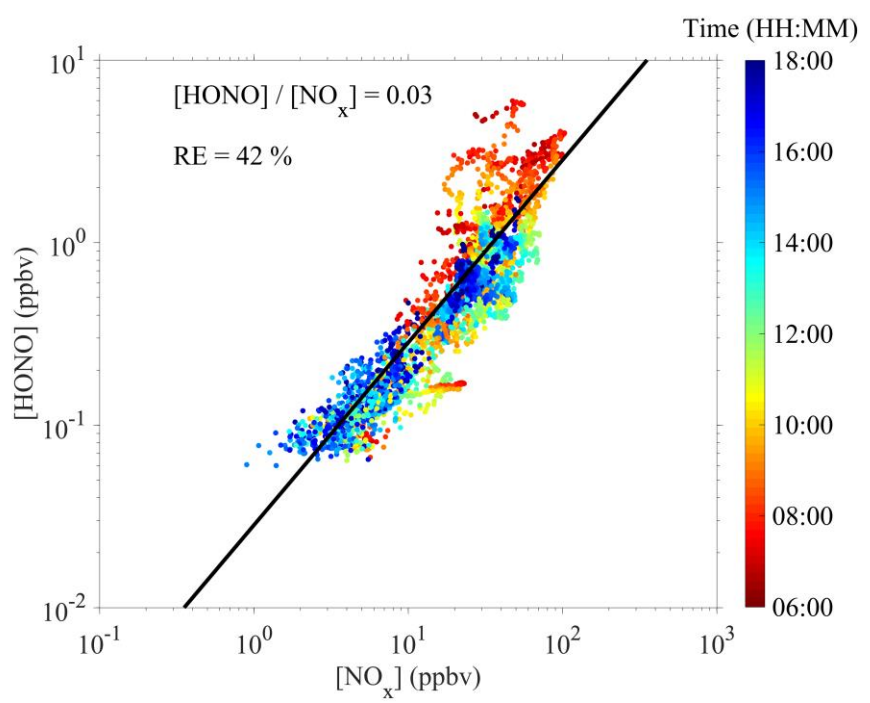
**Figure 4.** Comparison of measured  $[\text{H}_2\text{SO}_4]$ ,  $[\text{H}_2\text{SO}_4]_{\text{N2}}$ ,  $[\text{H}_2\text{SO}_4]_{\text{N7}}$  and  $[\text{H}_2\text{SO}_4]_{\text{Petäjä et al.}}$  on 10 March, 2018 with a time resolution of 5 min.



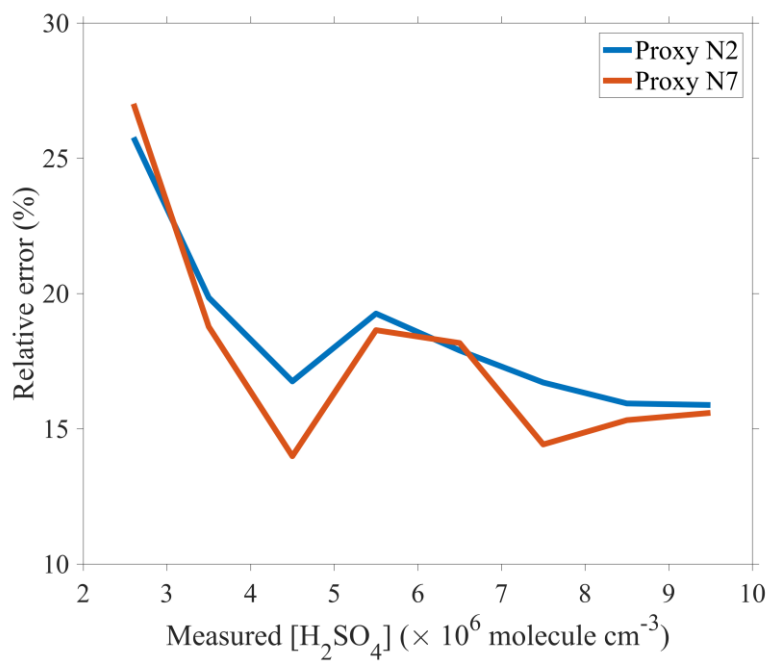
**Figure 1a**



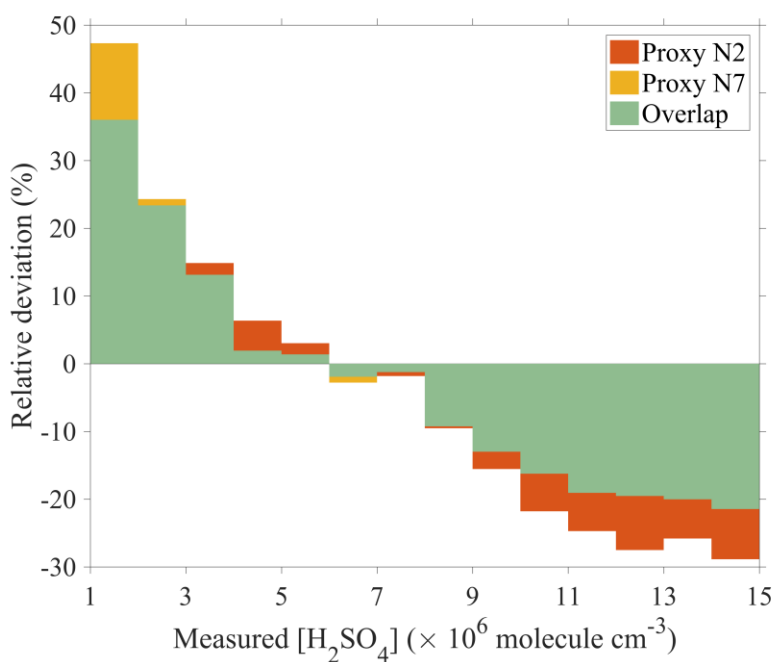
**Figure 1b**



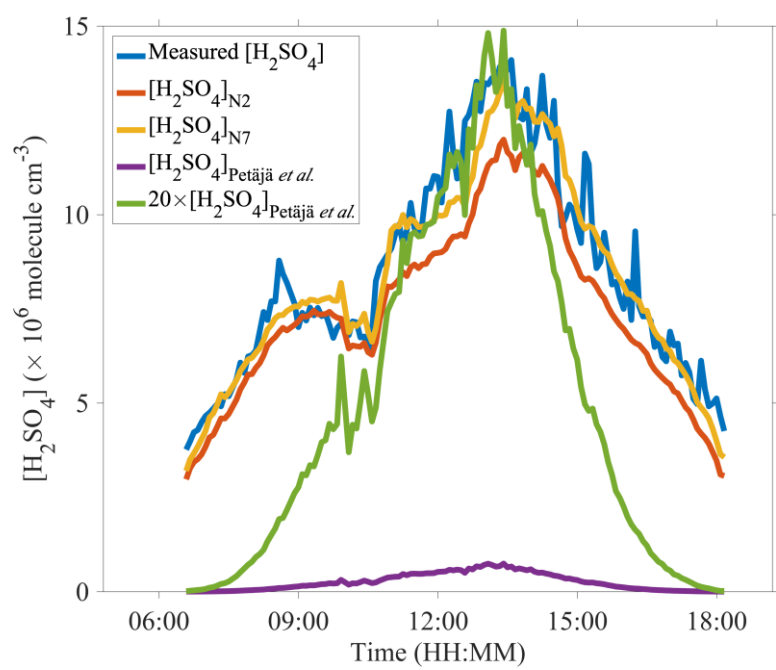
**Figure 2**



**Figure 3a**



**Figure 3b**



**Figure 4**