

Reply to Interactive comment on “Chemical Transformation of α -Pinene derived Organosulfate via Heterogeneous OH Oxidation: Implications for Sources and Environmental Fates of Atmospheric Organosulfates” by Rongshuang Xu et al.

Anonymous Referee #2

Interactive comment on Atmos. Chem. Phys. Discuss., <https://doi.org/10.5194/acp-2021-953-RC2>, 2022

10 *Xu et al. presents laboratory studies where they investigated the heterogenous oxidation of an alpha-pinene organosulfate surrogate, sodium 2-hydroxy-2,6,6-trimethylbicyclo[3.1.1]heptan-3-yl sulfate, or apOS-249, using an oxidation flow reactor. The reaction and products were measured with an UHPLC-ESI-QToFMS, HPLC-ESI-QTRAP-MS, and IC. The authors observed various products they provided chemical formula and compared against ambient and lab studies where these compounds had been observed before. Further, they determined the heterogenous OH oxidation of apOS-249 and found the lifetime to be comparable to the lifetime of aerosol (~10 days). They explored potential reaction mechanisms. Finally, they investigated the potential amount of inorganic sulfate (SO₄) produced from the heterogenous reaction of apOS-249 with OH. The paper is generally well written with interesting results that would be of importance for the community that reads ACP. I recommend publication after the authors address some of the comments below.*

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We thank the reviewer for his/her thoughtful comments. The referee’s comments are below in italics followed by our responses in normal font.

Comment #1:

1. In general, more information concerning the oxidation flow reactor (OFR) is needed to better understand and replicate the experiments. The following details needs to be included:

1a) How was the aerosol and gases introduced into the system? How were they sampled from the OFR?

1b) What material is the OFR made of?

30 *1c) Was this made in laboratory or purchased from a company?*

1d) What was the temperature for the experiments?

1e) How was OH reactivity determined during the experiments?

1f) What were the losses within the OFR? Was this included in the calculations?

Author Response:

40 Thanks for the suggestions. We have revised the experimental method in the manuscript and provided more details in the Supporting Information (**Sect. S1 Experimental Details** in SI). A simplified schematic diagram of our experimental setup was also added in the Supporting Information as **Scheme S1**. Please see our responses to specific questions below, followed by corresponding changes made in manuscript and SI (highlighted in blue).

1a) How was the aerosol and gases introduced into the system? How were they sampled from the OFR?

Response: We have prepared a simple schematic diagram of our experimental setup. apOS-249 aerosols were first generated by passing its solution through a constant output atomizer (TSI Model 3076) using 3 L min⁻¹ of nitrogen (N₂). Before entering the reactor, the aerosol stream was mixed with nitrogen (N₂), oxygen (O₂), and ozone (O₃) to make up a total flow of ~5 L min⁻¹, corresponding to a residence time of ~156 s. Aerosols leaving the reactor were collected into Teflon filters (2.0 μ m pore size, Pall Corporation) at a sampling flow rate of 3 L min⁻¹ conducted by an air sampling pump (Gilian 500, Sensidyne) for 30 min.

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1b) What material is the OFR made of?

Response: The OFR was made of aluminum with a volume of approximately 13 L (18-inch length and 8-inch inner diameter) (Kang et al., 2007).

1c) Was this made in laboratory or purchased from a company?

Response: The OFR (also called Potential Aerosol Mass, PAM) was designed and obtained from Prof. William Brune's group at Penn State.

1d) What was the temperature for the experiments?

Response: The experiment was conducted at 298.0 ± 0.5 K.

1e) How was OH reactivity determined during the experiments?

10 **Response:** As mentioned in the manuscript, OH was generated via photolysis of O₃ in the presence of water vapor. O₃ was generated by passing O₂ through an O₃ generator (ENALY 1000BT-12). The OH reactivity was determined by measuring the decay of sulfur dioxide (SO₂) in independent calibrating experiments (Teledyne SO₂ analyzer, Model T100) based on the reaction rate constant between gas-phase OH radicals and SO₂ ($= 9.0 \times 10^{-13}$ molecule⁻¹ cm³ s⁻¹) at 298 K (Kang et al., 2007) and was represented by the OH exposure, which is equivalent to the product of gas-phase OH concentration and the residence time. The OH exposure ranged from 0 to 17.4×10^{11} molecules cm⁻³ s in this study.

1f) What were the losses within the OFR? Was this included in the calculations?

20 **Response:** The OFR was designed with a small surface-to-volume ratio to minimize the aerosol wall loss (Kang et al., 2007; Lambe et al., 2011). The aerosol transmission efficiency for aerosol diameter larger than 150 nm was reported to be greater than 80% (Lambe et al., 2011). In our study, the wall loss was expected to be small as the aerosol diameter was measured to be 181.3 nm and the aerosols with a diameter larger than 150 nm accounted for a significant fraction of total aerosol number and mass. This wall loss factor was not corrected in the calculations. However, we expect this would not affect the determination of reaction kinetics (i.e. *k*) and inorganic sulfate yield. This is because concentration ratios (e.g. *I/I*₀) were used and the effect of wall loss would be cancelled out in the calculations.

References:

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- Kang, E., Root, M. J., Toohey, D. W., and Brune, W. H.: Introducing the concept of Potential Aerosol Mass (PAM), *Atmos. Chem. Phys.*, 7, 5727–5744, <https://doi.org/10.5194/acp-7-5727-2007>, 2007.
 - Lambe, A. T.; Ahern, A. T.; Williams, L. R.; Slowik, J. G.; Wong, J. P. S.; Abbatt, J. P. D.; Brune, W. H.; Ng, N. L.; Wright, J. P.; Croasdale, D. R.; Worsnop, D. R.; Davidovits, P.; Onasch, T. B.: Characterization of Aerosol Photooxidation Flow Reactors: Heterogeneous Oxidation, Secondary Organic Aerosol Formation and Cloud Condensation Nuclei Activity Measurements, *Atmos. Meas. Tech.*, 4, 445–461, <https://doi.org/10.5194/amt-4-445-2011>, 2011.

Revision in manuscript:

40 Page 5, Line 7: “Heterogeneous OH oxidation of α OS-249 aerosols was carried out using a 13-L aluminium OFR at 50 ± 2.0 % RH and 298.0 ± 0.5 K. Experimental details together with schematic diagram (**Scheme S1**) are given in the Supporting Information. Briefly, α OS-249 was first dissolved in deionized water (0.1 wt %) followed by a 30-min sonication. Aqueous aerosols were generated by passing the solution through an atomizer (TSI Model 3076) using 3 L min^{-1} of nitrogen (N₂). The aerosol stream was then directly mixed with ozone (O₃), wet/humidified nitrogen (N₂) and oxygen (O₂) to control the RH. A total flow of $\sim 5 \text{ L min}^{-1}$ was fed into the reactor, corresponding to a residence time of ~ 156 s (Xu et al., 2020a).”

50 Page 5, Line 19: “The OH exposure, a product of gas-phase OH radical concentration and the residence time, was in the range of $0\text{--}17.4 \times 10^{11}$ molecule cm⁻³ s. It was determined by measuring the decay of sulfur dioxide (SO₂) in independent calibrating experiments (Teledyne SO₂ analyzer, Model T100) based on the reaction rate between gas-phase OH radicals and SO₂ ($= 9.0 \times 10^{-13}$ molecule⁻¹ cm³ s⁻¹) at 298 K (Kang et al., 2007).”

Revision in SI:

Page 2, Line 1: “

1. Experimental Details

The heterogeneous OH oxidation of α pOS-249 aerosols was conducted using an OFR with a volume of ~ 13 L (18-inch length, 8-inch inner diameter) at 50 ± 2.0 % RH and 298.0 ± 0.5 K. As shown in Scheme S1, aqueous α pOS-249 aerosols were first generated by passing its solution through a constant output atomizer (TSI Model 3076) using 3 L min^{-1} of nitrogen (N_2). Before entering the reactor, the aerosols were directly mixed with dry/wet nitrogen (N_2), oxygen (O_2) and ozone (O_3) to make up a total flow of $\sim 5 \text{ L min}^{-1}$, corresponding to a residence time of ~ 156 s. The relative humidity (RH) inside the reactor was maintained by varying the mixing ratio of dry and humidified N_2 . The RH and temperature were measured by a RH-temperature sensor (Vaisala, HM40).

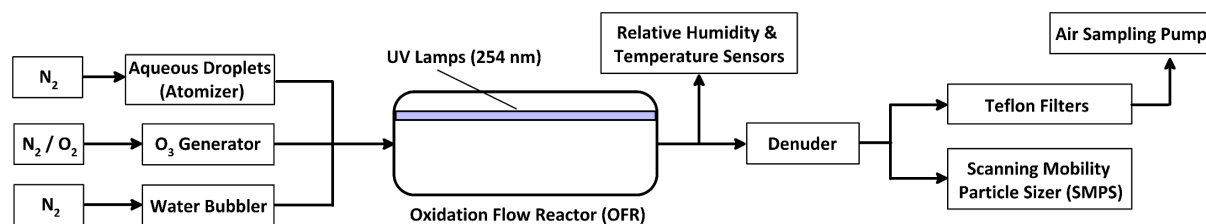
10 Inside the reactor, OH was generated via photolysis of O_3 with UV light at 254 nm in the presence of water vapor. The O_3 was generated by passing O_2 through an O_3 generator (ENALY 1000BT-12). The concentration of gas-phase OH radical was varied by changing the O_3 concentrations, monitored by an O_3 analyzer (2B technologies, Model 202). The OH exposure, a product of gas-phase OH radical concentration and the residence time, ranged from 0 – 17.4×10^{11} molecule cm^{-3} s and was determined by measuring the decay of sulfur dioxide (SO_2) (Teledyne SO_2 analyzer, Model T100) in independent calibrating experiments in the absence of α pOS-249 aerosols based on the reaction rate between gas-phase OH radicals and SO_2 ($= 9.0 \times 10^{-13}$ molecule $^{-1}$ $\text{cm}^3 \text{ s}^{-1}$) at 298 K (Kang et al., 2007). Furthermore, SO_2 calibration experiments in the presence of α pOS-249 aerosol were also conducted to investigate the effects of the aerosols on the generation and concentration of gas-phase OH radicals inside the reactor. A variation of ~ 10 % in the determination of OH exposure was observed over the experimental conditions.

20 The aerosol stream leaving the reactor then passed through an annular Carulite catalyst denuder (manganese dioxide/copper oxide catalyst; Carus Corp.) and an activated charcoal denuder to remove residual O_3 and other gas-phase species. 3 L min^{-1} of the stream was sampled onto the Teflon filters ($2.0 \mu\text{m}$ pore size, Pall Corporation) by an air sampling pump (Gilian 500, Sensidyne) for 30 min, with a total gas sampling volume of ~ 90 L. Duplicate filters were collected from each of oxidation experiments for subsequent chemical analysis. After collection, filters were immediately stored at -20°C in the dark and analysed within 3 months.

30 Part of the remaining stream was introduced into a scanning mobility particle sizer (SMPS, TSI, CPC Model 3775, Classifier Model 3081) to measure the size distribution of aerosols. The size distribution was sampled from 16 to 604.3 nm and scans were repeated every 180 s (sampling flow of $\sim 0.3 \text{ L min}^{-1}$ and sheath flow of 3 L min^{-1}). The aerosol mass loading was determined from measured volume concentration assumed for spherical aerosols with a unit density since the density of α pOS-249 is not available. As the sodium salts of the organosulfates ($\text{R-OSO}_3\text{Na}$) usually have density larger than 1.0 g cm^{-3} (e.g. 1.60 g cm^{-3} of $\text{CH}_3\text{SO}_4\text{Na}$; 1.46 g cm^{-3} of $\text{C}_2\text{H}_5\text{SO}_4\text{Na}$; Chemistry Dashboard), the reported aerosol mass loadings are considered as low limits. Before oxidation, the mean surface weighted diameter for aerosol distribution was about 181.3 ± 0.5 nm with a geometric standard deviation of 1.3 and the aerosol mass loading was measured to be $\sim 2000 \mu\text{g m}^{-3}$.

40 The OFR was designed with a small surface-to-volume ratio to minimize the aerosol wall loss (Kang et al., 2007; Lambe et al., 2011). The aerosol transmission efficiency for aerosol diameter larger than 150 nm was reported to be greater than 80% (Lambe et al., 2011). In our study, the wall loss was expected to be small as the aerosol diameter was measured to be 181.3 ± 0.5 nm and the aerosols with a diameter larger than 150 nm accounted for a significant fraction of total aerosol number and mass. This wall loss factor was thus not corrected in the calculations. However, we expect this would not significantly affect the determination of reaction kinetics (i.e. k) and inorganic sulfate yield. This is because concentration ratios (e.g. I/I_0) were used and the effect of wall loss would be cancelled out in the calculations.

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Scheme S1. Schematic diagram for experimental setup of the heterogeneous OH oxidation.’

Comment #2:

2. Other information within the methods that would improve the paper:

- 2a) Was a drier used after atomization?
- 2b) How were O₃ and gas-phase species removed prior to sampling? What was the particle loss through this method to remove gas-phase species?
- 2c) How was the SMPS operated? What SMPS system was used?
- 10 2d) What assumption was used to assume an aerosol mass loading of 2000 $\mu\text{g m}^{-3}$? Was this mass concentration too high for heterogenous oxidation (e.g., OH-limited at the beginning due to too high aerosol mass loading compared to OH concentration)?
- 2e) Description of the aerosol collection/filters are needed -- size, type, and pore size of filter, were they cleaned before use, how backgrounds were collected, stability of products on filters, and any impactors that may have been used.
- 2f) A simple diagram or an actual photo of the experimental set-up would be beneficial.

Author Response:

20 We agree with the reviewer’s suggestions. We have prepared point-to-point responses below. Changes have also been made in the manuscript and **Experiment Details** section of the SI provided above.

2a) Was a drier used after atomization?

Response: In our experiments, aqueous droplets generated by the atomizer did not pass through a diffusion dryer and were directly mixed with gases such as humidified nitrogen, oxygen, ozone before entering the flow tube reactor.

2b) How were O₃ and gas-phase species removed prior to sampling? What was the particle loss through this method to remove gas-phase species?

30 **Response:** An annular Carulite catalyst denuder and an activated charcoal denuder was used to remove O₃ and gas-phase species from the aerosol stream, respectively. We have measured the aerosol loss before and after the denuders under 50 % RH. Only 8.50 % and 2.04 % of differences were observed in aerosol number and mass concentration, respectively. We also would like to note that the size measurements of the aerosols were carried out after the denuders (**Scheme S1**).

2c) How was the SMPS operated? What SMPS system was used?

40 **Response:** A scanning mobility particle sizer (SMPS, TSI, CPC Model 3775, Classifier Model 3081) was used to measure the aerosol size and number distribution. The size distribution was sampled from 16 to 604.3 nm and scans were repeated every 180 s (sampling flow of $\sim 0.3 \text{ L min}^{-1}$ and sheath flow of 3 L min^{-1}).

2d) What assumption was used to assume an aerosol mass loading of 2000 $\mu\text{g m}^{-3}$? Was this mass concentration too high for heterogenous oxidation (e.g., OH-limited at the beginning due to too high aerosol mass loading compared to OH concentration)?

Response: Thanks for the comment. The aerosol mass loading was determined from measured volume concentration assumed for spherical aerosols with a unit density. We don’t correct for the aerosol density since the density of $\alpha\text{POS-249}$ is not known. From the literatures, the sodium salts of the organosulfates (R–OSO₃Na) usually have the density larger than 1.0 g cm^{-3} (e.g. 1.60 g cm^{-3} of

CH₃SO₄Na; 1.46 g cm⁻³ of C₂H₅SO₄Na; Chemistry Dashboard). We would like to consider the reported aerosol mass loadings as low limits.

10 In our experiment, the gas-phase OH radicals were generated by the photolysis of O₃ under UV light (λ = 254 nm) illumination in the presence of water vapor. O₃ was generated by passing the O₂ through an ozone generator. The OH concentration was regulated by changing the O₃ concentration and was determined by measuring the decay of sulfur dioxide (SO₂) in independent experiments in the presence or absence of aerosols. In the absence of aerosols, the measured OH concentration ranged from 3.06×10^9 molecules cm⁻³ to 1.11×10^{10} molecules cm⁻³. We also found that the presence of aerosols did not significantly affect the generation of gas-phase OH radicals and the determination of OH exposure, with differences less than ~10 % at each oxidation level. This may suggest that the oxidation would not be limited by the OH concentration.

2e) Description of the aerosol collection/filters are needed -- size, type, and pore size of filter, were they cleaned before use, how backgrounds were collected, stability of products on filters, and any impactors that may have been used.

20 **Response:** Thanks for the comment. In our study, aerosols leaving the reactor were collected into Teflon filters (47mm, 2.0 μ m pore size, Pall Corporation) through filtration at a sampling flow rate of 3 L min⁻¹ using an air sampling pump (Gilian 500, Sensidyne) for 30 min. These filters were used as purchased while the filter holder was cleaned prior to collection.

For the background, before the experiments, the OFR was flushed using high purity gases (N₂ and O₂) and then exposed to a high concentration of OH for several hours until background aerosol mass loading was less than a few μ g m⁻³. Given the low concentration of aerosols in the background, the background signals were considered to be insignificant.

30 For the stability of products, after collection, these filters were immediately stored at -20°C in the dark and analyzed within 3 months. The extraction efficiency of α pOS-249 was determined to be ~85 %. This high recovery suggests that α pOS-249 is relatively stable in our sample preparation and extraction procedure. We did not know the recovery of the reaction products as their standards were not available. a recent study by Hughes et al. (2019) evaluated the stability of a range of OSs (e.g. methyl sulfate, hydroxyacetone sulfate, two α -pinene derived OSs: $m/z = 279$ (C₁₀H₁₅O₇S⁻), and $m/z = 281$ (C₁₀H₁₇O₇S⁻)) on filters frozen at -20°C over the course of one year and extracted via similar procedure and analyzing using HPLC-ESI-HRMS. They found that stored OSs samples showed no degradation during the one year of storage and thus suggested that OSs with different alkyl, carboxylate and hydroxyl functional groups likely remain stable after collection onto filters when stored frozen. In addition, Wang et al. (2017) reported that there was no degradation over two-years' storage for α pOS-249 as well as limonene OS ($m/z = 249$ (C₁₀H₁₇O₅S⁻)) and limonaketone OS ($m/z = 249$ (C₉H₁₅O₆S⁻)), which have similar carbon skeletons while possessing extra functional (ketone or double bond) groups. 40 Altogether, we would suggest that reaction products are likely stable after collection onto filters during the storage at -20°C. However, further study is warrant for the investigation of stability of the reaction products.

The following changes were made in the manuscript, and details can be referred to the SI as replied to Comment #1.

50 Page 5, Line 19: "The OH exposure, a product of gas-phase OH radical concentration and the residence time, was in the range of $0-17.4 \times 10^{11}$ molecule cm⁻³ s. It was determined by measuring the decay of sulfur dioxide (SO₂) in independent calibrating experiments (Teledyne SO₂ analyzer, Model T100) based on the reaction rate between gas-phase OH radicals and SO₂ ($= 9.0 \times 10^{-13}$ molecule⁻¹ cm³ s⁻¹) at 298 K (Kang et al., 2007). It acknowledges that the presence of aerosols did not significantly affect the generation of gas-phase OH radicals and the determination of OH exposure (less than ~10 %). The aerosol stream leaving the reactor passed through an annular Carulite catalyst denuder (manganese dioxide/copper oxide catalyst; Carus Corp.) and an activated charcoal denuder to remove residual O₃ and other gas-phase species. Aerosols were collected onto the Teflon filters (47mm, 2.0 μ m pore size, Pall Corporation) through filtration at a sampling flow rate of 3 L min⁻¹ using an air sampling pump

(Gilian 500, Sensidyne) for 30 min, with a total gas sampling volume of ~ 90 L. Duplicate filters were collected from each of oxidation experiments for subsequent chemical analysis. After collection, filters were immediately stored at -20 °C in the dark and analysed within 3 months. Part of the remaining stream was introduced into a scanning mobility particle sizer (SMPS, TSI, CPC Model 3775, Classifier Model 3081) to measure the size distribution of the aerosols. The aerosol mass was determined from measured volume concentration assumed for spherical aerosols with a unit density.”

10 Page 8, Line 10: “The high recovery of α pOS-249 suggests the sample preparation and extraction methods are effective. Wang et al. (2017) have also reported that there was no degradation for α pOS-249 after two-years’ storage at low temperature (-20°C). A recent study by Hughes et al. (2019) examined the stability of a range of OSs (e.g. methyl sulfate, hydroxyacetone sulfate, two α -pinene derived OSs: $m/z = 279$ ($C_{10}H_{15}O_7S^-$), and $m/z = 281$ ($C_{10}H_{17}O_7S^-$)) on filters frozen at -20°C over one year. The filters were extracted via similar procedure applied in this study and the extracts were analyzed by HPLC-ESI-HRMS. They found that the investigated OSs with different functional groups (e.g. alkyl, carboxylate, and hydroxyl groups) showed no degradation during the storage. Taken together, α pOS-249 and its oxidation products (i.e. OSs) which have similar carbon skeletons while possessing different functional groups (alcohol and/or ketone) are likely stable during the storage and pre-treatment processes for chemical analysis.”

20 2f) *A simple diagram or an actual photo of the experimental set-up would be beneficial.*

Response: Thanks for the suggestion. We have prepared a simple schematic diagram of our experimental setup and have added this figure to the supporting information (**Scheme S1**). Please see our response above to Comment #1.

Comment #3:

3) *As reviewer #1 mentioned, sample preparation and/or sampling of the products with ESI may lead to side reactions. Though many of the products may be hard to synthesize, have surrogates been used to investigate their stability on filter, during preparation, and during sampling? One thing that would help with this question is stating that α pOS-249 shows high extraction efficiency (pg 8, ln4-5) sooner and maybe discuss other organosulfates.*

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Author Response:

Thanks for this insightful comment. We agree with reviewer’s comment that sample preparation and/or sampling of the α pOS-249 and reaction products with ESI may bring potential issues for identification and quantification of OSs. With regards to the side reactions (i.e. reactions between organics and solvent) during preparation and electrospray processes, as replied to the concern of reviewer #1, these reactions are considered to be negligible based on the insignificant peak intensities of potential reaction products in our aerosol mass spectra. Detailed discussions have been given in our responses to the reviewer #1 (Comment #1).

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To our best knowledge, standards are not yet available for investigating the stability on filter, during preparation, and during sampling of the reaction products. However, we have discussed the stability of the products using their surrogates to our best knowledge (Please see our responses to the reviewer’s Comment #2e). For products’ stability on filters after collection, a recent study by Hughes et al. (2019) evaluated the stability of a range of OSs (e.g. methyl sulfate, hydroxyacetone sulfate, two α -pinene derived OSs: $m/z = 279$ ($C_{10}H_{15}O_7S^-$), and $m/z = 281$ ($C_{10}H_{17}O_7S^-$)) on filters frozen at -20°C over the course of one year and extracted via similar procedure and analyzing using HPLC-ESI-HRMS. They found that stored OSs samples showed no degradation during the one year of storage and thus suggested that OSs with different alkyl, carboxylate and hydroxyl functional groups remain stable after collection onto filters when stored frozen. According to the synthesis reference for α pOS-249 by Wang et al. (2017), there is no degradation during over two-years’ storage for standard α pOS-249 as well as limonene OS ($m/z = 249$ ($C_{10}H_{17}O_5S^-$)) and limonaketone OS ($m/z = 249$ ($C_9H_{15}O_6S^-$)), which have similar carbon skeletons while possessing extra functional (ketone or double bond) groups. Altogether, reaction products are expected to be stable after collection onto filters during the storage at -20°C.

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We agree with the reviewer's suggestion and discussed this issue in earlier section of the manuscript. The following information is added in the manuscript.

Page 8, Line 10: "The high recovery of α pOS-249 suggests the sample preparation and extraction methods are effective. Wang et al. (2017) have also reported that there was no degradation for α pOS-249 after two-years' storage at low temperature (-20°C). A recent study by Hughes et al. (2019) examined the stability of a range of OSs (e.g. methyl sulfate, hydroxyacetone sulfate, two α -pinene derived OSs: $m/z = 279$ ($\text{C}_{10}\text{H}_{15}\text{O}_7\text{S}^-$), and $m/z = 281$ ($\text{C}_{10}\text{H}_{17}\text{O}_7\text{S}^-$)) on filters frozen at -20°C over one year. The filters were extracted via similar procedure applied in this study and the extracts were analyzed by HPLC-ESI-HRMS. They found that the investigated OSs with different functional groups (e.g. alkyl, carboxylate, and hydroxyl groups) showed no degradation during the storage. Taken together, α pOS-249 and its oxidation products (i.e. OSs) which have similar carbon skeletons while possessing different functional groups (alcohol and/or ketone) are likely stable during the storage and pre-treatment processes for chemical analysis.

We note that organic compounds such as carbonyls and carboxylic acids could undergo reactions with methanol during extraction, storage, and possibly during the electrospray process (Bateman et al., 2008). For instance, Batman et al (2008) suggested carboxylic acids could react with methanol to form esters and with carbonyls to hemiacetals and acetals. We checked the presence and relative abundance of these potential products (to that of our precursor, α pOS-249) in our aerosol mass spectra. At the maximum OH exposure, only a few products that could be potentially formed from the reactions of α pOS-249 with methanol were detected and they had negligible intensities. This would suggest that the influence of methanol is not significant on the identification of the major reaction products."

Comment #4:

4) Section 3.1. These results are really interesting and important for the community. However, as has been studied more within the community, phase state, ionic strength, and "shell" formation are most likely extremely important parameters in the OH heterogenous reaction of α pOS-249. As this is a single phase experiment, there is likely little phase state and shell formation concerns (which might be useful to explore in future experiments including with changing RH). I strongly recommend the authors briefly put this experiment into this context and how these results are most likely upper limits. Similar thoughts in regards to ionic strength, especially as this may impact the reaction mechanism and what intermediate and final products are stable.

Author Response:

Thanks for the valuable comments and suggestions. We agree with the reviewer that the complex interplay between aerosol phase, morphology and ionic strength can significantly alter the heterogenous OH reaction kinetics and mechanisms and are warrant for further study. We have added the following discussion in **Sect. 3.1 Oxidation Kinetics** to address these issues.

Page 12, Line 4: "In this study, the heterogenous OH reactivity of aqueous α pOS-249 aerosols at a single RH (50 %) was investigated. In the atmosphere, the complex interplay between aerosol phase state (e.g. solid or aqueous), morphology and the types and concentrations of salts could significantly alter the heterogenous OH reaction kinetics and mechanisms under different environmental conditions (e.g. RH and temperature).

The aerosol physical state can play a key role in determining the heterogeneous kinetics and chemistry of pure organic aerosols (Koop et al., 2011; Shiraiwa et al., 2011, 2013; Chan et al., 2014). For instance, Chan et al. (2014) reported that aqueous succinic acid aerosols reacted about 40 time faster than in solid aerosols towards heterogeneous OH oxidation. These could be explained by the more rapid diffusion of succinic acid to the surface of aqueous droplets for oxidation than solid aerosols. Moreover, for aqueous droplets, aerosol water content can vary considerably, depending on atmospheric conditions. The change in aerosol-phase water and solutes concentrations would influence the reactivity by varying aerosol viscosity (Slade and Knopf, 2014; Chim et al., 2017; Marshall et al., 2016, 2018). For instance, oxidation kinetics in highly concentrated aqueous organic aerosols are found to be much slower than those in diluted ones. This is because aerosol viscosity generally increases with the solute concentration,

thereby slowing down the diffusion of organic molecules within the aerosol and lowering the overall reactivity.

10 Atmospheric aerosols are comprised of organic compounds, inorganic salts and many other species. To date, large uncertainty remains in how inorganic salts alter the heterogeneous kinetics and chemistry (McNeill et al., 2007, 2008; Dennis-Smith et al., 2012). A few laboratory studies have revealed that the presence of dissolved inorganic ions (e.g., ammonium sulfate) can reduce the heterogeneous OH reactivity of organic compounds but does not significantly alter the reaction mechanisms (Mungall et al., 2017; Kwong et al., 2018; Lam et al., 2019). More recently, Xu et al. (2020b) reported that the change in the heterogeneous reactivity strongly depend on the concentration of organic compounds and inorganic salts. They found that the rate of the reactions decreases when the organic-to-inorganic mass ratio (OIR) decreases. This could be explained by the colliding probability between OH radical and organic species at the aerosol surface becoming lower in the presence of salt, resulting in a smaller overall reaction rate. We also acknowledge that different inorganic ions could have different propensity for air-aerosol interface depending on their polarizability and interactions with other components (Jungwirth and Tobias, 2002; Gopalakrishnan et al., 2005). Jungwirth and Tobias (2002) investigated the preference of sodium cation and chlorine anion to the interface or bulk in sea salts aerosols based on polarizable MD simulation. They found that chlorine anion has a stronger propensity for the interface than sodium cation and is proportional to its polarizability. Therefore, the types and concentration of inorganic ions (e.g. ionic strength) within aerosol could potentially alter the overall heterogeneous kinetics and reaction pathways. To date, it remains an open question whether the salts alter heterogeneous reactivity chemically, physically, or both.

30 We would like to note that additional uncertainties in heterogeneous reactivity of organic compounds in the presence of inorganic salts could also arise when these organic–inorganic droplets undergo phase separation, depending on environmental conditions and aerosol composition (e.g. different types of inorganic salts, the average oxygen-to-carbon (O:C) elemental ratio of organic compounds, and OIR) (You et al., 2014; Qiu and Molinero, 2015; Freedman, 2017, 2020). These phase-separated droplets typically exhibit two distinct liquid phases: an inorganic-rich inner phase and an organic-rich outer phase. Different morphologies have also been observed (e.g. core–shell morphology, partially engulfed morphology and transitions between different types of morphology). Lam et al. (2021) recently reported that phase-separated organic–inorganic droplets have a slightly higher reactivity towards gas-phase OH radicals, compared to single-phase ones. As phase separation occurred, an uneven distribution of organic species within the droplets increased the collision probability between organic molecules and OH radicals at or near the droplet surface. Overall, further studies emphasized on the effects of aerosol composition, phase transition and separation, and morphologies on heterogeneous reactivity of OSs are needed to better understand their transformation rates and chemistry.”

40 Relevant references were also added into the reference list:

- Chan, M. N., Zhang, H., Goldstein, A. H., and Wilson, K. R.: Role of water and phase in the heterogeneous oxidation of solid and aqueous succinic acid aerosol by hydroxyl radicals, *J. Phys. Chem. C*, 118, 28978–28992, <https://doi.org/10.1021/jp5012022>, 2014.
- Chim, M. M., Chow, C. Y., Davies, J. F. and Chan, M. N.: Effects of relative humidity and particle phase water on the heterogeneous OH oxidation of 2-methylglutaric acid aqueous droplets, *J. Phys. Chem. A*, 121, 1666–1674, doi:10.1021/acs.jpca.6b11606, 2017.
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Comment #5:

5) Section 3.4. Though the authors show that sulfate formation is minimally important, there is a question if the sulfate they observe may be due to the hydrolysis of the primary, secondary, etc. products. The authors investigate and determine that apOS-249 is likely stable but it is currently not clear if the other products may be.

Author Response:

10 Thanks for this insightful comment. We agree with the reviewer that inorganic sulfate could also be generated from the hydrolysis of reaction products. This is because secondary and tertiary OS could readily undergo hydrolysis and may form upon OH oxidation based on our proposed reaction mechanisms. We have added this information in the manuscript.

Page 20, Line 22: “Furthermore, we cannot rule out the possibility of the formation of SO_4^{2-} could be from the hydrolysis of reaction products as some secondary and tertiary OS could readily undergo hydrolysis and may form upon oxidation.”