Atmos. Meas. Tech. Discuss., doi:10.5194/amt-2017-344-RC1, 2017 © Author(s) 2017. This work is distributed under the Creative Commons Attribution 4.0 License.



Interactive comment on "Measurement of formic acid, acetic acid and hydroxyacetaldehyde, hydrogen peroxide, and methyl peroxide in air by chemical ionization mass spectrometry: airborne method development" by Victoria Treadaway et al.

Anonymous Referee #1

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This study details the detection and quantification of a selection of important atmospheric molecules using a multi-reagent ion chemical ionization mass spectrometer (CIMS). The multi-reagent ion system reported here blends CO2 in air and CH3I in N2, with the primary reagent ions being O2-, CO2(O2)-, and I-. This is different from previously implemented multi-reagent ion systems, as the two reagent gases are added simultaneously and tuned such that I-, O2-, and CO2(O2)- ion cluster chemistries are operable. The multi-reagent system was successfully deployed in ambient air-borne and laboratory measurements. This novel twin-reagent CIMS technique is likely to be in-

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teresting to researchers working with CIMS instruments to detect important gas-phase atmospheric molecules. The less selectivity of O2- (and CO2(O2)-) CIMS coupled with the better understood I- CIMS has potential to improve the current understanding of atmospheric gas-phase chemistry. I recommend publication of the manuscript after some relatively minor issues (detailed below) are addressed.

âĂć Line 145: What is the reaction time between the sample gas and the reagent ions inside the ion-sample reaction cell?

aĂć Line 153: "This pressure was stated to provide the maximum yield of cluster ions and peak sensitivity..." Just checking, was this stated by the RXN cell manufacturer?

âĂć Line 165: Maybe mention the optimized mixing ratios of CO2 and pure air used for HP and MHP signals in addition to the reference to the O'Sullivan paper?

aĂć Figure 2: The figure presently does not provide a lot of information. What was the averaging time used to obtain the spectrum? Would a longer averaging time provide a less noisy spectrum with the relevant peaks clearly defined? I suppose the log scale was used to show the lower signals of I-(HP), I-(HFo), and I-(MHP) in the same figure. Maybe having a linear scale (so the highest peaks can be clearly shown), and a zoomed-in inset of these lower signals would make a better figure?

âĂć Trivial comment: Figure 4 has no a) and b) labels although it is referenced as such in the text.

åÅć Paragraph starting from line 399: Do you see an increase in signals of (H2O)nlclusters (where n is 2,3,4...) at above 1000 ppm water vapor mixing ratio? It could be that, at higher degrees of hydration of the I- anion, (HAc)I- formation becomes unfavorable (probably due to a steric hindrance to (HAc)I- formation, i.e. the multiple water molecules attached to I- make the formation of I-(HAc) difficult), causing a decrease in sensitivity. On the other hand, I-(HFo) formation might become more favorable when multiple water molecules are attached to I- (HFo being a smaller molecule might be additionally stabilized by a sequential evaporation of multiple water molecules), explaining the increase in I-(HFo) sensitivity at higher water vapor mixing ratios you report. In any case, the possible detection of water dimers, trimers, tetramers clustered to I- at high water vapor concentration should probably be commented upon.

åĂć Continuing on the same theme, I would think that the binding strength of (H2O)Icluster is weaker than the (HAc)I- cluster, so a ligand-exchange reaction between HAc and water, which is reaction 4 in your manuscript, is likely not the reason for the decrease in (HAc)I- signal at higher water vapor concentrations.

åĂć Line 402: "indicated the switching reaction equilibrium for HAc (4) behaved like that for HFo...". I might have misunderstood, but don't you observe an increase in the sensitivity of (HFo)I- with an increase in water vapor mixing ratio? Does that then not imply that (HFo)I-, unlike HAc, is not affected by a possible ligand exchange reaction with water (reaction 4)?

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