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4, C127-C129, 2011

Interactive Comment

Interactive comment on "Measurements of tropospheric HO₂ and RO₂ by oxygen dilution modulation and chemical ionization mass spectrometry" by R. S. Hornbrook et al.

Anonymous Referee #2

Received and published: 15 March 2011

General comments

This paper presents an improved method for measuring hydroperoxy and organic peroxy radicals developed and well characterized for both ground-based and airborne measurements. The method builds upon previously described CIMS techniques with improved separation between HO2 and HO2+RO2 modes and a relatively good time resolution. Overall this is a well written paper and I recommend it be published in AMT after revision and ask the authors to consider the following special comments in their revision. Full Screen / Esc

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Special Comments

- 1. P.389: it is mentioned that in the method by Edwards et al. (2003), it takes 30 minutes for the reagent gases to be flashed from the inlet. Does that mean that previous measurements by this method have time resolution at least of 30 minutes? It seems to me too long considering the relatively large volume metric flow of ∼10 L/min at 200 mbar. I wonder if this is a typo, i.e. it should be 30 seconds instead of 30 minutes.
- 2. P.391, it is hard for me to believe that 2.32 SLPM of air flow can be pulled through a 0.50 mm (even smaller than the inlet diameter of 2.7mm) diameter orifice between the dilution region and neutral chemistry region without a significant pressure drop (from the text, the pressures in the dilution region and the neutral chemistry region are kept the same at 200 mbar). It seems to me that either a larger orifice (larger than 0.5 mm diameter) is used and/or there is a significant pressure drop between the two regions. Please double check these numbers
- 3. P.392 bottom: Are there any formed OH radicals left at the point where the SO2 rear injector is located? Have the authors done any tests to ensure that almost all produced OH has been converted into HONO? If not, the peroxy radical signals might be over-corrected.
- 4. P.394 in Eq. (1): is f(NO3-) the counting frequency when SO2 mixture is added through the front or the rear injector? Or the average of the both?
- 5. P.396, please add Edwards et al. (2003) where the N2O actinometry experiment is mentioned.
- 6. P.398, in the bottom line; radical wall loss due to small flow rate is mentioned. Were there any tests done to ensure that there is no significant wall loss at a typical flow rate of 5 SLPM?
- 7. P.399, 2nd line, change "that" to "than".
- 8. P.400: humidified N2 or O2 is added in case of very dry sample air. How much

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humidity is typically added? How does this affect the sensitivity and/or conversion efficiencies of RO2?

- 9. P.403: a lower inlet pressure of 133 mbar is used for sample at high altitude. How are the calibration and RO2 conversion efficiencies handled? Any significant difference from the normal operation at 200 mbar?
- 10. P.404: are the vapor flow rate of liquid RO2 precursors set using the needle valve so that the OH reactivity of each RO2 precursor is typical (60-250 per second)?
- 11. P.414: +/-35% uncertainty: does it include the uncertainties due to RO2 signals in the HO2 measurements and the unequal alpha values of RO2 in the HO2+RO2 mode? How does this additional uncertainties affect the model intercomparison shown in Fig. 9?

Interactive comment on Atmos. Meas. Tech. Discuss., 4, 385, 2011.

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