Supplement of Observations of VOC emissions and photochemical products over US oil- and gas-producing regions using high-resolution $\text{H}_3\text{O}^+$ CIMS (PTR-ToF-MS)

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Comparison of aromatic species measured by iWAS/GC-MS and H$_3$O$^+$ ToF-CIMS. The benzene comparison has a slope of 1.04 and R of 0.97; Toluene slope is 1.13 and R = 0.98; and C8 aromatics slope of 0.86 and R = 0.94.
S1. Peak fitting errors: data quality assessment
At the resolution of the H$_3$O$^+$ ToF CIMS instrument (~4000 m/dm), peaks at the same nominal mass frequently overlap. To separate the contribution from overlapping peaks at the same nominal mass, the Tofware peak fitting software (Aerodyne Research Inc./Tofwerk AG) iteratively fits a set of pseudo-gaussian high-resolution peaks to the sum signal across that nominal mass. The number of peaks at a given nominal mass, the peak shape, and the exact m/z of each peak center are user-defined. However, problems can occur in circumstances where a peak is much smaller than its neighbor, or where the m/z separation between peaks is unusually small. Under these circumstances, the returned peak-fit intensity can be highly inaccurate. The magnitude of the error depends on the relative size, closeness in m/z space, mass calibration precision and accuracy, and peak shape of the overlapping peak, and so it can change over time. These nonrandom errors can create false variability in a particular ion mass. This peak fitting issue is well known for both CISM and AMS instruments but has not yet been completely addressed (Müller et al., 2011; Cubison and Jimenez, 2015; Stark et al., 2015; Corbin et al., 2015).

Although there is not yet a quantitative method to determine the extent and magnitude of peak overlap errors in every situation, Tofware provides qualitative assessment tools that rely on the dependence of peak-fitting error on mass calibration precision (Timonen et al., 2016). Several studies have suggested that mass calibration precision and accuracy are strong controls on peak fitting error (Corbin et al., 2015; Cubison and Jimenez, 2015). When applied to a particular high-resolution peak, the tool re-calculates the fitted peak intensity for a range of expected error in the mass calibration (±10 ppm). Peaks that are strongly affected by peak overlap error will have highly variable fitted peak intensity. All high-resolution peaks reported in this work are not strongly affected by peak fit error, i.e., the fitted peak intensity is independent of mass calibration uncertainty. Mass spectra and time-series fitted with mass calibration ±10 ppm are shown in Figure SI 5.

S2. Limit of detection calculation
The measured instrument signal (ion counts) is the product of the measurement time $t$, the sensitivity $C_f$, and the concentration $[X]$:

$$S = C_f[X]t$$

Instrument noise is related to the square root of the signal:

$$N = \sqrt{S} = \sqrt{C_f[X]t}$$

The signal to noise ratio is therefore:

$$\frac{S}{N} = \frac{C_f[X]t}{\sqrt{C_f[X]t}} = \sqrt{C_f[X]t}$$
We define the “limit of detection” as the concentration \([X]\) at which the signal to noise ratio (S/N) equals 3:
\[
\frac{S}{N} = 3 = \sqrt{C_f[X]_{LoD} t}
\]

By solving for \([X]_{LoD}\), it can be seen that the limit of detection is inversely proportional to the measurement time:
\[
[X]_{LoD} = \frac{9}{C_f t}
\]

This is different from noise, which decreases as the square root of the measurement time.

For several reasons, the measurement limit of detection is actually somewhat higher than the calculations above would suggest. First, there is some background count rate \(B\), and the VOC signal is determined from the difference between the ambient signal and the background signal. Second, because of additional error from high-resolution peak fitting, the noise is somewhat higher than the square root of the signal. To account for this additional error, a linear scaling factor \(\alpha\) is included in the calculation of noise. With the inclusion of the background \(B\) and scaling factor \(\alpha\), the signal to noise calculation is (Bertram et al., 2011):
\[
\frac{S}{N} = \frac{C_f[X] t}{\alpha \times \sqrt{C_f[X] t + 2B t}} = 3
\]

The detection limit \([X]\) can then be solved for using the quadratic equation. Detection limits given in this work were calculated using raw counts-per-second, without reagent ion normalization or a ToF duty-cycle correction applied.

**S3. Photochemical aging analysis**

The relative ratios of different VOCs can be used to establish relative photochemical aging. Using PTRMS measurements of C6-C10 aromatic species, this method has been applied to photochemical aging of VOCs in urban areas (de Gouw et al., 2005; Warneke et al., 2007) and in an oil and gas producing region (Koss et al., 2015). Benzene (C6 aromatic), has the slowest reaction rate with OH; as the cumulative OH exposure of an air mass increases, the ratios of other aromatics to benzene should decrease. The ratio of an aromatic VOC to benzene at time \(t\) can be written as:
\[
\left( \frac{VOC}{Benzene} \right)_t = \left( \frac{VOC}{Benzene} \right)_0 \times e^{(k_{benzene} - k_{VOC}) \int_0^t [OH] dt}
\]

Where \(\left( \frac{VOC}{Benzene} \right)_0\) is the ratio of the aromatic to benzene at time \(t=0\), \(k_{benzene}\) and \(k_{VOC}\) are the rate constants of benzene and the aromatic with OH, and \(\int_0^t [OH] dt\) is the integrated OH exposure. If the ratio at time \(t\), the emission ratio, and the OH rate constants are known, the integrated OH exposure can be calculated.
We restricted this analysis to the three plume transects in the southwest portion of the flight survey. These three plumes have similar VOC composition, are located close to one another, and can be reasonably be assumed to be the same air mass. The flight was from south to north, so the northmost transect was latest in the day. The ratios of C7-C10 aromatics decrease from south to north (Figure SI 2).

We calculated integrated OH exposure in each of the three transects using toluene, C8 aromatics, C9 aromatics, and C10 aromatics. OH rate constants for C8, C9, and C10 aromatics were taken from Koss et al. (2015). OH rate constants for benzene and toluene were taken from Atkinson and Arey (2003). The average of the highest 5% points in the southmost transect was used as the ratio at time t=0. We note that in this analysis, time t=0 is not necessarily the time of emission. We set time t=0 to be the age of the least aged air mass. The exact emission ratio is not relevant.
to the analysis of pyrrole presented in the manuscript, since we are mainly interested in the relative rate of aging.

The calculated OH exposures from C7-C10 aromatics were averaged to give the final OH exposure used in the analysis. The separately calculated OH exposures were within 40% of the mean.

The OH rate constant for pyrroline was calculated using an exponential best fit to the pyrroline:benzene ratio as a function of the integrated OH exposure.

![Figure SI 3. Ratio of pyrroline to benzene as a function of OH exposure.](image)

<table>
<thead>
<tr>
<th></th>
<th>OH rate constant, cm$^3$ cmolecule$^{-1}$ s$^{-1}$</th>
<th>Emission ratio pyrroline:benzene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original calculation</td>
<td>$1.72 \pm 0.14 \times 10^{-11}$</td>
<td>$0.126 \pm 0.006$</td>
</tr>
<tr>
<td>Using maximum OHdt calculated from all aromatics</td>
<td>$1.39 \pm 0.11 \times 10^{-11}$</td>
<td>$0.142 \pm 0.008$</td>
</tr>
<tr>
<td>Using minimum OHdt calculated from all aromatics</td>
<td>$1.78 \pm 0.17 \times 10^{-11}$</td>
<td>$0.096 \pm 0.004$</td>
</tr>
<tr>
<td>Actual concentration 2x lower</td>
<td>$1.72 \pm 0.14 \times 10^{-11}$</td>
<td>$0.252 \pm 0.012$</td>
</tr>
<tr>
<td>Actual concentration 2x higher</td>
<td>$1.72 \pm 0.14 \times 10^{-11}$</td>
<td>$0.063 \pm 0.003$</td>
</tr>
</tbody>
</table>
Data selection based on flight altitude for each flight.
Figure SI 5

High-resolution mass spectra for select nominal masses.

Key:

<table>
<thead>
<tr>
<th>Mass spectra</th>
<th>Time series</th>
</tr>
</thead>
<tbody>
<tr>
<td>sum HR fit</td>
<td>Report fit</td>
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<tr>
<td>measurement</td>
<td>Fit +10 ppm</td>
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<tr>
<td>isotopes</td>
<td>m/z cal</td>
</tr>
<tr>
<td>HR masses</td>
<td>Fit -10 ppm</td>
</tr>
</tbody>
</table>

**m/z 31**

- CH$_2$OH$^+$
- NOH$^+$
- CH$_2$OH$^+$
- N$_2$H$^+$
- C$_2$H$_7$^+

**m/z 33**

- CH$_4$OH$^+$
- OH$^+$

**m/z 35**

- H$_2$SH$^+$
- (OH)H$_2$(H$^+$HN)

**m/z 41**

- C$_3$H$_5^+$
- ArH$^+$
- NH$_2$(H$_3$O)$^+$

**m/z 43**

- CH$_2$O$_2$H$^+$
- C$_2$H$_7^+$
- C$_3$H$_7^+$

Reported fit +10 ppm m/z cal
Fit -10 ppm m/z cal
Figure SI 6

Distribution of acetaldehyde, acetone, MEK, and acetic acid during each flight.
PTR-ToF-MS measurement of m/z 137 C_{10}H_{16}H^{+} and m/z 81 C_{6}H_{9}+ during the UBWOS 2013 campaign in the Uintah oil and gas field, Utah. The right two panels show the ion ratios to benzene, which indicate whether they belong to primary or photochemically produced species. We note that frequently, m/z 137 C_{10}H_{16}H^{+} is interpreted as monoterpenes, and m/z 81 C_{6}H_{9}+ as a monoterpene fragment (e.g. Kim et al. (2010)). In both Utah (UBWOS 2013) and the Permian (SONGNEX 2015) these two masses are uncorrelated, and m/z 81 has much higher intensity than m/z 137.
Potential diamondoid ion masses in SONGNEX measurement (top), headspace of West Texas Crude (middle), compared to reported distribution of diamondoids (bottom) (Stout and Douglas, 2004). Diamondoid masses are the bold sticks in the mass spectra. The three distributions are somewhat different. This could be because: (1) the C_{10}H_{16}H^+ detected during SONGNEX is not a diamondoid; (2) inlet effects reduced the measured amount of higher molecular weight species; (3) distribution of diamondoids in other fuels can differ from that presented by Stout and Douglas (2004); (4) The higher molecular weight species have a shorter atmospheric lifetime.
Comparison of $\text{H}_3\text{O}^+$ ToF-CIMS masses m/z 77 $\text{C}_3\text{H}_6\text{O}_3\text{H}^+$, m/z 91 $\text{C}_3\text{H}_6\text{O}_3\text{H}^+$, and m/z 46 NO$_2^+$ to various reactive nitrogen species during the April 23 SONGNEX flight. Ten-second average data is shown for clarity. For the comparison to small alkyl nitrates, the $\text{H}_3\text{O}^+$ ToF-CIMS data has been averaged to the iWAS sampling times. Small differences exist between the CaRDS and Chemiluminescence measurements; these were averaged where both were available. The choice of NO$_x$ and NO$_y$ data source does not affect our conclusions.
H$_3$O$^+$ ToF CIMS m/z 46 NO$_2^+$ tracer measurements compared to PAN. The high m/z 46 during the 4-21 Bakken flight was within an aged biomass burning plume (identified by enhanced acetonitrile).
Figure SI 11

A: H$_3$O$^+$ ToF-CIMS m/z 46 compared to NO$_2$ from all flights, colored by the fraction of NO$_2$ accounted for by PAN. B: H$_3$O$^+$ ToF-CIMS m/z 77 compared to PAN from all flights, colored by relative humidity proxy m/z 37 H$_5$O$_2^+$:m/z 19 H$_3$O$. 
Comparison of the distribution of unsaturated nitrogen species $C_nH_{2n-1}NH^+$ between (a) oil- and gas-related VOCs measured during the April 23 Permian flight, and (b) a biomass burning plume encountered during the April 21 Bakken flight. The series is m/z 28 HCNH$^+$, m/z 42 C$_2$H$_3$NH$^+$, m/z 56 C$_3$H$_5$NH$^+$, m/z 70 C$_4$H$_7$NH$^+$, etc. The values shown are the average signal enhancement (ncps enhancement) in the boundary layer during the Permian flight, and the average signal enhancement (ncps enhancement) in the biomass burning plume compared to immediately before the plume during the Bakken flight.

**S4. Comparison of nitrate reactivity to literature**

The estimated average pyrroline nitrate reactivity is 15x10⁻³ s⁻¹ (assuming the same nitrate reaction constant as pyrrole).

For comparison, Stutz et al. (2010) measured nocturnal nitrate loss rates in Houston, TX in an area influenced by emissions from the petrochemical industry. Total nitrate loss frequency was typically around 20 to 60 x10⁻³ s⁻¹ during the middle of the night. VOC nitrate reactivity ranged from 0 to 50x10⁻³ s⁻¹ and was typically the dominant loss mechanism. Indirect loss via N₂O₅ uptake to aerosol was the second most important pathway and ranged from 0 to 40 x10⁻³ s⁻¹.

Crowley et al. (2011) measured nocturnal nitrate loss frequency in polluted air in Spain. Total nitrate loss frequency was on the order of 5x10⁻³ s⁻¹ to 50x10⁻³ s⁻¹. Loss of NO₃ to VOCs, and loss via N₂O₅ uptake to aerosol were both important loss mechanisms, each on the order of 5x10⁻³ s⁻¹.

Conversely, (Tsai et al., 2014) found that loss via N₂O₅ uptake to aerosol accounted for nearly all of the typically 50 x10⁻³ s⁻¹ NO₃ loss in the Los Angeles metropolitan area.
References


