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Evaluation of NO⁺ reagent ion chemistry for online measurements of atmospheric volatile organic compounds

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Abstract. NO⁺ chemical ionization mass spectrometry (NO⁺ CIMS) can achieve fast (1 Hz and faster) online measurement of trace atmospheric volatile organic compounds (VOCs) that cannot be ionized with H₃O⁺ ions (e.g., in a PTR-MS or H₃O⁺ CIMS instrument). Here we describe the adaptation of a high-resolution time-of-flight H₃O⁺ CIMS instrument to use NO⁺ primary ion chemistry. We evaluate the NO⁺ technique with respect to compound specificity, sensitivity, and VOC species measured compared to H₃O⁺. The evaluation is established by a series of experiments including laboratory investigation using a gas-chromatography (GC) interface, in situ measurement of urban air using a GC interface, and direct in situ measurement of urban air. The main findings are that (1) NO⁺ is useful for isomerically resolved measurements of carbonyl species; (2) NO⁺ can achieve sensitive detection of small (C₄-C₈) branched alkanes but is not unambiguous for most; and (3) compoundspecific measurement of some alkanes, especially isopentane, methylpentane, and high-mass (C₁₂-C₁₅) n-alkanes, is possible with NO⁺. We also demonstrate fast in situ chemically specific measurements of C_{12} to C_{15} alkanes in ambient air.

1 Introduction

Volatile organic compounds (VOCs) are central to the formation of ozone and secondary organic aerosol and can have direct human health effects. Attempting to understand the behavior of these species in the troposphere presents sev-

eral measurement challenges (Glasius and Goldstein, 2016). First, VOCs are highly chemically diverse. Second, many environmentally important species require measurement precision of better than 100 parts per trillion (ppt). Finally, numerous applications, such as eddy flux analyses or sampling from a mobile platform, require fast in situ measurements, with 1 min or faster time resolution.

 $\rm H_3O^+$ chemical ionization mass spectrometry ($\rm H_3O^+$ CIMS), more commonly known as proton-transfer-reaction mass spectrometry (PTR-MS), is a well-established approach to measuring VOCs (de Gouw and Warneke, 2007; Jordan et al., 2009b). In $\rm H_3O^+$ CIMS, air is mixed with hydronium ($\rm H_3O^+$) ions in a drift tube region. VOCs are ionized by transfer of the proton from $\rm H_3O^+$ to the VOC. These instruments are capable of VOC measurements that are fast, sensitive, and chemically detailed (Jordan et al., 2009b; Graus et al., 2010; Sulzer et al., 2014; Yuan et al., 2016).

Despite these advantages, $\rm H_3O^+$ CIMS has several limitations related to the reagent ion chemistry. For one, this technique generally cannot distinguish between isomers. For instance, this is a significant limitation when measuring aldehyde and ketone carbonyl isomers, which display very different behavior in the atmosphere. Separation of propanal and acetone with PTR-MS has been explored using collision-induced dissociation with an ion-trap mass analyzer, but this technique negatively affects the instrument time resolution and sensitivity (Warneke et al., 2005). Additionally, some proton transfer reactions are dissociative. Large hydrocarbons ($\rm C_8$ and larger) fragment into common small masses, making spectra difficult to interpret (Jobson et al., 2005; Er-

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ickson et al., 2014; Gueneron et al., 2015). Alcohols and aldehydes can lose H_2O , lowering the sensitivity to the protonated parent mass; their product ion masses then coincide with those of hydrocarbons, making independent measurement difficult (Španěl et al., 1997; Buhr et al., 2002). Furthermore, H_3O^+ CIMS is not sensitive to small ($\sim C_8$ and smaller) saturated alkanes, as their proton affinities are lower than or very close to that of water (Arnold et al., 1998; Gueneron et al., 2015). This is a serious limitation in studies of urban air or emissions from oil and natural gas extractions, where small alkanes can contribute a large fraction to the total gas phase carbon and chemical reactivity (Katzenstein et al., 2003; Gilman et al., 2013). Gas-chromatography (GC) techniques avoid many of these limitations but have much slower time resolution.

Use of NO⁺ reagent ion chemistry may address some of the limitations of H₃O⁺. Reaction of NO⁺ with various VOCs has been extensively studied using selected-ion flow tube methods (SIFT-MS). SIFT methods use a quadrupole mass filter between the ion source and ion-molecule reactor, which provides a very pure reagent ion source but limits the primary ion signal. SIFT studies have identified the major products of the reaction of NO⁺ with VOCs representative of many different functional groups (Španěl and Smith, 1996, 1998a, b, 1999; Španěl et al., 1997; Arnold et al., 1998; Francis et al., 2007a, b). Aldehydes and ketones are easily separable: ketones cluster with NO⁺, forming mass (m + 30)ions, whereas aldehydes react by hydride abstraction, forming mass (m-1) ions (where m is the molecular mass of the species). Rather than losing H₂O, as in H₃O⁺ CIMS, alcohols react by NO⁺ adduct formation or hydride abstraction. And finally, NO^+ can be used to detect alkanes: small (> C_4) branched alkanes and large (> C₈) n-alkanes react by hydride abstraction, forming mass (m-1).

The application of SIFT methods to atmospheric analysis has been limited by relatively poor sensitivity (Smith and Španěl, 2005; Francis et al., 2007b; de Gouw and Warneke, 2007), although better sensitivities have been reported in recent years (Prince et al., 2010). The adaptation of an existing CIMS instrument to use the SIFT technique requires extensive instrument modification or the purchase of an external SIFT unit (Karl et al., 2012). Several groups have experimented with low-cost adaptation of H₃O⁺ CIMS instruments to use NO⁺ chemistry. Knighton et al. (2009) adapted an H₃O⁺ CIMS instrument to measure 1,3-butadiene and demonstrated in situ detection of this species in the atmosphere. Jordan et al. (2009a) have developed a hollowcathode ion source capable of switchable reagent ion chemistry, and they demonstrated laboratory measurement with NO⁺ of several aromatics, chlorinated aromatics, and carbonyls, with sensitivities comparable to H₃O⁺ CIMS. The NO⁺ capability of the Jordan et al. instrument has been used in the laboratory by Inomata et al. (2013) to investigate detection of n-tridecane, by Agarwal et al. (2014) to measure picric acid, and by Liu et al. (2013) to investigate the behavior of methyl vinyl ketone (MVK) and methacrolein in a reaction chamber.

These studies suggest that an easy, low-cost adaptation of H₃O⁺ CIMS instruments to NO⁺ chemistry could greatly enhance our capability to measure VOCs in the atmosphere. However, the number of VOC species investigated to date is small and few field measurements have been reported. The ability of a modified H₃O⁺ CIMS instrument to separate carbonyl isomers in ambient air, and to measure small alkanes both in the laboratory and in ambient air, has not been evaluated. Finally, the lack of fragmentation of n-tridecane reported in Inomata et al. (2013) is intriguing, but the use of an NO⁺ CIMS instrument to measure similar high-mass alkanes in ambient air has not been demonstrated.

Here we evaluate the adaptation of an ${\rm H_3O^+}$ CIMS instrument to use ${\rm NO^+}$ reagent ion chemistry. We provide specifics on instrument setup and operating parameters. We report the sensitivity and spectral simplicity of ${\rm NO^+}$ CIMS, relative to ${\rm H_3O^+}$ CIMS, for nearly 100 atmospherically relevant VOCs, including a wide range of functional groups, and provide product ion distributions for several representative compounds. We demonstrate, interpret, and evaluate measurements of separate aldehyde and ketone isomers, light alkanes, and several other species in ambient air. Finally, we investigate measurement of high-molecular-mass alkanes using ${\rm NO^+}$. We extend the laboratory analysis of high-mass alkanes to ${\rm C_{12}\text{--}C_{15}}$ n-alkanes and demonstrate fast, in situ measurement of these species in ambient air.

2 Methods

2.1 Instrumentation

Two separate H₃O⁺ CIMS instruments (referred to hereafter as PTR-QMS and H₃O⁺ ToF-CIMS) were adapted to NO+ chemistry in this work. Both instruments consist of (1) a hollow cathode reagent ion source, (2) a drift tube reaction region, (3) an ion transfer stage that transports from the drift tube to the mass analyzer and allows differential pumping, and (4) a mass analyzer. Both instruments have nearly identical hollow cathode ion sources and drift tube reaction regions, described in detail in de Gouw and Warneke (2007). The PTR-QMS (Ionicon Analytik) uses ion lenses to transfer ions from the drift tube to a unit-mass-resolution quadrupole mass analyzer (Pfeiffer). This instrument is described further by de Gouw and Warneke (2007). The H₃O⁺ ToF-CIMS uses RF-only segmented quadrupole ion guides to transfer ions from the drift tube to a time-of-flight mass analyzer produced by Aerodyne Research Inc./Tofwerk with a mass resolution of 4000-6000 (Bertram et al., 2011). This instrument is described further by Yuan et al. (2016). A similar PTR-ToF instrument using quadrupole ion guides has also been recently described (Sulzer et al., 2014). ToF-CIMS data were analyzed using Tofwerk high-resolution peak-fitting software (Aerodyne Research Inc./Tofwerk AG). A description of the algorithm is given in DeCarlo et al. (2006).

A GC instrument was used both as an interface to the ToF-CIMS and as a separate instrument using an electron-impact quadrupole mass spectrometer. The GC collects VOCs in a liquid nitrogen cryotrap for a 5 min period every 30 min. VOCs are then injected onto parallel Al₂O₃/KCl PLOT and semi-polar DB-624 capillary columns to separate C₂-C₁₁ hydrocarbons and heteroatom-containing VOCs. When used as an interface to the ToF-CIMS, the column eluant was directed to the inlet of the ToF-CIMS, where it was diluted with 50 sccm of clean air with controlled humidity. When operated as a separate instrument, the column eluant was directed to an electron-ionization quadrupole mass spectrometer (EIMS) operated in selected-ion mode. The response of this GC-EIMS instrument to various VOCs has been well characterized over a long period of field and laboratory applications, and further operational details have been reported elsewhere (Goldan et al., 2004; Gilman et al., 2010, 2013).

2.2 Adaptation of H₃O⁺ to NO⁺ CIMS

Ideally, both H₃O⁺ and NO⁺ reagent ion chemistry can be utilized with a single instrument. The fewest possible number of hardware parameters were changed to facilitate fast switching and instrument stability.

To achieve generation of $\mathrm{NO^+}$ ions, the water reservoir was replaced with ultra-high-purity air. The source gas flow (5 sccm), the hollow cathode parameters, and the drift tube operating pressure (2.4 mbar) were not changed. To optimize the generation of $\mathrm{NO^+}$ ions relative to $\mathrm{H_3O^+}$, $\mathrm{O_2^+}$, and $\mathrm{NO_2^+}$, and the generation of the desired $\mathrm{VOC^+}$ ion products, the voltages of the intermediate chamber plates, V_{IC1} and V_{IC2} , and the drift tube voltage V_{DT} were adjusted. An instrument schematic showing the locations of V_{IC1} , V_{IC2} , and V_{DT} can be found in the Supplement (Fig. S1). Optimization was performed sampling dry air.

It has been demonstrated that the quadrupole ion guides of the ToF-CIMS can significantly change the measured distribution of reagent and impurity ions (Yuan et al., 2016). The PTR-QMS does not have that issue as strongly (modeled and measured cluster distributions are largely similar, as discussed by de Gouw and Warneke, 2007) and therefore we explored the effect of $V_{\rm IC1}$, $V_{\rm IC2}$, and $V_{\rm DT}$ on reagent ion distribution using the PTR-QMS. As the PTR-QMS and ToF-CIMS have nearly identical ion source and drift tube design, we assume that ion behavior in these regions is the same for the two instruments.

First, $V_{\rm DT}$ was held constant at 720 V (the original setting of the PTR-QMS instrument), and $V_{\rm IC1}$ and $V_{\rm IC2}$ were varied (Fig. 1). The settings of $V_{\rm IC1}$ (140 V) and $V_{\rm IC2}$ (80 V) were selected as a compromise between high NO⁺ ion count rate and low-impurity ion count rates. The major impurity ions are $\rm H_3O^+$, $\rm O_2^+$, and $\rm NO_2^+$, and it is desirable to limit the formation of these ions because they react with VOCs, compli-

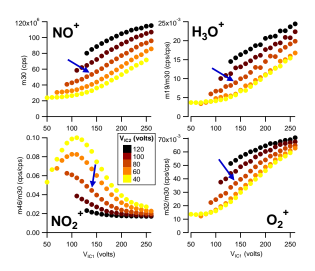


Figure 1. Dependence of NO^+ , H_3O^+ , NO_2^+ , and O_2^+ on intermediate chamber voltages. The arrow denotes the selected operating conditions. Experiment conducted in dry air (H_3O^+) is from residual water in the instrument and in commercial ultra-zero air.)

cating the interpretation of spectra. Next, several VOCs with different functional groups were introduced into the instrument, separately, and the drift tube electric potential scanned. A drift tube voltage of 350 V (electric field intensity relative to gas number density $E/N=60\,\mathrm{Td}$) was selected as a compromise between maximizing NO+ ion count rate, minimizing H₃O+, O₂+, and NO₂+, maximizing VOC ion count rates, minimizing alkane fragmentation, and promoting different product ions for carbonyls and aldehydes (Fig. 2). This setting results in about 10×10^6 cps (counts per second) of NO+ primary ions, while in typical PTR-MS settings we achieve about 30×10^6 cps of H₃O+ primary ions.

We note that the E/N of 60 Td used for the NO⁺ CIMS is much lower than that used in typical PTR-MS settings (circa 120 Td). In air, NO⁺ will react with water to produce H₃O⁺ and HNO₂ (Fehsenfeld et al., 1971). The electric field in the drift tube limits the formation of the NO⁺ · (H₂O)_n intermediaries in this reaction, promoting high NO⁺ count rates and VOC sensitivity. In PTR-MS, the drift field is used to prevent the formation of analogous H₃O⁺ · (H₂O)_n clusters. The bond energy of H₃O⁺ · (H₂O)_n clusters is significantly higher than that of NO⁺ · (H₂O)_n clusters (Keesee and Castleman, 1986); hence the need for a higher E/N in PTR-MS settings.

The remainder of the work detailed in this paper was performed using the ToF-CIMS with the settings as described here. The ToF-CIMS has the advantages of high mass resolution, fast time resolution, and simultaneous measurement of all masses. Further small adjustments were made to the ToF-CIMS quadrupole ion guide voltages using Thuner software (Tofwerk AG) to promote sensitivity to VOCs and separate carbonyl isomers. The two most important such adjustments decreased the electric potentials immediately upstream of each quadrupole ion guide (Fig. S2). These adjustments re-

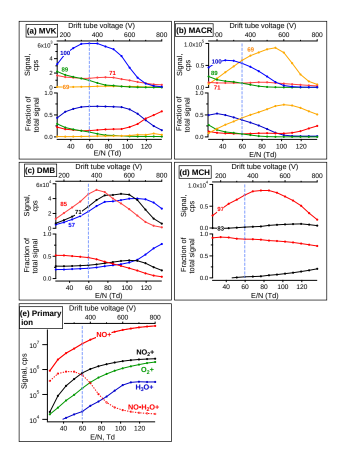


Figure 2. VOC and primary product ion dependence on drift tube voltage. Traces are labeled by the nominal product ion m/z in Th. (a) Methyl vinyl ketone; (b) methacrolein; (c) 2,2-dimethylbutane; (d) methylcyclohexane; (e) primary ions and clusters. The dashed line indicates the selected operating voltage. Experiment conducted in dry air (H_3O^+) is from residual water in the instrument and in commercial ultra-zero air.)

duced declustering at these locations, which improved the transmission of $VOC \cdot NO^+$ clusters.

3 Results and discussion

3.1 Laboratory experiments

3.1.1 Sensitivity and simplicity of the NO⁺ reagent ion chemistry

VOCs from several calibration cylinders (VOCs listed in Table S1 in the Supplement) were diluted with high purity air to mixing ratios of approximately 10 ppbv and introduced into the sampling inlet of the GC interface. Eluant from the column was directed into the ToF-CIMS as described above. A relative humidity of 20 % was used for this experiment. This humidity condition is similar to that expected for ambient measurements discussed in Sect. 3.2; this condition was cho-

sen to aid interpretation of ambient air data. Humidity effects are discussed in Sect. 3.1.5. Several species co-elute with another compound (m- and p-xylenes; myrcene and camphene; 1-ethyl,3-methylbenzene and 1-ethyl,4-methylbenzene); reported sensitivities and product ions are an average of the two co-eluting species.

Each VOC mixture was sampled twice, once with H₃O⁺ and once with NO⁺ reagent ion chemistry and instrument settings. Based on the results we evaluated the utility of NO⁺ CIMS relative to H_3O^+ CIMS using two metrics. The first metric is sensitivity for individual VOCs. To determine the sensitivity (S), the signals (cps) of all product ions were integrated over the width of the chromatographic peak and sensitivities for the measured VOCs using NO⁺ chemistry were calculated relative to the sensitivity using H₃O⁺ chemistry $(S_{NO+}/S_{H_3O^+})$. For several VOCs, we also calculated the relative sensitivity when only the most abundant product ion (the quantitation ion) is measured (Table 2b). Because only one concentration was sampled, this metric relies on sensitivity being linear with concentration. Linear sensitivity is a reasonable assumption for the NO⁺ and H₃O⁺ ToF-CIMS because separate multiple-point calibrations for select VOCs showed a linear response (Sect. 3.1.4, Fig. S10), H₃O⁺ CIMS has demonstrated linear sensitivity over a wide range of concentration (de Gouw and Warneke, 2007; Sulzer et al., 2014; Yuan et al., 2016), and the NO⁺ CIMS agrees well with an independent technique over a range of atmospheric concentrations (Sect. 3.2.2).

The second metric is the simplicity of spectra. In an ideal instrument, each VOC would produce only one product ion, and each ion mass would be produced by only one VOC. However, using NO⁺ and H₃O⁺ reagent ions, fragmentation of product ions does occur. As a metric for the complexity of the product ion distribution resulting from particular VOCs, we determined the fraction of the most abundant ion to the total signal from this VOC(F) and discuss (F_{NO^+}) relative to $(F_{H_3O^+})$. Figure S3 contains a comparison of $F_{\rm NO^+}$ and $F_{\rm H_3O^+}$ and an example product ion distribution. A larger value of this ratio means that NO⁺ reagent ion chemistry creates a simpler product ion distribution for that particular VOC. This metric does not indicate whether a particular product ion is produced by only one VOC. Uniqueness of product ions is discussed in Sect. 3.1.2. The NO⁺ CIMS product ion distributions of 25 atmospherically relevant VOCs are reported in Table 2.

Figure 3 summarizes the comparison between NO⁺ and H_3O^+ reagent ion chemistry for the two metrics. On the y axis the spectrum simplicity metric and on the x axis the sensitivity metric are shown.

Branched alkanes and most cyclic alkanes are detected with far greater sensitivity using NO^+ chemical ionization than with H_3O^+ chemical ionization. Aromatics and alkenes are detected slightly more sensitively, and, on average, ketones are detected slightly less sensitively. Alcohols are detected more sensitively, by at least a factor of 2, with the

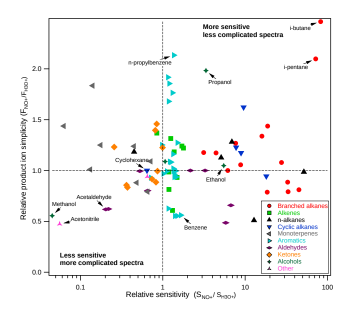


Figure 3. Comparison of production ion distribution and sensitivity of VOCs using NO^+ and H_3O^+ reagent ion chemistry, at a relative humidity of 20 %.

exception of methanol. The lower sensitivity to methanol is consistent with slower reaction kinetics reported in the literature (Španěl and Smith, 1997). Monoterpenes and acetonitrile are detected substantially less sensitively.

In comparing the simplicity of the product ion distribution between $\rm H_3O^+$ and $\rm NO^+$ chemistry, most branched and cyclic alkanes, ketones, and monoterpenes have a higher fraction of signal on a single product ion (simpler spectra). We also highlight that many alkyl substituted aromatics fragment substantially with $\rm H_3O^+$ chemistry but do not with $\rm NO^+$ chemistry. The few exceptions (notably, benzene) create more complicated spectra because an $\rm NO^+$ cluster product is also present (m+30).

3.1.2 Distribution of product ions

It is somewhat more difficult to predict the ionized VOC products of NO⁺ CIMS compared to H₃O⁺ CIMS, because NO⁺ has three common reaction mechanisms: charge transfer, hydride abstraction, and cluster formation. Groups of VOCs that have similar charge transfer and hydride abstraction enthalpies tend to react with similar ionization mechanisms (Fig. 4). Figure 4 uses thermodynamic information from Lias et al. (1988), as well as mechanistic information from this work (see Table S1 for a list of species) and from SIFT studies (Španěl and Smith, 1996, 1998a, b, 1999; Španěl et al., 1997; Arnold et al., 1998; Francis et al., 2007a, b). Charge transfer occurs when the reaction enthalpy is favorable, regardless of the hydride transfer enthalpy. When the charge transfer enthalpy is close to zero, then NO⁺ clustering occurs; when charge transfer is not favorable but hydride transfer is, then hydride transfer will occur. In terms

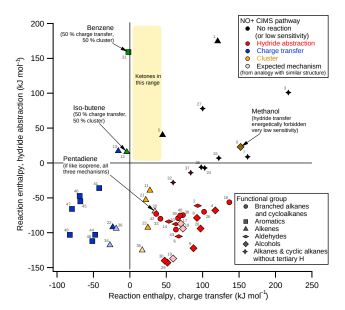


Figure 4. VOC–NO⁺ reaction mechanism dependence on charge transfer and hydride transfer reaction enthalpy. VOC identification is indicated by the small numbers and is listed in Table 1. Hydride abstraction enthalpies for ketones are not known but can be assumed to be positive based on structural considerations (lack of tertiary hydrogen). Ion thermodynamic information is available for several species whose reaction mechanism was not experimentally verified in this or previous work; an expected mechanism was determined by analogy with a VOC of similar structure. 17 is 1-butanol, by analogy with 1-propanol; 18 is 2-methylpropanol, by analogy with 1-propanol; 19 is 2-butanol, by analogy with 2-propanol; 20 is 1,4-pentadiene, by analogy with isoprene; 34 is 4-methyl-2-pentene, by analogy with 2-pentene; 35 is 3-methyl-1-pentene, by analogy with 1-hexene; 36 is 2,3,-dimethyl-1-butene, by analogy with isobutene.

of VOC families, this means that carbonyls participate in two mechanisms: ketones cluster with NO⁺, and aldehydes hydride transfer. Branched alkanes exclusively undergo hydride transfer. Aromatics undergo charge transfer and benzene also clusters; alcohols undergo hydride transfer, and alkenes charge transfer, cluster, or hydride transfer depending on the size of the molecule and the location of the double bond within the molecule.

Although Fig. 4 provides a general way to predict the possible mechanisms for a particular VOC, it provides no information about the distribution of the signal between different mechanisms or the degree of fragmentation. The distribution depends strongly on instrumental conditions, which include E/N settings in the ion–molecule reaction region (by far the most important effect), fragmentation and clustering in the ion optics, presence of impurity ions such as O_2^+ from the converted hollow cathode ion source, and relative humidity (Sect. 3.1.5).

In Fig. 5 the product ion distributions of several VOCs determined in this work are compared to three others using NO⁺. Studies by the University of Leicester used a much

Table 1. VOC species in Fig. 4 and their charge transfer and hydride transfer reaction enthalpies.

ID no.	Species name	Hydride transfer enthalpy (kJ mol ⁻¹)	Charge transfer enthalpy (kJ mol ⁻¹)
0	methanol	22.98	152.05
1	ethene	174.58	120.59
2	acetaldehyde	-61.32	93.20
3	ethane	100.98	217.65
4	ethanol	-68.02	117.31
5	propene	40.17	44.96
6	propanal	-105.32	67.15
7	propane	8.88	161.69
8	n-propanol	-78.72	92.23
9	i-propanol	-122.22	87.41
10	methacrolein	-87.62	63.29
11	1-butene	-39.39	27.59
12	isobutene	15.88	-4.24
13	2-butenes	17.18	-15.82
14	butanal	-84.02	53.64
15	n-butane	6.98	122.14
16	isobutane	-56.32	136.61
17	1-butanol	-87.02	70.04
18	2-methylpropanol	-94.02	72.94
19	2-butanol	-137.02	59.43
20	1,4-pentadiene	-69.32	34.35
21	1-pentene	-53.02	21.80
22	2-pentene	-92.02	-23.54
23	3-methyl-1-butene	-92.52	24.70
24	cyclopentane	-7.22	102.84
25	n-pentane	-6.02	98.02
26	isopentane	-70.02	101.88
27	neo-pentane	77.98	99.95
28	1-pentanol	-94.02	97.05
29	3-methyl-2-butanol	-143.02	51.71
30	3-pentanol	-140.02	47.85
31	benzene	159.08	-1.93
32	cyclohexane	-28.02	59.43
33	methylcyclopentane	-80.02	42.06
34	4-methyl-2-pentene	-117.82	-27.40
35	3-methyl-1-pentene	-125.22	16.98
36	2,3,-dimethyl-1-butene	-94.02	-18.72
37	n-hexane	-94.02 -13.92	83.55
38	2-methylpentane	-74.72	72.94
39	2,3-dimethylbutane	-74.72 -79.22	66.18
40	3-methylpentane	-75.42	69.08
41	toluene	-75.42 -36.02	-42.06
42	methylcyclohexane	-30.02 -73.02	36.27
43	1,2-dimethyl-cyclopentane	-75.02 -95.52	63.29
43 44	ethylbenzene	-93.32 -103.02	-47.66
45	•		
43 46	o-xylene	-55.02 -47.22	-67.92
	m-xylene		-68.88
47 48	p-xylene	-65.92	-79.50 -51.52
48	isopropylbenzene	-111.92	
49 50	3-ethyltoluene	-103.12	-82.41
50	acetone		43.03
51	butanone		24.70
52	2-pentanone		11.19
53	3-pentanone		4.44
54	MVK		37.24

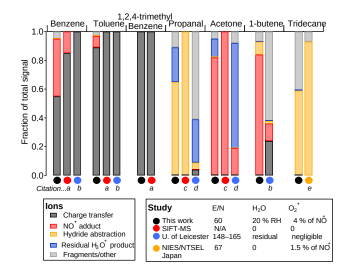


Figure 5. Comparison of product ion distributions between four sets of instrumental and environmental conditions. References: (a) Španěl and Smith (1998b); (b) Blake et al. (2006); (c) Španěl et al. (1997); (d) Wyche et al. (2005); (e) Yamada et al. (2015).

higher E/N ratio in the drift tube, leading to higher fragmentation and lower NO⁺ adduct formation compared to this work (Wyche et al., 2005; Blake et al., 2006). Investigation of higher-mass alkanes by Yamada et al. (2015) used similar E/N but achieved lower contaminant O_2^+ , which is a likely explanation for the higher degree of fragmentation of tridecane seen in this work. In SIFT-MS studies, without an electric field, fragmentation is minimized and preselection of NO⁺ primary ions eliminates contaminant H_3O^+ and O_2^+ and therefore SIFT product ion distributions are generally simpler. These differences highlight the importance of selection of drift tube operating conditions and instrument characterization.

3.1.3 Alkane fragmentation

Small (C₄-C₁₀) branched alkanes cannot be measured by H₃O⁺ CIMS. With NO⁺ CIMS, these VOCs are detectable but generally fragment to produce several ionic fragments that are common to different species. These masses (for example, m/z 57 C₄H₀⁺) are produced by many different compounds and are likely not useful for chemically resolved atmospheric measurements. A few masses (e.g., m/z 71 C₅H $_{11}^{+}$ and m/z 85 C₆H₁₃⁺) are only produced by a few compounds and were therefore targeted for further investigation in ambient air measurements. Conversely, cyclic alkanes fragment very little. Fig. S4 shows the product ion distributions of several representative aliphatic compounds. We note that the major product ions of cyclic alkanes (M-H) are the same with H₃O⁺ and with NO⁺ chemistry. However, the mechanism is different: NO⁺ ionizes by hydride abstraction, while H₃O⁺ ionizes by protonation followed by loss of H2 (Midey et al., 2003). The H₃O⁺ ionization mechanism has a secondary

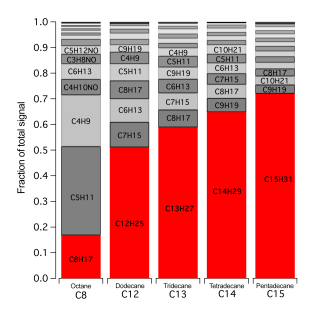


Figure 6. Large $(C_{12}-C_{15})$ n-alkane product ion distribution, using relative humidity of 20 %. The expected largest mass resulting from hydride abstraction (m-1) is highlighted in red. N-octane (C_8) is shown for comparison.

channel consisting of protonation followed by elimination of CH₄ or C_nH_{2n} (Midey et al., 2003). The difference in ionization mechanism is a likely explanation for the lower degree of fragmentation observed using NO⁺ chemistry.

Compared to small (C₈ and smaller) alkanes, large (C₁₂ and higher) n-alkanes show little fragmentation, with at least 50% of the total ion signal accounted for by the expected parent mass (m-1; Fig. 6). Additionally, the degree of fragmentation decreases with increasing carbon chain length. It is quite difficult to measure these compounds with H₃O⁺ CIMS because they fragment extensively and are not detected sensitively (Erickson et al., 2014). NO⁺ CIMS could provide a fast, sensitive, chemically specific measurement of these compounds. It should be mentioned that large n-alkanes (C_{10} and larger) are not measurable with the GC interface. Dodecane $(C_{12}H_{26})$, tridecane $(C_{13}H_{28})$, tetradecane $(C_{14}H_{30})$, and pentadecane (C₁₅H₃₂) were sampled directly with the NO⁺ ToF-CIMS and product ions were identified by correlation with the expected major product ion (m-1). The NO⁺ ToF-CIMS sensitivity to pentadecane was determined using a permeation source (Veres et al., 2010). Contaminant O_2^+ could potentially reduce the measured parent ion ([M-H]⁺) through fragmentation; an alkane measurement corrected for O₂⁺ interference would have higher sensitivity and a simpler product ion distribution (e.g., Yamada et al., 2015).

3.1.4 Instrument response factor for select compounds

A calibration factor was determined for various VOCs by (1) direct calibration, (2) estimation from sensitivity relative to H_3O^+ CIMS, or (3) estimation from correlation with GC-

Table 2. Sensitivities and detection limits of NO⁺ ToF-CIMS for various VOCs. Additional product ions not used to establish sensitivity are listed in italic. The H_3O^+ ToF-CIMS detection limits in the farthest right column are calculated from separate H_3O^+ ToF-CIMS calibrations as described in Yuan et al. (2016).

	ed directly with NO ⁺ CIMS; ambient (20 %) relative humic Ion formula (% of total signal)				Back-	Noise	NO ⁺ sensitivity		NO ⁺ 1 s	H ₃ O ⁺ CIMS 1 s
VOC species	Formula	Mech- anism	(% of total signal)	Exact m/z (Th)	ground cps	scale factor α	ncps/	cps/	detection limit	detection limit
Methanol	CH ₄ ONO ⁺	M+NO ⁺	(12 %)	62.024	0.70	1.23	0.07	0.67	19 ppb	0.397 ppb
	$CH_4OH^{+^a}$ $CH_7O_2^{+^a}$	$M+H^+$ $M+H_3O^+$	(49%) (39%)	33.034 51.044						
Acetonitrile	C ₂ H ₃ NNO ⁺	M+NO ⁺	(48 %)	71.024	0.49	1.33	3.7	24	540 ppt	45 ppt
	$C_2H_3NH^{+a}$	$M+H^+$	(44%)	42.034						
	$C_2H_6NO^{+a}$	$M + H_3 O^+$	(8%)	60.044						
Acetaldehyde	C ₂ H ₃ O ⁺	M-H ⁻	(60 %)	43.018	33	1.33	29	146	268 ppt	195 ppt
	$C_2H_5O_2^+$	M - H + H_2O	(13%)	61.028						
	$C_2H_4OH^{+a}$	$M+H^+$	(11%)	45.034						
	$C_2H_4ONO^+$	$M+NO^+$	(9%)	74.024						
Acetone	C ₃ H ₆ ONO ⁺	$M+NO^+$	(82 %)	88.039	19	1.16	51	376	73 ppt	97 ppt
	$C_3H_6OH^{+a}$	$M+H^+$	(13%)	59.049						
Isoprene	$C_5H_8^+$	M^+	(46 %)	68.062	0.76	1.34	44	286	48 ppt	162 ppt
	$C_5H_8NO^+$	$M+NO^+$	(17%)	98.060						
	$C_5H_7^+$	M- H	(7%)	67.054						
MEK	C ₄ H ₈ ONO ⁺	M+NO ⁺	(86 %)	102.055	3.4	1.33	98	767	23 ppt	45 ppt
	$C_4H_8OH^{+a}$	$M+H^+$	(8%)	73.065						
Benzene ^b	$C_6H_6^+$	M^+	(55%)	78.046	2.3	1.37	57	391	43 ppt	
	C ₆ H ₆ NO ⁺	M+NO ⁺	(40 %)	108.044	2.5	1.72	36	292	74 ppt	96 ppt
	sum				4.8	1.59	93	683	34 ppt	
Toluene	C ₇ H ₈ ⁺	M ⁺	(89 %)	92.062	3.5	1.33	110	825	22 ppt	47 ppt
	$C_7H_8^{\circ}NO^+$	$M+NO^+$	(8%)	122.060						
o-Xylene	$C_8H_{10}^+$	M^+	(94%)	106.078	1.3	1.51	121	972	17 ppt	40 ppt
	$C_8H_{10}^{10}NO^+$	$M+NO^+$	(5%)	136.076						
1,2,4- Trimethylbenzene	C ₉ H ₁₂ ⁺	M ⁺	(100 %)	120.093	0.86	1.75	125	1068	17 ppt	45 ppt
n-Pentadecane ^c	$C_{15}H_{31}^{+}$	M-H ⁻	(72 %)	211.242	2.7	1.83	48	512	46 ppt	-
	$C_9H_{19}^{+31}$	fragment	(3%)	127.148						
	$C_{10}H_{21}^{+}$	fragment	(3%)	141.164						
	$C_8H_{17}^{+21}$	fragment	(3%)	113.132						

EIMS (Table 2). Direct calibrations were performed by mixing a known concentration of a VOC from either a permeation cell (pentadecane) or a calibration gas cylinder (other VOCs) into an ambient humidity ($\sim 20\,\%$) high-purity air dilution stream. Calibration factors estimated from sensitivity relative to H_3O^+ CIMS were calculated using H_3O^+ ToF-CIMS calibration factors and results from laboratory GC-CIMS experiments (Sect. 3.1.1). Calibration factors for H_3O^+ ToF-CIMS were determined in previous work (Yuan et al., 2016). These calibration factors were multiplied by the relative peak areas determined in Sect. 3.1.1 to obtain estimated NO $^+$ ToF-CIMS calibration factors. (An example

chromatogram and calculation is shown in Fig. S5.) Calibration factors estimated from correlation with GC-EIMS were calculated from the slope of NO⁺ ToF-CIMS measurements against GC-EIMS measurements in ambient air (discussed in further detail in Sect. 3.2.2).

In the following discussion we use two metrics of instrument response: cps and normalized cps (ncps). cps is the raw ion count rate of the instrument. Two operations were applied to cps measurements to obtain ncps. First, a duty cycle

Table 2. Continued.

(b) Sensitivity estimated via sensitivity relative to H_3O^+ CIMS; 20 % relative humidity												
		Product ions						NO ⁺ sensitivity		1 .	H ₃ O ⁺	
	H ₃ O ⁺ cps/		Mech-	(% of total	Exact m/z	Relative (NO ⁺ cps /	Back- ground	Noise scale	ncps/	cps/	NO ⁺ 1 s detection	CIMS 1 s detection
VOC species	ppb	Formula	anism	signal)	(Th)	H_3O^+ cps)	cps	factor α	ppb	ppb	limit	limit
Ethanol	119	$C_2H_5O^+ C_2H_7O_2^+$	M-H ⁻ <i>M-H+H</i> ₂ <i>O</i>	(80 %) (15 %)	45.033 63.044	6.2	149	1.37	127	738	105 ppt	1627 ppt
Methyl- cyclohexane	27	$C_7H_{13}^+$ $C_6H_{11}^+$	M-H ⁻ fragment	(98 %) (2 %)	97.101 83.086	17	6.6	1.32	53	448	50 ppt	943 ppt
MVK	539	C ₄ H ₆ ONO ⁺	M+NO ⁺	(100 %)	100.039	0.38	4	1.71	24	202	112 ppt	85 ppt
Pentanone	770	$C_5H_{10}ONO^+ C_5H_{10}OH^{+a}$	$M+NO^+$ $M+H^+$	(83 %) (7 %)	116.071 87.080	1.18	4.4	1.32	97	906	21 ppt	47 ppt
α-Pinene	262	C ₁₀ H ₁₆ ⁺	M ⁺	(59 %)	136.125	0.28	0.39	1.69	7.3	73	233 ppt	67 ppt

92.062 93.070 137.132

(c) Sensitivity estimated via correlation with GC-EIMS; ambient (20%) relative humidity

fragment fragment

		Product	ions			Back-	Noise	NO ⁺ Se	ensitivity	NO^+ 1 s
VOC species	Formula	Mech- anism	(% of total signal)	Exact m/z (Th)	Correlation with GC (R^2)	ground cps	scale factor α	ncps/ ppb	cps/	detection limit
Propanal	C ₃ H ₅ O ⁺	M-H ⁻	(65 %)	57.033	0.928	11	1.40	170	1057	26 ppt
	$C_3H_7O_2^{+^a}$	M - H + H_2O	(17%)	75.044						
	$C_3H_6OH^{+a}$	$M+H^+$	(7%)	59.049						
Methacrolein +	C ₄ H ₅ O ⁺	M-H ⁻	(64 %)	69.033	0.984	4.1	1.37	48	325	60 ppt
crotonaldehyde	$C_4H_6ONO^+$	$M+NO^+$	(16%)	100.039						
	$C_3H_5^+$	fragment	(10%)	41.039						
Isopentane	C ₅ H ₁₁ ⁺	M-H ⁻	(82 %)	71.086	0.888	23	1.36	101	706	49 ppt
	$C_3H_7^+$	fragment	(11%)	43.054						
Methylcyclo- pentane	$C_6H_{11}^+$	M-H ⁻	(99 %)	83.086	0.961	7.4	1.34	154	1225	18 ppt
C ₅ aldehydes	C ₅ H ₉ O ⁺	M-H ⁻	(49 %)	85.065	0.936	9.8	1.38	119	904	28 ppt
	$C_4H_9^+$	fragment	(22%)	57.070						
	$C_5H_{11}O_2^{+a}$	M - H + H_2O	(19%)	103.075						
2- and 3-	$C_6H_{13}^+$	M-H ⁻	(82 %)	85.101	0.978	16	1.34	122	981	30 ppt
methylpentane	$C_3H_7^{+3}$	fragment	(10%)	43.054						
	$C_4H_{+9}^{+}$	fragment	(4%)	57.070						
Hexanal	$C_6H_{11}O^+$	M-H ⁻	(49 %)	99.080	0.945	10	1.47	160	1270	22 ppt
	$C_6H_{13}O_2^{+^a}$	M - H + H_2O	(23%)	117.091						
	$C_5H_{11}^+$	fragment	(15%)	71.086						
Styrene	$C_8H_8^+$	M^+	(100%)	104.062	0.949	0.62	1.47	112	966	15 ppt
Benzaldehyde	C ₇ H ₅ O ⁺	M-H ⁻	(100%)	105.033	0.923	12	1.37	75	621	43 ppt

 $[\]overline{a}$ Product from residual H_3O^+ . b Both product ions can be unambiguously assigned to benzene. We therefore report also the counting statistics and limit of detection for the sum of the two ions. c For technical reasons, pentadecane sensitivity was determined in dry air.

correction was applied (Chernushevich et al., 2001):

$$I_{\rm corr} = {\rm cps} \times \sqrt{\frac{m/z_{\rm reference}}{m/z}},$$
 (1)

where $I_{\rm corr}$ is the duty-cycle-corrected ion count rate and $m/z_{\rm reference}$ is an arbitrary reference mass (in this work $m/z_{\rm reference} \equiv 55$). The duty-cycle correction accounts for differences in ion residence time in the extraction region of

the ToF and eliminates a mass-dependent sensitivity bias. Then, measurements were normalized to the duty-cycle-corrected NO⁺ (primary ion) measurement, which typically has count rates on the order of 10⁶ above that of VOCs:

$$ncps = 10^6 \frac{I_{corr}}{NO_{corr}^+}.$$
 (2)

The normalization removes variability due to fluctuations in the ion source and detector. In calculating limits of detection, we use duty-cycle-uncorrected cps, as this best reflects the fundamental counting statistics of the instrument. In reporting ambient air measurements, we use ncps. The ncps measurement reduces several significant instrumental biases and better reflects VOC abundances in air.

Limits of detection at 1 Hz measurement frequency were calculated by finding the mixing ratio at which the signal-to-noise ratio (S/N) is equal to 3. The calculation can be expressed by (Bertram et al., 2011; Yuan et al., 2016)

$$\frac{S}{N} = 3 = \frac{C_f[X]_{\text{lod}}t}{\alpha \times \sqrt{C_f[X]_{\text{lod}}t + 2Bt}},$$
(3)

where C_f is the instrument response factor, in cps per ppb; $[X]_{lod}$ is the limit-of-detection mixing ratio of species X in ppb; t is the sampling period of 1 s; α is the scaling factor of noise compared to expected Poissonian counting statistics; and B is the background count rate in cps. The scaling factor α is generally greater than 1 because high-resolution peak overlap and fitting algorithms create additional noise (Cubison and Jimenez, 2015). For comparison, H_3O^+ ToF-CIMS limits of detection, using the same ToF-CIMS instrument, are included where available.

Aliphatics and aromatics are generally detected quite sensitively. Aromatics have sub-100 ppt detection limits and are detected slightly more sensitively with NO⁺ CIMS than with $\rm H_3O^+$ CIMS, with NO⁺ detection limits generally about 30 % lower. Aliphatic species are detected with quite low detection limits (less than 50 ppt) and with substantially better sensitivity than $\rm H_3O^+$: the detection limit of methylcyclohexane using NO⁺ is a factor of 27 lower than with $\rm H_3O^+$.

Aldehydes and ketones also have detection limits of around 100 ppt or less, with the exception of acetaldehyde (lod = 355 ppt). The higher detection limit of acetaldehyde is due to a somewhat higher instrumental background and a lower response factor that is consistent with reaction kinetics (Španěl et al., 1997). Methanol has a very high detection limit (19 ppb); this is expected from the anomalously low rate constant of the methanol–NO⁺ reaction (Španěl and Smith, 1997). In contrast, ethanol is detected far more sensitively with NO⁺ than with $\rm H_3O^+$, with a detection limit of 105 ppt (compared to 1600 ppt for $\rm H_3O^+$).

3.1.5 Humidity dependence

Humidity-dependent behaviors of primary ions and selected VOCs (acetaldehyde, acetone, isoprene, 2-butanone, benzene, toluene, o-xylene, and 1,3,5-trimethylbenzene) were

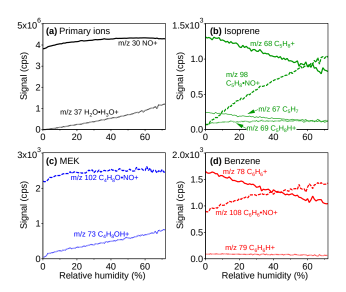


Figure 7. Humidity dependence of primary ions and selected VOCs. (a) NO⁺ and water clusters; (b) isoprene; (c) methyl ethyl ketone (MEK); (d) benzene.

determined by diluting a VOC calibration standard into humidified air to reach approximately 10 ppb mixing ratio, then sampling directly with the NO⁺ ToF-CIMS. Air temperature was 27 °C. Product ion and signal dependencies on humidity for selected primary ions and VOCs are shown in Fig. 7 (additional species are included in Fig. S6). As relative humidity increases, NO⁺ (m/z 30) remains relatively constant, while protonated water and protonated water clusters (especially m/z 37, $H_5O_2^+$) increase. As the abundance of H_3O^+ in the drift tube increases, one might expect to see increased products of VOC reaction with H₃O⁺ with a corresponding decrease in NO⁺ products. Although an increase of H₃O⁺ product is seen for some species (e.g., MEK), it is not universally true. For many species, the major effect is that the NO⁺ adduct product increases relative to other NO⁺ product ions. This effect is especially intense for isoprene, where the isoprene–NO⁺ cluster $(m/z 98, C_5H_8NO^+)$ increases by a factor of 10 from 0 to 70% relative humidity. A similar humidity effect, observed during SIFT measurements of alkenes, has been reported previously by Diskin et al. (2002), who attributed the effect to better stabilization of excited intermediary $(NO^+ \cdot R)^*$ ions by H_2O . A full investigation of this effect is beyond the scope of this paper. In lieu of a complete theoretical understanding of humidity effects, we suggest that an experimental humidity correction could be applied as in Yuan et al. (2016).

3.2 Measurements of urban air

3.2.1 GC-NO⁺ CIMS measurements

Measurement of ambient air using the GC interface allowed us to determine which compounds in ambient air produce

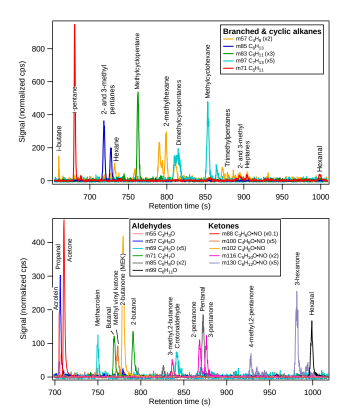


Figure 8. Example GC-CIMS chromatogram of ambient air sample. Masses have been split between two panels for clarity. Top: select masses corresponding to branched and cyclic alkanes. Bottom: select masses corresponding to aldehydes and ketones.

which masses. This is the essential link between laboratory measurements of calibration standards and interpretation of ambient NO⁺ ToF-CIMS measurements. Ambient air from outside the laboratory was sampled from 27 to 30 October 2015 through an inlet 3 m above ground level and directed through 10 m of 1/2 in. diameter Teflon tubing at a flow rate of 17 standard $L \min^{-1}$ (residence time approximately 4 s). The GC interface subsampled this stream. Eluant from the column was directed into the NO⁺ ToF-CIMS as described in Sect. 2.1. The laboratory is in an urban area (Boulder, CO) and the inlet was located near a parking lot and loading dock. Absolute instrument background (including the GC interface) was determined by sampling zero air at the beginning and end of the 3-day measurement period. Instrument performance and stability, retention times of selected compounds, and instrument background were checked at least once per day by sampling a 56-component hydrocarbon calibration standard.

Figure 8 shows several masses from a typical chromatogram. In this chromatogram, it is clear, for instance, that the majority of signal from m/z 83 (C₆H₁₁⁺) can be attributed to one compound (methylcyclopentane). In contrast, m/z 57 (C₄H₉⁺) is produced from many different compounds with comparable intensities. Aldehydes and ketones appear

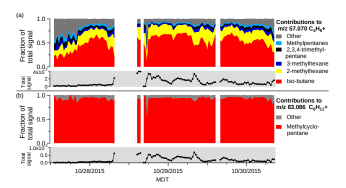


Figure 9. Contributions to two masses based on GC-CIMS measurements of ambient air. "Total signal" is normalized counts per chromatogram. (a) m/z 57 C₄H₉⁺; (b) m/z 83 C₆H₁₁⁺.

to be well separated, as expected from the laboratory experiments. Figure 9 summarizes the contributions of different VOCs to several ions $(m/z\,57,\,C_4H_9^+)$ and $m/z\,83\,\,C_6H_{11}^+)$ during the entire 3-day measurement period. $M/z\,57\,\,(C_4H_9^+)$ has contributions from many different VOCs, and the relative proportions are highly variable. Conversely, $m/z\,83\,\,(C_6H_{11}^+)$ is mostly attributable to methylcyclopentane during the majority of the 3-day measurement period. $M/z\,57\,\,(C_4H_9^+)$ does not provide a useful measurement of alkanes, while $m/z\,83\,\,(C_6H_{11}^+)$ may possibly provide a useful measurement of methylcyclopentane. Corresponding figures for other masses can be found in the supplemental information (Figs. S7–S9). Table 3 summarizes our assessment of key ions.

3.2.2 NO⁺ CIMS vs. GC-EIMS measurement comparison

Measurements using the GC interface do not provide any information about the fast time response capability of the NO⁺ ToF-CIMS. Additionally, not all compounds detectable by NO⁺ CIMS and present in ambient air can be transmitted through the GC interface. Simultaneous GC-EIMS and NO⁺ ToF-CIMS measurements were conducted to investigate fast NO⁺ measurements, determine whether there are any significant interferences to key NO⁺ masses, and explore NO⁺ CIMS response to VOCs not transmittable through the GC interface.

Ambient air was sampled into the laboratory as described in the previous section. The GC-EIMS and the NO⁺ ToF-CIMS were run as separate instruments and subsampled the 17 SLPM flow at the same point. Measurements were taken from 4 to 6 November 2015. The GC-EIMS instrument was operated on a 30 min schedule. GC-EIMS instrument background was determined from zeros taken at the beginning and end of the 3-day measurement period. The 56-component hydrocarbon calibration standard was sampled once per day. The NO⁺ ToF-CIMS measured at 1 Hz frequency. Instru-

Table 3. Assessment of significant product ions investigated by GC-NO⁺ CIMS and parallel GC-EIMS and NO⁺ CIMS measurement of ambient air. Masses in bold can be unambiguously assigned to a single VOC or a structurally related, correlated group of VOCs.

			Correlation with parallel GC-EIMS		
Ion formula (Th)	Exact mass	Assessment from series GC-NO ⁺ ToF-CIMS	R ²	Slope (ppbv ppbv ⁻¹)	
C ₃ H ₅ ⁺	41 039	Several non-correlated species			
$\mathbf{C}_2\mathbf{H}_3\mathbf{O}^+$	43.018	Acetaldehyde	0.942	0.892	
$C_3H_7^+$	43.054	Several non-correlated species			
$C_2H_5O^+$	45.033	Ethanol	0.998		
C ₂ H ₃ O ⁺ C ₃ H ₇ ⁺ C ₂ H ₅ O ⁺ C ₄ H ₆ ⁺	54.046	Propyne ¹			
C ₄ H ₈ ⁺ C ₃ H ₅ O ⁺ C ₄ H ₉ ⁺ C ₃ H ₇ O ⁺	56.062	Several non-correlated species			
$C_3H_5O^+$	57.033	Propanal	0.928		
$C_4H_9^+$	57.070	Several non-correlated species			
$C_3H_7O^+$	59.049	Interference from acetone;			
		If accounted for, sum of C ₃ alcohols			
CH ₄ NO ₂ ⁺	62.024	Methanol, but poor sensitivity	0.904	1.25	
$C_5H_6^+$	66.046	Interference from benzene;			
		if accounted for, cyclopentadiene			
$C_4H_4O^+$	68.026	Furan ²			
$C_5H_8^+$	68.062	Possibly isoprene ³			
$C_4H_5O^+$	69.033	${f Methacrolein}+{f crotonaldehyde}^4$	0.984		
$C_5H_8^+$ $C_4H_5O^+$ $C_5H_9^+$	69.070	Several non-correlated species			
C ₅ H ₁₀ ⁺ C ₄ H ₇ O ⁺ C ₅ H ₁₁ ⁺ C ₄ H ₉ O ⁺ C ₆ H ₆ ⁺	70.078	Possibly the sum of 2-pentenes ³			
$C_4H_7O^+$	71.049	Several non-correlated species			
$C_5H_{11}^+$	71.086	Isopentane	0.888		
$C_4H_9O^+$	73.065	Several non-correlated species			
$C_6H_6^+$	78.046	Benzene ⁵	0.987	0.847	
$C_5H_6O^+$	82.041	Possibly the sum of 2- and 3-methylfuran ³			
$\mathbf{C}_{6}\mathbf{H}_{11}^{+}$ $\mathbf{C}_{5}\mathbf{H}_{9}\mathbf{O}^{+}$	83.086	Methylcyclopentane	0.961		
$\mathbf{C}_{5}\mathbf{H}_{9}\mathbf{O}^{+}$	85.065	Sum of C ₅ aldehydes	0.936		
$C_6H_{13}^+ C_4H_8NO^+$	85.101	Sum of 2- and 3-methylpentane	0.978		
	86.060	Several non-correlated species			
$C_5H_{11}O^+$	87.080	C ₅ alcohols and ethers; significant			
a w wo+	00.020	interference from minor carbonyl product ions	0.050	1.10	
$C_3H_6NO_2^+$ $C_2H_4NO_3^+$	88.039	Acetone	0.978	1.13	
$C_2H_4NO_3^{-1}$	90.019	Possibly acetic acid (chromatography too poor to determine)			
$\mathbf{C}_{7}\mathbf{H}_{8}^{+}$	92.062	Toluene	0.999	0.810	
$C_7H_{13}^+$	97.101	Sum of C ₇ cyclic alkanes	0.917		
$C_6H_{11}O^+$	99.080	Hexanal	0.945		
$C_7H_{15}^+$	99.117	Possibly the sum of 2- and 3-methylhexane,			
С.Н.NO+	100 020	but poor sensitivity	0.050		
$C_4H_6NO_2^+ C_5H_{10}NO^+$	100.039	MVK Describly the sum of C terminal alleges	0.950		
$C_5H_{10}NO$	100.076	Possibly the sum of C ₅ terminal alkenes, but poor sensitivity			
$C_4H_8NO_2^+$	102.055	MEK	0.971	0.843	

ment zeros were taken for a 2 min period once every hour (example time series with zeros in Fig. S10). Calibration gas from a 10-component hydrocarbon standard was sampled for

2 min once every 3 h. At the end of the 3-day measurement period, both instruments were disconnected from the ambient air line and sampled air from inside the laboratory for 1.5 h

Table 3. Continued.

			Correlation with parallel GC-EIMS			
Ion formula (Th)	Exact mass	Assessment from series GC-NO ⁺ ToF-CIMS	\mathbb{R}^2	Slope (ppbv ppbv ^{−1})		
$C_8H_8^+$	104.062	Styrene (vinyl benzene)	0.949			
$\mathbf{C}_{7}\mathbf{H}_{5}^{\circ}\mathbf{O}^{+}$	105.033	Benzaldehyde	0.923			
$C_8H_{10}^+$	106.078	Sum of C ₈ aromatics	0.952	0.746		
$C_6H_6NO^+$	108.044	Benzene ⁵				
$C_8H_8^+$ $C_7H_5O^+$ $C_8H_{10}^+$ $C_6H_6NO^+$ $C_8H_{15}^+$	111.117	Possibly the sum of C ₂ alkyl-substituted cyclohexanes ⁶	0.761			
$C_7H_{13}O^+$	113.096	Heptanal ²				
$C_8H_{17}^+$	113.132	Possibly the sum of methylheptanes,				
1,		but poor sensitivity				
$C_5H_{10}NO_2^+$	116.071	Sum of C ₅ ketones	0.945			
$C_9H_{10}^+$	118.078	Possibly the sum of methylstyrene isomers ³				
$C_9H_{12}^{+}$	120.093	Sum of C ₉ aromatics; scatter possibly due to disparity in response factors	0.600			
$C_8H_{15}O^+$	127.112	Octanal ²				
$C_6H_{12}NO_2^+$ $C_{10}H_{14}^+$	130.086	Possibly the sum of C ₆ ketones ³				
$C_{10}H_{14}^{+}$	134.109	Possibly the sum of C ₁₀ aromatics				
$C_{10}H_{16}^{+7}$	136.125	Monoterpenes plus unknown interference;	0.584			
. 10		possibly adamantane from vehicle exhaust				
$C_7H_{14}NO_2^+$	144.102	Heptanone ²				

¹ Cross-comparison with independent GC-EIMS not possible due to chromatographic quantitation ion overlap with neighboring peaks.

(three GC samples) to investigate the NO⁺ ToF-CIMS response to air with a VOC composition substantially different from urban air.

For all comparisons between the two instruments, the 1 Hz NO⁺ ToF-CIMS measurements were averaged over the 5 min GC-EIMS collection period. The NO⁺ ToF-CIMS was calibrated using air with ambient humidity (approximately 20%) for the 10 species listed in Table 2a, and no further humidity correction was applied. Correlations between independent GC and calibrated CIMS measurements generally show a high correlation coefficient ($R^2 > 0.9$) and slopes close to 1 (examples in Fig. 10a, b). This demonstrates that an adapted NO⁺ CIMS instrument retains sensitive measurement of atmospherically important species such as aromatics that are often targeted using PTR-MS and in addition can detect compounds such as isopentane, sum of 2- and 3methylpentane, methylcyclopentane, and sum of C₇ cyclic alkanes (Fig. 10c-f) that are usually not detected with PTR-MS. Slopes for calibrated VOCs, and correlation coefficients (R^2) for all VOCs investigated, are included in Table 3. The good agreement also indicates that humidity dependence of sensitivity is likely not a severe effect for most species; however, addressing and quantifying this effect should be a priority for future work.

To assess the ability of the NO⁺ ToF-CIMS to separate ketones and aldehydes, we explore measurements of propanal and acetone. The separate measurement of these two species is a good test case because the two peaks are chromatographically well resolved on the GC-EIMS, there are few isomers of C₃H₆O (of which acetone and propanal are likely the only atmospherically relevant species), and independent measurements of these two species are interesting for scientific reasons: aldehydes are generally much more reactive with OH than their ketone isomers and may have significantly different behavior in the atmosphere (Atkinson and Arey, 2003).

A time series of propanal and acetone is shown in Fig. 11a. The two compounds have clearly different behavior in the atmosphere: there is fast (seconds to minutes), high variability in the acetone measurement that is not seen in the propanal measurement, and the longer-term (\sim hours) variability of acetone and propanal is not the same. The fast, high spikes in acetone may come from local sources such as exhaust from chemistry labs in the building. The acetone comparison between the GC-EIMS and the NO⁺ ToF-CIMS has a slope of 1.13, a correlation coefficient R^2 of 0.978, and negligible

² Cross-comparison with independent GC-EIMS not possible due to EIMS quadrupole selected-ion scan window restrictions. ³ Concentrations too low in ambient air to determine. ⁴ Winter urban air sampled was likely influenced by local domestic biomass burning; crotonaldehyde may be a smaller fraction of signal in other environments. ⁵ Benzene correlation using sum of m108 $C_6H_6NO^+$ and m78 $C_6H_6^+$. ⁶ With exclusion of single outlier, $R^2 = 0.831$.

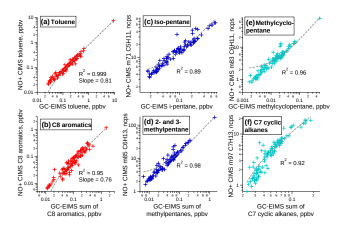


Figure 10. Correlations between VOCs measured with GC-EIMS and NO⁺ ToF-CIMS in ambient air. The 1 Hz NO⁺ ToF-CIMS measurement is averaged to the 5 min GC collection period. Orthogonal least-squares linear best fits (ODR best fit) are shown with dashed lines. The lines appear curved due to log-scale axes. For several compounds (e.g., methylcyclopentane, 2- and 3-methylpentane), the single high outlier pulls the best fit slightly away from the data points at low mixing ratios. (a) Toluene. (b) C₈ aromatics: sum of ethylbenzene, o-xylene, m-xylene, and p-xylene. (c) Isopentane. (d) Sum of 2-methylpentane and 3-methylpentane. (e) Methylcyclopentane. (f) C₇ cyclic alkanes: sum of methylcyclopentane, ethylcyclopentane, and dimethylcyclopentane.

offset. The comparison between the GC and CIMS propanal measurements has an R^2 of 0.928 (Fig. 11b, c).

Several episodes occurred with elevated high-mass n-alkane masses (m/z 169 $C_{12}H_{25}^+$, dodecane; m/z 183 $C_{13}H_{27}^+$, tridecane; m/z 197 $C_{14}H_{29}^+$, tetradecane; m/z 211 $C_{15}H_{31}^+$, pentadecane). Two examples are shown in Fig. 12. The episodes show high temporal and compositional variability. The inlet was downwind from a parking lot and next to a loading dock and electric power generator for the building, and it is likely that the elevated C_{12} – C_{15} alkanes are from any or all of these sources. An ambient air measurement of these species is particularly interesting because they have been implicated in efficient secondary organic aerosol production from diesel fuel exhaust (Gentner et al., 2012).

4 Summary and conclusions

In summary, an ${\rm H_3O^+}$ ToF-CIMS (PTR-MS) instrument was easily and inexpensively converted into an NO⁺ CIMS by replacing the reagent source gas and modifying the ion source and drift tube voltages. The usefulness of NO⁺ CIMS for atmospheric VOC measurement was then evaluated by (1) using a GC interface to determine product ion distributions for nearly 100 VOCs and compare the sensitivity and simplicity of spectra to ${\rm H_3O^+}$ CIMS; (2) measuring ambient air with a GC interface, to map product ions to their VOC precursors and determine which ions may be useful for chem-

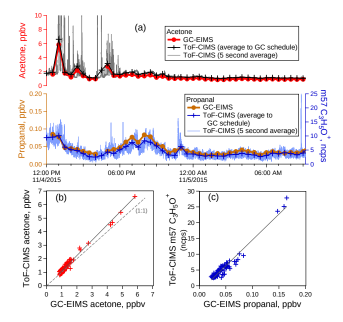


Figure 11. (a) Time series of acetone and propanal measurements from NO⁺ ToF-CIMS and GC-EIMS in ambient air. Measurements shown include the GC-EIMS measurement (5 min sample every 30 min, circle markers), the NO⁺ ToF-CIMS measurement averaged over the 5 min GC sampling period (cross markers), and the NO⁺ ToF-CIMS measurement averaged to a 5 s running mean. (b) Correlation between NO⁺ ToF-CIMS and GC-EIMS measurement of acetone. (c) Correlation between NO⁺ ToF-CIMS and GC-EIMS measurement of propanal.

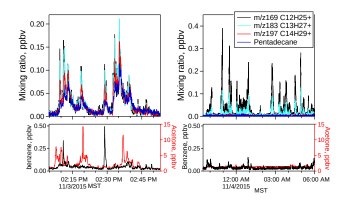


Figure 12. Episodes with elevated high-mass alkane masses in ambient air. Mixing ratios for m/z 169 $C_{12}H_{25}^+$ (dodecane), m/z 183 $C_{13}H_{27}^+$ (tridecane), and m/z 197 $C_{14}H_{29}^+$ (tetradecane) are shown in approximate ppbv, assuming the same instrument calibration factor as pentadecane. Additional VOC species (benzene, acetone) are shown in the bottom panels for context.

ically specific measurement; and (3) measuring ambient air directly, to evaluate chemical specificity and investigate fast (1 Hz) time measurement of new compounds. Additionally, the NO^+ CIMS response to C_{12} – C_{15} n-alkanes and to vari-

able humidity was determined in some detail. Further work is needed to better understand the humidity dependence.

NO⁺ CIMS is a valuable technique for atmospheric measurement because it can separate small carbonyl isomers, it can provide fast and chemically specific measurement of cyclic and a few important branched alkanes (notably, isopentane and methylpentane) that cannot be detected by PTR-MS, it can measure alkyl-substituted aromatics with less fragmentation than H₃O⁺ CIMS, and it can detect larger (C₁₂-C₁₅) alkanes. With NO⁺ CIMS significant fragmentation of most small alkanes does occur, making them difficult to measure quantitatively. There are also interferences on many alcohols (with the exception of ethanol) and butanal. Additionally, it is worth considering that VOC · NO⁺ cluster formation moves certain species into a higher mass range. This may be a drawback because the number of possible isobaric compounds increases with mass, and it may be more difficult for high-resolution peak-fitting algorithms to separate species of interest from isobaric interferences (example in Fig. S11). Finally, because there are three different ionization mechanisms (hydride transfer, charge transfer, and NO⁺ adduct formation), it may be difficult to determine which VOC precursors correspond to particular ions. NO⁺ CIMS may be an extremely useful supplementary approach for specific applications such as studying secondary organic aerosol precursors in vehicle exhaust, investigating emissions from oil and natural gas extraction, identifying additional species in complex emissions such as biomass burning, measuring emissions of oxygenated consumer products and solvents in urban areas, and investigating photochemistry of biogenic VOCs.

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