Calibration of Hydroxyacetonitrile (HOCH₂CN) and Methyl 1

isocyanate (CH₃NCO) Isomers using I⁻ Chemical Ionization 2

- **Mass Spectrometry (CIMS)** 3
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Abstract. The toxic reduced nitrogen compound methyl isocyanate (CH₃NCO, MIC) has been 17 reported present in wildfire and bio-mass burning emissions, agricultural fumigation plumes, and 18 indoor air. Its isomer, hydroxyacetonitrile (HOCH2CN, glycolonitrile, HAN) has not been 19 observed in the Earth's atmosphere to date. In this study, absolute sensitivity calibrations for these 20 isomers using two I⁻ chemical ionization mass spectrometry (I-CIMS) instruments, time-of-flight 21 (ToF) and quadrupole (Quad) instruments, commonly used in laboratory and field measurements, 22 were performed, for the first time, over a range of ion-molecule reactor temperature (10-40°C) 23 and I(H2O)^{-/I-} ratio (0.01-1). This study demonstrates that I-CIMS, under typical operating 24 conditions, is not sensitive to MIC with limits of detection (LOD) of >860 and >570 ppb for the 25 ToF and Quad I-CIMS instruments, respectively. Both I-CIMS instruments are, however, highly 26 sensitive to the HAN isomer with 0.3 and 3 ppt LODs for the ToF-CIMS and Quad-CIMS 27 instruments, respectively. The present results show that several recent field studies using I-CIMS 28 instrument detection have mis-attributed the C2H3NO to MIC. This study proposes that HAN, 29 rather than MIC, was most likely the C₂H₃NO isomer observed in those field studies, although the 30 31 source chemistry for HAN remains uncharacterized. This study demonstrates the importance of applying absolute calibration standards in the identification and quantification of isomeric 32 33 compounds.

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40 1 Introduction

41 The identification and quantitative measurement of atmospheric trace species are critical elements in the development 42 of air quality models and the establishment of environmental policies. Methyl isocyanate (CH₃NCO, MIC) and hydroxyacetonitrile (HOCH2CN, glycolonitrile, HAN) (Wolfsie, 1960) are toxic reduced nitrogen isomers present in 43 44 an extreme range of environments. MIC has been reported in wildfire and bio-mass burning, e.g., Koss et al. (2018), and building fire emissions (Blomqvist et al., 2003), agricultural fumigation plumes (Woodrow et al., 2014), and 45 indoor air (Bekki et al., 2018), particularly where there is cigarette smoke (Moldoveanu, 2010), cooking emissions, 46 47 and chlorine-disinfectant use. HAN has been observed spectroscopically in interstellar space (see Zhao et al. (2021) 48 and references within), while measurements in the Earth's atmosphere have not been reported to date. The atmospheric degradation of MIC leads to the formation of isocyanic acid (HNCO) (Papanastasiou et al., 2020), another 49 50 toxic reduced nitrogen compound. The atmospheric chemistry of the HAN isomer is not presently characterized.

51 The implementation of I⁻ chemical ionization mass spectrometry (I-CIMS) instruments, in particular high-52 resolution time-of-flight mass spectrometer instruments (I-CIMS ToF-MS), has enabled high sensitivity laboratory 53 and ground-based field measurements as well as high spatial and temporal resolution in airborne field studies. 54 However, a general limitation of CIMS measurements is the inability to distinguish isomers, e.g., MIC and HAN 55 C2H3NO isomers, without using pre-separation techniques such as gas chromatography. Field measurements of MIC have been reported in outdoor (Priestley et al., 2018) and indoor (Mattila et al., 2020a; Mattila et al., 2020b; Wang et 56 57 al., 2022) environments using (I-CIMS) instruments, with sensitivity calibration estimated using voltage-scan techniques (Mattila et al., 2020b; Wang et al., 2022) and analogy to the sensitivity of other VOC species (Priestley et 58 59 al., 2018).

60 The primary objective of this work was to establish sample handling and I-CIMS sensitivity calibration protocols 61 for the C2H3NO MIC and HAN isomers. Absolute pressure and FTIR measurements were used to establish MIC 62 standards, while diffusion and infusion methods, coupled with total reactive nitrogen (Nr) methods, were used for the quantification of HAN standards. Calibration measurements are presented for I-CIMS time-of-flight (ToF-CIMS) and 63 64 quadrupole CIMS (Quad-CIMS) mass spectrometry instruments over a range of I(H₂O)⁻/I⁻ ratios and ion-molecule 65 reactor (IMR) temperatures. The results from this work can be applied to similar instruments used in laboratory and 66 field studies, although instrument sensitivities may vary somewhat depending on instrument configuration and operating conditions. This work demonstrates that the recent field measurements reported by Priestley et al. (2018), 67 68 Mattila et al. (2020a), and Wang et al. (2022) have mis-attributed their measured I-CIMS signal, which was assigned 69 to MIC, when it was most likely due to HAN.

70 2 Experimental Methods

A primary goal of this work was the development of measurement protocols for hydroxyacetontrile (HOCH₂CN, HAN) and methyl isocyanate (CH₃NCO, MIC) using iodide chemical ionization mass spectrometry (I-CIMS) with a time-of-flight mass spectrometer (ToF-CIMS) (Lee et al., 2014) and quadrupole mass spectrometer (Quad-CIMS) (Neuman et al., 2000) instruments. We have focused on calibration measurements for ToF-CIMS and Quad-CIMS instruments for which calibrations for MIC and HAN have not been performed previously. An emphasis in this work was placed on MIC and HAN sample handling and quantification, i.e., knowing what the sample concentration is at the instrument inlet.

78 MIC and HAN samples for calibration measurements were prepared using dilute gas mixtures prepared using 79 absolute pressure measurements, diffusion sources, and infusion sources that are described in detail below. Absolute 80 concentrations were determined using FTIR spectroscopy (MIC) and a total reactive nitrogen (Nr) instrument (HAN). 81 Calibration measurements were performed with ToF- and Quad-CIMS instruments to evaluate differences in 82 sensitivity for instruments with different configurations, e.g., ion-molecule reactor (IMR) geometries, ionization 83 sources, and ion focusing optics. In addition, sensitivities were determined for a range of IMR temperature (10-40°C) and I(H2O)'/I' ratios, commonly used in laboratory and field studies, which are known to affect I-CIMS sensitivity 84 (see Robinson et al. (2022)). The MIC and HAN calibration methods, Nr instrument and methods, and ToF- and 85 86 Quad-CIMS instruments are described below.

87 2.1 Methyl isocyanate source and calibration

88 Methyl isocyanate (MIC) is a stable volatile liquid at room temperature with a vapor pressure of ~467 hPa (1 hPa = 89 0.75 Torr) at 25°C. For the present study, a MIC sample was obtained commercially in pure form. Dilute gas-phase 90 methyl isocyanate (MIC) samples were prepared manometrically at a total pressure of 1066 hPa in a 12 L Pyrex bulb 91 with He bath gas. The MIC mixing ratio was ~3.5%. The MIC mixing ratio was also determined by Fourier transform 92 infrared spectroscopy using the absorption cross section data measured in Papanastasiou et al. (2020) from this 93 laboratory. Dilute high-pressure gas standards were prepared by diluting glass bulb standards into aluminum cylinders 94 and filled with zero air. Methyl isocyanate bulb and aluminum cylinder standards were quantitatively added to the 95 calibrated zero air flow sampled by the I-CIMS instruments.

96 2.2 Hydroxyacetonitrile (HAN) source and calibration methods

HAN is a stable semi-volatile compound, vapor pressure of 1.33 hPa at 336 K (NIOSH), that is available commercially
as a 70 wt% mixture with H₂O. Over the course of this study, liquid-to-gas-phase diffusion and infusion methods
were applied for the delivery of gas-phase HAN to the N_r calibration and I-CIMS instruments.

100 The diffusion source is described in detail elsewhere (Roberts et al., 2010; Williams et al., 2000). Basically, a capillary diffusion cell regulated vapor from the liquid sample into the gas stream feeding into the Nr and I-CIMS 101 102 instruments. The concentration of source compound is regulated by varying the total gas flow, i.e., dilution factor. 103 This source method requires an independent determination of the compound concentration in the gas stream. The 104 infusion method has been used previously in our laboratory and is described in detail elsewhere (Bernard et al., 2017; 105 Bernard et al., 2018). Basically, a syringe pump is used to deliver a constant liquid flow (0.01-0.3 µL min⁻¹ in our 106 experiments) of a HAN sample into the gas stream feeding into the N_r and I-CIMS instruments. The region of the 107 liquid-gas interface was heated (40 to 60°C) to ensure uniform volatilization and minimize potential sample condensation. Measurements were performed using the commercially available stock HAN solution and a HAN 108 109 sample diluted in acetone, prepared off-line. The mixing ratio of HAN in the gas flow can in principle be calculated 110 using the calibrated gas and liquid flow rates, the density of the compound, and its liquid mixing ratio, although in this work, the HAN infusion source was calibrated using a total reactive nitrogen, Nr, measurement. 111

112 2.3 Total reactive nitrogen, Nr, measurement

113 The infusion source gas-phase concentration of HAN was determined by a total reactive nitrogen, Nr, measurement.

114 Total reactive nitrogen is defined as all reduced and oxidized nitrogen-containing compounds with the exception of

115 N2 and N2O. The Nr instrument has been demonstrated for both gas-phase and particle-phase Nr, and is described in 116 detail elsewhere (Stockwell et al., 2018). In this study, the nitrogen containing compounds in a gas-phase sample are 117 first catalytically converted on a 750°C Pt catalyst to NO and NO2. The NO2 is subsequently converted to NO on a 118 molybdenum oxide catalyst at 350°C. The NO then reacts with an excess of O₃ to form NO₂, which is detected by 119 chemiluminescence (Williams et al., 1998). The Nr instrument was calibrated using commercial dilute mixtures of 120 NO (5.18 ppm) and HCN (9.5 ppm) in a N2 bath gas. HCN and NO calibrations performed over the course of the 121 study agreed to within 3%. The total flow through the Nr instrument was set to 1.048 sLPM (standard liter per minute) 122 and the total zero air flow from the infusion source was 2.148 sLPM. The excess flow passed through an exhaust line. 123 Considering uncertainties in the standards and flow rates, the 2σ uncertainty in the Nr calibration of the HAN source 124 was estimated to be 15%. The HAN infusion source was calibrated by measuring the Nr concentration as a function 125 of the liquid injection flow rate. Over the course of the study, multiple calibration experiments were performed using 126 independently prepared HAN samples.

127 2.4 Iodide Chemical Ionization Mass Spectrometry (I-CIMS)

128 I-CIMS has the ability to measure sub-part per trillion (ppt) gas-phase concentrations of organic acids, halogens, 129 oxidized organic compounds, and N₂O₅ at up to 10 Hz resolution (Huey, 2007; Neuman et al., 2000; Veres et al., 130 2020). This relatively soft ionization technique usually yields an I⁻ cluster ion with the intact analyte molecule. I-131 CIMS is highly selective, with the sensitivity to an analyte dependent on the binding enthalpy of the compound with 132 I. Time-of-flight mass spectrometers typically contain ion focusing quadrupoles that can also impart changes to the 133 sensitivity of analytes due to mass-dependent ion transmission and collision-induced dissociation (Robinson et al., 134 2022). The I-CIMS instruments used in the present study have been described in detail previously and only pertinent 135 details are described below.

136 2.4.1 Time-of-Flight Chemical Ionization Mass Spectrometer (ToF-CIMS)

137 The ToF-CIMS is setup with a pressure-controlled inlet, ion molecule reactor (IMR), small segmented quadrupole 138 (SSQ), big segmented quadrupole (BSQ), and time-of-flight (ToF) mass analyzer (Veres et al., 2020). A Kr lamp 139 provided vacuum ultraviolet radiation at 124 and 117 nm that photoionized CH₃I to produce I⁻ in the IMR 140 (Breitenlechner et al., 2022; Ji et al., 2020). Analytes (A) react with I⁻ or I(H₂O)⁻ to form adducts:

141	$I^- + A$	$\leftrightarrow [\text{I-A}]^{-}$	(1)
142	$[I-H_2O]^- + A$	\leftrightarrow [I-A] ⁻ + H ₂ O	(2)

with I-C₂H₃NO⁻ detected at m/z 183.9265. Key instrument conditions that impact the sensitivity for an analyte are the
IMR temperature and I(H₂O)⁻/I⁻ ratio (Lee et al., 2014; Robinson et al., 2022; Veres et al., 2020). In this study, the
IMR was temperature controlled between 20 and 40°C and the I(H₂O)⁻/I⁻ ratio was dynamically controlled at 0.01,
i.e., dry, and over the range 0.4 to 0.6, which represents a typical range of operating conditions of laboratory and field
instruments. The ToF-CIMS inlet was pressure controlled at 130 hPa and the flow into the inlet was controlled to 6
sLPM. The IMR was pressure controlled to 44 hPa (Zhang and Zhang, 2021). The pressure in the SSQ and BSQ
were 1.7 and 0.013 hPa, respectively. The sum of I⁻ + I(H₂O)⁻ was typically 5MHz in these experiments.

150 2.4.2 Quadrupole Chemical Ionization Mass Spectrometer (Quad-CIMS)

151 The Quad-CIMS instrument setup consisted of a critical orifice inlet (600 μ m dia.) combined with an IMR, ion

focusing lenses, and a quadrupole mass analyzer (Neuman et al., 2000). I⁻ ions were generated by passing a 0.1%
 CH₃I in N₂ mixture through a ²¹⁰Po radioactive source. The IMR was temperature controlled between 10 and 30°C.

The $I(H_2O)/I$ ratio was adjusted between 0.01–1.0 by controlling the humidity of the gas flow into the CIMS, which

ranged from 4.2 to 4.5 sLPM. This system did not have an intermediate pressure zone between the IMR and mass

spectrometer chamber, common to many I-CIMS instruments, that would serve as a collisional-dissociation chamber.

157 The pressure in the IMR was maintained at 37 hPa by varying the pumping speed. The mass resolution of the Quad-

158 CIMS instrument is ~200 and that of the ToF-CIMS is ~5000. The sum of $I^- + I(H_2O)^-$ was typically 500 kHz in these

159 experiments.

160 2.5 Materials

Synthetic air (zero grade), N₂ (UHP, 99.999%), He (UHP, 99.999%) gases and CH₃I (99%) and acetone (99%) were
used as provided. Standard dilute mixtures of NO (5.18 ppm in N₂) and HCN (9.5 ppm in N₂) were obtained
commercially. Hydroxyacetonitrile (HAN, ~70% in H₂O, CAS RN: 107-16-4) and methyl isocyanate (MIC, 97+%,
CAS RN: 624-83-9) samples were obtained commercially. The MIC sample contained a <3% trimethylchlorosilane
(CAS RN: 75-77-4) inhibitor. The HAN and MIC samples were degassed in freeze—pump—thaw cycles. Samples
were stored in a chemical refrigerator in vacuum sealed Pyrex reservoirs prior to use.

A dilute (3.48%) gas mixture of MIC in He was prepared manometrically in a 12 L Pyrex bulb. Fourier transform
 infrared spectroscopy (FTIR) measurements (1 cm⁻¹ resolution, 425 cm pathlength) using a previously determined
 absorption spectrum from our laboratory (Papanastasiou et al., 2020) confirmed the mixing ratio, to within 10% of the

170 manometric preparation. Standard dilute solutions of HAN in acetone were prepared volumetrically using 1.0–2.0 μL

171 of the commercial HAN solution and 5.0 or 10.0 mL of acetone. The standard solutions were stored in a chemical

- 172 refrigerator in vacuum sealed Pyrex reservoirs. Samples used in the calibration infusion experiments were extracted
- 173 from the standard solution with a gas-tight 10 or 100 μ L syringe.

174 3. Results and Discussion

175 3.1 Methyl isocyanate (CH₃NCO, MIC)

176 ToF-CIMS and Q-CIMS measurements with MIC mixing ratios of up to 860 ppb yielded no measurable signal for the

177 I-MIC⁻ or I(H₂O)-MIC⁻ adducts above the background (1 σ background noise level of 3 ncps). Measurements were

178 performed over the range of CIMS conditions described in the experimental section. The sensitivity upper-limits

- 180 per mixing ratio unit increase of the compound). Our measurements indicate that MIC mixing ratios of greater than
- 181 l ppm may be required to generate a detectable I-CIMS signal, if MIC can be detected by I-CIMS at all.

Table 1. ToF-CIMS and Quad-CIMS instrument sensitivity (S) and limits of detection (LOD) measured in this work
 for hydroxyacetonitrile (HOCH₂CN, HAN) and methyl isocyanate (CH₃NCO, MIC) for typical field operating

184 conditions with an IMR temperature of 30°C, I(H₂O)/I⁻ ratio of 0.55, and 1 s integration. See Fig. 3 for sensitivity

185 dependence on operating conditions.

	CH ₃ NCO (MIC)		HOCH ₂ CN (HAN)	
Instrument	S	LOD	S	LOD

	(ncps ppb ⁻¹)	(ppb)	(ncps ppt ⁻¹)	(ppt)
ToF-CIMS	< 0.009	>860	19.6 ± 0.5	0.3
Quad-CIMS	< 0.044	>570	25.8 ± 0.7	3

186 The ToF- and Quad-CIMS instruments used in the present study were found to be insensitive to the detection of CH₃NCO (MIC). The lack of MIC sensitivity for both instruments suggests that the I⁻ cluster with MIC is not 187 thermodynamically stable in the IMR. The ion focusing or ion optics in the ToF-CIMS and the Quad-CIMS are, 188 189 however, quite different. The ToF-CIMS SSQ was set to 1.7 hPa, so collisions occur in this region, resulting in some 190 collisional dissociation. The BSQ focuses ions and can also result in fragmentation of I clusters. The ion optics of 191 the Quad-CIMS uses only static electric fields in a low-pressure region ($<1.3 \times 10^{-4}$ hPa) and does not have sufficient 192 frequency to dissociate I' clusters. Given the lack of sensitivity of both ToF- and Quad-CIMS instruments to MIC, it 193 is unlikely that other common I-CIMS instrument configurations have the sensitivity to detect MIC at atmospherically 194 relevant mixing ratios.

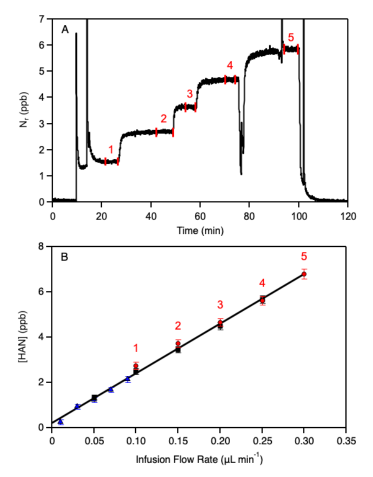
195 3.2 Hydroxyacetonitrile (HOCH₂CN, HAN)

The diffusion source, with the commercial 70% HAN/H₂O solution, proved to be an unsuccessful HAN delivery method. This method produced high gas-phase concentrations of an HCN impurity as identified by I-CIMS. Although HCN was a minor sample impurity, <1%, its high vapor pressure (862 hPa at 295 K (Perry and Porter, 1926)) and low Henry's law coefficient (~9 M Atm⁻¹ (Burkholder et al., 2019)), resulted in a [HCN]/[HAN] gas-phase mixing ratio of greater than 1000, i.e., HAN was not detectable by I-CIMS using this source. The high gas-phase HCN concentration, using the diffusion method, precluded quantitative calibration of HAN concentration using the total reactive nitrogen calibration method.

203 The infusion source did not produce detectable gas-phase HCN above the detection limit of the mass spectrometers, 204 i.e., the HCN impurity level was much less than 1%. Therefore, HCN did not influence the absolute HAN calibration 205 determination using the Nr instrument. The region around the infusion source was heated to 50°C when using the 206 commercially available HAN stock solution. Although this source worked, it didn't provide a stable HAN signal to 207 within 10%. This source, using a dilute HAN/acetone mixture with the region of the infusion source heated to 45°C, i.e., slightly below the boiling point of acetone, yielded more stable HAN signals with variations of a few percent. 208 209 Measurements performed at temperatures greater than 45°C yielded reasonable results, but the signal was less stable 210 due most likely to boiling of the acetone solvent. 211 Calibration of hydroxyacetonitrile (HAN) solutions by total reactive nitrogen (Nr) is shown in Fig. 1. The infusion

method produces a stable source of HAN with a high signal-to-noise ratio for a typical calibration experiment. The total HAN concentration was set by adjusting the injection flow rate. Individual solutions were calibrated multiple times, as shown in **Fig. 1**. The 2σ uncertainty of the fit precision of [HAN] vs. infusion flow rate was determined to

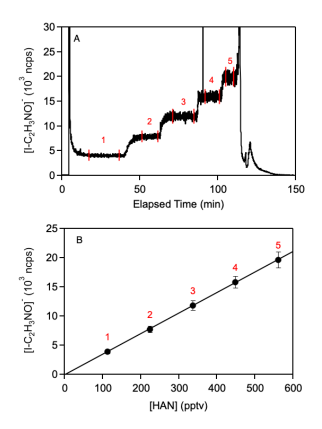
 $\label{eq:215} \be 4\%. \ The small positive intercept may be due to minor unidentified N_r impurities.$



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Figure 1: Calibration of the hydroxyacetonitrile (HOCH₂CN, HAN) infusion source at 23°C using the total nitrogen, N_r, instrument
 (see text for details of the N_r instrument). (a) Background-corrected time series of a representative calibration experiment. The
 data within the numbered vertical lines were averaged and correspond to the numbered points in panel b. (b) Calibration of HAN
 concentration, taking the Nr measured concentration, example in panel A, to be equal to the HAN concentration, as a function of
 infusion source flow rate. Different symbols represent independent calibration experiments. The line is an unweighted linear least squares fit of all the data. Error bars represent 2σ measurement precision.

Calibrations of the HAN signal on the ToF-CIMS and Quad-CIMS instruments were made by varying the infusion
flow rate, see representative data for the Quad-CIMS in Fig. 2. The obtained ToF- and Quad-CIMS sensitivity for
HAN given in Table 1 was obtained using the infusion method with the dilute HAN/acetone samples. For typical
field operating conditions with an IMR temperature of 30°C, I(H₂O)/I⁻ ratio of 0.55, and 1 s integration the HAN
sensitivity of the ToF I-CIMS was determined to be ~20 ncps ppt⁻¹.



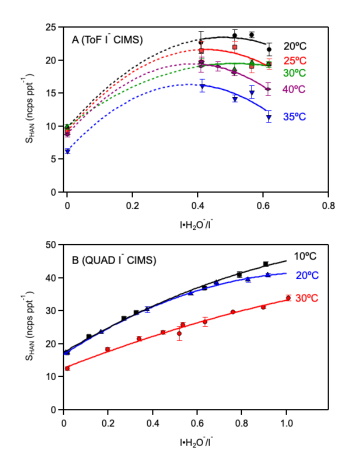
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Figure 2: Calibration of the Quad-CIMS instrument using the hydroxyacetonitrile (HOCH₂CN, HAN) infusion source at 23°C (see text for details of the Quad-CIMS instrument). (a) Background-corrected time series of a representative Quad-CIMS calibration experiment (IMR temperature = 20° C, $I(H_2O)/I^{-} = 0.57$). The data within the numbered vertical lines were averaged and correspond to the numbered points in panel B. (b) Quad-CIMS HAN calibration curve where the HAN concentration was determined from the N_r calibration of the infusion source, e.g., see Fig. 1. The line is an unweighted linear least-squares fit of the data. Error bars represent 2 σ measurement precision.

235 The HAN concentration from the infusion source at the instrument inlet was varied, for each IMR temperature and 236 I(H₂O)^{-/I} ratio, to determine the HAN sensitivity temperature and I(H₂O)^{-/I} dependence shown in Fig. 3. The ToF-237 CIMS and Quad-CIMS instruments are both highly sensitive to HAN, but displayed slightly different temperature and 238 I(H₂O)^{-/I⁻} dependencies. As the temperature increased, the HAN sensitivity decreased consistent with the [I-HAN]⁻ 239 adduct being less stable at higher temperatures. For the ToF-CIMS instrument, the HAN sensitivity decreased a factor 240 of ~1.5-2 between 20 and 40°C at the highest I(H2O)'/I' ratio included in this study. For the Quad-CIMS instrument, 241 a ~25% decrease in HAN sensitivity was observed at a I(H2O)^{-/1} ratio of 1 when increasing the IMR temperature from 242 10 to 30°C. The I(H₂O)⁻/I⁻ dependency for HAN sensitivity was fit reasonably well with a quadratic dependence on 243 $I(H_2O)^{-1}$: A + B($I(H_2O)^{-1}$) - C($I(H_2O)^{-1}$)², where the constant term represents HAN clustering with I⁺, reaction 1; 244 the linear term represents HAN undergoing a ligand switching reaction with I(H₂O)⁻, reaction 2; and the quadratic 245 term may represent a lack of reactivity of higher-order H2O clusters, I(H2O)n where n >1 or a shift in the

246 $(\text{I-HAN})^{-} + \text{H}_2\text{O} \longleftrightarrow (\text{I-H}_2\text{O})^{-} + \text{HAN}$

equilibrium. 247



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249 Figure 3: Hydroxyacetonitrile (HOCH2CN, HAN) calibration factors for the ToF and QUAD mass spectrometers used in this 250 work as a function of the $I(H_2O)/\Gamma$ ratio and ion-molecule reactor, IMR, temperature. Error bars represent 2σ precision of the 251 linear calibration fits. Lines are empirical polynomial fits to guide the eye. (a) HAN calibration factors measured for the ToF-252 CIMS. Symbols represent IMR temperatures of 20°C (black circles), 25°C (red squares), 30°C (green triangles), 35°C (blue upside-

down triangles), and 40°C (purple diamonds). (b) Quad-CIMS HAN calibration data. Symbols represent IMR temperatures of: 10° C (black squares), 20° C (blue triangles), 30° C (red circles). 253

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255 There is a plausible explanation for the differences in the Quad-CIMS and ToF-CIMS instrument I(H₂O)⁻/I⁻ sensitivity dependence. First, there are differences in these instruments in the ion focusing downstream of the IMR. 256 257 The ToF-CIMS contains an SSQ at 1.73 hPa that imparts an electric field and acts as a collisional dissociation chamber. 258 The subsequent BSQ also imparts an electric field that can dissociate weakly bound I⁻ clusters. The I(H₂O)/I⁻ is 259 dynamically controlled in the ToF-CIMS by what is observed at the detector. Therefore, the I(H₂O)⁻ (and I(H₂O)⁻) 260 ion counts are not representative of the I(H2O)⁻/I⁻ ratio in the IMR, i.e., the I(H2O)⁻/I⁻ ratio in the IMR is being under-261 estimated, although we were unable to quantify the dependence. The Quad-CIMS, which has ion lenses at lowpressure downstream of the IMR, 1.3-0.13) × 10⁻³ hPa, should have only minor collisional dissociation, if any. 262 263 Therefore, the I(H₂O)⁻/I⁻ ratio measured by the Quad-CIMS is, most likely, close to the actual I(H₂O)⁻/I⁻ in the IMR. In conclusion, it is recommended that instruments used to quantify HAN be calibrated under actual operating 264 conditions. The protocols presented in this work can be used for the calibration of HAN. 265

266 4 Conclusions

267 In this study, protocols for determining I-CIMS instrument sensitivity for CH₃NCO (MIC) and HOCH₂CN (HAN), 268 two stable toxic C2H3NO isomers (Panda et al., 2023), were developed. Calibration of ToF-CIMS and Quad-CIMS 269 instruments for HAN were performed over a range of instrument conditions commonly used in laboratory and field 270 experiments, including: ion molecule reactor (IMR) temperature (10-40°C) and I(H2O)^{-/I-} ratios between 0.1 and 1. 271 Both I-CIMS instruments were found to be highly sensitive to HAN with 0.3 and 3 ppt limits of detection (LOD) for 272 the ToF-CIMS and Quad-CIMS instruments, respectively, for measurements with the IMR at 30°C, I(H₂O)/I⁻ ratio of 0.55, and 1 s integration (see Table 1). The instruments were insensitive to MIC with LODs of >860 and >570 ppb 273 274 for the ToF and Quad I-CIMS instruments, respectively. A weak negative temperature dependence and systematic 275 positive I(H2O)'/I' ratio dependence was observed for HAN. The ToF and Quad I-CIMS instruments have similar 276 normalized sensitivities, reflecting the similar chemistry in the ion molecule reaction regions. The ToF instrument 277 has much lower LOD because the reagent ion concentration was approximately 10 times greater than that in the Quad 278 instrument and the ToF has far better mass resolution. The results from this study should, to a first approximation, 279 translate to other laboratory and field I-CIMS instruments.

Our work demonstrates that the previous field studies of Priestley et al. (2018), Mattila et al. (2020a; 2020b), and Wang et al. (2022), which used I-CIMS detection methods, mis-attributed the C₂H₃NO mass signal as methyl isocyanate, MIC, and also provides evidence for the observation of HAN, a previously unrecognized species in these environments. Our results suggest that HAN was observed, but do not imply that MIC was not present in the environments studied by Priestley et al., Mattila et al., and Wang et al. Since I-CIMS is not sensitive to MIC, alternative measurement methods, such as proton transfer CIMS, would be required to identify the presence of MIC.

286 Our work indicates that HAN is likely to be present in the troposphere. Iver et al. (2016) and Hyttinen et al. (2018)

287 provide an explanation for the significant difference in the I-CIMS sensitivity for MIC (CH₃NCO) and HAN

288 (HOCH₂CN), due to the stability of I- cluster binding energies. That is, the H-bonding with the HO group in HAN

289 leads to a stable I- cluster, while MIC would not form a stable I- cluster.

The heterogeneous and gas-phase atmospheric chemistry of HOCH₂CN (HAN) are, however, not presently characterized. Here, we postulate that in addition to primary HAN emissions, e.g., from wildfires, that HAN would

be formed heterogeneously in clouds, or on hydrated aerosol, via the liquid-phase reaction:

 $\label{eq:HCN} 293 \qquad \mathrm{HCN} + \mathrm{H_2CO} \Leftrightarrow \mathrm{HOCH_2CN}$

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- 295 The partitioning of HAN between the liquid- and gas-phase will depend on its Henry's law coefficient, which has not
- 296 been measured to date. Sander (2023) reports an estimated Henry's law coefficient value of ~130 M atm⁻¹, using a
- 297 quantitative structure-property relationship, which implies partitioning of HAN into the gas-phase. HCN and H₂CO
- are ubiquitous in the atmosphere with elevated concentrations in wildfire plumes, which may lead to a significant
- 299 enhancement of the HAN concentration following a wildfire plume exposure to clouds. Our work will aid future
- 300 laboratory and field studies to identify the atmospheric source chemistry of HAN as well as its atmospheric loss
- 301 processes and degradation products.
- 302 Data availability. NA
- Author contributions. ZF undertook the experimental measurements and contributed to the first draft and writing of the paper. AC performed the MIC infrared measurements and initial ToF MIC measurements. JAN performed the
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