1 Response of protonated, adduct, and fragmented ions in Vocus proton-

2 transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS)

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17 Abstract

Volatile organic compounds (VOCs) affect secondary pollutant formation via active 18 chemistry. Proton-transfer-reaction mass spectrometry (PTR-MS) is one of the most 19 important techniques to study the highly variable spatial and temporal characteristics 20 of VOCs. The response of protonated, adduct, and fragmented ions in PTR-MS in 21 changing instrument settings and varying relative humidity (RH) requires rigorous 22 characterization. Herein, dedicatedly designed laboratory experiments were conducted 23 to investigate the response of these ions for 21 VOCs, including 12 oxygenated VOCs 24 and two nitriles, using the recently developed Vocus PTR-MS. Our results show that 25 26 the focusing ion-molecule reactor (FIMR) axial voltage increases sensitivity by three to four orders of magnitude but does not significantly change the fractions of protonated 27 ions. Reducing the FIMR pressure, however, substantially increases fragmentation. 28 Applying a high radio frequency (RF) amplitude radially on FIMR can enhance 29 sensitivity by one to two orders of magnitude without affecting the protonated ion 30 fractions. The change in big segmented quadrupole (BSQ) amplitude mainly affects 31 32 sensitivity and protonated ion fraction by modifying ion transmission. The relationship between sensitivity and proton-transfer reaction rate constant is complicated by the 33 influences from both ion transmission and protonated ion fraction. The protonated ions 34 35 of most VOCs studied (19 out of 21) show less than 15% variations in sensitivity as RH increases from ~5% to ~85%, except for some long-chain aldehydes which show a 36 positive RH variation of up to 30%. Our results suggest that the Vocus PTR-MS can 37 reliably quantify the majority of VOCs under ambient conditions with varying RH. 38 However, caution is advised for small oxygenates such as formaldehyde and methanol 39 40 due to their low sensitivity, as well as for long-chain aldehydes for their slight RH dependence and fragmentation. 41

42 1 Introduction

Atmospheric volatile organic compounds (VOCs) affect atmospheric chemistry by 43 44 forming secondary pollutants such as tropospheric O₃ (Shao et al., 2016) and secondary organic aerosols (SOA) (Shrivastava et al., 2017). In addition to their low mixing ratios 45 (parts per billion by volume, ppbv, or even lower), the spatial and temporal variabilities 46 of atmospheric VOCs pose another analytical challenge to the study of their 47 atmospheric occurrence, sources, and fates. Mass spectrometric (MS) techniques based 48 on ion-molecule reactions (IMR) (Španěl and Smith, 1996) or specifically proton-49 transfer reactions (PTR) (Hansel et al., 1995; Lindinger et al., 1998) in a selected ion 50 flow tube (SIFT) have been developed to provide fast-responding measurements of 51 VOCs. These techniques, especially the PTR-MS, have been widely used in VOC 52 measurements in outdoor and indoor environments (Salazar Gómez et al., 53 2021;Sekimoto and Koss, 2021;Pagonis et al., 2019;Pleil et al., 2019;Claflin et al., 54 2021;Schripp et al., 2014;Jensen et al., 2021). 55

Quantification of VOCs (denoted as M) by PTR-MS relies heavily on their proton-56 transfer reactions with the hydronium ion H_3O^+ (R1). In early SIFT-MS studies where 57 reagent ions include a multitude of $H_3O^+(H_2O)_n$ (n =0, 1, 2, 3...) ion series (Španěl and 58 Smith, 2000), proton-transfer reactions with more hydrated ($n \ge 1$) hydronium ions 59 (R2a) are also important for species with proton affinity (PA) larger than water clusters. 60 In addition, ligand switching reactions (R2b) and association reactions (R3, with N 61 being N₂ or O₂) are also common, leading to $[MH + H_2O]^+$ instead of MH⁺. Under these 62 circumstances, the quantification of VOCs might be heavily influenced by water vapor 63 concentration, or relative humidity (RH), of the sample. For instance, acetone 64 65 concentrations in exhaled air were overestimated by 13% even using both protonated (MH^+) and water adduct ($[MH + H_2O]^+$) ions for quantification, when water vapor 66 varied in the range of $(1 - 10) \times 10^{12}$ molecules cm⁻³ (Španěl and Smith, 2000). A later 67 study (Smith et al., 2001) showed that quantification of other oxygenated VOCs 68 (OVOCs) such as ethyl acetate, diethyl ether, methanol, ethanol, and propanol by SIFT-69 MS also suffered from RH dependence to various degrees. 70

 $M + H_3O^+ \rightarrow MH^+ + H_2O \tag{R1}$

$$M + H_3O^+(H_2O)_n \to MH^+ + (H_2O)_{n+1}$$
 (R2a)

$$M + H_3O^+(H_2O)_n \rightarrow [MH + H_2O]^+ + (H_2O)_n$$
 (R2b)

$$MH^{+} + H_2O + N \rightarrow [MH + H_2O]^{+} + N$$
(R3)

For PTR-MS that normally uses the MH⁺ for quantification, RH dependence was 71 72 also widely reported. For instance, Warneke et al. (2001) reported that the sensitivity of benzene in PTR-MS decreased significantly with the increase of RH, while Steinbacher 73 et al. (2004) suggested a slight decrease with the increase of RH. Quantification of 74 biogenic volatile organic compounds (BVOCs) was also reported to be slightly affected 75 76 by RH (Kari et al., 2018). The RH dependence stems from the change of reagent ion distribution, i.e., among H_3O^+ and $H_3O^+(H_2O)_n$ (n ≥ 1), which can lead to 77 overestimation or underestimation of VOCs if such dependence is strong because 78 ambient RH is deemed highly variable. Therefore, RH-dependent calibrations for VOC 79 measurements using PTR-MS were normally recommended (de Gouw and Warneke, 80 2007;Inomata et al., 2008;Sinha et al., 2009;Vlasenko et al., 2010;Cui et al., 81 2016;Michoud et al., 2018). 82

Another complication in VOC measurements using SIFT-MS or PTR-MS is that, due to the nucleophilicity of the oxygen atom, protonated OVOCs would dehydrate, forming fragmented ions (R4). This reaction often occurs in heavy alcohols, aldehydes, and carboxylic acids (Španěl et al., 1997;Ŝpaněl and Smith, 1998;Hartungen et al., 2004;Baasandorj et al., 2015).

88
$$MH^+ \rightarrow [MH - H_2O]^+ + H_2O$$
 (R4)

In addition, cleavage on the C-C bond of the protonated ion (R5) is also possible, especially for alkyl-substituted VOCs under high-energy conditions (e.g., a high E/N ratio, which is the reduced electric field parameter with E being the electric field and N the number density of the gas in the drift tube).

93 $MH^+ \rightarrow [MH - C_xH_y]^+ + C_xH_y$ (R5)

For instance, at an E/N ratio of 120 Townsend (Td), substituted monocyclic aromatic compounds such as ethylbenzene and propylbenzene start to fragment into a benzenium ion ($C_6H_7^+$) (de Gouw et al., 2003;Gueneron et al., 2015).

A newly designed focusing ion-molecule reactor (FIMR) was used for PTR-MS,
termed Vocus, and has been shown to have little RH dependence for the protonated ion
because of the high concentration of water vapor introduced into the FIMR (Krechmer

et al., 2018). The concentration of hydronium ion (H_3O^+) in the FIMR is high enough 100 to maintain at a constant level and dominate over other side reactions, thereby 101 minimizing the RH dependence for VOC measurement. Yet, the formation of adduct 102 ions and fragmented ions in Vocus PTR-MS as a function of RH has not been fully 103 scrutinized, hindering a complete understanding of the ion chemistry in the Vocus PTR-104 MS and potential cross interference when measuring ambient air with complex VOC 105 mixtures. Herein, we conducted experiments on the effects of instrumental settings and 106 RH variations on the quantification of 21 VOCs, including 12 OVOCs and 2 nitriles, 107 108 using a Vocus PTR-MS. Response of protonated ions (MH⁺), adduct ions ([MH + H_2O^{+}), and fragmented ions ($[MH - H_2O^{+}]$ or $[MH - C_xH_y]^{+}$) of these VOCs was 109 investigated as a function of instrumental setting and RH. Results are interpreted based 110 on the PA values and/or proton-transfer reaction rate constants (k_{ptr}) . Some caveats on 111 using the Vocus PTR-MS to measure VOCs, especially OVOCs, are also provided. 112

113 2 Methodology

114 **2.1 Instrument settings**

115 Experiments were performed with a Vocus proton-transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS, Vocus 2R, TOFWERK AG, Thun, Switzerland), 116 117 hereinafter referred to as Vocus. The Vocus consists of (i) a discharge ion source, (ii) a focusing ion-molecule reactor (FIMR), (iii) a big segmented quadrupole (BSQ), (iv) a 118 119 series of direct current (DC) optics that further focus and accelerate the primary beam (PB), and (v) a time-of-flight (ToF) mass analyzer (Krechmer et al., 2018). The ion 120 121 source is a plasma discharge composed of two conical surfaces. Water vapor is supplied by purging 20 to 30 mL of milli-Q water and is ionized by plasma discharge. The 122 reagent ions pass through a ring offset from the central axis so that the photons 123 generated by the discharge cannot enter. The drift tube was improved by replacing the 124 stacked ring electrodes of the traditional PTR-MS with a FIMR, which is a glass tube 125 with a resistive coating on the inner surface and a quadrupole with a radio frequency 126 (RF) electric field applied. The FIMR increases ion transmission by a factor of 7 to 9 127 and sensitivity by more than one order of magnitude (Krechmer et al., 2018). Moreover, 128 the mean kinetic energy of H_3O^+ is increased by three times, and the formation of more 129 hydrated hydronium ions is reduced, suppressing RH dependence for most VOCs 130 measured (Krechmer et al., 2018). Meanwhile, the mean kinetic energy of VOCs 131

measured is not significantly increased, thereby minimizing fragmentation (Krechmer et al., 2018). The ToF mass analyzer offers a mass resolving power of 12,000 at a massto-charge ratio (m/Q) of 107 Thomson (Th).

In our experiments, sample air was drawn into the instrument using 0.5 m long perfluoroalkoxy (PFA) Teflon tubing of ~0.5 m length and 1/4 " outer diameter, with a flow rate of 0.5 L·min⁻¹. Most of the sample air was directed to the exhaust, while the actual flow into the FIMR was around 0.15 L·min⁻¹. In typical experiments, the FIMR was operated at a pressure of 2.0 mbar and a temperature of 373.15 K. The axial and radial voltages were normally set to be 625 and 500 V, respectively, unless stated otherwise.

We also performed experiments by varying the instrument settings such as FIMR 142 axial voltage (V) and FIMR pressure (p), both of which affect the E/N ratio, as well as 143 144 RF and BSQ amplitudes to investigate how protonated, adduct, and fragmented ions respond to those changes. These experiments were performed under dry (RH ~5%) 145 146 conditions, and the concentrations were approximately 12 ppbv for most VOCs (except for β -caryophyllene at about 1.2 ppby). The instrument settings were varied by: 1) 147 changing the FIMR axial voltage from 260 to 700 volts, 2) changing the pressure in 148 FIMR from 1.5 to 3.5 mbar; 3) changing the RF amplitude from 13 to 500 volts (with 149 p of 2.0 or 3.5 mbar); 4) changing the BSQ voltage from 50 to 300 volts (with p of 2.0 150 or 3.5 mbar). The other instrument settings were fixed as the default values while 151 changing the tested ones. Specifically, RF amplitude was at 500 volts and BSQ 152 amplitude was at 300 volts when changing E (i.e., V) and N (i.e., p), and an E/N ratio 153 of 142 Td was used when changing RF and BSQ amplitudes (Table S1). 154

155 **2.2 Experimental setup**

The VOCs (Table S2 and Figure 1) in mixtures from two cylinders were separately 156 delivered to the dilution and/or RH control setup (Figure S1). Dilution air was generated 157 from a zero-air generator (Environics series 7000, Environics Analytics Group Ltd., 158 Canada). Gas cylinder I (Table 1, Apel-Riemer Environmental Inc., US, valid for 12 159 months) contains mainly hydrocarbons, while gas cylinder II (Table 1, Linde Gases, 160 US, valid for 12 months) contains mainly OVOCs and nitriles. Most VOCs in the 161 cylinders are at approximately 1000 ppbv, except for β -caryophyllene that is at 162 approximately 100 ppbv. Table S2 shows their CAS numbers, m/Q values of the 163

164 protonated ions (MH⁺), as well as PA and k_{ptr} values. According to their functional 165 groups, the 21 VOCs are grouped into 9 categories, and Figure 1 shows their structures. 166 Note that although n-butanal and methyl ethyl ketone are isomers, they are in different 167 cylinders and measured separately, thus will not interfere with each other.

Dry experiments were performed by diluting the VOCs from the cylinders with dry zero 168 air with 7 concentrations from 0 to approximately 22 ppbv (or approximately 2 ppbv 169 for β -caryophyllene). For each concentration step, measurements lasted for about half 170 an hour for gas cylinder I with hydrocarbons but about two hours for gas cylinder II 171 172 with OVOCs and nitriles. The 2-hour stabilization time for cylinder II, which contained mainly OVOCs and nitriles, should be sufficient because even with half-an-hour 173 stabilization time for cylinder I, the overlapping species acetone and acetaldehyde 174 showed deviations of less than $\pm 1.5\%$ in sensitivity. Other measures such as minimizing 175 the length of the Teflon tube at the inlet (less than 30 cm) were also undertaken to allow 176 fast establishment of equilibrium-state concentrations for OVOCs. In RH-dependent 177 experiments (Figure S1), dilution was made by RH-conditioned air produced from a 178 179 humidity generator (OHG-4, Owlstone, US). The accuracy of the RH sensor (RH-USB Probe, Omega) is within 4% of RH. Nine RH ramping steps from ~5% to ~85% with 180 181 approximately 10% intervals were used, and the VOCs were set with 4 concentrations from 0 to approximately 12 ppbv (or approximately 1.2 ppbv for β -caryophyllene). 182 After the initial equilibration of 0.5 hours under dry conditions (RH ~5%), each RH 183 ramping lasted for 15 min. Triplicate experiments were performed for the highest 184 concentration. 185



Figure 1. Names, structures, and grouping of the 21 VOCs in this study. These VOCs were prepared in two gas standard cylinders (I and II) with mixing ratios of ~1000 ppbv (~100 ppbv for β -caryophyllene), balanced by N₂. Acetaldehyde and acetone are present in both gas standard cylinders, with concentrations differing by <5%. n-Butanal and methyl ethyl ketone are isomers but are in different gas standard cylinders.

192 2.3 Data analysis

The Vocus data were analyzed with the manufacturer-supplied software package 193 Tofware (v3.2.3) based on Igor Pro (Wavemetrics). Peak fitting was performed using 194 Tofware routines and the measured m/Q values of the pronated ions (MH⁺) are shown 195 in Table S2 together with their exact m/Q values. In addition to MH⁺, we also looked 196 for adduct ions ($[MH + H_2O]^+$), and fragmented ions ($[MH - H_2O]^+$ or $[MH - C_xH_y]^+$), 197 since those ions are also anticipated for VOCs, especially OVOCs, in PTR-MS 198 measurements (Pagonis et al., 2019). Although the VOCs were in mixtures and showed 199 ensemble mass spectra, we constructed the mass spectrum for each VOC by plotting 200 201 their identified adduct or fragmented ions alongside the protonated ions, as shown in Figures S2-S4 in the SI. Their percentage contributions are presented in Figure 2 and 202 numerically in Table S4. 203



204

Figure 2. The average signal percentages of protonated, adduct, and fragmented ions at a concentration of ~12 ppbv (~1.2 ppbv for β -caryophyllene).

For both dry and RH-dependent experiments, from the 1-minute averages from 1-207 Hz datasets, the last five points were averaged to obtain stable signals. Figure S5 shows 208 the time series of selected hydrocarbons (groups A1 and A2, aromatic hydrocarbons 209 and terpenoids, Figure S5a) and OVOCs (groups B2 and B5, long-chain and aromatic 210 aldehydes, Figure S5b) during dry experiments with concentration stepping. Figure S6 211 shows the time series of protonated ions (MH⁺), adduct ions ($[MH + H_2O]^+$), and 212 fragmented ions $([MH - H_2O]^+ \text{ and/or } [MH - C_xH_y]^+ \text{ for n-butanal, pentanal and}$ 213 hexaldehyde, whose adduct and fragmented ions contributed substantially to the total 214 signals (Figure 2 and Table S4). Some dealkylated fragments ($[MH - C_xH_v]^+$) of long-215 chain aldehydes (e.g., n-butanal and pentanal) might overlap with the protonated ions 216 ([MH]⁺) of unsaturated aldehydes (i.e., acrolein and methacrolein). Yet since the 217 218 intensities of the former are expected to be low by analogy with that of hexaldehyde (~2%, Figure 2), the ions at those m/Q values are only considered as the protonated ions 219 of unsaturated aldehydes (the latter). 220

Similar to RH-dependent experiments, the duration of each instrument setting experiment was 15 min. The signal intensities and the ratio of MH^+ , $[MH + H_2O]^+$ and $[MH - H_2O]^+$ and/or $[MH - C_xH_y]^+$ to all for two typical hydrocarbons (α -pinene and 1,2,4-trimethylbenzene) and two OVOCs (acetone and hexaldehyde) as the axial voltage and pressure (which both affects the E/N ratio) in the FIMR, RF amplitude, and BSQ amplitude varied are shown in Figures S7 to S10. The reduced electric field parameter (E/N ratio) was estimated by comparing the signal fraction of fragment of α pinene in Materić et al. (2017), in which detailed examination on fragment signal fraction at different E/N ratio was performed (Figure S11).

230 2.4 Sensitivity estimation

The formation of the protonated ion MH⁺ via reaction R1 is desirable for quantification of VOCs, which is described by the kinetics of the proton-transfer reaction (de Gouw and Warneke, 2007;Yuan et al., 2017):

234
$$[MH^+] = [H_3O^+]_0 \quad (1 - e^{-k[M]\Delta t}) \quad (\text{Eq. 1})$$

where $[MH^+]$ is the number concentration of the protonated ion, $[H_3O^+]_0$ that of the 235 236 initial hydronium ion, k is the rate constant of R1 (k_{ptr}), [M] is the number concentration of the target VOCs in the sample air, and Δt is the reaction time in the FIMR. Two 237 conditions allow simplification of Eq. 1 to Eq. 2 below for easy quantification of VOCs: 238 1) The term $k[M]\Delta t$ is much smaller than 1, such that R1 can be considered essentially 239 first-order; and 2) H₃O⁺ is not significantly depleted and remain more or less constant 240 after the FIMR. Compared with traditional ion sources, the Vocus ion source produces 241 sufficient H₃O⁺ (Krechmer et al., 2018). Ambient levels of ppbv (or less) for [M] (~10¹⁰ 242 molecule cm⁻³) generally fulfill such requirements, given that k_{ptr} is on the order of 10⁻ 243 9 cm³ molecule⁻¹ s⁻¹ and Δt of 10⁻⁴ s (Ellis and Mayhew, 2014). Therefore, 244

245
$$[MH^+] = [H_3O^+]k[M]\Delta t \text{ (Eq. 2)}$$

where $[H_3O^+]$ is the mixing ratio of hydronium ions after the FIMR (i.e., being detected in the mass spectrometer). Then,

248
$$[M] = \frac{I_{MH^+}}{I_{H_30^+}} \frac{1}{k\Delta t} \quad (Eq. 3)$$

where I_{MH^+} and $I_{H^{3O^+}}$ are signal intensities of the protonated ion and the hydronium ion, respectively.

In general, the sensitivity (S) of PTR-MS for quantification of VOCs is defined as the ratio between the signal intensity I_{MH^+} normalized by 10⁶ cps (counts per second) of I_{H3O^+} and 1 ppbv (10⁻⁹ mol mol⁻¹) of VOCs, i.e.:

254
$$S = \frac{\frac{I_{MH^+}}{I_{H_3O^+}} \times 10^6}{\frac{[M]}{N} \times 10^9} \quad (\text{Eq. 4})$$

where N is the number density of air in the FIMR. The sensitivity S is thus expressed
as a normalized signal per ppbv, having a unit of neps ppbv⁻¹. Combining Eq. 3 and Eq.
4 yields,

$$S = 10^{-3} \times N\Delta t \times k (\text{Eq. 5})$$

where $10^{-3} \times N \Delta t$ is specific to the instrumental settings. Eq. 5 dictates that S should have a linear relationship with the proton-transfer reaction rate constant (k_{ptr}) if the instrument settings are fixed and can be utilized to predict S if k_{ptr} values are known (Ellis and Mayhew, 2014).

In reality, however, quantification of VOCs using MH^+ from PTR-MS measurements is complicated by 1) formation of adduct (e.g., with H₂O) and fragmented (e.g., dehydration) ions, and 2) discriminated transmission for MH^+ ions with different *m/Q* values (de Gouw and Warneke, 2007; Yuan et al., 2017). The fraction of MH^+ in all related ions (f_{MH^+}) and the relative transmission efficiency ($T_{MH^+}/T_{H_3O^+}$) are used to account for these two effects, respectively:

269
$$S = 10^{-3} \times N\Delta t \times \frac{T_{MH^+}}{T_{H_30^+}} \times f_{MH^+} \times k \text{ (Eq. 6)}$$

In our study, the sensitivity is expressed as the slope of signal intensity (in counts per second, cps) vs. concentration (in ppbv), having a unit of cps ppbv⁻¹ (Figure S12). Signal normalization to H_3O^+ (ncps) was not adopted because the signal of H_3O^+ (m/Q= 19 Th) was substantially suppressed with low transmission (see below) for those ions with small m/Q values (but too high intensities) to minimize ion currents.

275 **3 Results and Discussion**

276 **3.1 Effects of instrumental settings on the ion signals**

An increase of the E/N ratio from 48 to 142 Td manifested by the increases of the FIMR axial voltage (V, with front from 226 to 666 volts and back keeps at 34 volts) led to drastic increases of MH⁺ signal intensity by three to four orders of magnitude for all VOCs studied (Figure 3a). Such increases were also observed for adduct and

fragmented ions (Figures S7 - S10), albeit to different extents. It has been shown that 281 increasing the axial voltage in the FIMR can substantially increase protonated ion 282 signals, which is due mainly to three reasons (Krechmer et al., 2018;de Gouw and 283 Warneke, 2007). First, a high FIMR axial voltage can accelerate the ions and thus 284 reduce their residence time, thereby preventing diffusional loss. Second, the high 285 voltage in FIMR significantly increases the concentration of all reagent ions (Figure 286 S13). Lastly, at high voltage, reactions between some OVOCs (such as acetone) and 287 H_3O^+ leads to more protonated ions (MH⁺) instead of adduct ions ([MH + H₂O]⁺) 288 289 through ligand switching reaction (R2b). The last effect was believed to be less significant for species that do not readily react with H₃O⁺(H₂O)_n (such as aromatics and 290 terpenoids). 291



Aromatic hydrocarbons 🧧 Terpenoids 🥅 Small aldehydes 🥅 Long-chain aldehydes 1 Ketones
Unsaturated aldehydes Aromatic aldehydes 🔲 Nitriles 🔲 Methanol

Figure 3. The ratio of the logarithm of intensity (panels a, b, and c) and the difference 293 of fractional signal of the protonated ion (MH⁺) among all ions (panels d, e, and f), 294 when changing axial voltage (V) or FIMR pressure (p) (panels a and d), RF amplitude 295 (panels b and e), and BSQ amplitude (panels c and f). The ratios were taken after taking 296 the logarithm of the signal intensities of MH⁺ at the right-hand side of the instrument 297 setting (after the arrow) to that at the left-hand side of the instrument setting stated in 298 the panel label; likewise, the fractional differences are the fractions of the MH⁺ signal 299 300 among the protonated, fragmented, and adduct signals under these two instrumental 301 settings.

To investigate whether the desired MH⁺ is indeed more favorably formed by 302 suppressing R2b under high axial voltages, we plot the differences in the signal 303 fractions of MH⁺ between axial voltages of 666 volts and 226 volts (Figure 3d). The 304 results show that about one-third of the 21 VOCs do not have significant differences in 305 the signal fractions: most VOCs in this one-third have the MH⁺ as the sole or 306 307 dominating ion observed (cf., Figure 2). Meanwhile, there are about one-third showing negative differences (i.e., decreasing MH⁺ fractions) of up to 0.2, including the two 308 hydrocarbons and two OVOCs shown in panel d of Figures S7 – S10. The remaining 309 310 one-third show positive differences (i.e., increasing MH⁺ fractions) of up to 0.3, mainly for unsaturated or aromatic aldehydes, as well as nitriles (Figure 3d). A closer 311 inspection of the fractional changes as axial voltage increases for acetone (Figure S9d) 312 and hexaldehyde (Figure S10d) reveals that the fractions of both MH^+ and $[MH + H_2O]^+$ 313 decrease, while those of fragmented ions $([MH - H_2O]^+ \text{ and/or } [MH - C_xH_y]^+)$ increase. 314 The relative decreases of the signal fractions of MH⁺ (8% for acetone and 51% for 315 hexaldehyde) are, however, much lower than those of $[MH + H_2O]^+$ (57% for acetone 316 317 and 80% for hexaldehyde). These observations suggest that while fractions of both MH^+ and $[MH + H_2O]^+$ decrease, the decreases of the adduct ion $([MH + H_2O]^+)$ are more 318 significant, supporting the third reason that relatively more H_3O^+ (instead of 319 $H_3O^+(H_2O)_n$) at higher axial voltages to react with these OVOCs. However, as the 320 voltage increases, all ion signals are increasing (Figure S7a-S10a). This observation 321 illustrates that the ion acceleration and diffusion prevention should be the primary 322 reason for signal enhancement at high axial voltages. Nevertheless, the signal fractions 323 of the MH⁺ do not change substantially (within 30%) as the FIMR axial voltage 324 325 increases, making quantification reliable even for species with high signal contributions from adduct and fragmented ions (e.g., long-chain aldehydes, group B2, cf., Figure 2). 326 327 The increase of E/N ratio by decreasing FIMR pressure from 3.5 to 1.5 mbar increases signal intensities (Figure S14a) of MH⁺ by less than one order of magnitude, 328 or even decreases those for some OVOCs such as long-chain aldehydes. The changes 329 in signal fractions of MH⁺ (Figure S14d), on the other hand, are more than those when 330 changing axial voltages, especially for ketones, unsaturated aldehydes, aromatic 331 aldehydes, and nitriles. The increase of pressure in the PTR reactor also favors the 332 formation of reagent clusters $H_3O^+(H_2O)_n$, which leads to the formation of adduct ions 333 (Wang et al., 2020). For OVOCs acetone and hexaldehyde, the fractions of MH⁺ do 334

increase when FIMR pressure was decreased from 3.5 to 2.5 mbar (E/N ratio from 162 335 to 95), which is accompanied by significant decreases of the adduct ion $[MH + H_2O]^+$ 336 (Figure S9j and Figure S10j). This observation suggests less formation of adduct ions 337 at lower FIMR pressures. Further decrease of FIMR pressure to 1.5 mbar, however, 338 results in slight decreases of MH⁺ fractions, in lieu of increases of fragmented ions [MH 339 $-H_2O^{\dagger}$ and $[MH - C_xH_v]^+$ (Figure S9j and Figure S10j); for hydrocarbons α -pinene 340 and 1,2,4-trimethylbenzene (Figure S7j and Figure S8j) that only have MH⁺ and [MH 341 $-C_xH_y$ ⁺, continuous decreases of MH⁺ fractions and increases of [MH - C_xH_y]⁺ are 342 343 observed for the whole range of FIMR pressure tested (3.5 to 1.5 mbar). A recent study using Vocus PTR-MS to measure organic peroxides also observed that less fragmented 344 ions were formed under higher FIMR pressure (Li et al., 2022), presumably due to the 345 efficient transfer of excess kinetic energy by frequent collisions at higher pressures. A 346 medium FIMR pressure of 2.0 mbar was chosen to have relatively low fractions of both 347 348 adduct ions ($[MH + H_2O]^+$) and fragmented ions ($[MH - H_2O]^+$ and $[MH - C_xH_v]^+$).

The radial RF electric field in the FIMR is unique for the Vocus PTR-MS, which 349 350 can (1) collimate the ions towards the central axis (especially heavier ions) and (2) increase the kinetic energy of the ions (especially for lighter ions) (Krechmer et al., 351 352 2018). These effects led to 1 to 1.5 orders of magnitude higher signals for MH^+ at 2.0 mbar FIMR pressure (Figure 3b) and 1.5 to 2 orders of magnitude at 3.5 mbar (Figure 353 S14b) when the RF amplitude was changed from 13 to 500 volts. The additional 354 enhancement of signal intensity at a higher FIMR pressure (i.e., 3.5 mbar as compared 355 to 2.0 mbar) can be attributed to a longer residence time of the reagent ions (Krechmer 356 et al., 2018). The more pronounced increase of kinetic energy for lighter ions (e.g., 357 H_3O^+) than heavier ions [i.e., clusters $H_3O^+(H_2O)_n$] might imply the favorable 358 formation of the protonated ion MH⁺ rather than adduct ions. The fractions of MH⁺ for 359 different RF amplitudes do not change significantly (within ±0.2) either at 2.0 mbar 360 (Figure 3e) or 3.5 mbar (Figure S14e). This observation thus suggests that adding the 361 RF can increase signal intensities by 1-2 orders of magnitude but does not affect the 362 fractional signal for MH⁺, making it beneficial for accurate quantification. 363

The BSQ amplitude above 100 volts does not change the signal intensities significantly (Figure 3c and Figure S14c, as well as Figures S7– S10). The BSQ ion guide provides a high-pass band filter to reduce the number of ions (thus signal intensity) of low m/Q values (especially for reagent ions, H₃O⁺, with high ion currents),

preventing the fast degradation of the microchannel plate (MCP) detector (Krechmer et 368 al., 2018). This bandpass filter leads to lower ion transmission efficiency (<1) for ions 369 with smaller m/Q values, which is discussed below. Therefore, the signal reduction 370 when the BSQ amplitude increased from 100 to 300 volts is more obvious for small 371 analytes such as formaldehyde, acetonitrile, and methanol (Figure 3c and Figure S14c). 372 For other analytes whose fragmented ions have m/Q values of less than 60 Th, the signal 373 fractions of MH⁺ would also be affected (Figure 3f and Figure S14f). For example, the 374 intensities of fragmented ions $[MH - H_2O]^+$ (CH₃CCH₂⁺, m/Q = 41 Th) and $[MH - H_2O]^+$ 375 C_xH_y ⁺ (CH₃CO⁺, m/Q = 43 Th) for acetone had substantial decreases when BSQ 376 amplitude was higher than 200 volts (Figure S9c and Figure S10). The protonated ion 377 MH⁺ (CH₃COCH₃H⁺, m/Q = 59 Th) and adduct ion (CH₃COCH₃H₃O⁺, m/Q = 77 Th), 378 however, remained less unaffected. This effect leads to noticeable changes in signal 379 fractions of MH⁺ (maximum 0.4) for small analytes such as ketones, unsaturated 380 aldehydes, nitriles, as well as methanol (Figure 3f and Figure S14f) as the BSQ 381 amplitude changes from 100 to 300 volts. 382

383 **3.2 Sensitivity and transmission of protonated ions**

We calculated the sensitivity and transmission of protonated ions (MH⁺) of the 21 384 VOCs studied when the instrument was under the optimized conditions. Table 1 shows 385 the sensitivities (cps ppbv⁻¹), as slopes of MH⁺ signals vs. mixing ratios (average value 386 from 0 to 22 ppbv at dry condition, except for β -caryophyllene to 2 ppbv) and the limit 387 of detection (LOD, 3σ). Panels a and b in Figure 4 show the sensitivity vs. k_{ptr} for all 388 21 VOCs, while panels c and d show the transmission efficiencies calculated from the 389 division of the sensitivity vs. k_{ptr} ratio for each VOC by the slope fitted in Figure 4a and 390 4b. 391

Most VOCs had sensitivities above 1000 cps ppbv⁻¹, except 1) formaldehyde (A2-392 1) and methanol (C2-1), whose MH^+ ions have m/Q values much lower than 60 Th; and 393 2) β -caryophyllene (A2-3) that came with a very low concentration range. In addition 394 395 to its low m/Q values that limit the transmission, the backward reaction is also an important reason for the low sensitivity. For instance, formaldehyde has a low PA value 396 (712.5 kJ mol⁻¹) that is not much higher than that of water (691.0 kJ mol⁻¹) and has been 397 shown to have a high tendency of backward reaction of R1 (Inomata et al., 398 399 2008; Vlasenko et al., 2010; Warneke et al., 2011). The two compounds in group B5

400 (aromatic aldehydes), benzaldehyde and m-tolualdehyde, had the highest sensitivities 401 of >12000 cps ppbv⁻¹. This might be due to their high PA values (>830 kJ mol⁻¹), which 402 are among the highest except those of terpenoids (Table S2).

Table 1. Sensitivity (slope), intercept, and limit of detection (LOD) based on 3 standard

deviations (σ) and cylinder numbers (No.). Results were obtained from measurements

of 0 - 22 ppbv for all VOCs except for β -caryophyllene (up to ~ 2 ppbv). The cylinder

406 numbers for the VOCs studied are also shown here.

Group ^a	Name	Cylinder No.	Label	Sensitivity (cps ppbv ⁻¹)	3σ LOD (pptv), 5 s
A1	Benzene	Ι	A1-1	2596	35
	Toluene	Ι	A1-2	5724	2
	m-Xylene	Ι	A1-3	8669	3
	1,2,4-Trimethylbenzene	Ι	A1-4	8951	1
A2	Isoprene	Ι	A2-1	2140	16
	α-Pinene	Ι	A2-2	4046	2
	β-Caryophyllene	Ι	A2-3	723	1
B1	Formaldehyde	Π	B1-1	c	c
	Acetaldehyde	I&II	B1-2	2096 ^b	283
B2	n-Butanal	Π	B2-1	1114	343
	Pentanal	Π	B2-2	1465	63
	Hexaldehyde	Π	B2-3	1595	35
B3	Acetone	I&II	B3-1	9932 ^b	127
	Methyl ethyl ketone	Ι	B3-3	9636	51
B4	Acrolein	Ι	B4-1	7224	16
	Methacrolein	Ι	B4-2	6090	13
B5	Benzaldehyde	Π	B5-1	16089	1
	m-Tolualdehyde	Π	B5-2	12893	1
C1	Acetonitrile	Ι	C1-1	1511	6
	Acrylonitrile	Ι	C1-2	9275	1
C2	Methanol	Ι	C2-1	c	c

407 Notes:

408 a, A1: aromatic hydrocarbons, A2: terpenoids, B1: small aldehydes; B2: long-chain aldehydes, B3:

409 ketones, B4: unsaturated aldehydes, B5: aromatic aldehydes, C1: nitriles, C2: methanol; 410 b, the average value from gas cylinders I and II, with a deviation of less than $\pm 1.5\%$;

410 b, the average value non gas cynders r and n, whera devia 411 c, low sensitivity and high LOD due to low transmission.



413 Figure 4. Sensitivity as a function of k_{ptr} for (a) MH⁺ and (b) all ions. Linear fitting was 414 performed only for MH⁺ sensitivity. Species included in the fitting were those with m/Q415 value > 60 Th (Table S2) and signal percentage of MH^+ (f_{MH^+}) > 75% (Table 1). The 416 grey-shaded area is bounded by $0.5 \times$ Slope and $2 \times$ Slope. The fitted curves in panel b 417 are the same as in panel a and are for reference only. Panels c and d are transmission 418 curves from MH⁺ only and from the sum of all ions, respectively. The sigmoidal curve 419 for MH⁺ (same for c and d) was fitted from species except for β -caryophyllene, α -pinene, 420 n-butanal, pentanal, and hexaldehyde whose fragmentation was significant. Note that 421 only the m/Q values of MH⁺ was used in the x axes of panel c and d, although panel d 422 contains information of adduct and fragmented ions that have m/Q values different from 423 that of MH⁺, which does not consider the differences in m/Q values of adduct and 424 425 fragmented ions.

It was shown that the sensitivities for different VOCs in PTR-MS can be calculated 426 from the kinetics of the proton-transfer reactions (Warneke et al., 2003;Sekimoto et al., 427 2017;Cappellin et al., 2012). For Vocus, Krechmer et al. (2018) also pointed out that 428 the relationship between sensitivity and k_{ptr} can be established and used to calculate the 429 sensitivity for other compounds. We herein compare the relationship between 430 sensitivity and k_{ptr} (Figure 4a). In our data, the uncertainties for sensitivity were 431 conservatively taken as the maximum percentage uncertainty (5.3%) of fitted slopes. 432 Values of k_{ptr} were calculated as averages of both modeled and experimental results 433 434 found from literature (Table S3), with uncertainties propagated from an estimated percentage error of 15% for both modeled (Zhao and Zhang, 2004) and experimental 435 values. The anticipated linear relationship of sensitivity vs. k_{ptr} is not easily visible, 436 most likely due to the formation of fragments/adducts for some VOCs and low 437 transmission efficiencies for others. However, a relatively improved linear relationship 438 was found if we limit the VOCs to 1) m/Q values for MH⁺ > 60 Th, and 2) a fraction of 439 MH⁺ ion in all ions (including adduct/fragmented ions) larger than 75% (cf, Figure 2). 440 With these limitations, the fitted linear line gives a slope of $(2.8 \pm 0.3) \times 10^{12}$ cps ppbv⁻ 441 ¹ molec s cm⁻³, approximately 38% lower than that $[(4.5 \pm 0.4) \times 10^{12} \text{ cps ppbv}^{-1} \text{ molec}$ 442 s cm⁻³] of Krechmer et al. (2018). Pearson's R (R_{Pr}) is 0.77. A grey area is also shown 443 by two lines of $2 \times \text{slope}$ and $0.5 \times \text{slope}$, which includes approximately half (ten) of 444 the VOCs studied. Those that fall out of the grey area to the lower region are mainly 445 compounds in groups B2 (long-chain aldehydes, purple), C1 and C2 (nitriles or 446 methanol, black), B1 (small aldehydes, red), and A2 (terpenoids, green). Using the total 447 signals of all ions (protonated, adduct, and fragmented), Figure 4b shows the 448 improvements for compounds in group B2 only, while others (especially those in C1, 449 C2, and B1) do not move up to the grey area. 450

We also calculated the transmission efficiencies (Figure 4c and 4d) from the 451 division of the sensitivity vs. k_{ptr} ratio for each VOC by the slope fitted in Figure 4a. It 452 is shown that compounds in groups B1, C1, and C2 are mainly on the rising range of 453 the sigmoidal curve, while the three long-chain aldehydes in group B2 (purple) are well 454 below the curve if only MH⁺ ions were used; these three long-chain aldehydes move up 455 to transmission > 0.5 when all ions are considered (Figure 4d). Of particular interest is 456 the compound α -pinene, whose transmission was < 0.5 when only the MH⁺ ion was 457 used, but it increases to about 1.5 when the fragmented ion was considered. Note that 458

in Figure 4d, the m/Q value of MH⁺ was used in the x axis for the summed ion signals, which might lead to certain bias as the fragmented and adduct ions have different m/Qvalues. Nevertheless, the above analysis suggests that both the formation of adduct/fragmented ions and transmission affect the relationship between sensitivity and k_{ptr} , which needs precaution when using predicted sensitivity directly from k_{ptr} . In addition, one needs to be cautious about the prediction of transmission efficiency with m/Q greater than 150.

466 **3.3 RH dependence of ion signals**

One of the most important reasons for RH dependence is that the distribution of 467 reagent ions might vary with ambient RH, especially when the abundance of H₃O⁺ in 468 the PTR reactor is not high. While the Vocus has been shown to have abundant enough 469 H_3O^+ (Krechmer et al., 2018), whether it can substantially minimize RH dependence 470 for most VOCs deserves scrutinization. Figure 5a-h shows the relative sensitivity, 471 defined as the relative change of sensitivity of MH⁺ ion vs. VOC concentration under 472 473 different RH conditions. Among the nine groups of VOCs studied, seven groups show almost flat relative sensitivities within the RH range of ~5% to ~85% (298 K), with the 474 exceptions of long-chain aldehydes (group B2) that show increasing sensitivities as RH 475 increases (Figure 5d) and methanol (group C2) showing large variations (Figure 5h). 476 Some other compounds, such as β -caryophyllene (A2-3, Figure 5b) and formaldehyde 477 (B1-1, Figure 5c) also show either relatively large uncertainties or fluctuations, which 478 can be ascribed to their low intensities (cf., Table 1). 479



480

Figure 5. The dependence of the MH⁺ signals on RH for the VOCs studied. Panels a-h: the relative sensitivity was calculated as the slope (Sensitivity) under all conditions to that at the dry (RH<5%) condition. Panel i: the percent change of relative sensitivity per 10% RH increase. See Figures S15-S17 for other ion signals.

Figure 5i shows the RH dependence of the MH^+ ions, defined as the percentage change of sensitivity per 10% RH increase, for all 21 VOCs studied. Aromatic hydrocarbons (group A1) show negative RH dependence as in previous studies using PTR-MS with a drift tube (Warneke et al., 2001;Steinbacher et al., 2004). While previous studies reported decreases in benzene sensitivity by 16 – 56% from dry to humid conditions (up to 100% RH), our results show a decrease of less than 1.1% per 10% RH increase (i.e., <11% in the whole RH range) with a somewhat narrower RH

range (up to ~85% RH). In addition, two out of the three terpenoids (group A2) also 492 show slightly negative RH dependence, and the other one (α -pinene) shows very small 493 positive RH dependence (Figure 5i). These hydrocarbons (aromatics and terpenoids) in 494 groups A1 and A2 have relatively low k_{ptr} values (mostly $<2.5 \times 10^{-9}$ cm³ molec⁻¹ s⁻¹, 495 Table S3), and their RH dependence shows a fairly good correlation with the PA value 496 497 (Figure 6a, purple circles and squares). This observation suggests that there might be a thermodynamic reason behind the noticeable decrease of sensitivity for hydrocarbons 498 such as benzene as RH increases. Since hydrocarbons such as benzene and toluene do 499 500 not readily react with H₃O⁺(H₂O) (Warneke et al., 2001), R1 is the main reaction to 501 form MH⁺. As water vapor concentration increases at high RH, the reverse reaction of R1 might be important for compounds with low PA and low k_{ptr} values (Inomata et al., 502 2008). 503



Figure 6: The RH dependence of MH^+ ion plotted against PA, color-coded by (a) k_{ptr} and (b) f_{MH}^+ .

504

507 Long-chain aldehydes (group B2) have the largest RH dependence of 1-4% 508 positive deviation per 10% RH increase for the MH⁺ ions. The RH dependence of the 509 $[MH + H_2O]^+$ ions (Figure S15g) is even much higher (1.4–8.5% positive deviation per 510 10% RH increase). Interestingly, the trends of RH dependence for the MH⁺ ions and 511 that for the $[MH + H_2O]^+$ ions for long-chain aldehydes are exactly opposite (Figure 5i

and Figure S15g); that is, pentanal > n-butanal > hexaldehyde. The reason behind this 512 513 observation is out of the scope of this study. The dominating $[MH - H_2O]^+$ ions for long-chain aldehydes (Figure 2), however, show much less ($\pm 1\%$ per 10% RH increase) 514 RH dependence (Figure S16i). Other carbonyl compounds (groups B1, B3, B4, and B5) 515 also show positive deviations as RH increases (less than 1.5% increase in sensitivity 516 per 10% RH increase), albert to various degrees (Figure 5i). Similar to long-chain 517 aldehydes, their $[MH + H_2O]^+$ ions also show a large positive deviation (Figure S16g), 518 and $[MH - H_2O]^+$ ions show little RH dependence (Figure S16e). These carbonyl 519 520 compounds have medium PA values (Figure 6a) except formaldehyde. If we exclude formaldehyde (with an extremely low PA value, the up triangle to the far left in Figure 521 6b) and long-chain aldehydes (low percentages of MH⁺ ions, <25%, down triangles in 522 Figure 6b), the RH dependence of other carbonyl compounds shows a slightly 523 decreasing trend of RH dependence vs. PA values (Figure 6b, up triangles, left triangles, 524 525 right triangles, and double triangles). These observations might hint on the relationships between the RH dependence of carbonyl compounds and reagent ion distribution as 526 well as reaction direction for R1-R4, which is different from those for pure 527 hydrocarbons (groups A1 and A2). Finally, the RH dependence of MH⁺ for compounds 528 529 in groups C1 (nitriles) and C2 (methanol) is within +1% per 10% RH increase (Figure 5i). 530

Overall, in the whole RH range studied (~5% to ~85%), the RH dependence of 531 MH⁺ ions for the 21 VOCs studied is less than 30%, with most compounds (except 532 group B2, long-chain aldehydes) less than 15%. For $[MH + H_2O]^+$ ions (mainly for 533 carbonyl compounds), strong RH dependence was observed (Figure S15), being 1.4 -534 8.5% per 10% RH increase, or 8.9 - 63.2% from ~5% to ~85% RH. The dehydrated 535 ions ($[MH - H_2O]^+$), however, show the smallest RH dependence (±1% per 10% RH 536 increase) among all the ions (Figure S16). Fragmented ions with decarbonization ([MH 537 $-C_xH_y$ ⁺) show mainly negative RH dependence, generally less than 3% per 10% RH 538 539 increase (Figure S17).

540 **4 Conclusions**

541 We investigated the response of protonated, adduct, and fragmented ions of 21 542 atmospherically relevant VOCs in a Vocus PTR-MS as instrument setting and RH 543 condition vary. For the two ways of increasing the E/N ratio, increasing the FIMR axial

voltage can substantially (by three to four orders of magnitude) increase sensitivity but 544 does not change the fractions of the MH⁺ ions (mostly within 30%); reducing the FIMR 545 pressure, however, does not enhance sensitivity much but can lead to more substantial 546 fragmentation. Therefore, a high FIMR axial voltage of 600 - 700 volts and a medium 547 pressure of around 2.0 mbar are recommended. Increasing the RF amplitude of FIMR 548 can increase sensitivity by 1 to 1.5 orders of magnitude at 2.0 mbar and 1.5 to 2 orders 549 of magnitude at 3.5 mbar, and it does not change the MH⁺ ion fractions (within 20%). 550 Therefore, a high RF amplitude of 500 V is recommended. Increasing the BSQ 551 552 amplitude does not increase the sensitivity much but changes the MH⁺ ion fractions of small ions substantially by changing the transmission efficiency. The choice of this 553 instrument setting mainly relies on what ions (i.e., those reagent ions with too high an 554 abundance) one wants to filter out. Our choice is at 300 V, which gives a 50% 555 transmission at about 55 Th. 556

557 The relationship between sensitivity and k_{ptr} is strongly affected by two factors: 1) whether the MH⁺ ion has a high transmission efficiency, and 2) whether the MH⁺ ion 558 is the dominating ion. If so, a fairly good correlation ($R_{Pr} = 0.77$) was observed for the 559 VOCs studied. The transmission curve is also more reasonably resembling the sigmoid 560 561 function only if all the ions (protonated, adduct, and fragmented) are considered. The low transmission efficiencies of formaldehyde and methanol result in extremely low 562 sensitivities of these two small OVOCs, although a low PA value is another reason for 563 564 the former.

As RH increases from $\sim 5\%$ to $\sim 85\%$, the MH⁺ ions for 19 out of the 21 VOCs 565 studied have sensitivity variation of less than 15%, but long-chain aldehydes have 566 positive RH dependence of up to 30%. The RH dependence of $[MH + H_2O]^+$ ions for 567 long-chain aldehydes is stronger, while that of the dominating $[MH - H_2O]^+$ ions is 568 limited. Therefore, the signal distributions among protonated, adduct, and fragmented 569 ions are also affected by RH variation. Together with their relatively high background 570 signals (especially for n-butanal, Figure S5b), quantification of long-chain aldehydes 571 in the ambient environment using Vocus requires special attention. It is also worth 572 573 noting that hydrocarbons generally show slight negative RH dependence, probably due to their relatively low k_{ptr} values, although such RH dependence does not affect 574 quantification significantly; their RH dependence has a fairly good correlation with 575 their PA values, hinting a thermodynamic reason behind this trend. 576

577

578 **Competing interests**

579 The authors declare that they have no conflict of interest.

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585 **References**

- Baasandorj, M., Millet, D. B., Hu, L., Mitroo, D., and Williams, B. J.: Measuring acetic and formic acid
 by proton-transfer-reaction mass spectrometry: sensitivity, humidity dependence, and quantifying
 interferences, Atmos. Meas. Tech., 8, 1303-1321, <u>https://doi.org/10.5194/amt-8-1303-2015</u>, 2015.
- Cappellin, L., Karl, T., Probst, M., Ismailova, O., Winkler, P. M., Soukoulis, C., Aprea, E., Märk, T. D.,
 Gasperi, F., and Biasioli, F.: On Quantitative Determination of Volatile Organic Compound
 Concentrations Using Proton Transfer Reaction Time-of-Flight Mass Spectrometry, Environmental
 Science & Technology, 46, 2283-2290, https://doi.org/10.1021/es203985t, 2012.
- Claflin, M. S., Pagonis, D., Finewax, Z., Handschy, A. V., Day, D. A., Brown, W. L., Jayne, J. T., Worsnop,
 D. R., Jimenez, J. L., Ziemann, P. J., de Gouw, J., and Lerner, B. M.: An in situ gas chromatograph with
 automatic detector switching between PTR- and EI-TOF-MS: isomer-resolved measurements of indoor
- air, Atmospheric Measurement Techniques 14, 133-152, <u>https://doi.org/10.5194/amt-14-133-2021</u>, 2021.
- 597 Cui, L., Zhang, Z., Huang, Y., Lee, S. C., Blake, D. R., Ho, K. F., Wang, B., Gao, Y., Wang, X. M., and
- Louie, P. K. K.: Measuring OVOCs and VOCs by PTR-MS in an urban roadside microenvironment of
 Hong Kong: relative humidity and temperature dependence, and field intercomparisons, Atmos. Meas.
 Tech., 9, 5763-5779, <u>https://doi.org/10.5194/amt-9-5763-2016</u>, 2016.
- 601 de Gouw, J., and Warneke, C.: Measurements of volatile organic compounds in the earths atmosphere 602 using proton-transfer-reaction mass spectrometry, Mass Spectrom Rev, 26, 223-257, 603 https://doi.org/10.1002/mas.20119, 2007.
- de Gouw, J. A., Goldan, P. D., Warneke, C., Kuster, W. C., Roberts, J. M., Marchewka, M., Bertman, S.
- B., Pszenny, A. A. P., and Keene, W. C.: Validation of proton transfer reaction-mass spectrometry (PTRMS) measurements of gas-phase organic compounds in the atmosphere during the New England Air
 Quality Study (NEAQS) in 2002, Journal of Geophysical Research: Atmospheres, 108, https://doi.org/10.1029/2003JD003863, 2003.
- Ellis, A. M., and Mayhew, C. A.: Chemical Ionization: Chemistry, Thermodynamics and Kinetics, in:
 Proton Transfer Reaction Mass Spectrometry, 25-48, 2014.
- Gueneron, M., Erickson, M. H., VanderSchelden, G. S., and Jobson, B. T.: PTR-MS fragmentation
 patterns of gasoline hydrocarbons, International Journal of Mass Spectrometry, 379, 97-109,
 https://doi.org/10.1016/j.ijms.2015.01.001, 2015.
- Hansel, A., Jordan, A., Holzinger, R., Prazeller, P., Vogel, W., and Lindinger, W.: Proton transfer reaction
 mass spectrometry: on-line trace gas analysis at the ppb level, International Journal of Mass Spectrometry
 and Ion Processes, 149-150, 609-619, <u>https://doi.org/10.1016/0168-1176(95)04294-U</u>, 1995.
- Hartungen, E. v., Wisthaler, A., Mikoviny, T., Jaksch, D., Boscaini, E., Dunphy, P. J., and Märk, T. D.:
- Proton-transfer-reaction mass spectrometry (PTR-MS) of carboxylic acids: Determination of Henry's law constants and axillary odour investigations, International Journal of Mass Spectrometry, 239, 243-248, https://doi.org/10.1016/j.ijms.2004.09.009, 2004.
- Inomata, S., Tanimoto, H., Kameyama, S., Tsunogai, U., Irie, H., Kanaya, Y., and Wang, Z.: Technical
 Note: Determination of formaldehyde mixing ratios in air with PTR-MS: laboratory experiments and
 field measurements, Atmospheric Chemistry and Physics, 8, 273-284, <u>https://doi.org/10.5194/acp-8-273-</u>
 2008, 2008.
- Jensen, A., Liu, Z., Tan, W., Dix, B., Chen, T., Koss, A., Zhu, L., Li, L., and de Gouw, J.: Measurements
 of Volatile Organic Compounds During the COVID-19 Lockdown in Changzhou, China, Geophysical
 Research Letters, 48, e2021GL095560, https://doi.org/10.1029/2021GL095560, 2021.
- Kari, E., Miettinen, P., Yli-Pirilä, P., Virtanen, A., and Faiola, C. L.: PTR-ToF-MS product ion
 distributions and humidity-dependence of biogenic volatile organic compounds, International Journal of
 Mass Spectrometry, 430, 87-97, https://doi.org/10.1016/j.ijms.2018.05.003, 2018.
- Krechmer, J., Lopez-Hilfiker, F., Koss, A., Hutterli, M., Stoermer, C., Deming, B., Kimmel, J., Warneke,
 C., Holzinger, R., Jayne, J., Worsnop, D., Fuhrer, K., Gonin, M., and de Gouw, J.: Evaluation of a New
 Reagent-Ion Source and Focusing Ion–Molecule Reactor for Use in Proton-Transfer-Reaction Mass
- 634 Spectrometry, Anal Chem, 90, 12011-12018, <u>https://doi.org/10.1021/acs.analchem.8b02641</u>, 2018.
- Li, H., Almeida, T. G., Luo, Y., Zhao, J., Palm, B. B., Daub, C. D., Huang, W., Mohr, C., Krechmer, J.
 E., Kurtén, T., and Ehn, M.: Fragmentation inside proton-transfer-reaction-based mass spectrometers
 limits the detection of ROOR and ROOH peroxides, Atmos. Meas. Tech., 15, 1811-1827,
 https://doi.org/10.5194/amt-15-1811-2022, 2022.
- 639 Lindinger, W., Hansel, A., and Jordan, A.: On-line monitoring of volatile organic compounds at pptv
- 640 levels by means of proton-transfer-reaction mass spectrometry (PTR-MS) medical applications, food
- control and environmental research, International Journal of Mass Spectrometry and Ion Processes, 173,
 191-241, <u>https://doi.org/10.1016/S0168-1176(97)00281-4</u>, 1998.
- 643 Materić, D., Lanza, M., Sulzer, P., Herbig, J., Bruhn, D., Gauci, V., Mason, N., and & Turner, C.: Selective

- reagent ion-time of flight-mass spectrometry study of six common monoterpenes, International Journal
 of Mass Spectrometry, 421, 40-50, <u>https://doi.org/10.1016/j.ijms.2017.06.003</u>, 2017.
- 646 Michoud, V., Sauvage, S., Léonardis, T., Fronval, I., Kukui, A., Locoge, N., and Dusanter, S.: Field
- measurements of methylglyoxal using proton transfer reaction time-of-flight mass spectrometry and
 comparison to the DNPH-HPLC-UV method, Atmospheric Measurement Techniques, 11, 5729-5740,
 https://doi.org/10.5194/amt-11-5729-2018, 2018.
- Pagonis, D., Sekimoto, K., and de Gouw, J.: A Library of Proton-Transfer Reactions of H3O+ Ions Used
 for Trace Gas Detection, J Am Soc Mass Spectr, 30, 1330-1335, <u>https://doi.org/10.1007/s13361-019-</u>
 02209-3, 2019.
- Pleil, J. D., Hansel, A., and Beauchamp, J.: Advances in proton transfer reaction mass spectrometry
 (PTR-MS): applications in exhaled breath analysis, food science, and atmospheric chemistry, J Breath
 Res, 13, <u>https://doi.org/10.1088/1752-7163/ab21a7</u>, 2019.
- Salazar Gómez, J. I., Sojka, M., Klucken, C., Schlögl, R., and Ruland, H.: Determination of trace
 compounds and artifacts in nitrogen background measurements by proton transfer reaction time-of-flight
 mass spectrometry under dry and humid conditions, J Mass Spectrom, 56, e4777,
 https://doi.org/10.1002/jms.4777, 2021.
- Schripp, T., Etienne, S., Fauck, C., Fuhrmann, F., Märk, L., and Salthammer, T.: Application of protontransfer-reaction-mass-spectrometry for Indoor Air Quality research, Indoor Air, 24, 178-189,
 https://doi.org/10.1111/ina.12061, 2014.
- Sekimoto, K., Li, S.-M., Yuan, B., Koss, A., Coggon, M., Warneke, C., and de Gouw, J.: Calculation of 663 the sensitivity of proton-transfer-reaction mass spectrometry (PTR-MS) for organic trace gases using 664 of 665 molecular properties, International Journal Mass Spectrometry, 421, 71-94, https://doi.org/10.1016/j.ijms.2017.04.006, 2017. 666
- Sekimoto, K., and Koss, A. R.: Modern mass spectrometry in atmospheric sciences: Measurement of
 volatile organic compounds in the troposphere using proton-transfer-reaction mass spectrometry, J Mass
 Spectrom, 56, <u>https://doi.org/10.1002/jms.4619</u>, 2021.
- Shao, P., An, J., Xin, J., Wu, F., Wang, J., Ji, D., and Wang, Y.: Source apportionment of VOCs and the
 contribution to photochemical ozone formation during summer in the typical industrial area in the
 Yangtze River Delta, China, Atmospheric Research, 176-177, 64-74,
 https://doi.org/10.1016/j.atmosres.2016.02.015, 2016.
- Shrivastava, M., Cappa, C. D., Fan, J., Goldstein, A. H., Guenther, A. B., Jimenez, J. L., Kuang, C.,
 Laskin, A., Martin, S. T., Ng, N. L., Petaja, T., Pierce, J. R., Rasch, P. J., Roldin, P., Seinfeld, J. H.,
 Shilling, J., Smith, J. N., Thornton, J. A., Volkamer, R., Wang, J., Worsnop, D. R., Zaveri, R. A., Zelenyuk,
- A., and Zhang, Q.: Recent advances in understanding secondary organic aerosol: Implications for global climate forcing, Reviews of Geophysics, 55, 509-559, https://doi.org/10.1002/2016RG000540, 2017.
- Sinha, V., Custer, T. G., Kluepfel, T., and Williams, J.: The effect of relative humidity on the detection of
 pyrrole by PTR-MS for OH reactivity measurements, International Journal of Mass Spectrometry, 282,
 108-111, https://doi.org/10.1016/j.ijms.2009.02.019, 2009.
- Smith, D., Diskin, A., Ji, Y. F., and Spanel, P.: Concurrent use of H3O+, NO+, and O-2(+) precursor ions
 for the detection and quantification of diverse trace gases in the presence of air and breath by selected
 ion-flow tube mass spectrometry, INTERNATIONAL JOURNAL OF MASS SPECTROMETRY, 209,
 81-97, https://doi.org/10.1016/S1387-3806(01)00478-X, 2001.
- Spaněl, P., and Smith, D.: SIFT studies of the reactions of H3O+, NO+ and O+2 with a series of volatile
 carboxylic acids and esters, International Journal of Mass Spectrometry and Ion Processes, 172, 137-147,
 https://doi.org/10.1016/S0168-1176(97)00246-2, 1998.
- Španěl, P., and Smith, D.: Selected ion flow tube: a technique for quantitative trace gas analysis of air
 and breath, Medical and Biological Engineering and Computing, 34, 409-419,
 <u>https://doi.org/10.1007/BF02523843</u>, 1996.
- Španěl, P., Ji, Y., and Smith, D.: SIFT studies of the reactions of H3O+, NO+ and O2+ with a series of
 aldehydes and ketones, International Journal of Mass Spectrometry and Ion Processes, 165-166, 25-37,
 https://doi.org/10.1016/S0168-1176(97)00166-3, 1997.
- Španěl, P., and Smith, D.: Influence of water vapour on selected ion flow tube mass spectrometric
 analyses of trace gases in humid air and breath, Rapid Communications in Mass Spectrometry, 14, 18981906, https://doi.org/10.1002/1097-0231(20001030)14:20<1898::AID-RCM110>3.0.CO;2-G, 2000.
- 698 Steinbacher, M., Dommen, J., Ammann, C., Spirig, C., Neftel, A., and Prevot, A. S. H.: Performance
- characteristics of a proton-transfer-reaction mass spectrometer (PTR-MS) derived from laboratory and
 field measurements, International Journal of Mass Spectrometry, 239, 117-128,
 https://doi.org/10.1016/j.ijms.2004.07.015, 2004.
- Vlasenko, A., Macdonald, A. M., Sjostedt, S. J., and Abbatt, J. P. D.: Formaldehyde measurements by
 Proton transfer reaction Mass Spectrometry (PTR-MS): correction for humidity effects, Atmospheric

- 704 Measurement Techniques, 3, 1055-1062, https://doi.org/10.5194/amt-3-1055-2010, 2010.
- 705 Wang, Y., Yang, G., Lu, Y., Liu, Y., Chen, J., and Wang, L.: Detection of gaseous dimethylamine using
- vocus proton-transfer-reaction time-of-flight mass spectrometry, Atmospheric Environment, 243, 117875,
 <u>https://doi.org/10.1016/j.atmosenv.2020.117875</u>, 2020.
- 708 Warneke, C., van der Veen, C., Luxembourg, S., de Gouw, J. A., and Kok, A.: Measurements of benzene
- and toluene in ambient air using proton-transfer-reaction mass spectrometry: calibration, humidity
 dependence, and field intercomparison, International Journal of Mass Spectrometry, 207, 167-182,
- 711 <u>https://doi.org/10.1016/S1387-3806(01)00366-9</u>, 2001.
- Warneke, C., De Gouw, J. A., Kuster, W. C., Goldan, P. D., and Fall, R.: Validation of atmospheric VOC
 measurements by proton-transfer-reaction mass spectrometry using a gas-chromatographic preseparation
 method, Environ Sci Technol, 37, 2494-2501, https://doi.org/10.1021/es026266i, 2003.
- 715 Warneke, C., Veres, P., Holloway, J. S., Stutz, J., Tsai, C., Alvarez, S., Rappenglueck, B., Fehsenfeld, F.
- C., Graus, M., Gilman, J. B., and de Gouw, J. A.: Airborne formaldehyde measurements using PTR-MS:
 calibration, humidity dependence, inter-comparison and initial results, Atmos. Meas. Tech., 4, 2345-2358,
- 718 <u>https://doi.org/10.5194/amt-4-2345-2011</u>, 2011.
- Yuan, B., Koss, A. R., Warneke, C., Coggon, M., Sekimoto, K., and de Gouw, J. A.: Proton-Transfer Reaction Mass Spectrometry: Applications in Atmospheric Sciences, Chemical Reviews, 117, 13187-
- 721 13229, <u>https://doi.org/10.1021/acs.chemrev.7b00325</u>, 2017.
- 722Zhao, J., and Zhang, R.: Proton transfer reaction rate constants between hydronium ion (H3O+) and723volatileorganiccompounds,AtmosphericEnvironment,38,2177-2185,
- 724 <u>https://doi.org/10.1016/j.atmosenv.2004.01.019</u>, 2004.
- 725