

1 **Characterization of offline analysis of particulate matter with**
2 **FIGAERO-CIMS**

3 Jing Cai^{1,2,†}, Kaspar R. Daellenbach^{1,2,3,‡,*}, Cheng Wu^{4,5}, Yan Zheng⁶, Feixue Zheng¹, Wei Du^{1,2}, Sophie L. Haslett⁴,
4 Qi Chen⁶, Markku Kulmala^{1,2}, Claudia Mohr^{4,*}

5 ¹ Aerosol and Haze Laboratory, Beijing Advanced Innovation Center for Soft Matter Science and Engineering,
6 Beijing University of Chemical Technology, Beijing 100029, China

7 ² Institute for Atmospheric and Earth System Research, Faculty of Science, University of Helsinki, Helsinki 00014,
8 Finland

9 ³ Laboratory of Atmospheric Chemistry, Paul Scherrer Institute, Villigen, Switzerland.

10 ⁴ Department of Environmental Science, Stockholm University, Stockholm, 11418, Sweden

11 ⁵ Department of Chemistry and Molecular Biology, Atmospheric Science, University of Gothenburg, Gothenburg,
12 SE-412 96, Sweden

13 ⁶ State Key Joint Laboratory of Environmental Simulation and Pollution Control, Beijing Innovation Center for
14 Engineering Science and Advanced Technology, College of Environmental Science and Engineering, Peking
15 University, Beijing, 100871, China

16 [†] These authors contributed equally to this work.

17 Correspondence to: kaspar.dallenbach@helsinki.fi and claudia.mohr@aces.su.se

18 **Abstract:** Measurements of the molecular composition of organic aerosol (OA) constituents improve our
19 understanding of sources, formation processes, and physicochemical properties of OA. One instrument providing
20 such data at a time resolution of minutes to hours is the Chemical Ionization time-of-flight Mass Spectrometer
21 with Filter Inlet for Gases and AEROsols (FIGAERO-CIMS). The technique collects particles on a filter, which
22 are subsequently desorbed, and the evaporated molecules are ionized and analyzed in the mass spectrometer.
23 However, long-term measurements using this technique and/or field deployments at several sites simultaneously,
24 require substantial human and financial resources. The analysis of filter samples collected outside the instrument
25 (offline) may provide a more cost-efficient alternative and makes this technology available for the large number
26 of particle filter samples collected routinely at many different sites globally. Filter-based offline use of the
27 FIGAERO-CIMS limits this method albeit to particle-phase analyses, likely at reduced time resolution compared
28 to online deployments. Here we present the application and assessment of offline FIGAERO-CIMS, using Teflon
29 and Quartz fiber filter samples that were collected in autumn 2018 in urban Beijing. We demonstrate the feasibility
30 of the offline application with “sandwich” sample preparation for the identified over 900 organic compounds with
31 (1) high signal-to-noise ratios, (2) high repeatability, and (3) linear signal response to the filter loadings.
32 Comparable overall signals were observed between the Quartz fiber and Teflon filters for 12-h and 24-h samples,
33 but with larger signals for semi-volatile compounds for the Quartz fiber filters, likely due to adsorption artifacts.
34 We also compare desorption profile (thermogram) shapes for the two filter materials. Thermograms are used to
35 derive volatility qualitatively based on the desorption temperature at which the maximum signal intensity of a
36 compound is observed (T_{max}). While we find that T_{max} can be determined with high repeatability ($\pm 5.7^\circ\text{C}$), from
37 the duplicate tests for one filter type, we observe considerable differences in T_{max} between the Quartz and Teflon
38 filters, warranting further investigation into the thermal desorption characteristics of different filter types. Overall,
39 this study provides a basis for expanding OA molecular characterization by FIGAERO-CIMS to situations where
40 and when deployment of the instrument itself is not possible.

41 **1. Introduction**

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42 Molecular information on organic aerosol (OA) composition is important for understanding the role that OA
43 plays in the atmosphere regarding its impacts on air quality, human health, and the climate (Daellenbach et al.,
44 2020; Huang et al., 2014; Cappa et al., 2012; Yao et al., 2018; Riipinen et al., 2012). Such data can be obtained
45 from offline filter collection and analysis in the laboratory using optical (e.g. Fourier transform infrared
46 spectroscopy, FTIR) and magnetic (e.g. Nuclear magnetic resonance spectroscopy, NMR) spectroscopy or, more
47 commonly, high-resolution mass spectrometer methods, which include gas/liquid chromatography coupled to
48 mass spectrometry (GC/LC-MS), ultrahigh-performance liquid chromatography coupled to Orbitrap mass
49 spectrometry and electrospray ionization mass spectrometry (ESI-MS) (Noziere et al., 2015). In contrast, online
50 mass spectrometers provide direct and in-situ information on particles' molecular composition, e.g. the filter inlet
51 for gases and aerosols coupled to a high-resolution time-of-flight chemical ionization mass spectrometer
52 (FIGAERO-HR-ToF-CIMS, Aerodyne Research Inc., US, hereafter FIGAERO-CIMS (Lopez-Hilfiker et al.,
53 2014)) or the extractive electrospray ionization time-of-flight mass spectrometer (EESI-MS) (Lopez-Hilfiker et
54 al., 2019). Since the particle-phase measurement by FIGAERO-CIMS is filter-based, it has the potential to be used
55 for offline analysis. Briefly, in the FIGAERO, particles are collected on a Teflon® (hereafter Teflon) filter and
56 analyzed via thermal desorption. When coupled to a high-resolution time-of-flight chemical-ionization mass
57 spectrometer (hereafter CIMS), molecular composition information of inorganic and organic aerosol compounds
58 that evaporate at temperatures up to 200 °C can be achieved. Having the advantage of combining molecular
59 composition and volatility information, the FIGAERO-CIMS has been widely used for measuring OA compounds
60 in many different environments including e.g. forests (Lopez-Hilfiker et al., 2016; Lee et al., 2016; Lee et al., 2018;
61 Mohr et al., 2019), rural and urban areas (Le Breton et al., 2019; Huang et al., 2019b; Cai et al., 2022), indoor air
62 (Farmer et al., 2019), and cooking emissions (Masoud et al., 2022).

63 Both online and offline techniques have their advantages and disadvantages and are associated with artefacts
64 (Turpin and Lim, 2001; Turpin et al., 2000). Both online and offline techniques have their advantages and
65 disadvantages and are associated with artefacts (Turpin and Lim, 2001; Turpin et al., 2000). Offline techniques
66 are an easy alternative to demanding online in-situ approaches requiring large human and financial resources.
67 Moreover, one collected filter can be used for different analysis methods and purposes. However, the offline
68 approaches are susceptible to sample handling and storage artefacts. The condensation and re-evaporation of
69 vapors, and potential reactions on the filter during sampling and storage can result in both positive and negative
70 sampling biases (Turpin et al., 2000; Cheng et al., 2009). Online instruments generally allow for measurements at
71 higher time resolution, which is an advantage when studying rapid atmospheric processes, and no sample storage
72 is needed before analysis. However, the deployment of the FIGAERO-CIMS outside the laboratory requires a
73 well-equipped site that is easily accessible. In addition, long-term maintenance of these complex mass
74 spectrometers needs substantial human and financial resources. Therefore, deployments are often achieved only
75 for short periods (i.e. campaigns lasting from a couple of weeks to months), which limits the application of this
76 technique for monitoring and simultaneous measurements at multiple sites. Furthermore, FIGAERO gas-phase
77 measurements have to be interrupted regularly for particle-phase analysis in online usage, which could be a
78 problem for measurements requiring high time resolution data (e.g. chamber studies). Using the FIGAERO-CIMS
79 for analyzing filters collected elsewhere ("offline application") may therefore provide a valid alternative for long-
80 term monitoring or simultaneous measurements at multiple sites. Whereas the online FIGAERO-CIMS technique
81 typically uses Teflon filters to reduce interferences from the gas phase. Quartz fiber filters are widely used for
82 offline sampling of OA due to their high melting point and insolubility in water and typical organic solvents
83 (Watson and Chow, 2002; Tao et al., 2017; Schauer et al., 2002; Gustafson and Dickhut, 1997). Up to now, only
84 a few studies have used the FIGAERO-CIMS in offline mode with Teflon filters (Siegel et al., 2020; Huang et al.,
85 2019a), and an in-depth characterization of the method is missing. The performance of Quartz fiber filters in
86 FIGAERO-CIMS needs to be assessed and compared to Teflon filters.

87 Here, we describe the application of FIGAERO-CIMS in offline mode for the analysis of particles deposited on
88 Teflon and Quartz fiber filters in urban Beijing during the autumn and winter of 2018. The filter deposition time
89 varies from 30 min to 24 h. We assess the performance of FIGAERO-CIMS for offline characterization of OA as

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91 well as inorganic compounds and discuss background determination, reproducibility, and linearity of response for
92 the two filter types. We describe filter handling and offline analysis procedures and show the comparison of signals
93 from different mass loadings collected on both filter types. The utility of the FIGAERO for offline use is
94 demonstrated in this study. The potential to broaden its application for OA component measurements in future
95 research is also discussed. We note, however, that it is not the scope of this paper to discuss aspects of offline
96 FIGAERO-CIMS that also apply to its online deployment, such as e.g. general percentage of recovery from the
97 filter or calibrations.

98 **2. Methods**

99 **2.1 Filter sampling**

100 The sampling site is situated on the west campus of the Beijing University of Chemical Technology (BUCT, 39°
101 56'31" N, 116°17'50" E). BUCT is located near the West Third Ring Road of Beijing, surrounded by residential
102 areas. A more detailed description of the sampling site can be found elsewhere (Cai et al., 2020; Kontkanen et al.,
103 2020; Liu et al., 2020; Yao et al., 2020; Fan et al., 2021; Guo et al., 2021). From November to December 2018,
104 samples of fine particulate matter with an aerodynamic diameter of up to 2.5 μm (PM_{2.5}) were collected by a four-
105 channel sampler (TH-16A, Tianhong Co., China) with a sampling flow rate of 16.7 L min⁻¹, installed on the rooftop
106 of a five-floor building (~20m above ground). Both Teflon (Zefluor® PTFE membrane, 1 μm pore size, 47 mm
107 diameter, Pall Corp., US) and Quartz fiber filters (7202, 47 mm diameter, Pall Corp., US) were collected
108 simultaneously at separate channels. The four parallel channels of the sampler had a shared PM₁₀ cyclone inlet and
109 were equipped with 4 independent PM_{2.5} cyclones and auto flow controllers for each channel. All channels were
110 measuring the same size range of particles. A sizing effect from the interactions between different channels can
111 therefore be neglected. The setup of filter type for each channel was as follows: Channel 1, Teflon (12 h for or 0.5
112 h); Channel 2, Quartz (12 h or 0.5 h); Channel 3, Teflon (24 h or 2.5 h); Channel 4, Quartz filters (24 h or 2.5 h).
113 This is listed in Table 1. The flow rate was regularly calibrated individually for each channel during the sampling
114 process.

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115 To investigate the influence of filter mass loadings and collection time on the signal response, the following filter
116 samples were taken: (1) 5 pairs of samples (Teflon/Quartz fiber filters, Channels 1 and 2) with 30 min deposition
117 time on Dec 15, 2018 between 14:00 to 16:30 (Table 1). At the same time, an additional pair of Teflon/Quartz
118 samples were deposited for 2.5 hours using the other two separate channels of the sampler (Channels 3 and 4). (2)
119 12-h samples of Quartz/Teflon filters (Channels 1 and 2) from Oct 26 to Oct 30 and Nov 3 to Nov 24 (here only
120 the Quartz filters from Nov 3 to Nov 16 were analyzed (in total 27 pairs of samples), shown in Table 1). (3) 24-h
121 Quartz/Teflon samples (Channels 3 and 4) from Oct 26 to Oct 30 and Nov 3 to Nov 25 (here only one pair of
122 Teflon/Quartz filters was analyzed, shown in Table 1). During the last sampling period, high PM_{2.5} and relative
123 humidity (RH) conditions prevailed (Nov 3: 181 $\mu\text{g m}^{-3}$, 60%, and Nov 13: 227 $\mu\text{g m}^{-3}$, 75%), and the channel of
124 the 24-h sampling Teflon filter got clogged. Thus, only one pair of 24-h Teflon/Quartz samples from this period
125 was analyzed (Table 1).

126 Detailed information on the sampling protocol is listed in Table 1. Three pairs (Teflon/Quartz) of field blank
127 samples were also collected during the sampling period. Before sampling, Teflon filters were baked for 2 hours at
128 200 °C, which is much longer than the typical desorption time for FIGAERO-CIMS online usage (Ylisirniö et al.,
129 2021), and Quartz filters for 4.5 hours at 550 °C (Liu et al., 2016) in order to minimize contamination. After
130 sampling, samples were put in filter holders wrapped in pre-baked aluminum foils, individually sealed in a sealed
131 bag and stored in a freezer at -20 °C for 7 months until being analyzed in the laboratory.

132 To calculate the OA mass loadings of the samples, an online Time-of-Flight-Aerosol Chemical Speciation
133 Monitor (Aerodyne Research Inc., US, hereafter ToF-ACSM) equipped with a PM_{2.5} lens and standard vaporizer
134 was operated during the sampling period at the same site. Details of the ToF-ACSM settings can be found in Cai

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137 et al. (2022). The OA loading on each filter (OA_{filter}) was determined relying on the OA concentrations from the
 138 co-located TOF-ACSM (OA_{ACSM}), the offline filter sampling flow rate (16.7 L min^{-1}), the sampling time, the
 139 surface of the entire offline filter sample (A_{filter}), and the analyzed offline filter sample (A_{punch}) (Equation 1):

$$OA_{filter} = \frac{A_{punch}}{A_{filter}} \times OA_{ACSM} \times \text{Sampling flow rate} \times \text{Sampling time} \quad (1)$$

141 Table 1: Testing objectives, filter deposition dates and times, flows, filter material (T = Teflon, Q = Quartz fiber),
 142 filter mass loadings of OA, number of samples, and number of sample repeats (filter punches) for the same filter.

| Testing objective | Sampling date | Sampling time | Filter material | OA loading [μg] per punch (punch diameter, area) | Number of samples/repeats |
|--|------------------------------------|---------------|-----------------|---|---|
| (1) Baseline subtraction tests, (2) reproducibility tests, (3) filter type comparison | Dec 15 | | | | |
| | 14:00 – 16:30 (30 min-interval) | 30 min | T & Q | 1.7×10^{-2} – 2.0×10^{-2} (2 mm, 0.031 cm^2) | 1/1 |
| | Dec 15: 14:00 – 16:30 | 2.5 h | T & Q | 9.1×10^{-2} (2 mm, 0.031 cm^2) | 1/3 for repeats |
| (1) Reheating tests, (2) filter type comparison | Nov 8 21:30 – Nov 9 9:00 | 12 h | T & Q | 6.5×10^{-1} (2 mm, 0.031 cm^2) | 1/1 |
| Reheating tests | Nov 12 21:30 – Nov 13 9:00 | 12 h | Q | 0.75 (2 mm, 0.031 cm^2) | 1/1 |
| Reheating tests | Nov 13 21:30 – Nov 14 9:00 | 12 h | Q | 1.2 (2 mm, 0.031 cm^2) | 1/1 |
| (1) Filter type comparison, (2) different ramping protocols for 2 mm punch, (3) linearity response for signals from different filter punch areas | Nov 24 9:30 – 9:00 25 | 24 h | T & Q | 1.2 (2 mm, 0.031 cm^2) | 1/3 for repeats and 1/3 for different ramping protocols |
| | | | | 2.7 (3 mm, 0.071 cm^2) | 1/1 |
| | | | | 4.8 (4 mm, 0.13 cm^2) | 1/1 |

| | | | | | |
|--|-----------------|------|---|--|------|
| | | | | 15 (7 mm, 0.38 cm ²) | |
| Comparison of 12-h signals to ToF-ACSM | Nov 3 to Nov 16 | 12 h | Q | $5.0 \times 10^{-2} - 1.2$ (2 mm, 0.031 cm ²) | 27/1 |

144

145

146 **2.2 Offline application of FIGAERO-CIMS**147 **2.2.1 Measurement approach**148 **2.2.1.1 FIGAERO-CIMS setup**

149 The molecular composition of OA collected on the filter samples was characterized with FIGAERO-CIMS using
 150 iodide (I⁻) as the reagent ion. In typical online FIGAERO-CIMS operation, particles are collected on a filter
 151 (Zefluor[®] Teflon filters) with a sampling time of a few minutes to hours and then thermally desorbed by a flow of
 152 temperature-controlled ultra-pure nitrogen (99.999 %) immediately following deposition. The thermally desorbed
 153 compounds are charged by clustering with I⁻, which is typically generated through the exposure of methyl iodide
 154 to an X-ray or radioactive source for FIGAERO-CIMS (Po²¹⁰ in our study). In this study, we used the FIGAERO-
 155 CIMS in the laboratory to analyze filter samples collected earlier in the field. These samples were placed manually
 156 one by one in the dedicated filter holder of the FIGAERO-CIMS and the desorption procedure was started (see
 157 2.2.1.3).

158 **2.2.1.2 Sample preparation and test design**

159 Since the total particle mass collected on one filter was generally too large to be analyzed at once in its entirety
 160 by FIGAERO-CIMS (due to the risk of titration of the reagent ion), we only analyzed small circular punches of
 161 the collected filters. The default punching area was 3.1×10^{-2} cm² (punch diameter $d=2$ mm). In addition, to test the
 162 linearity of response to sample mass loadings, punch areas for the same filter were varied between 3.1×10^{-2} cm²
 163 ($d=2$ mm) and 0.38 cm² ($d=7$ mm), resulting in variation in mass loadings by a factor of 10 (shown in Table 1).
 164 Since the filter punches were too small for the filter holder of the FIGAERO, we put them between two pre-baked
 165 originally sized ($d=25$ mm) Zefluor[®] Teflon filters (“sandwich technique”, Fig. 1a). Field blanks were prepared
 166 analogously.

167 The OA mass loadings of the filter punches were estimated with the co-located ToF-ACSM in this study (details
 168 shown in Table 1). To test the performance of the method, we did the following tests (Fig. 1, Table 1): (1) reheating
 169 a few filters to determine backgrounds (see section 2.2.4), (2) assess different background subtraction methods,
 170 (3) reproducibility of signals from the same filter (section 3.4), (4) the linearity of signal response from different
 171 punching areas from the same filter (section 3.4), (5) comparing signals from different ramping protocols (section
 172 2.2.1.3), (6) comparison between and offline FIGAERO-CIMS and online ToF-ACSM (section 3.5), (7) signals
 173 from different filter types (section 3.6), and (8) thermograms from different types of filters (section 3.7).

174 **2.2.1.3 Temperature ramping protocols**

175 Reagent ion depletion is undesired as it can create non-linearities in the instrument response (Koss et al., 2018;
 176 Zheng et al., 2021). To avoid reagent ion depletion in FIGAERO-CIMS, the concentration of sample ions entering

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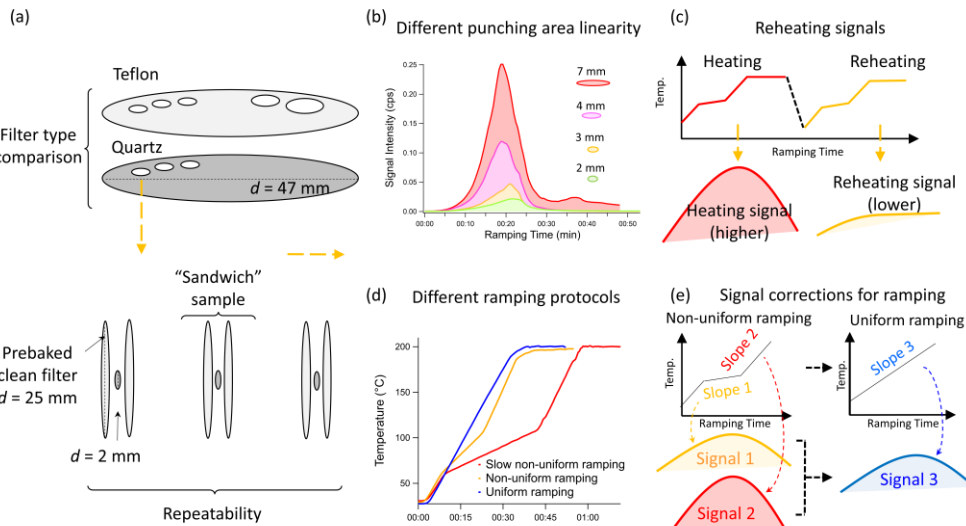
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182 the instrument is controlled, typically by modifying the particle mass loading on the filter and/or the heating rate.
183 While the particle mass loading can be varied easily when operating the FIGAERO-CIMS online through
184 adjustment of sampling time and flow, in offline mode with pre-collected samples this can only be modified by
185 the fraction of filter surface analyzed. For our Beijing filter samples, even when using the smallest punch sizes
186 ($3.1 \times 10^{-2} \text{ cm}^2$), mass loadings of especially nitric acid (HNO_3) were still high enough to lead to titration of the
187 reagent ion. We note that this can also be an issue for online measurements in presence of high nitrate
188 concentrations, e.g. in highly polluted areas. In order to reduce reagent ion depletion between 60 °C to 105 °C
189 desorption temperature, where HNO_3 exhibits a maximum signal, we used a heating protocol with a non-uniform
190 temperature ramping procedure. Instead of ramping from room temperature to 200 °C with a constant heating rate,
191 we divided the temperature ramp into several periods: (1) from room temperature (~25 °C) to 60 °C in 8 min
192 ($4.4 \text{ }^{\circ}\text{C min}^{-1}$), (2) from 60 °C to 105 °C in 15 min ($3 \text{ }^{\circ}\text{C min}^{-1}$), (3) from 105 °C to 200 °C in 12 min ($7.9 \text{ }^{\circ}\text{C min}^{-1}$).
193 The ramp period was followed by a 20-minute soaking period (200 °C) to allow signals to go to background
194 levels. We called this temperature ramping protocol non-uniform temperature ramping and used it as the default
195 desorption procedure in this study. The maximum reagent ion depletion achieved in this way was ~35% for the
196 samples with the highest mass loadings on a 2 mm punch, which was mostly used in this study. We also tested
197 two alternative heating protocols:

- 198 1) Slow non-uniform temperature ramping: Same as the non-uniform ramping protocol, but with (2)
199 slowed down to $1.5 \text{ }^{\circ}\text{C min}^{-1}$. The total heating time for this protocol was 70 minutes, and the
200 maximum reagent ion depletion was ~ 20%.
- 201 2) Uniform temperature ramping: The temperature was increased from room temperature to 200 °C
202 in 31.5 min ($5.7 \text{ }^{\circ}\text{C min}^{-1}$). Including the 20 min soak, the total heating was 51.5 minutes, and the
203 maximum reagent ion depletion was around 50%. In order to limit reagent ion depletion, the
204 heating rate was 1.8–3.5 times slower than typical rates used for online FIGAERO-CIMS
205 applications ($10\text{--}20 \text{ }^{\circ}\text{C min}^{-1}$ (Thornton et al., 2020)).

206 The 3 temperature ramping protocols are displayed in Fig. 1d. As different heating rates lead to different
207 thermogram shapes and T_{\max} for individual compounds, we developed a correction method in an effort to be able
208 to compare desorption-derived volatility for the different ramping protocols. This will be further discussed in
209 section 3.3.



212 **Figure 1.** Schematic of the tests conducted in this study, (a) sample preparation using punching areas of different sizes of the
 213 Teflon and Quartz fiber filters and squeezing them between two original-sized filters for analysis, (b) signal intensities of
 214 different punching areas from the same sample with the same analytical procedure, (c) reheating tests by conducting two
 215 consecutive heating cycles, (d) different temperature procedures, and (e) signal intensity correction from non-uniform ramping
 216 to uniform ramping.

217 2.2.2 Data analysis

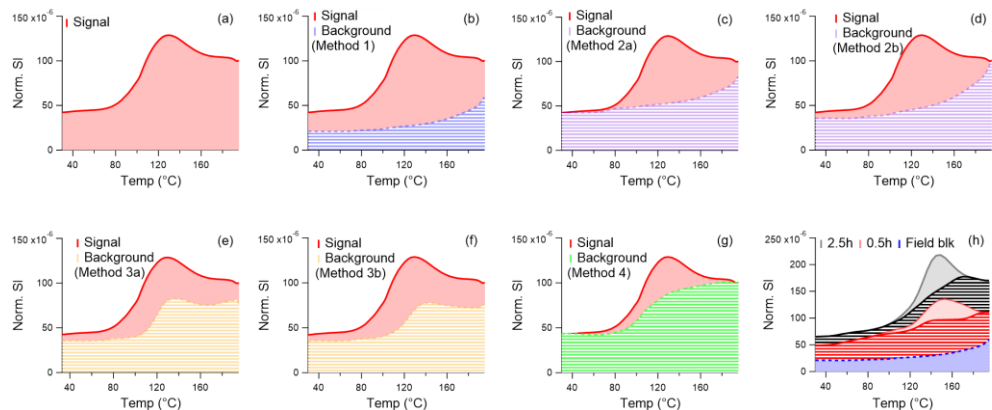
218 In this study, FIGAERO-CIMS data were analyzed with the Tofware package (v.3.1.0, Tofwerk, Switzerland,
 219 and Aerodyne, US) within the Igor Pro software (v.7.08, Wavemetrics, US). Mass accuracies of low- to high-mass
 220 species (~130 to 500 Da) were within ± 10 ppm for all the samples. A total of ~1,200 peaks were found in the range
 221 of 46 and 500 Da, of which 916 were identified as organic species. Detailed information about the identified
 222 chemical compounds can be found in Cai et al. (2022). The total signal of a compound per filter sample, defined
 223 as the integrated signals (I_s), calculated by first normalizing by the signals of the primary ions (I) and then
 224 integrating the entire thermogram (ramping and soaking, normalized by the signals of I). Signals of the first 1.5
 225 min of ramping and the last 1.5 min of soaking periods were excluded in order to remove potential interference
 226 from switching to and from the heating status. In this study, we use the term CHOX to represent all organic
 227 compounds identified by FIGAERO-CIMS, $C_{x \geq 1}H_{y \geq 1}O_{z \geq 1}X_{0-n}$, detected as clustered with I , with X being different
 228 atoms including N, S, Cl, or a combination of them.

229 2.2.3 Background subtraction

230 The background in offline FIGAERO-CIMS is a combination of instrument background and field blank. The
 231 field blanks provide information on sampling and handling artefacts, while the instrument background is mainly
 232 from (1) the desorption of semi-volatile or low-volatile compounds adsorbed on instrument surfaces (such as the
 233 ion-molecular reaction region (IMR)), and (2) impurity of the reagent ion precursors and carrier gases. Thus,
 234 instrument background signal can vary for different samples and depending on instrument status. For FIGAERO-
 235 CIMS online deployments, frequent blank measurements and calibrations are recommended (Bannan et al., 2018;

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237 Thornton et al., 2020). The common method for online FIGAERO-CIMS of placing an additional filter upstream
 238 of the FIGAERO filter is impossible for offline pre-sampled filters. Given 1) the large variation of the filter sample
 239 loadings ($\sim 1 \times 10^{-2}$ μg – 1.2 μg), which influences the number of compounds that can potentially adsorb to
 240 instrument surfaces, 2) the general scarcity of field blanks in offline mode compared to background filter samples
 241 in online FIGAERO-CIMS, and 3) that the instrument background can be influenced by instrument history very
 242 different from the offline sample due to the temporal separation of sample and analysis, choosing an appropriate
 243 instrumental and field blank determination method is crucial and challenging for offline FIGAERO-CIMS analysis.
 244 Here we describe and discuss performance of 6 different background subtraction methods (schematically shown
 245 in Fig. 2):

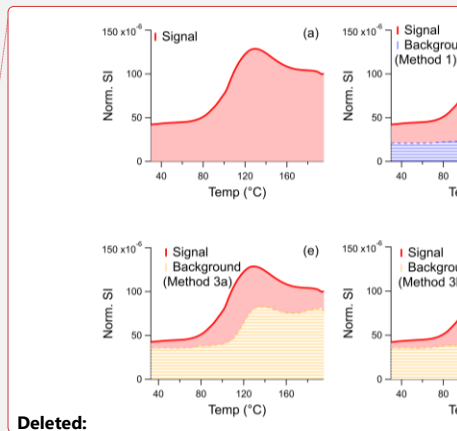


246
 247 **Figure 2.** Schematic of a compound's signal and background thermograms for different background determination methods.
 248 The x-axis is the temperature during ramping, and the y-axis is the signal intensity normalized by the primary ion (I^*). (a) total
 249 sample signal of a model compound without blank subtraction, (b) Method 1: field blank only, (c) Method 2a: scaling field
 250 blank to the start of ramping, (d) Method 2b: scaling field blank to the end of soaking, (e) Method 3a: reheating of a subset of
 251 filters, and using the average signal ratio from reheated and heated filters as background signal for all filters (individual
 252 compound-based), (f) Method 3b: reheating of a subset of filters, and using an exponential fit to the entire mass range of the
 253 average signal ratio from reheated and heated filters as background signal for all filters, (g) Method 4: thermal baseline using
 254 a spline algorithm, and (h) one 0.5-h and one 2.5-h sample with blank-subtraction. Ideally, the I_s of the 2.5-hour collection
 255 sample ($I_{s,2.5h}$) would be close to the sum of the 5 paralleled 0.5-hour collection sample ($I_{s,0.5h}$).

256 **Method 1:** Background is the average integrated signal intensity (I_s , the integrated signal of the thermograms
 257 shown in Fig. 2a) of field blanks ($\bar{I}_{s,field\ blk,i}$), which are three in our case (Fig. 2b). The integrated background-
 258 subtracted signal for compound i ($I_{s,blksub,i}$) is then $I_{s,i} - \bar{I}_{s,field\ blk,i}$.

259 **Method 2:** Background is field blank average ($\bar{I}_{s,field\ blk,i}$, see Method 1) scaled to the ratio of ambient sample
 260 and field blank signals during a reference period (ref period) – either prior to the start of heating (the first 1.5 to 3
 261 min of the ramping procedure before the temperature starts to increase, Method 2a or at the end of the soaking (the
 262 last 1.5 to 3 min of the soaking period, Method 2b). Method 2 corrects for variation in instrument background that
 263 is not necessarily related to the sample to be analyzed. The integrated background-subtracted signal for compound
 264 i ($I_{s,blksub,i}$) is then

$$I_{s,blksub,i} = \int I_{sample,ij} - \int I_{field\ blk,ij} \times \frac{\int_{ref\ period} I_{s,i,ambient}}{\int_{ref\ period} I_{s,i,field\ blk}} \quad (2)$$



267 By using Method 2a, it is assumed that the signal measured before heating, but with the filter already in place, is
268 due to instrument background, which can vary between the measurement of a sample filter and a blank filter (Fig.
269 2c). However, this method may lead to underestimation of the sample signal for compounds that already evaporate
270 at room temperature.

271 By using Method 2b, it is assumed that the signal measured at the end of soaking is due to instrument background,
272 which can vary between the measurement of a sample filter and a blank filter. The variation in instrument
273 background is taken into account at maximum heating temperature (200 °C) and thus elevated temperature of
274 surfaces downstream of the filter, and at the end of the soaking period when presumably all material that can
275 evaporate from the filter has evaporated (shown in Fig. S1).

276 **Method 3:** In this method (Siegel et al., 2021), the instrument background is assessed by heating the same filter
277 twice, assuming that during the first heating cycle, all detectable material has evaporated, and that what is measured
278 in a reheating cycle is the instrument background signal. Ideally, reheating would be done for each sample
279 individually. Since this was not done for our dataset, the instrument background determined based on a few reheatings
280 (3 in our case, the details of the reheating samples are shown in Table 1) had to be extrapolated to all samples
281 (Method 3a and 3b). It is clearly shown that the signals from the reheating cycle are much lower than those from
282 the first heating (Fig. S1) without a clear peak in thermograms for both filter types, suggesting sampled compounds
283 were well desorbed in the original heating cycle. Simple reheating does not consider the field blanks, which need
284 to be subtracted in addition.

285 For Method 3a we assumed that the ratio of the integrated signal of the second heating cycle (heating C2) and first
286 heating cycle (heating C1) of the same filter is influenced by volatility and therefore compound-dependent. Here
287 we used the average ratio from 3 reheating tests done for this dataset (Fig. S2). The distribution of the ratios is
288 shown in Fig. S3. The $I_{S_{blksub},i}$ was then calculated following Eq. 2, where the instrument background is the fraction
289 of the sample signal established from the re-heating, and added to the signal from the field blank, which is
290 calculated in the same way.

$$291 I_{S_{blksub},i} = \left(I_{S_{sample,i}} - I_{S_{sample,i}} \times I_{S_{i,(\text{heating C2},i)}} \right) \\ 292 - \left(I_{S_{field blk,i}} - I_{S_{field blk,i}} \times I_{S_{i,(\text{heating C2},i)}} \right) \quad (3)$$

293 For Method 3b, we assumed that the ratio of heating C2 to heating C1 exhibits a signal dependency (relatively
294 higher background for compounds with lower signal), calculated using an exponential fit to the data from the 3
295 reheat tests (Fig. S4) using Eq. (4) with the constants A, B, and C. The field blanks are calculated in the same way.
296 Then the $I_{S_{blksub}}$ can be calculated as in Eq. (3)

$$297 I_{S_{i,(\text{heating C2},i)}} = A + B \times \exp(I_{S_{sample,i}} + C) \quad (4)$$

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298 **Method 4:** Thermal baseline subtraction. In this method, we determined for every thermogram of each compound
299 a background thermogram termed thermal baseline ($I_{S_{thbsl}}$). The thermal baseline was computed using a spline
300 algorithm initially developed by Wang et al. (2018) for determining the background concentration of a pollutant
301 using its concentration time series (by determining the spline of background from varying time intervals).
302 Thermogram data were pre-averaged to 1.8 mins (corresponding to 4 data points of the original time resolution of
303 27s) to reduce noise for the thermal baseline computation. Field blanks were handled in the same way (shown in
304 Fig. S5). Thus, the blank-subtracted signal $I_{S_{blksub}}$ of a compound i is:

$$305 I_{S_{blksub},i} = I_{S_{sample,blksub,i}} - I_{S_{field blk,blksub,i}} \\ 306 = \left(\int I_{S_{sample,i,j}} - I_{S_{sample,thbsl,i}} \right) - \left(\int I_{S_{field blk,i,j}} - I_{S_{field blk,thbsl,i}} \right) \quad (5)$$

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312 $I_{sample, thbsl,i}$ and $I_{field blk, thbsl,I}$ represent the thermal baseline of compound i for samples and field blanks, respectively.

313 2.2.4 Thermograms and T_{max} recovery

314 The amount of compounds coming off the filter at a certain temperature varies as a function of temperature
315 ramping rates, resulting in different thermogram shapes and T_{max} (shown in Fig. 1d). This is especially important
316 in our case for the non-uniform ramping protocols. In an attempt to make the different cases comparable for
317 qualitative volatility studies, we developed a thermogram correction where the blank-subtracted signal as a
318 function of temperature for each compound i is re-distributed to constant temperature intervals (Eq. (6)):

$$319 I_{thermocorrected,i,j} = \int_{T-\Delta t}^T I_{sample,blksub,i,j} dT \quad (6)$$

320 Considering the ~ 2 °C variation in thermogram reproducibility reported from an online FIGAERO-CIMS study
321 (Lopez-Hilfiker et al., 2014), the temperature interval ΔT used in this study is 3°C.

322

323 3. Results

324 3.1 Assessment of the background: Signal comparison between different blank subtraction methods

325 To assess the influence of the 6 background methods on the resulting signal, Quartz fiber filter samples from 5
326 different 0.5-h samples (OA: $\sim 2.0 \times 10^{-2}$ µg for each punch) and a 2.5 h sample collected in parallel (OA: 9.1×10^{-2}
327 µg) were used, and the sum of their background-subtracted integrated signals (I_{blksub}) compared (Fig. 2 h). Without
328 background subtraction, the sum of the signals from the five 0.5-h samples was generally higher than the I_s of the
329 2.5-h sample (shown in Fig. 3a). An exception to this is HNO_3 , which has the highest signal of all compounds and
330 therefore is the least influenced by background. The higher I_s for the sum of the five 0.5-h samples is likely because
331 of the low signal-to-noise ratio compared to the 2.5-h sample. Subtracting only the field blank (Method 1) therefore
332 yielded the same result (Fig. 3b). Scaling the heating baseline (Method 2a and 2b) led to a better agreement between
333 the sum of the five 0.5-h and the 2.5-h samples (Figs. 3c and d). Compounds with high abundance generally fall
334 on a 1:1 line (slope range 0.5–2) by using these two background subtraction methods. With the thermal baseline
335 subtraction method (Method 4), results were comparable between 2.5-h and five 0.5-h samples. For the approach
336 using filter reheating (Method 3), there was a lesser agreement between the sum of the 0.5-h samples and the 2.5-h
337 sample (Figs. 3e and 3f). We speculate that this could be improved with a reheating cycle for every sample. For
338 future offline FIGAERO-CIMS analyses, we recommend carefully determining the background. Following our
339 assessment of blank determination methods, we suggest regular collections of field blanks and scaling their signal
340 (Methods 2a/b), and if field blanks are not available, computing a thermal baseline (Method 4). If using the
341 reheating approach as in a previous study with FIGAERO-CIMS in offline mode (Siegel et al., 2021), the
342 background should be determined by conducting reheating desorption cycles for each sample and blank
343 individually.

344 In general, as expected, high mass loadings are less sensitive to the various background subtraction methods due
345 to the higher signal-to-noise ratio (for example, 12-h/24-h sampling with OA loading of ~ 1 µg, Fig. S6). Besides
346 filter loadings, baseline levels can also be influenced by the properties of compounds (e.g. stickiness) and
347 instrument geometry. In summary, of all background subtraction methods shown here, Methods 2a, 2b, and 4
348 achieved the best agreement in signal intensities between the sum of 0.5-h and 2.5-h samples (Fig. S7). With these
349 methods, 82% to 93% of high-signal compounds (25% highest signal) fell into a signal ratio of ~ 1 (0–2, Fig. S8).
350 This shows the importance of correctly assessing the instrument background, especially for compounds with low
351 signal.

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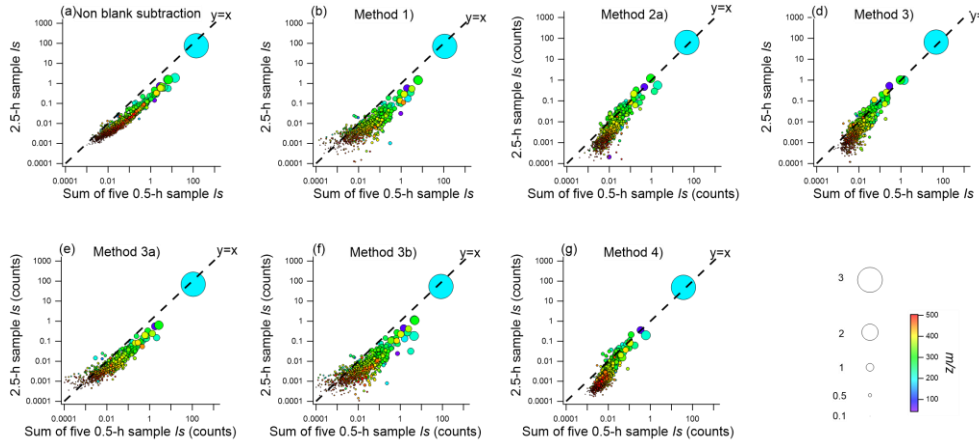
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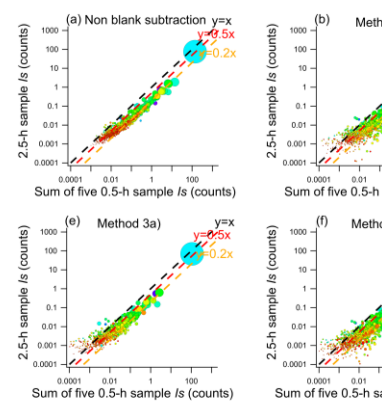
360
361 **Figure 3.** Comparison of the integrated signals (I_s) for the 2.5-h versus sum of 0.5-h samples (a) without blank subtraction,
362 with blank subtraction using (b) Method 1, (c) Method 2a, (d) Method 2b, (e) Method 3a, (f) Method 3b, (g) Method 4. The
363 size of dots is proportional to the 4th root of integrated signal intensities of compounds, and they are color-coded by the ions'
364 m/z (mass-to-charge ratio).

365 In this study, we applied Method 2b in the following discussions due to its better performance for the compounds
366 with both higher ($I_s > 0.1$ counts) and lower signal ($I_s < 0.01$ counts, Fig. 3d). First, we examined the signal-to-noise
367 ratios for offline FIGAERO-CIMS, defined as the ratio of the blank-subtracted signal to the standard deviation
368 (STDs) of the background determined using method 2b per compound. Most of the identified compounds are
369 above the estimated detection limit (3 times STDs of the backgrounds) for both filter types (87% and 87% of
370 CHOX peaks for both 24-h Quartz and Teflon filters, OA loadings of $1.2 \mu\text{g}/3.1 \times 10^2 \text{ cm}^2$ (2 mm punch)). For the
371 12-h samples (OA loadings of $0.58 \mu\text{g}/3.1 \times 10^2 \text{ cm}^2$ (2 mm punch)), 84% and 70% of CHOX compounds were
372 above the detection limit for Quartz and Teflon filters, respectively (Fig. S9). This varies for different filter
373 loadings and punch areas.

374 **3.2 Reproducibility of signal**

375 We performed reproducibility tests using three 2-mm punches from the same 24-h and 2.5-h samples of both
376 Teflon and Quartz filters and checked the signal response with the non-uniform temperature ramping procedure.
377 The comparisons of the blank-subtracted CHOX I_s for the 24-h and 2.5-h sample punches for both filter types are
378 displayed in Fig. 4 and Fig. S10, respectively.

379 In Figs. 4a and 4b, we plotted the compounds' signal from one punch versus their average signal from all 3
380 punches for the Teflon and Quartz filters, respectively. We observe a high correlation between the individual and
381 average signals (Spearman correlation coefficients R_{sp} are 0.95–0.96 and 0.97–0.99 for Teflon and Quartz filters,
382 respectively). For each CHOX compound, we also computed the relative error (standard deviation/average signals
383 ($\text{Std}(I_s)/\text{Avg}(I_s)$) for the three punches) versus the average signal (Figs. 4c, 4d). The relative error for a CHOX
384 compound was 9% for Quartz and 18% for Teflon (median relative errors) for 24-h samples (Figs. 4c, 4d). The
385 relative error decreased with higher signal intensities (Figs. 4c, 4d), especially for the Quartz filters, suggesting
386 that abundant compounds are measured more precisely than less abundant compounds. This trend is less apparent
387 for Teflon filters, which is likely caused by less reproducibility for high I_s compounds. Possible explanations could
388 be uneven distribution of particulate mass on the filter or larger uncertainties in the punching process for Teflon
389 filters due to the extension of the material. 86% and 94% of all CHOX compounds for Teflon and Quartz filters,
390 respectively, had >3 times higher signals than the variability from the duplicate tests (Fig. S9). For the 2.5-h filter



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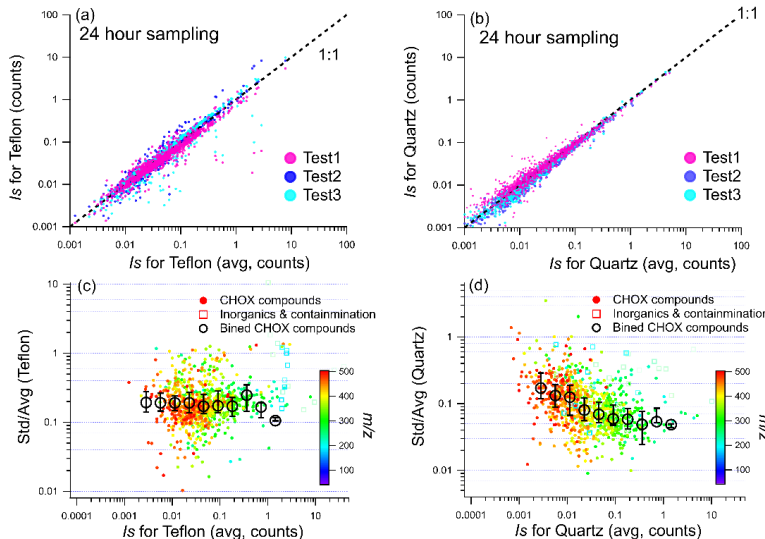
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396 samples (Fig. S10), the relative error is higher compared to the 24-h samples (25% for Quartz, and 31% for Teflon).
 397 This is likely due to the lower OA loadings (9.1×10^{-2} $\mu\text{g/punch}$) of the 2.5-h sample compared to the 24-h sample
 398 ($1.2 \mu\text{g/punch}$), which leads to higher uncertainties for blank subtraction and peak fitting. Still, the analytical
 399 reproducibility is acceptable, even for samples with OA loadings as low as $\sim 0.1 \mu\text{g}$. The relative error between
 400 repeats reported here is slightly larger (~9% and 18% for $\sim 1 \mu\text{g}$ OA/punch for Quartz and Teflon filters, and 25%
 401 for Quartz, 31% for Teflon for $\sim 0.1 \mu\text{g}$ OA/punch) compared to the variability in signal for online FIGAERO-
 402 CIMS (5–10% for $1 \mu\text{g}$ OA, (Lopez-Hilfiker et al., 2014)).

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403
 404 **Figure 4.** Comparison of the integrated signals from duplicate tests of the same 24-h sample for (a) Teflon and (b) Quartz
 405 fiber filters. The relative error (I_s ratio of standard deviation/average) value of the 3 duplicate tests as a function of I_s for (c)
 406 Teflon and (d) Quartz filters. In (c) and (d), CHOX compounds are shown as dots, inorganics as well as contaminants as
 407 squares colored by the m/z . The black cycles in (c) and (d) represent median values of signal intensity bins (with log I_s intervals
 408 of 0.3 for the I_s range of 0 to 2) and error bars represent the 25th and 75th percentile of binned values of $\text{Std}(I_s)/\text{Avg}(I_s)$ for
 409 CHOX.

410 3.3 Comparison of signal for different temperature ramping protocols

411 Here we compare the signal from different ramping protocols for the punches from the same 24-h Quartz and
 412 Teflon filters (Table 1). Since as suggested in the section 2.2.2, the I_s were calculated by the integration of the
 413 normalized signals (normalized to the primary ion (I)), which to some extent compensates for reagent ion
 414 depletion. The signal of the field blanks is largely dominated by instrument background (i.e. there is no distinct
 415 peak in the thermogram (Fig. S1e) thus the I_s of the field blanks is highly influenced by integration time. Since
 416 the field blanks were only analyzed with non-uniform ramping, the I_s for slow non-uniform and uniform ramping
 417 protocols were assumed as the I_s of non-uniform scaled by their integration time ratios.

418 The comparison of the background-subtracted I_s of all identified compounds from different ramping protocols
 419 for a pair of 24-h Quartz and Teflon filters each is shown in Fig. 5. Since the integrated signals of the compounds
 420 within a mass spectrum are log-normally distributed (shown in Fig. S11a and S11b), a linear fit would be strongly
 421 biased by high-signal compounds such as HNO_3I or $\text{C}_6\text{H}_{10}\text{O}_5\text{I}$. Thus, we calculated the correlation coefficients of

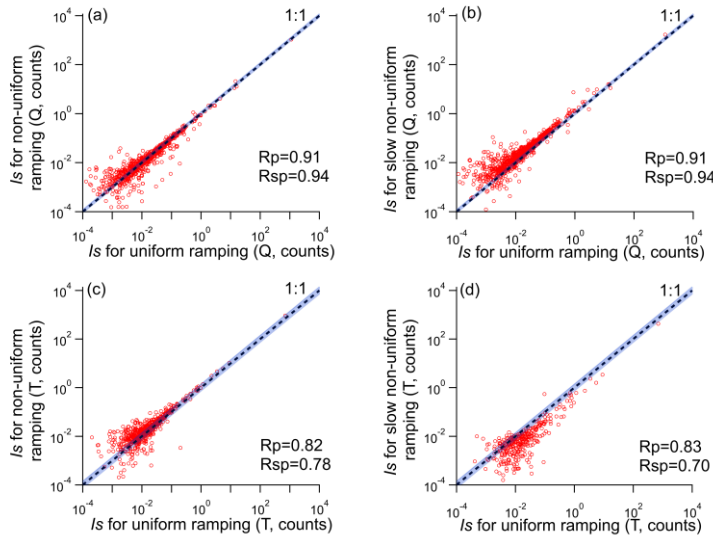
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426 the log-transformed signal intensities in the comparison. The Pearson correlation coefficients (R_p) and Spearman
 427 correlation coefficients (R_{sp}) are as follows: for Quartz filters $R_p = 0.91$, $R_{sp} = 0.94$ for non-uniform vs uniform,
 428 and $R_p = 0.91$, $R_{sp} = 0.94$ for slow non-uniform vs uniform, and for Teflon filters $R_p = 0.82$, $R_{sp} = 0.78$ for non-
 429 uniform vs uniform, and $R_p = 0.83$, $R_{sp} = 0.70$ for slow non-uniform vs uniform protocols.

430 These numbers suggest that the Quartz samples were less affected by different temperature ramping protocols
 431 than the Teflon samples. We also note that Teflon samples exhibited lower reproducibility than Quartz samples
 432 (see section 3.2). The lowest R_p and R_{sp} were observed for the comparison between the slow non-uniform ramping
 433 and the uniform ramping procedure for Teflon filters (Fig. 5d). Possible explanations could be the higher
 434 background and thus lower signal-to-noise ratios for Teflon filters in the low ramping rate region ($1.3\text{ }^{\circ}\text{C min}^{-1}$ for
 435 the range of $60\text{ }^{\circ}\text{C}$ to $105\text{ }^{\circ}\text{C}$) of the slow non-uniform ramping protocol. Thus, care needs to be taken when using
 436 very slow heating rates and backgrounds need to be carefully assessed, especially for Teflon filters.



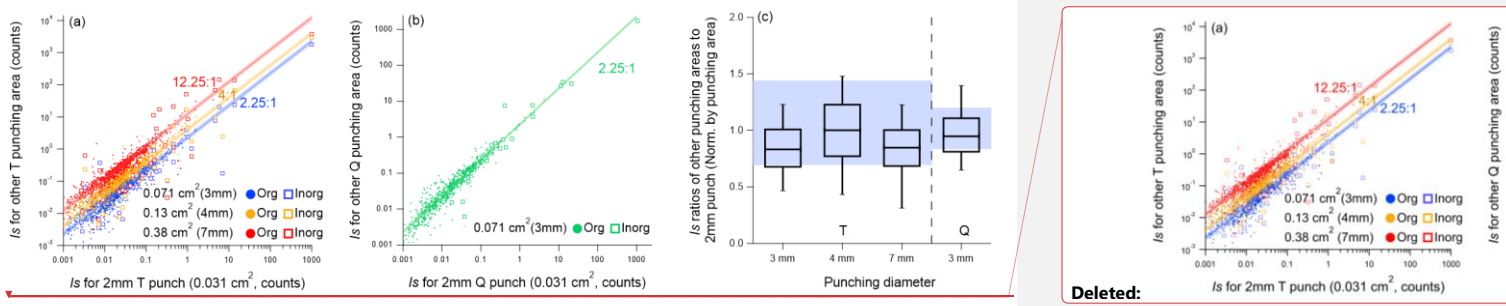
437

438 **Figure 5.** Comparison of I_s from the different temperature ramping protocols of the 24-h Quartz (Q) and Teflon (T) filter
 439 samples, (a) non-uniform and uniform ramping (Quartz sample), (b) slow non-uniform and uniform ramping (Quartz sample),
 440 (c) non-uniform and uniform ramping (Teflon sample), (d) slow non-uniform and uniform ramping (Teflon sample). The blue
 441 shaded areas represent the relative error of signal assessed in the reproducibility tests of the 24-h samples (18% for Teflon
 442 and 9% for Quartz filters). The upper and lower limits for the reproducibility-based variation are calculated as $(1+18\%)/(1-18\%)$
 443 and $(1-18\%)/(1+18\%)$, respectively. The upper and lower limits for the I_s distribution of Quartz caused by
 444 reproducibility are calculated as $(1+9\%)/(1-9\%)$ and $(1-9\%)/(1+9\%)$, respectively.

445 For further analyses, we use the results from the non-uniform temperature ramping protocol, which represents a
 446 good balance between the influence of background due to low signal-to-noise ratios, and I^- depletion. The good
 447 agreement between offline FIGAERO-CIMS and ToF-ACSM discussed in Section 3.5 further implies that such a
 448 ramping protocol is suitable for the OA loadings observed in our study.

449 **3.4 Linearity of signal response**

450 To assess the linearity of signal response to the amount of sample collected on the filter, we used punches with
 451 varying areas from one single filter. We used punch diameters of 2, 3, 4, and 7 mm for a Teflon filter and 2 mm
 452 and 3 mm for a Quartz filter. The analytical protocol was kept constant between the individual sample punches
 453 (non-uniform ramping protocol and method 2b for background subtraction). The mass loadings of the analyzed
 454 filter punches ranged from 1.2 to 15 μg OA (2.2 to 27 μg PM_{2.5}) for the Teflon filter and from 1.2 to 2.7 μg OA
 455 (2.2 to 5.0 μg PM_{2.5}) for the Quartz filter (Table 1). The blank-subtracted I_s from the different punching areas for
 456 the Quartz and Teflon filters is shown in Fig. 6. Overall, the offline FIGAERO-CIMS approach responds linearly
 457 to changes in filter mass loadings. The integrated signal ratios of CHOX are consistent with their respective area
 458 ratios (Figs. 6a, 6b), within uncertainty. In Fig. 6c we also plot the signal ratios of the 2 mm punch to the other
 459 punches, normalized by punching area (where 1 signifies perfect linearity). These ratios are generally in the range
 460 of possible variability caused by the relative error from the reproducibility tests.



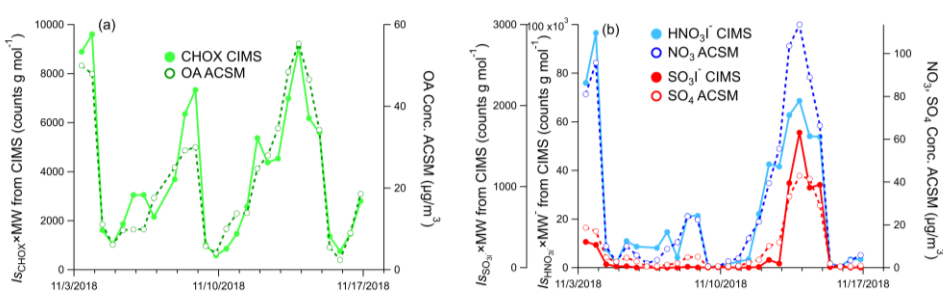
462 **Figure 6.** Comparison of the I_s between signals from punches (a) with 3 mm, 4 mm, 7 mm, and 2 mm in diameter for the
 463 same Teflon filter, and (b) with 3mm and 2 mm in diameter for the same Quartz filter. The lines in (a) and (b) represent the
 464 punching area ratios. The shaded areas in (a) and (b) represent the area ratio plus/minus the relative errors (9% for Quartz,
 465 and 18% for Teflon) from the reproducibility tests. (c) Distribution of I_s ratios normalized by the punching area ratios (3 mm,
 466 4 mm, and 7 mm to 2 mm diameter punches for Teflon, 3 mm to 2 mm diameter punches for Quartz). Within each box,
 467 the median (middle horizontal line), 25th and 75th percentiles (lower and upper ends of the box), and 10th and 90th percentiles
 468 (lower and upper whiskers) are shown. The shaded area in (c) represents the possible distribution of the I_s ratios due to the
 469 relative error established from the 24-h sample reproducibility tests (18% for Teflon and 9% for Quartz filters). The upper
 470 and lower limits for the Teflon I_s ratio distribution are calculated as $(1+18\%)/(1-18\%)$ and $(1-18\%)/(1+18\%)$, respectively.
 471 The upper and lower limits for the Quartz I_s ratio distribution are calculated as $(1+9\%)/(1-9\%)$ and $(1-9\%)/(1+9\%)$,
 472 respectively.

473 For compounds with very high signals, the response I_s ratio can deviate from the punch area ratio, not least also
 474 due to the varying degree of reagent ion depletion. The highest I⁻ depletions were ~35%, ~60%, ~68%, and ~70%
 475 for 2mm, 3mm, 4mm, and 7mm punches, respectively. For e.g. the highest inorganic (HNO₃I⁻) and organic
 476 (C₆H₁₀O₅I⁻) ions, the I_s from a 7mm punch is only 30% and 67%, respectively, of what would be expected based
 477 on punching area ratios (7mm to 2mm). For smaller punches (4 mm/3 mm), 75%/80% and 105%/107% of the
 478 expected HNO₃I⁻ and C₆H₁₀O₅I⁻ signals, respectively, are detected. This indicates that for reduced amounts of
 479 desorbing material provided by smaller filter fractions, the amount of reagent ion is sufficient during the whole
 480 ramping process (lowest I⁻/C₆H₁₀O₅I⁻ signal ratio: $\sim 10^3$). In other words, if titration of reagent ion can be avoided
 481 as much as possible (e.g. I⁻/target ion signal ratio: $\sim 10^3$) the I_s responds linearly to concentration changes. In this
 482 study, titration is non-apparent for OA loadings of $< 5 \mu\text{g}$ and I⁻ signals of ~ 1 million. Therefore, it is recommended
 483 to calculate OA loadings of the samples prior analysis to determine the punching sizes in offline FIGAERO-CIMS
 484 analysis.

487 **3.5 Comparison between offline FIGAERO-CIMS and *in-situ* ToF-ACSM**

488 In the following, we compare the time series of the signals of the signals from offline FIGAERO-CIMS from Quartz filters and
 489 the corresponding chemical components from online ToF-ACSM measurement. The comparison between the total
 490 signal of all identified CHOX compounds and OA concentrations from the ToF-ACSM is displayed in Fig. 7a.
 491 Here, the FIGAERO-CIMS signals of five polyols ($C_8H_{18}O_5I$, $C_{10}H_{22}O_6I$, $C_{12}H_{26}O_7I$, $C_{14}H_{30}O_8I$, $C_{16}H_{34}O_9I$)
 492 were excluded, which were contaminants from the lab due to their inexplicably high I_s in 3 of the 27 12-h samples
 493 and the usage of diethylene glycol (DEG) in the lab. To compare with the $PM_{2.5}$ component concentrations from
 494 the ToF-ACSM, for each 12-h filter, we compute the sum of integrated signals (I_s , signal integration over the
 495 entire thermogram, counts) multiplied by their molecular weight (MW, g mol⁻¹) of all compounds from FIGAERO-
 496 CIMS for comparison to the corresponding $PM_{2.5}$ component concentrations from the ToF-ACSM. Even though
 497 I is selective towards oxygenated organic compounds, the total MW-weighted CHOX signal measured by offline
 498 FIGAERO-CIMS in this study highly correlates with OA measured by the ToF-ACSM ($R_p = 0.94$), which is
 499 known to be dominated by secondary organic aerosols (SOA) (Cai et al., 2020; Kulmala et al., 2021; Jia et al.,
 500 2008).

501 The time series of the 12h- I_s for HNO_3I and SO_3I measured by offline FIGAERO-CIMS correlate well with the
 502 NO_3 and SO_4 concentrations from ToF-ACSM ($R_p = 0.94$ and 0.95, Fig. 7b). The signal of HNO_3I in the particle
 503 phase measured by FIGAERO-CIMS is as an indicator of particulate nitrate and organonitrate (Lee et al., 2016),
 504 and the signal of SO_3I is related to inorganic sulfate and sulfur-containing organics (Ye et al., 2021; Cao et al.,
 505 2019). Following the same method, after calibrations, the quantified CHOX mass concentrations of offline
 506 FIGAERO-CIMS were found to be highly correlated with OA and SOA from ToF-ACSM in another dataset at
 507 the Peking University campus (PKU) in Beijing, indicating offline FIGAERO-CIMS analysis can be quantitative
 508 with proper calibrations (shown in Fig. S12 (Zheng et al., 2021)). Like other offline sampling methods, the offline
 509 FIGAERO-CIMS method may be affected by artefacts from sampling and storage of the filters. Both positive
 510 (absorption of gaseous OA), and negative artefacts (volatilization of collected OA), may occur during the sampling
 511 and storage, even if filters were stored frozen (Cheng et al., 2009). However, the signals from FIGAERO-CIMS
 512 correlate generally well with major components measured by TOF-ACSM, suggesting that those artefacts can be
 513 considered minor in our study, at least in terms of bulk PM constituents (Figure 7).



515
 516 **Figure 7.** Comparison of the time series of the integrated signals of inorganic and organic compounds from 12-h samples (2
 517 mm punches) analyzed by offline FIGAERO-CIMS, and chemical components measured *in-situ* by ToF-ACSM, (a) total
 518 CHOX from FIGAERO-CIMS and OA from ToF-ACSM, (b) HNO_3I from FIGAERO-CIMS and NO_3 from ToF-ACSM,
 519 (c) SO_3I from FIGAERO-CIMS and SO_4 from ToF-ACSM. To compare with the $PM_{2.5}$ component concentrations from the
 520 ToF-ACSM, the I_s of each compound from FIGAERO-CIMS was multiplied by their molecular weight (MW) in (a) and (b).
 521 Note that FIGAERO-CIMS and ToF-ACSM data are on different axes

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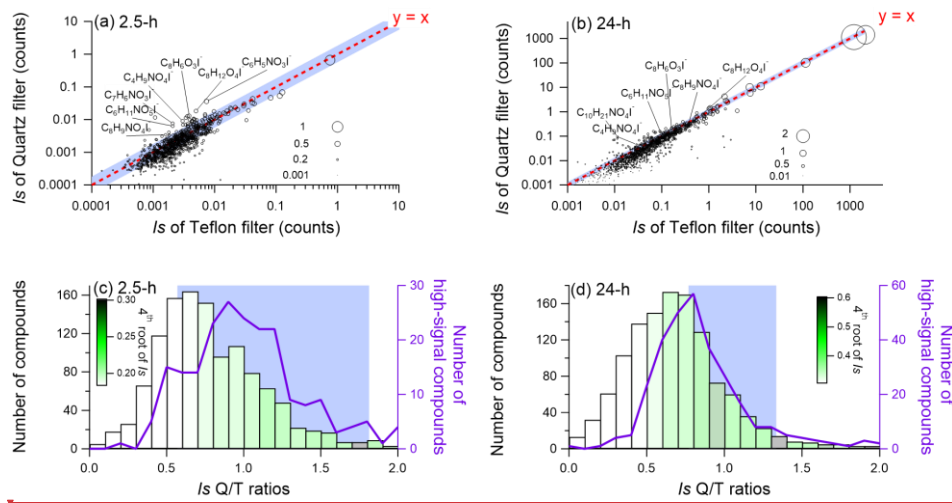
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533 **3.6 Comparison of Quartz and Teflon filters**

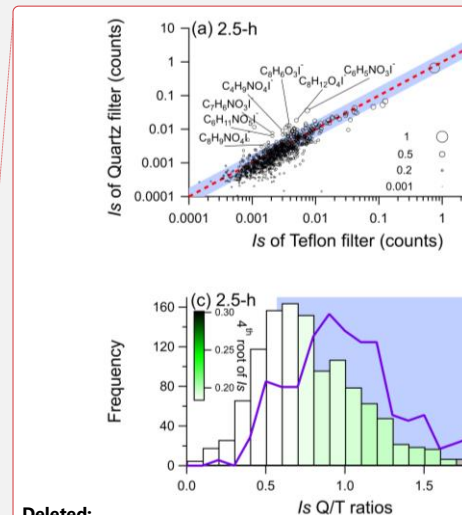
534 In the following, we compare the I_s from simultaneously collected Quartz and Teflon filter samples (collection
 535 times 2.5 h, 12 h, and 24 h, see Table 1). Fig. 8a and b show the comparison of the average I_s of compounds (3
 536 samples each) for both filter types, with 2.5h (OA loading of $9.1 \times 10^{-2} \mu\text{g}$) and 24h (OA loading of $1.2 \mu\text{g}$)
 537 collection times. The mass spectra show an overall similar pattern, we observe a non-negligible difference,
 538 especially for the 2.5h samples (Fig. 8a). The log-transformed signals from Quartz and Teflon samples correlate
 539 better for 24-h samples ($R_p = 0.96$, $R_{sp} = 0.95$, Fig. S11d) than for the 2.5-h samples ($R_p = 0.88$, $R_{sp} = 0.87$, Fig.
 540 S11c). In addition, the signal observed for Quartz filter samples is generally slightly lower than for Teflon filter
 541 samples (Fig. 8c, d). Compounds with high Quartz/Teflon-signal ratios are in general semi- or low volatile
 542 compounds (operationally defined as having a $T_{\max} < 60^\circ\text{C}$). These compounds tend to be in the CHO and especially
 543 CHON category and exhibit a higher degree of unsaturation (e.g. $\text{C}_8\text{H}_6\text{O}_3\text{I}$, $\text{C}_6\text{H}_5\text{NO}_3\text{I}$ and $\text{C}_7\text{H}_6\text{NO}_3\text{I}$). They can
 544 be aromatics or their thermal fragmentation products (Liu et al., 2019). Due to the high surface area of the Quartz
 545 filters, semi- or low volatile compounds are more easily adsorbed than on Teflon filters, potentially resulting in
 546 higher positive artefacts. Compounds with low Quartz/Teflon-signal ratios tend to have overall low signal. Despite
 547 the application of a blank determination method that takes instrument backgrounds into account (Method 2b),
 548 higher residuals were still observed for the lower signal compounds, especially for the Teflon filters (as seen also
 549 for the 2.5-h and 0.5-h sample comparison (Fig. 3d). In contrast, compounds with a higher signal tend to be in the
 550 range of Q/T ratios expected based on the observed variability from the reproducibility tests (shown in Fig. 8c and
 551 8d).

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552
 553 **Figure 8. Comparison of the integrated signal intensities of all identified compounds for the Quartz fiber and Teflon filter**
 554 **samples for (a) 2.5-h samples, and (b) 24-h samples. The size of symbols in (a) and (b) is proportional to the 4th root of the**
 555 **signal intensity of each compound from the Quartz filter. Frequency distribution (number of compounds) per signal ratio of**
 556 **Quartz/Teflon for all compounds (green bars), and high-signal compounds (highest 25% signal compounds) only (purple lines)**
 557 **for 2.5-h samples (c), and 24-h samples (d). The bars in (c) and (d) are colored by the average of the 4th root of the**
 558 **signal intensity of the Quartz filter. The blue shaded area in each panel represents the possible distribution of I_s ratios of**
 559 **Quartz/Teflon from the relative errors from the duplicate tests of 2.5-h (25% for Quartz and 31% for Teflon) and 24-h (9%**
 560 **for Quartz and 18% for Teflon) samples. The upper and lower limits for the 2.5-h Quartz/Teflon I_s ratios were calculated as**
 561 **(1+25%)/(1-31%) and (1-25%)/(1+31%), respectively. The upper and lower limits for the 24-h Quartz/Teflon I_s ratios were**
 562 **calculated as (1+9%)/(1-18%) and (1-9%)/(1+18%), respectively.**



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Deleted: Comparison of the integrated signal intensities of all identified compounds for the Quartz fiber and Teflon filter samples, (a) 2.5-h samples, (b) 24-h samples. The size of symbols in (a) and (b) is proportional to the 4th root of the signal intensity of each compound from the Quartz filter. The distribution of I_s ratios of Quartz/Teflon from the relative errors from the duplicate tests of 2.5-h (25% for Quartz and 31% for Teflon) and 24-h (9% for Quartz and 18% for Teflon) samples. The upper and lower limits for the 2.5-h Quartz/Teflon I_s ratios were calculated as $(1+25\%)/(1-31\%)$ and $(1-25\%)/(1+31\%)$, respectively. The upper and lower limits for the 24-h Quartz/Teflon I_s ratios were calculated as $(1+9\%)/(1-18\%)$ and $(1-9\%)/(1+18\%)$, respectively. The T_{\max} was corrected.

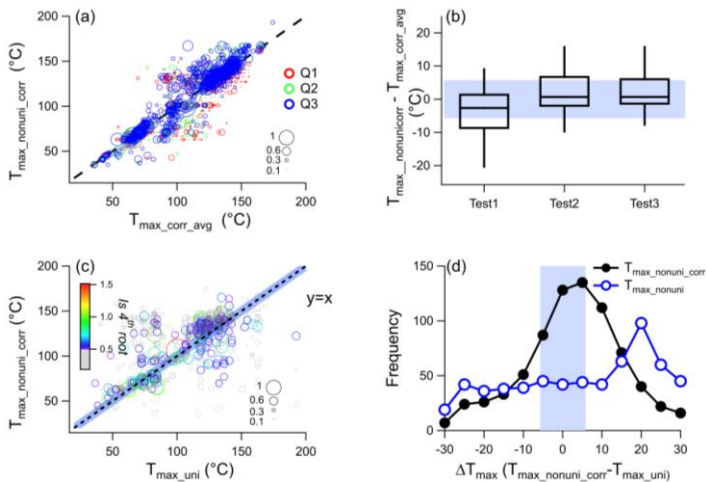
586 **3.7 T_{\max} : Influence of temperature ramping protocol and filter type**

587 Non-uniform ramping of the temperature due to reagent ion titration is more likely needed when the FIGAERO-
 588 CIMS is run in offline mode compared to online mode, where sampling times and resulting filter mass loadings
 589 can be adjusted more easily. We have therefore developed a method (see section 2.2.4) to recover T_{\max} from non-
 590 uniform ramping protocols, i.e. to make it comparable to T_{\max} from uniform ramping protocols. Compared to the
 591 raw thermograms, the shape of the corrected thermograms is more similar to that of the uniform protocol (Fig. [S13](#)
 592 and [S14](#)), since the thermograms were re-gridded to the same temperature intervals (3 °C).

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593 Firstly, we tested the variation of T_{\max} from the three duplicate tests of the Quartz filters using the non-uniform
 594 ramping protocol and thermogram correction (Fig. 9a). After correction, the corrected T_{\max} ($T_{\max_nonuni_corr}$) from
 595 individual tests was highly correlated with their average ($T_{\max_corr_avg}$, $R_p = 0.87$ – 0.93). The median value of the
 596 difference between $T_{\max_nonuni_corr}$ of duplicate tests and their average for all compounds ranges from -2.7–0.7 °C
 597 (shown in Fig. 9b). The majority of compounds (52%–70%) have a T_{\max} difference within 5 °C, close to the value
 598 reported in previously (~2°C, (Lopez-Hilfiker et al., 2014)). The median standard deviation of the difference
 599 between the corrected T_{\max} of individual tests ($T_{\max_nonuni_corr}$) and their average ($T_{\max_corr_avg}$) from all compounds
 600 is 5.7 °C, which is defined as the variation of T_{\max} for duplicate tests.



601

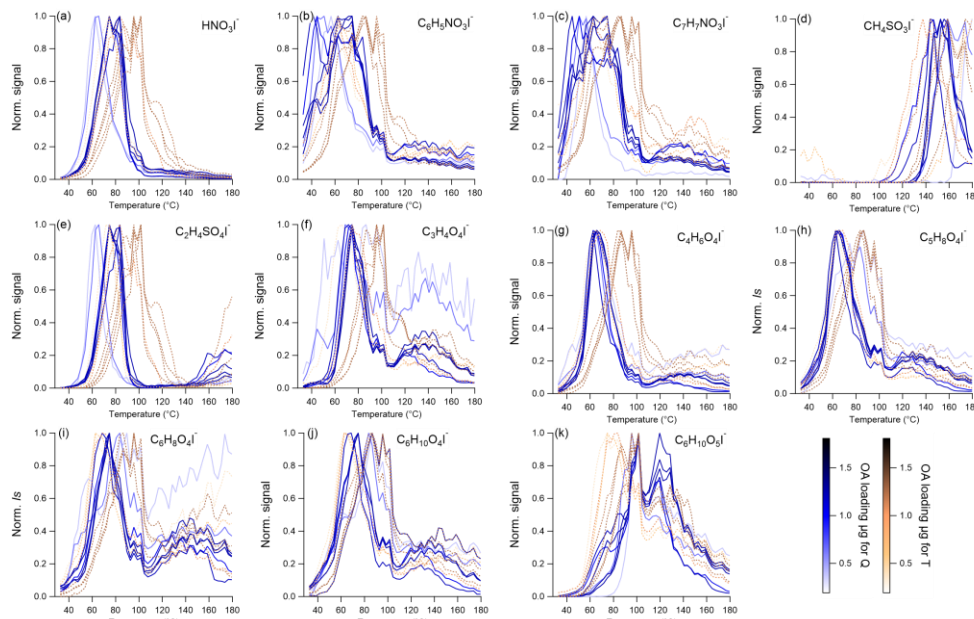
602 **Figure 9.** (a) Comparison of $T_{\max_nonuni_corr}$ from the 3 duplicate tests and their average ($T_{\max_corr_avg}$), (b) distribution of the
 603 difference between the 3 triplicate tests and the $T_{\max_corr_avg}$, (c) comparison of T_{\max} from the corrected non-uniform ramping
 604 and uniform ramping protocol (T_{\max_uni}), (d) histogram of ΔT_{\max} between T_{\max} from the uniform ramping protocol (T_{\max_uni})
 605 and non-uniform with ($T_{\max_nonuni_corr}$)/without (T_{\max_nonuni}) correction. The size of symbols in (a) and (b) is proportional to the
 606 4th root of the integrated signal intensity. The 4th root of the signal intensity <0.5 is shown in grey. The uniform ramping
 607 protocol test and 3 duplicate non-uniform ramping protocol tests were conducted for the same 24-h Quartz filter (Nov 23 to
 608 24). The shaded area in (b), (c), and (d) represents T_{\max} variation (± 5.7 °C) from the duplicate tests.

609 We take the uniform sampling protocol (see Fig. 1d) as the basis since this is the commonly used protocol for
 610 FIGAERO-CIMS in online mode. The comparison of T_{\max} from the corrected non-uniform and the uniform
 611 ramping protocols is shown in Fig. 9c. Generally, after correction for the non-uniform ramping, the Pearson
 612 correlation coefficient of $T_{\max_nonuni_corr}$ and T_{\max_uni} is higher ($R_p = 0.60$) compared to the uncorrected ones with
 613 the uniform protocol ($R_p = 0.20$, T_{\max_nonuni} vs T_{\max_uni}). The correlation coefficients were even higher (0.72 and
 614 0.84) for the 400 and 100 compounds with the highest signal intensity. In Fig. 9d we plot the frequency distribution

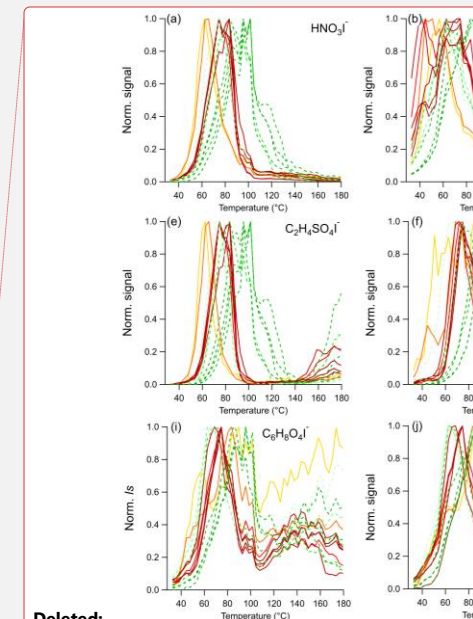
617 of the differences between the corrected T_{\max} ($T_{\max_nonuni_corr}$) and T_{\max} from the uniform protocol (T_{\max_uni}) for each
 618 CHOX compound in the spectrum. For 73% of the compounds, the difference in T_{\max} between the two ramping
 619 protocols lies between -15 and 15 °C, and 41 % of compounds exhibit a difference of $0 - \pm 5$ °C.

620 In the next step, we compared the volatility derived from T_{\max} for Quartz fiber and Teflon filters. We selected a
 621 number of inorganic and organic compounds, based on their high average signals for the whole sampling period,
 622 for comparison of thermograms from 12-h and 24-h Teflon and Quartz filters sampled in parallel (Table S1, Fig.
 623 10). Compounds include HNO_3I , $\text{CHON}(\text{C}_6\text{H}_5\text{NO}_3\text{I}, \text{C}_7\text{H}_7\text{NO}_3\text{I})$ and $\text{CHOS}(\text{CH}_4\text{SO}_3\text{I}, \text{C}_2\text{H}_4\text{SO}_4\text{I})$ compounds
 624 as well as CHO compounds with $\text{C}_{\text{num}} \geq 3$ ($\text{C}_3\text{H}_4\text{O}_4\text{I}$, $\text{C}_4\text{H}_6\text{O}_4\text{I}$, $\text{C}_5\text{H}_8\text{O}_4\text{I}$, $\text{C}_6\text{H}_8\text{O}_4\text{I}$, $\text{C}_6\text{H}_{10}\text{O}_4\text{I}$, $\text{C}_6\text{H}_{10}\text{O}_5\text{I}$).
 625 Compounds with $\text{C}_{\text{num}} < 3$ (e.g. $\text{CH}_2\text{O}_2\text{I}$) were excluded due to possible gas-phase interference and more likely
 626 influenced by thermal decomposition. Some compounds exhibited similar thermogram shapes for the two types of
 627 filters, such as $\text{C}_6\text{H}_{10}\text{O}_5\text{I}$ and $\text{CH}_4\text{SO}_3\text{I}$, while for some other species, the thermograms were different. Taking
 628 $\text{C}_3\text{H}_4\text{O}_4\text{I}$ as an example, a bimodal thermogram shape with peaks around 100 °C and 150 °C was observed for the
 629 Quartz filter, while only a unimodal peak around 90 °C was observed for the Teflon filter. The different
 630 thermogram shapes of individual compounds for the different filter types might warrant further investigation with
 631 a focus on the role of filter type properties (such as pore size, thickness, absorption, and hydrophobic/hydrophilic
 632 properties).

633



634
 635 **Figure 10.** Normalized thermograms for Teflon (T, dashed lines) and Quartz (Q, solid lines) filters of, (a) HNO_3I , (b)
 636 $\text{C}_6\text{H}_5\text{NO}_3\text{I}$, (c) $\text{C}_7\text{H}_7\text{NO}_3\text{I}$, (d) $\text{CH}_4\text{SO}_3\text{I}$, (e) $\text{C}_2\text{H}_4\text{SO}_4\text{I}$, (f) $\text{C}_3\text{H}_4\text{O}_4\text{I}$, (g) $\text{C}_4\text{H}_6\text{O}_4\text{I}$, (h) $\text{C}_5\text{H}_8\text{O}_4\text{I}$, (i) $\text{C}_6\text{H}_8\text{O}_4\text{I}$, (j) $\text{C}_6\text{H}_{10}\text{O}_4\text{I}$,
 637 (k) $\text{C}_6\text{H}_{10}\text{O}_5\text{I}$. The thermograms were first corrected (section 2.2.4) and then normalized to signals in T_{\max} and colored by the
 638 OA mass loading. The sampling information of the thermograms presented here is listed in Table S1.



640 In addition, we found that compounds with higher mass loadings appeared to have a higher T_{max} (e.g. $C_2H_4SO_4I^-$
641 and $C_2H_7NO_3I^-$, shown in Fig 10), consistent with previous findings using Teflon filters (Huang et al., 2018;
642 Ylisirniö et al., 2021). The variability in T_{max} induced by varying PM loadings is within 5°C for 29% of compounds,
643 and within 15°C for 54% of all compounds for Quartz filters, and 35% and 57% of compounds, respectively, for
644 Teflon samples. The higher T_{max} variation for different OA loading samples compared to the duplicate samples
645 (+5.7°C, Fig.9 b) is likely caused by other factors, such as particle viscosity, the particles on the filter, and/or mass
646 loadings on the filter (Huang et al., 2018; Ylisirniö et al., 2021; Wu et al., 2021; Graham et al., 2022). The T_{max}
647 variation due to filter type ($R_p=0.27$) is much larger than the one induced by filter loadings. Thus, the direct
648 comparison of T_{max} between Quartz and Teflon filters is not feasible, warranting further research.

649 4. Discussion

650 This study introduces methods and assesses the performance of using the FIGAERO-CIMS in offline mode, i.e.
651 to analyze particulate matter collected temporally and locally distant from the instrument on filter samples (Quartz
652 and Teflon). Such an approach greatly enhances the capabilities of the FIGAERO-CIMS for analyzing atmospheric
653 samples, as it enables the probing of the air at locations where and on occasions when *in-situ* deployments are
654 difficult.

655 Due to the difficulties in background determination for offline FIGAERO-CIMS, in this study, we propose
656 different background determination methods, which were further assessed by the comparison between samples
657 from 5 different 0.5-h samples and a 2.5-h sample collected in parallel. We applied non-uniform temperature
658 ramping to avoid reagent ion titration and a background scaling method taking interference of variable instrument
659 backgrounds into account. In general, the offline FIGAERO-CIMS approach using the methods presented in this
660 study can be used for providing OA composition information with typical offline sampling times (e.g. 12h and
661 24h) samples: (1) the reproducibility of integrated signal intensity is within $\pm 20\%$ for both filter types (18% for
662 Teflon and 9% for Quartz), (2) detected signals respond linearly to changes in the samples' mass loadings, (3) the
663 signals of $CHOX$ and SO_3I^- , HNO_3I^- correlated well with corresponding $PM_{2.5}$ chemical component concentrations
664 of OA, SO_4 , and NO_3 measured by ToF-ACSM ($R_p=0.94$ to 0.95), (4) the log-transformed mass spectra are highly
665 correlated ($R_p>0.9$) between Quartz and Teflon filters for typical offline sampling times (e.g. 12h and 24h), and
666 for high-signal compounds the $I_{SO_3I^-}$ ratios between Quartz and Teflon filters are generally within reproducibility
667 variation. Overall, this highlights the possibility of using widely available and stored Quartz filters to identify
668 $CHOX$ molecular composition with FIGAERO-CIMS.

669 T_{max} retrieved from corrected thermograms of desorption with non-uniform ramping protocols are comparable to
670 T_{max} from uniform ramping protocol for high signal intensity compounds ($R_p = 0.72$ – 0.84). More than 50% of
671 compounds have T_{max} values that are reproducible within 5 °C for duplicate tests ($R_p = 0.87$ – 0.93) of the same
672 sample, and for >50% of compounds, T_{max} varies within 15 °C for different mass loadings. Yet, T_{max} is strongly
673 affected by the filter material (Teflon vs Quartz) leading to a large discrepancy in T_{max} between Quartz and Teflon
674 samples ($R_p = 0.27$), hindering direct comparisons and warranting further research.

675 In summary, using FIGAERO-CIMS to analyze offline samples is a useful and simple way to investigate OA
676 molecular composition, but care needs to be taken for T_{max} analyses. This opens broad applications to study OA
677 molecular composition, sources, and formation processes at several sites simultaneously and in long-term
678 deployments.

679 Author contributions

680 JC, KRD, CM, and MK designed the research. JC, FXZ, and WD collected the samples at the BUCT site. JC, CW,
681 SH, KRD, and CM analyzed the samples and interpreted the data. ZY and CQ analyzed the samples collected at

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683 the Peking University campus site. CM, KRD, and MK supervised this research. JC, KRD, and CM wrote the
684 manuscript with contributions from all co-authors. All authors have given approval to the final version of this
685 manuscript.

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693

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