



# **Retention During Freezing of Raindrops, Part II: Investigation of Ambient Organics from Beijing Urban Aerosol Samples**

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### Abstract

The freezing of hydrometeors incurs certain water-soluble organic compounds dissolved in the supercooled cloud droplets to be released into the gas phase. This may lead to the vertical redistribution of

- 15 substances that become available for new particle formation in the upper troposphere. Drop freezing experiments were performed on the Mainz Acoustic Levitator (M-AL) using aqueous extracts of ambient samples of Beijing urban aerosol. The retention coefficients of over 450 compounds were determined. Most nitroaromatics and organosulfates were fully retained along with the aliphatic amines (AA) and higher-order amines and amides while sulfides, lipids, aromatic hydrocarbons, and long chain compounds
- 20 are among the most unretained and incidentally the fewest species observed. The findings here also indicate that  $NO<sub>x</sub>$  and  $SO<sub>x</sub>$  chemistry, particularly anthropogenically related, enhances the retention of the resulting secondary organic aerosols (SOA). A positive correlation between polarity and freezing retention along with a negative correlation with vapor pressure and freezing retention was observed. No sigmoidal relationship with effective Henry's law constant was observed which differs with the
- 25 parameterizations of riming retention presented in current literature, which is justified by the lower surface-to-volume ratio of the large drop size investigated.





### 1 Introduction

Atmospheric organic matter (OM) plays a critical role in climate regulation directly through radiation scattering and indirectly through cloud condensation nucleation which impacts Earth's energy 30 balance through radiative forcing (Fofie et al., 2018; Liu et al., 2018). These effects are controlled by factors such as their optical properties, size, and the hygroscopicity (Dusek et al., 2006; Sun et al., 2021), which can change based on the proportions of primary organic aerosols (POA)—directly emitted aerosols—and secondary organic aerosols (SOA)—aerosols formed from the oxidation products of volatile organic compounds (VOC) as part of new particle formation (NPF) (Hallquist et al., 2009; Liu et 35 al., 2021; Riva et al., 2019). Convective systems have been suggested to support NPF in the outflow region by reducing existing particle concentrations, facilitating cold temperatures, and transporting

reactive gases into regions with high actinic fluxes (Clarke et al., 1998; Zheng et al., 2021)

Nucleation-mode particles—with sizes in the lower tens of nm—have consistently been observed in concentrations of up to  $10^4 \text{ cm}^3$  from aircraft in the upper troposphere (Andreae et al., 2018; Andrés

- 40 Casquero-Vera et al., 2020; Clarke et al., 1999; Heitto et al., 2024; Weigel et al., 2011; Williamson et al., 2019). These measurements significantly exceed the corresponding concentrations in the planetary boundary layer and indicate that the main source of such ultra-fine particles in the upper troposphere is in situ NPF rather than their direct transport from the boundary layer (Bardakov et al., 2021). The traditional explanation for this phenomenon has been that the reduction of existing aerosol particles in deep
- 45 convective clouds eliminates removal processes for small particles and condensable vapors, supporting NPF (Clarke et al., 1998). However, Williamson et al. (2019) also showed that even without these conditions, such as in tropical convection, these newly formed particles can still be found. They then argue that most models underestimate available organic matter at high altitudes and as a result predict less NPF in these regions. If this NPF is the result of an overlooked mechanism of organic matter transport, it
- 50 is then critical to elucidate this mechanism so to constrain uncertainty around the influence of high altitude NPF from convective outflows (Bardakov et al., 2021).





Publications by Borchers et al. (2024) and Jost et al., (2017) have demonstrated a potential mechanism for organic matter transport in mixed-phase clouds. They describe how organic compounds that are exchanged between the gas and aqueous phase in cloud droplets can either be trapped in the ice

- 55 phase during freezing—washing them out by precipitation—or return to the gas phase by volatilization. This revolatilization incurred by the freezing process leads to a vertical redistribution and has the potential to explain the occurrence of organic matter at high altitudes in regions with deep convection. Alternatively, earlier publications by Pruppacher and Klett (2010) and Snider and Huang (1998) have suggested that complete sublimation of ice particles can also transport 'retained' compounds trapped in
- 60 the ice phase and release them at high altitudes.

In convective clouds the main formation pathway of precipitation-size ice particles is riming, i.e. the freezing of supercooled µm-sized cloud droplets on the surface of a mm-sized ice particle. Thus, riming retention is an important process for the vertical redistribution of water-soluble organic compounds (WSOC). Convective clouds with warm bases favor the formation of mm-sized drops by

- 65 collision-coalescence (Lamb and Verlinde, 2011), which subsequently can be uplifted in the updraft to regions with temperatures below 0°C. Once beyond the freezing level they can freeze and thereby release dissolved matter into the gas phase. This identifies freezing retention of mm-sized drops as a potential contributor to the vertical redistribution of WSOCs and was experimentally investigated in the present study.
- 70 The proportion of a substance that remains in the ice during this phase change is described by the retention coefficient R, which indicates the percentage of the trapped substance with a value between 0 and 1 (Bela et al., 2018; Iribarne and Pyshnov, 1990; Snider et al., 1992; Stuart and Jacobson, 2004). A species' retention is influenced by its chemical properties, such as its dimensionless effective Henry's law solubility constant  $(H^*)$ , as well as the physical properties of the droplet such as temperature, liquid water
- 75 content, droplet size, and ventilation. Substances with a small H\* are more likely to return to the gas phase during riming, which results in a lower retention coefficient. Additionally, these external and

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physical conditions of the droplet disproportionally influence the retention for these small H\* substances (Jost et al., 2017; Stuart and Jacobson, 2003, 2004)

- Current experimental studies to determine retention coefficients for atmospheric constituents and 80 relevant SOA precursors have focused on inorganic species, small organics, or single component mixtures with significantly higher than natural concentrations. Additionally, current studies have only examined retention in droplets within natural cloud size range rather than raindrop sizes. The few studies that look at complex mixtures are limited to compounds of similar families and only a handful of species (Borchers et al., 2024). Naturally occurring atmospheric constituents that are observed in rainwater are present as
- 85 complex mixtures of potentially thousands of species (Seymore et al., 2023). To get closer to observing the retention of compounds in their natural conditions, this study presents measurements of retention coefficients for a real, complex mixture of WSOC extracted from filter samples taken in an urban environment.

#### 90 2 Methods

#### 2.1 Sampling Location and Method

A high-volume sampler (HiVol) was run with quartz fiber TSP filters over three nights between March 3 and 5, 2022 in Beijing, China (40.0426° N, 116.4197° E) for an approximate sample volume of 550 m<sup>3</sup> between the hours of 21:00 to 9:00. These filters were sealed in aluminum foil and stored at -20 $\degree$ C

95 until analysis. Aqueous extracts of these filters were prepared by taking 1/4 of each filter, combining them (in total 3/4 of a 203 x 254 mm filter area) in 30 ml Milli-Q water, and then extracting with an orbital shaker for 15 min. The same was performed for a blank sample; a total 3/4 of unsampled filter area was extracted in 30 ml Milli-Q water for 15 min. These extracts were filtered through a 0.2 µm PTFE filter. 10 ml of the prepared extract was reserved for Ultra-High Performance Liquid Chromatography High-





100 Resolution Orbitrap Mass Spectrometry (UHPLC-HRMS) analysis and stored at 3 °C while the remaining 20 ml was sent to Institute for Atmospheric Physics at the Johannes Gutenberg University of Mainz, Germany for freezing experiments.

### 2.2 Mainz Acoustic Levitator (M-AL)

105 Freezing experiments were performed in an acoustic levitator (APOS BA 10, tec5 GmbH). This allows contact-free single-drop levitation maintained by a standing ultrasonic wave. This setup and its relevant physical influences are described in detail by Diehl et al., (2014), Szakáll et al., (2021), and in part 1 of this publication series by Gautam et al. (2024). For the freezing experiment, the M-AL is placed inside a walk-in cold room where the ambient temperature was set to -15<sup>o</sup>C. The M-AL is surrounded by a 110 protective acrylic housing to prevent any disturbance from air motion which may cause unsteady temperature conditions, unstable levitation, or carry unwanted ice-nucleating particles onto the drop surface. Air temperature in the M-AL was measured by a PT100 sensor and an infrared thermometer (KT 19.82 II, Heitronics) was used to monitor drop surface temperature.

The aqueous filter extract was injected with a syringe into the M-AL node to form a single free-

- 115 floating drop with a diameter of  $2 \pm 0.1$  mm. The drop was allowed to freeze without the introduction of artificial freezing nucleator. Freezing time was approximately 90 seconds on average but not longer than 3 min. Once the drop was fully frozen, it was removed from the M-AL and stored in a polytetrafluoroethylene (PTFE) vial at  $-20\degree$ C until analysis. Enough drop to reach the minimum viable sample volume for analysis, 50 µl, were collected to produce a single sample (approximately 12 drops).
- 120 Two full samples were collected for UHPLC-HRMS analysis. The blank filter extract was also used in the freezing experiment to produce two more travel blank samples for comparative analysis and background subtraction.





#### 2.3 UHPLC-HRMS analysis

- 125 In addition to the M-AL frozen extract and the travel blanks, 100 µl of the reserved extract and Milli-Q solvent was analyzed by UHPLC-HRMS. Chromatographic separation was performed (Vanquish Flex, Thermo Fisher Scientific Inc.) on a reversed phase column (Cortecs Solid Core T3, 2.7  $\mu$ m, 150  $\times$  3 mm, with the corresponding VanGuard Cartridge, Waters Corp.). Samples were ionized in negative and positive mode using a heated electrospray ionization source (HESI-II Probe, Thermo Fisher Scientific
- 130 Inc.) and then detected with a high-resolution hybrid quadrupole-Orbitrap mass spectrometer (Q Exactive Focus, Thermo Fisher Scientific Inc.). The chromatographic settings and gradient are as follows: LC solvent A: ultrapure water with 0.1% formic acid; LC solvent B: methanol with 0.1% formic acid; Flow rate 400 µl min<sup>-1</sup>; pre-column heater and post-column cooler 40 °C; Gradient: 0 min 1% B; 1 min 1% B; 15 min 99% B; 16.5 min 99% B; 17.5 min 1% B; 20 min 1% B. The MS settings were at fullMS scan
- 135 (*m/z* 50-750; resolution 70k) along with data-dependent MS2 in discovery mode (resolution at 17.5k) for acquiring fragmentation spectra of the largest peaks.

#### 2.4 Non-targeted Analysis and Property Estimation

Compound identification confidence is communicated here using the convention described in

- 140 Schymanski et al., (2014). The raw UHPLC-HRMS files were processed on Compound Discoverer 3.3 (Thermo Fisher Scientific). This software aligned chromatographic peaks of interest with a maximum shift of 0.1 min in retention time and a mass tolerance of  $\pm 2$  ppm. Mass traces with retention times less than 1.8 min were excluded as they are not considered to be chromatographically separated. Ions were detected if the peak intensity was at least  $5\times10^5$  counts for [M−H]<sup>-</sup> for negative mode or [M+H]<sup>+</sup> and
- $145$  [M+Na]<sup>+</sup> for positive mode. In addition to the mass-to-charge ratio of the detected ion, at least one corresponding isotopologue had to be measured. The tolerance between the measured and calculated intensity of the isotopologue was less than 30 %. These unknown compounds were then grouped with a





retention time tolerance of 0.1 min to produce a merged MS feature and those of them with a sample-toblank ratio smaller than 5 were marked as background and removed from the dataset. Any compounds

- 150 that did not appear in the reserved sample of filter extract were also removed from the dataset. A peak quality score was given on a scale of 0 to 10—with 10 being a perfect chromatographic peak—for each mass trace based on its peak shape qualities, e.g. peak jaggedness, modality. For all mass traces with a peak quality higher than 6 in all samples, a predicted composition for each mass trace was calculated within  $\pm 2$  ppm with the allowed elements of carbon (C), hydrogen (H), nitrogen (N), oxygen (O), and
- 155 sulfur (S). Compounds were grouped together as CHO, CHNO, CHOS, and CHNOS. It is important to note that phosphorus (P) containing species were not considered for this study. All level 5 (L5) or higher compounds including any mass traces that did not fit a predicted composition within  $\pm 2$  ppm were used for calculating retention coefficients using their integrated MS signals. Level 4 (L4) or higher compounds with determined compositions were used for Van Krevelen and Kroll analysis to highlight the validity of
- 160 the dataset as a real, complex mixture of urban-influenced WSOC. To aid the visualization of MS data, a Van Krevelen diagram cross-plots the H:C ratio as a function of the O:C ratio while a Kroll diagram cross-plots the estimated average carbon oxidation state as a function of the number of carbon atoms. An estimated vapor pressure at 298 K was then calculated for the elemental composition based on the parameterization by Li et al. (2016).
- 165 The predicted compound list was then matched against the mzCloud database (HighChem LLC, 2013-2021) for comparing  $MS<sup>2</sup>$  spectra. If a compound had at least one positive match with the predicted compound in either database as well as a peak quality score above 8 in the reserved extract, this level 3 (L3) or higher tentative candidate was selected to be used for calculating its effective Henry's law constant (K<sub>H</sub>, mol Pa<sup>-1</sup> m<sup>-3</sup>). These properties were predicted using the HENRYWIN<sup>TM</sup> model as part of
- 170 the EPI Suite<sup>TM</sup> package which provides the values at 298 K (US EPA, n.d.). If the EPI Suite<sup>TM</sup> was able to find an experimental value based on a CAS lookup match, those values were used over the model





prediction. This is only applicable to the minority of identified species. The effective Henry's law solubility was converted to a dimensionless effective Henry's law constant  $(H^*)$  using the equation:

$$
H^* = K_H * \overline{R}T \tag{1}
$$

175 Where  $\bar{R}$  is the ideal gas constant (8.3144626 m<sup>3</sup> Pa K<sup>-1</sup> mol<sup>-1</sup>) and T is temperature (K). This conversion allows for a dimensionless comparison and considers dissociation and hydration effects. This calculation, however, requires structural information about a compound. As a result, performing this calculation on a L3 tentative structural candidate can be specious or misleading. Regardless, H\* for structural isomers using HENRYWIN™ typically differs less than 3 orders of magnitude with an overall average of

180 approximately 1.5. This accuracy is sufficient for this analysis (Isaacman-Vanwertz and Aumont, 2021).

#### 2.3 Retention Calculation and Tracer Corrections

The signals of species that also appeared in the travel blanks were first subtracted from the M-AL samples to remove their ambient signal but remained in the dataset as they exceeded the sample-to-blank

- 185 ratio of 5 and could not be considered background. A naturally occurring tracer was selected from the dataset for both positive and negative mode to correct for any dilution, evaporation, and desorption that may occur. To be an ideal tracer, the compound should be fully retained during freezing and have an adequate MS signal. For this work, the peak quality was required to be higher than 8 in the reserved extract and for there to be a positive database match for the predicted composition. For negative mode,
- 190 this tracer was 4-Nitrophenol (C6H5NO3, 139.0269 *m/z*, 9.7 min, L3). This was chosen as previous experiments by Borchers et al., (2024) have identified this compound to have a retention coefficient of  $1.01 \pm 0.07$  during riming experiments, where desorption and evaporation effects are likely to be more influential than in the present experiment due to enhanced ventilation and much smaller droplet size. For positive mode, xylitol  $(C_5H_{12}O_5, 152.0685 m/z, 1.6 min, L3)$  was selected as the tracer as its H<sup>\*</sup> has been
- 195 determined to be higher than  $10^8$  (Compernolle and Müller, 2014), which according to Borchers et al.,





(2024) and Jost et al., (2017) indicate its retention coefficient can be safely assumed to be 1, even under the higher exchange conditions of wind tunnel experiments. Regardless of the accuracy of the L3 assigned structure, these tracers represent species with retentions very close to 1 that allow for a reference to that value to compare between samples and make corrections for non-freezing mass exchange.

200 The equation for calculating the retention coefficient is adopted from Borchers et al., (2024) and Jost et al., (2017) and used here as

$$
R = \frac{s_{\text{substance}}^{\text{sample}} / s_{\text{substance}}^{\text{RES}}}{s_{\text{tracer}}^{\text{sample}} / s_{\text{tracer}}^{\text{RES}}}
$$
(2)

where the numerator describes the ratio between the peak area of the substance of interest in the ice sample ( $S_{\text{substance}}^{\text{sample}}$ ) and in the reserved extract sample ( $S_{\text{substance}}^{\text{RES}}$ ). The denominator describes the same

- 205 ratio but for the tracer ( $S_{\text{tracer}}^{\text{sample}}$  /  $S_{\text{tracer}}^{\text{RES}}$ ). Since no dilution effects are involved in the measurement and all samples are measured in the same aqueous matrix, signal ratios can be compared directly without a calibration curve, provided that detector response is linear within the given range of measurement. Given that ions below the threshold intensity of linearity were excluded from measurement, dynamic mass calibration of the HRMS was performed prior to measurement, and that HRMS instruments of this
- 210 generation show linear dynamic ranges of at least five orders of magnitude (Kaufmann and Walker, 2017), it is reasonable to assume linearity over the measurement range. As ventilation and evaporation effects are quite low in the M-AL (Szakáll et al., 2021), their effects are compensated for by the tracer. Compensating for desorption effects is more complex. As desorption is thought to be driven mainly by linear diffusion and thus enhanced by increased ventilation, species with higher than ambient
- 215 concentrations as well as species with higher vapor pressures are thought to be disproportionally influenced. However, since the M-AL has little ventilation to enhance desorption, HiVol filter sampling already bias against high vapor pressure species (Bidleman et al., 2020), and ambient filter sampling is closer to ambient concentrations relative to previously simulated single component mixtures, desorption





effects are only compensated for by the tracer and nonuniform desorption effects are considered

#### 220 negligible.

Further data analysis was performed with MATLAB ver. R2023a. Distribution modeling was performed using the Distribution Fitter from the Statistics and Machine Learning Toolbox 12.5, based on the Stable Distribution and t Location Scale Distribution models provided in *Univariate Stable Distributions* (2020) and *Univariate Continuous Distributions* (2015) (Nolan, 2020; Yee, 2015).

## 225 3 Results and Discussion

#### 3.1 Dataset Description

In the negative mode, from over 2800 MS features measured, 548 significant, non-background detected compounds were found. 208 met the peak quality constraints and were then used for analysis. 196 compounds (94%) had successfully assigned compositions and 77 were then selected for additional 230 property calculations.

In the positive mode, 342 significant, non-background detected compounds were found from over 1800 features. 250 met the peak quality constraints and were then used for analysis. 218 of those compounds (87%) had successfully assigned compositions. 84 were then selected for additional property calculations. Comparatively fewer compounds were assigned compositions in the positive mode as

235 phosphorous containing species were not considered. These can represent almost a third of positively ionizable species in rainwater WSOC (Seymore et al., 2023) so it likely makes up a significant portion of species variety that is not considered.







Figure 1. (A) Scatter plot of HPLC Retention time (min) vs *m/z* ratio; (B) van Krevelen diagram of O/C 240 ratio vs H/C ratio; (C) Kroll diagram of number of C atoms against the average oxidation state of C; Left panels are for negative ionization mode (−)HESI, right panels are positive ionization mode (+)HESI. Area of the marker indicates relative intensity in the reserved extract while color denotes compositional class of the assigned compound: Green for CHO, blue for CHNO, red for CHOS, yellow for CHNOS, magenta for CHN, cyan for CH.

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Figure 1 illustrates that the dataset is indicative of a typical urban influenced WSOC profile of a dilute sample. In the negative mode, the most significant signals are several nitrophenols and nitroaromatics; notably C<sub>6</sub>H<sub>5</sub>NO<sub>3</sub> (139 *m/z*, 9.7 min, L3) and C<sub>6</sub>H<sub>4</sub>N<sub>2</sub>O<sub>5</sub> (184 *m/z*, 10.3 min, L3) are





tentatively identified as 4-nitrophenol and 2,4-dinitrophenol respectively. These nitrophenols ionize 250 efficiently in (−)HESI which explains in part their prominence in Figure 1.B where they cluster around 0.6-0.9 H/C and are indications of biomass and fossil fuel burning emissions (Taneda et al., 2004). The prominent CHOS compounds in the negative mode are alkylorganosulfates, notably  $C_5H_{12}O_4S$  (168  $m/z$ , 8.4 min, L3) and C8H18O4S (210 *m/z*, 12.9 min, L3), which are typical markers of secondary processed automobile and shipping traffic OM.(Blair et al., 2017; Qi et al., 2021) These can most easily be seen in 255 Figure 1.B above 2 H/C. Some of the other notable CHOS compounds below 2 H/C are terpene-derived

organosulfates such as camphorsulfonic acid  $(C_{10}H_{16}O_4S, 232 m/z, 9.4 m in, L2)$ , which also demonstrate secondary processing under urban conditions (Iinuma et al., 2007; Surratt et al., 2007).

The most significant positive mode signals in Figure 1 come from caprolactam  $(C_6H_{11}NO, 114)$  $m/z$ , 7.0 min, L2) and several coumarin derivatives (C<sub>9</sub>H<sub>6</sub>O<sub>2</sub>, 146  $m/z$  7.4 min, L2; C<sub>9</sub>H<sub>8</sub>O<sub>4</sub>, 162  $m/z$ , 10.9

260 min, L3; C9H8O3, 164 *m/z*, 8.7 min, L3; C10H8O4, 192 *m/z*, 9.0 min, L3). Caprolactam is a cyclic amide and indicative of industrial emission influence as it is primarily used for manufacturing synthetic fibers but also used in numerous other manufacturing activities. Caprolactam is a monitored compound on the hazardous air pollutants list by the United States Environmental Protection Agency (U. S. Environmental Protection Agency, n.d.). Coumarin species are known brown carbon components and have biomass 265 burning sources as well as potential secondary pathways (Xing et al., 2023).

The several other prominent CHNO compounds are mostly amines, e.g.  $C_{11}H_{23}NO_2$  (201  $m/z$ , 3.6) min, L3), C9H11NO2 (165 *m/z*, 3.7 min, L3), DL-Stachydrine (C7H13NO2, 143 *m/z*, 1.8 min, L2), etc. This is consistent with known amine-nitrate aerosol formation during winter months where there are sources of amine salts and semi-volatile organic amine compounds, particularly in areas with high agricultural and

270 combustion emissions (Price et al., 2016). The other prominent CHNO compounds are tentatively identified as amides such as C12H18N2O (206 *m/z*, 6.6 min, L3), C10H14N2O (178 *m/z*, 4.2 min, L4), C<sub>3</sub>H<sub>4</sub>N<sub>4</sub>O<sub>2</sub> (128 *m/z*, 1.8 min, L<sub>3</sub>), etc; which can either be further secondary products of AA (Price et al., 2014) or the result of anthropogenic emissions (Li et al., 2022; Schollée et al., 2017). These amine and





amides tend to have lower retention times and can been seen in the lower cluster in Figure 1.A. The CHO

- 275 species present are generally either aromatics or aliphatic acids and separate out as so in the van Krevelen diagram in Figure 1.B, with aromatics below 1.5 H/C and acids above such as C<sub>12</sub>H<sub>24</sub>O<sub>3</sub> (216 *m/z*, 14.2) min, L3) and C9H10O3 (166 *m/z*, 9.1 min, L3). Very few biogenic CHO species are present as there are very few CHO species within 1.5-1.8 H/C that would indicate humics, ligins, or other raw biomass markers (Qian et al., 2013). This is consistent with the winter season sampling. Further characterization of
- 280 the nonaromatic CHO is difficult to generalize, as there are a variety of sugars, ethers, alcohols, and acids that represent various possible biogenic species, terpenoids, and terpene derivatives. For example, xylitol (possibly arabitol) (C5H12O5, 152 *m/z*, 1.6 min, L3), hexitol (C6H14O6, 182 *m/z*, 1.6 min, L3), cinnamic acid (C<sub>9</sub>H<sub>8</sub>O<sub>2</sub>, 148 *m/z*, 8.7 min, L3), phthalates such as dimethyl phthalate (C<sub>10</sub>H<sub>10</sub>O<sub>4</sub>, 194 *m/z*, 10.9 min, L2) and phthalic acid (C<sub>8</sub>H<sub>6</sub>O<sub>4</sub>, 166 *m/z*, 7.6 min, L3), as well as succinic acid (C<sub>4</sub>H<sub>2</sub>O<sub>3</sub>, 98 *m/z*, 2.6 min,
- 285 L3), levoglucosan (C<sub>6</sub>H<sub>10</sub>O<sub>5</sub>, 162 *m/z* , 4.5 min, L2) and farnesol (C<sub>15</sub>H<sub>26</sub>O, 222 *m/z*, 14.6 min, L3) are all species potentially identified in the dataset.

A few CH and CHN compounds were found only in the positive mode, primarily AA and a couple of aromatic hydrocarbons. Combined, these represent less than 11% of the positive mode compounds. Only one CHS species was identified  $(C_{18}H_{12}S, 260 \frac{m}{z}$ , 10.0 min, L4) but was not used for analysis as it 290 was below peak quality requirements. No CH, CHN, or CHS species were found in the negative mode.

#### 3.2 Retention Coefficients

Table 1. Mean and Median Retention Coefficients by Compound Class and Heteroatom Group









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Figure 2. Histograms of Retention Coefficients for all measured L5 and above compounds that met peak quality constraints; (A) (−)HESI, (B) (+)HESI

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Figure 3. Merged Histogram for (−)HESI and (+)HESI of all Retention Coefficients measured fit with Stable and t Location-Scale distribution parameterizations of statistical density.





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Table 3. Parameters for t Location-Scale Distribution Fit presented in Figure 3

Mean	0.9442	
<b>Log Likelihood</b>	210.775	
Parameter	Value	<b>Std. Error</b>
Mu(u)	0.94415	0.00566
Sigma $(\sigma)$	0.08665	0.00630
Nu(v)	2.02867	0.26969







Figure 4. Probability plot of Retention Coefficients with the Stable and t Location-Scale distribution 310 parameterizations and a Normal distribution

The histograms presented in Figure 2 and Figure 3 illustrate the distribution of retention coefficients determined for this dataset. Each histogram shows a peak at 0.96, 0.93, and 0.94 for (−)HESI, (+)HESI, and the full dataset respectively with average retentions all at 0.95 with standard deviations of 315 0.21 and 0.53 respectively. These values and the values for each compound class are presented in Table 1. Visually the distributions in Figure 2 appear nonnormal, suggesting a true distribution is being measured. Additionally, Figure 4 shows the data deviates strongly from a normal distribution. Both Shapiro-Wilk and Shapiro-Francia tests indicate nonnormality (p-values: 0.4052, 0.3940 (−)HESI; 0.5698, 0.5611 (+)HESI respectively).





320 Combining these distributions and filtering out outliers, the dataset is fitted with two distributions to model the dataset: Stable and t Location-Scale. The parameters for these fits can be found in Tables 2 and 3. These distributions are functional estimations of the statistical density of retention coefficients based on the empirical measurements in this experiment. The Stable distribution appears to model the data most accurately as the parameter errors are lower than the t Location-Scale and the data points in 325 Figure 4 lie closer to the Stable distribution curve.



Figure 5. Boxplot of Retention Coefficients by Composition Class; (A) (−)HESI, (B) (+)HESI

330 The means of the composition classes in (−)HESI vary little, generally less than 5% from each other. In contrast, the means of the composition classes in (+)HESI vary more, up to 40%. The deviations and ranges of values are also wider in (+)HESI, from 0.21 to 0.82 as seen in Table 1. Visually this can be seen in Figure 5. For both ionizations, CHNOS tends to be the highest retained along with CHN in (+)HESI. CHNOS represents more of the heaviest species in the sample set while CHN is entirely AA. In 335 (+)HESI, CHNOS, CHOS, and CH represent the smallest portion of the dataset—less than 6%—and some of the most variably retained species. For CH, this follows with the variability in hydrocarbon

aqueous solubility, however this variability is more likely explained for CHNOS, CHOS as well with the





smaller sample set bias. Notably, CHO has lower retention with a wider distribution than CHNO in both ionizations. In (−)HESI, CHNO is mostly nitroaromatics while in (+)HESI, CHNO is mostly amines and

- 340 amides. CHO represents a more similar distribution of organic acids and terpenoids in both positive and negative mode, with more nonpolar species represented in (+)HESI. The lower retention among CHO may then be based on its distribution of organic acids versus terpenoids. This data suggests that nitrate species and amines/amides have similar retentions. It is known that  $NO<sub>x</sub>$  removal is enhanced by aqueous phase reactions (Daito et al., 2000) and that organic nitrogen represents an important fraction of WSOC
- 345 (Saxena and Hildemann, 1996; Zhang et al., 2002), but this data may also indicate that nitrogen chemistry on CHO species enhances their retention in hydrometeors. Other UHRMS studies of rainwater have suggested similar explanations for nitrogen uptake in hydrometeors and rainwater organic nitrogen's high bioavailability (Seymore et al., 2023).



Figure 6. Violin plots of Retention Coefficients by Composition Class; (A) (−)HESI, (B) (+)HESI

The distributions of CHNO and CHOS in (−)HESI seen in Figure 6 show a coincidence of the mean and median along with strong symmetry around the mean. This indicates visually that they appear to be normally distributed, suggesting that the true retentions for the whole of these compound classes 355 may be close to 1. Both Shapiro-Wilk and Shapiro-Francia tests indicate normality (p-values: 0.1920,





0.1922) for CHNO but not for CHOS (7.7595  $\times$  10<sup>-9</sup>). It therefore would be reasonable to assume a normal distribution for CHNO, but CHOS cannot be rigorously stated as normally distributed. CHO and CHNOS show unique distributions with a significant number of values within  $-\sigma$  and below, indicating there are certain CHO species that are not retained during freezing.

- 360 For (+)HESI, the samples sizes for CHNOS, CHOS, and CH are too small to make meaningful descriptions of their distributions. For CHO, CHN, CHNO, the distributions are visually nonnormal and also do not pass any statistical test for normality. The distributions also appear less smooth than their negative mode counterparts, likely a result of previously discussed ionization variability in (+)HESI. Notably, (+)HESI shows a few species with very low retention specifically within the CHO, CHOS, and
- 365 CH groups. Specifically, these are C14H22 (190 *m/z*, 14.5 min, L3), C15H26O (222 *m/z*, 14.6 min, L3), and what appears to be a phenyl-sulfide species ( $C_{16}H_{18}OS$ , 258  $m/z$ , 8.0 min, L4). This is the only identified organosulfide within the dataset. It would be speculative to say that its low retention may indicate that organosulfides as a class are unretained and thus unlikely to appear in the dataset. Its low retention likely has more to do with its low polarity.  $C_{14}H_{22}$  and  $C_{15}H_{26}O$  as long chain, nonpolar species demonstrate that

370 species with lower aqueous solubilities are also likely to have low retentions.

Concerning heteroatoms, the distributions and ranges of retentions are quite similar among all groups. Oxygen-containing species appear to have a slightly wider distribution which is mostly weighted by the CHO class. Nitrogen-containing species have a smaller standard deviation than the O or S containing species, indicating fewer species with variable retentions and more fully retained compounds. This further

375 suggests that nitrogen inclusion enhances retention (see also Figure S1).

### 3.3 Correlation of Retention Coefficients with Chemical Properties







Figure 7. Retention Coefficient as a function of HPLC Retention time; (A) (−)HESI, (B) (+)HESI; Color denotes compositional class of the assigned compound, as used in Fig. 1: Green for CHO, blue for CHNO, red for CHOS, yellow for CHNOS, magenta for CHN, cyan for CH. Dashed line shows linear fit.

- 385 The determined molecular weight (MW) shows little correlation linearly with retention (as seen in Figure S2). In (−)HESI, there is a weak positive correlation, suggesting larger compounds are more likely to be retained. This is likely related to lower vapor pressures associated with larger MW species in the negative mode. An F-test against the constant value model indicates that this correlation is not significant (p-value: 0.0857). However in (+)HESI, there's a very weak negative correlation for MW 390 which suggests the opposite. An F-test against the constant value model indicates that this correlation with MW is also not significant (p-value: 0.1440). This trend in the positive mode is likely driven more by polarity, as Figure 1.A also demonstrates that larger species in (+)HESI tend to have higher HPLC retention times and are therefore more nonpolar. The plot in Figure 8 further demonstrates this with a stronger negative correlation between the HPLC retention time and the retention.
- 395 Figures 7 shows correlations for retention coefficients with HPLC retention time. With reversephase HPLC, retention time is a direct proxy for molecular polarity, i.e. shorter retention times indicate higher polarity and longer retention times indicate more nonpolar species. Both (−)HESI and (+)HESI show significant negative correlation between retention and retention time and therefore polarity; an F-

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- test against the constant value model shows p-values of 0.00193 for (−)HESI and  $1.44 \times 10^{-6}$  for (+)HESI. 400 This indicates that nonpolar species are likely to be unretained and this appears to be especially true for the previously discussed long chain species, such as  $C_{14}H_{22}$  (190  $m/z$ , 14.5 min, L3) and  $C_{15}H_{26}O$  (222)  $m/z$ , 14.6 min, L3) compounds. In Figure 7, a few compound classes separate distinctly by polarity, particularly the CHOS and CHNO in (−)HESI as well as CHO and CHNO in (+)HESI. These polarity differences in these classes may be the driving force in the difference of the retention between CHOS and
- 405 CHNO in (−)HESI, but unlikely for CHO and CHNO in (+)HESI as CHO spans a much wider range of retentions that cannot be explained solely by polarity.



Figure 8. Retention Coefficient as a function of Estimated Vapor Pressure; Color denotes compositional class of the assigned compound: Green for CHO, blue for CHNO, magenta for CHN, cyan for CH, yellow 410 squares for values taken from literature. Dashed line shows linear fit.

Further chemical property correlations with retention could be made for the species with estimated chemical properties. The first plot, Figure 8, uses the measured retention coefficient to plot against calculated vapor pressure (VP). It demonstrates a significant negative correlation with VP; an F-





- 415 test against the constant value model gives a p-value of  $1.42 \times 10^{-4}$ . It is relevant to note that the majority of species measured are considered semi-volatile (vapor pressure:  $10<sup>-9</sup>$  to 10 Pa; SVOC) with few low volatility (LVOC) and intermediate volatility organic compounds (IVOC) (Weschler and Nazaroff, 2008). LVOC and IVOC are bias against in the sampling method as many LVOCs are highly oxygenated compounds which may be less sensitive compared to other compounds in UHPLC-Orbitrap MS (Wang et
- 420 al., 2024) and most IVOCs are revolatilized during sampling (Bidleman et al., 2020). VP is also the property most associated with desorption effects, likely contributing to some of the negative trend seen in Figure 8. However, it is mostly associated with IVOC and less with SVOC so desorption alone is not likely to fully explain this correlation.



Figure 9. Effective Henry's Law Constant H\* versus Retention Coefficient; (A) (−)HESI, (B) (+)HESI; Color denotes compositional class of the assigned compound: Green for CHO, blue for CHNO, magenta for CHN, cyan for CH. Dashed line shows linear fit.

430 Demonstrated in Figure 9, retention shows little or no dependency on H\* under the present experimental conditions. The linear correlations are not significant for neither (−)HESI or (+)HESI; with p-values of 0.9270 and 0.3530 respectively for the F-test against the constant value model. The very slight





positive correlation shows agreement with Stuart and Jacobson's (2003, 2004) observation that high H\* species are more likely to be 100% retained but does not show sigmoidal behavior as modeled by Jost et

435 al., 2017 (plotted for reference in Figure 10). This could be the result of differing physical experimental parameters, such as larger droplet size (2 mm versus 20 micron). However, directly comparing the known literature values for retention coefficients with the observations made here does not immediately indicate the systems are incompatible or exclude the comparison.



440 Figure 10. Effective Henry's Law Constant H\* versus Retention Coefficient; Color denotes compositional class of the assigned compound: Green for CHO, blue for CHNO, magenta for CHN, cyan for CH; Yellow squares denote values taken from Borchers et al., (2024), Jost et al., (2017), and Von Blohn et al., (2011, 2013). Dashed line shows linear fit. Solid red line gives the parameterization by Jost et al., (2017).





Figure 10 compares the data presented in Figure 9 against the literature values presented by 445 Borchers et al., (2024), Jost et al., 2017, and Von Blohn et al., (2011, 2013). This comparison does not appear incongruous, i.e. no discernible difference can be seen between wind tunnel experiments and these observations. While the measurements presented by Borchers et al., (2024), Jost et al., (2017), and Von Blohn et al., (2011, 2013) are physically dissimilar experiments to this study—i.e. wind tunnel experiments, small droplets of micrometer size, high ventilation conditions—their observations are

- 450 congruent with this experiment. The literature values include only three species with  $H^* < 10^4$  and the lowest H\* of them, pinandiol, is excluded from the parameterization presented by Borchers et al., (2024) as it was perceived as an outlier. Without more measurements of compounds with  $H^* < 10^4$  under wind tunnel conditions, it is difficult to determine if the nonsigmoidal behavior seen in this experiment is the result of different physical parameters—specifically the lower surface-to-volume ratio—or if the
- 455 sigmoidal behavior described by Borchers et al., (2024) and Jost et al., (2017) is an overfit of a limited dataset. Evidence presented by Gautam et al., (2024) in part 1 of this publication series makes a compelling case for the dominance of physical parameters at these drop sizes. Critical to these experiments, Gautam et al., (2024) observed the formation of an ice shell, which inhibited any further expulsion of dissolved substances during freezing.

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### 4 Conclusions

This study presents the measurement of the retention coefficients for real, complex WSOCs from urban particulate matter for direct drop freezing under raindrop size conditions. The overall distribution of the retention of WSOCs forms a real, nonnormal distribution up to 1. Looking at the individual

465 compound classes of organics, the data shows that they may have different distributions of retention coefficients. Most negatively ionizable CHNO and CHOS compounds appear to be fully retained, indicating that nitroaromatics and organosulphates are favorable to be retained. Slight positive

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correlations between MW, polarity, and  $H^*$  are seen with retention along with a negative correlation with VP. No sigmoidal relationship with H\* was observed. This is likely the result of the lower surface-to-area

- 470 ratio for this drop size and the ice shell formation observed by Gautam et al., (2024) in part 1 of this publication series. However without further measurements of single component solutions for compounds with  $H^*$  < 10<sup>4</sup> under wind tunnel conditions, specifically for small cloud droplet sizes, it is difficult to determine if the nonsigmoidal behavior seen in this experiment is solely the result of physical parameters or if the sigmoidal behavior described by other studies is an overfit of a limited dataset.
- 475 Sulfides, lipids, aromatic hydrocarbons, and long chain compounds are among the most unretained and incidentally the fewest species observed. These are also among the most nonpolar species observed, which is presumably the dominant factor in that regard. CHO species show the highest variability for their measured retentions, most likely related to the distributions of polarity and VPs among the sugars, organic acids, and terpenoids seen here.
- 480 AA don't follow the trends associated with polarity and VP but are among the most highly retained species. The explanation for this is possibly in its structural properties, which cannot be easily determined using this analytical method. AA solubility in water is largely determined by the dimension and structure of the alkyl substituents, such that AA with longer chains are less soluble than AA with shorter chains and AA with branched substituents are less soluble than AA with linear groups with the 485 same number of carbons (Badocco et al., 2015). Polarity and hydrogen bonding are also known contributors to AA solubility but this is not unique to AA. Hydrogen bonding potential may enhance retention along oxygen functionalities such as among nitro and sulphate species. Hydrogen bonding alone, however, is unlikely to fully explain the high retentions among nitro and sulphate species. Specifically, nitro species have weak hydrogen bonding potential (Shugrue et al., 2016) and as a result is
- 490 likely influenced mostly by the increased polarity imparted by the nitro substituent or its dissociation.

The data suggests that nitrogen and sulfur inclusion generally increase a species' ability to be retained. Overall, this insulates that the products of  $NO<sub>x</sub>$  and  $SO<sub>x</sub>$  reactions from anthropogenic emissions





enhances the retention of these SOA species, reducing the likelihood of reaching the upper atmosphere. Further on this, other studies have demonstrated that  $NO<sub>x</sub>$  removal is enhanced by aqueous phase

- 495 reactions (Daito et al., 2000). These findings may also indicate that this  $NO<sub>x</sub>$  chemistry on CHO species enhances their retention in hydrometeors, potentially by increasing its polarity or solubility. Other UHRMS studies of rainwater have suggested similar explanations for nitrogen uptake in hydrometeors and rainwater organic nitrogen's high bioavailability (Seymore et al., 2023). Additionally, correlations with VP and polarity show that lower VP species and more polar species tend to be retained. Atmospheric
- 500 chemical processing generally tends to oxidatively degrade large nonpolar species into more watersoluble, less volatile species (Iavorivska et al., 2016). Specifically, aqueous phase droplet chemistry is known to facilitate condensed phase SOA formation from highly volatile species (McNeill, 2015). However, many freshly aged terpene products increase in volatility or see only small decreases in VP with oxidation for their early generation products (Bilde and Pandis, 2001; Kurtén et al., 2018; Wu et al.,
- 505 2021). This insulates that many freshly oxidized SOA precursors may have a lower potential to be retained than aged organics and may generally suggest that freshly oxidized SOA precursors are more likely to reach the upper atmosphere than primary organics or aged SOA.

The use of UHPLC-HRMS has allowed for the study of ambient WSOC retention rather than single component or limited mixture experiments from previous studies. The experiment in this paper

- 510 demonstrates the viability of UHPLC-HRMS analysis for ambient WSOC and shows the need for further complex mixture study regarding retention. Future study on retention within hydrometeors should include complex mixture analysis under the physical conditions most similar to the atmosphere, i.e. wind tunnel experiments, smaller droplets, increased ventilation. These studies would also be improved with more distinguishable tracers with known retentions and more sophisticated corrections for desorption. Further
- 515 studies on single component solutions of species with  $H^* < 10^4$  under atmospherically similar physical conditions would also allow for stronger conclusions from the comparison of the retentions measured in this study with other experiments performed under wind tunnel conditions.





The experiment presented here also cannot distinguish between species incorporated within ice crystal structure and those phase-separated but physically constrained to the hydrometeor, potentially

- 520 between crystal grain boundaries or on the particle surface. That distinction is also not atmospherically relevant regarding the net transport of organics into the upper troposphere. While this method aims to demonstrate the retention for real WSOC, this method is still sample method biased against higher volatility species and likely features other sampling bias typical for HiVol filter based measurements such as filter extraction bias and solvation effects. These measurements also present the distribution of
- 525 retention coefficients for the variety of species present and not necessarily the mass distribution of species potentially present in the atmosphere. Corrections for species abundancy must first be made in order to apply this data to organic transport models.

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### Author Contributions

JS, MG, MS, AT participated in designing the experiments; LZ provided the samples; JS, JM prepared the samples for experiments; MG performed the experiments; JS, JM, AV conducted the analytical





measurements; JS analyzed the data and wrote the manuscript draft; MG, MS, AT, JM, AV, TH reviewed

540 and edited the manuscript.

# Competing Interests

The contact author has declared that none of the authors has any competing interests.

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