

# 1 Investigating small ion number size distributions: 2 insight into cluster formation and growth

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## 12 Abstract

13 Small ions, consisting mostly of charged molecular clusters with mobility diameters below 2 nm,  
14 exist continuously in the atmosphere. Here, we studied small ion number size distributions  
15 measured with Neutral cluster and Air Ion Spectrometer measurements in Hyytiälä, Finland and  
16 Beijing, China. We found that in Hyytiälä, there is a strong positive relationship between the  
17 concentration and diameter of small ions of both polarities and highly oxidized organic molecule  
18 (HOM) and sulfuric acid concentrations, and that the relationship with the former is especially  
19 strong. The relationship between the negative sulfuric acid cluster ions and the small ion number  
20 size distribution in Hyytiälä was found to be more complex, but overall positive. In contrast to  
21 Hyytiälä, we found that in Beijing the small ion number size distribution does not have a clear  
22 relationship with sulfuric acid or oxidized organic molecule (OOM) concentration. We found that  
23 the small ion size distribution in Hyytiälä behaved as expected with respect to varying coagulation  
24 sink (CoagS), with concentrations of the smallest ions decreasing most with increasing CoagS.  
25 Surprisingly, the small ion size distribution in Beijing did not vary significantly with varying  
26 CoagS. However, in both locations, the impact of growth on the small ion number size distribution  
27 during periods of intense cluster formation and new particle formation is clearly seen. Our results  
28 show that while in Hyytiälä the growth of small ions to larger diameters is limited by the  
29 concentrations of sulfuric acid and OOMs, in Beijing there are additional factors required for the  
30 small ions to grow.

## 31 1 Introduction

32 Atmospheric aerosol particles influence the Earth's climate (e.g., Quaas et al., 2009; Boucher et al.,  
33 2013; Schmale et al., 2021; Li et al., 2022) and can have adverse effects on human health (e.g.,  
34 Shiraiwa et al., 2017; Arfin et al., 2023). These influences have commonly been related to  
35 properties, such as the mass or number concentration of an atmospheric aerosol population, its size  
36 distribution, or its chemical composition (Shiraiwa et al., 2017; Atkinson et al., 2015; Finlay, 2021).  
37 The electric charging state of atmospheric aerosols has attracted much less interest, although this  
38 property may have large influences on the dynamics of atmospheric aerosol populations (Harrison  
39 and Carslaw, 2003; Fdez-Arroyabe et al., 2022), thereby affecting many other important aerosol

40 properties. The presence of charges also makes it possible to measure low aerosol concentrations at  
41 high resolution in both time and particle size (Mirme and Mirme, 2013; Mirme et al., 2024).

42 Charged atmospheric particles, or more broadly ions, include charged aerosol particles, charged  
43 molecular clusters, and even large molecules having a charge. Ions with electrical mobility  
44 diameters roughly below 2 nm in diameter are classified as small ions, and consist of charged  
45 molecular clusters, while ions above 2 nm consist of charged aerosol particles (Tammet, 1995; Ehn  
46 et al., 2010). Of these charged aerosol particles, those with diameters between 2 and 7 nm are  
47 referred to as intermediate ions (Tammet, 1995).

48 Atmospheric ions are created through ionization of molecules the atmosphere. Most important of  
49 these ionization sources are cosmic ray radiation, gamma radiation, and radon decay (Harrison and  
50 Tammet, 2008). Small ions are constantly present in the troposphere as molecules are ionized and  
51 subsequently grow to small ions (Harrison and Tammet, 2008; Hirsikko et al., 2011). The lifetime of  
52 small ions is short at around 100 s, and their chemical composition depends on the atmospheric  
53 trace gas concentrations and their chemistry (Harrison and Tammet, 2008; Ehn et al., 2010; Shuman  
54 et al., 2015). In contrast, intermediate ions are typically detected mainly during the occurrence of  
55 atmospheric new particle formation (Tammet et al., 2014, Tuovinen et al., 2024), or during snowfall  
56 or rain (Hirsikko et al., 2007; Tammet et al., 2014). New particle formation (NPF) is considered to  
57 occur when constantly existing stable clusters, neutral or charged, start to grow to larger sizes by  
58 uptake of precursor vapors such as sulfuric acid and organic compounds with low volatilities  
59 (Kulmala et al., 2006; Kulmala et al., 2007; Lehtipalo et al., 2018; Kirkby et al., 2023).

60 A recent study by Kulmala et al. (2024a) presented the use of a novel cluster ion counter (CIC) for  
61 measuring small and intermediate ion concentrations to study local-scale NPF and to derive other  
62 parameters such as condensation sink (CS). The information gained by these measurements can be  
63 used further to study the complex climate-biosphere feedbacks (Kulmala et al., 2020; Kulmala et  
64 al., 2024b). These recent advances have motivated us to take a deeper look at the small ion size  
65 distribution.

66 The concentration of small ions depends on the ionization rate and the losses of small ions due to  
67 ion-ion recombination, coagulation with larger aerosol particles, and deposition (Tammet et al.,  
68 2006; Hörrak et al., 2008). The size of small ions depends on their chemical composition and age as  
69 the ions grow through chemical reactions and condensation of vapors, or through coagulation with  
70 neutral clusters. By investigating small ion number size distributions, we can learn more about these  
71 chemical and dynamical processes.

72 In this study, we combine ion number size distribution data measured by Neutral cluster and Air Ion  
73 Spectrometer (NAIS; Manninen et al., 2009; Mirme and Mirme, 2013) with concentrations of low-  
74 volatility vapors and ion clusters measured by mass spectrometer instruments to identify how, and  
75 why, the size distribution of small ions changes and evolves. Data from two different contrasting  
76 locations, Hyytiälä, Finland and Beijing, China (Kulmala et al., 2025), are used. First, we will study  
77 if the variation of the small ion size distribution with season is considerable. Secondly, we will  
78 quantify the potential relationship of organic low-volatility vapors and sulfuric acid on the size and  
79 number of small ions. Third, we will analyze the impact of coagulation scavenging on small ion size  
80 distribution. Fourth, we will analyze the small ion size distribution as a function of intensity of NPF  
81 to reveal how the small ion size distribution changes as the clusters grow. Finally, some case studies

82 are presented. With these, we aim to identify the most important processes impacting the small ion  
 83 number size distribution, and to evaluate the role of these processes in driving the growth of small  
 84 ions to intermediate ions.

## 85 **2 Background and methods**

### 86 **2.1 Evolution of small ion size distribution**

87 Typically, the parameter of interest when considering small ions is their total number concentration  
 88 and its temporal evolution. The changes in the small ion number concentration can be described by  
 89 the simplified air ion balance equation:

$$\frac{dN^{\pm}}{dt} = Q - CoagS N^{\pm} - \alpha N^{\pm} N^{\mp} - S N^{\pm} \quad (1)$$

90 Here,  $N^{\pm}$  is the concentration of one polarity, while  $N^{\mp}$  is the concentration of the other polarity.  
 91 The first term on the right-hand side of the equation describes the source rate of the ions, where  $Q$  is  
 92 the ionization rate of air molecules. The second term, where CoagS stands for coagulation sink,  
 93 tells the loss rate of small ions due to coagulation on larger aerosol particles. The third term tells the  
 94 loss rate of ions due to ion-ion recombination, where  $\alpha$  is the ion recombination coefficient. The  
 95 final term describes other losses of the ions, including deposition, and  $S$  is the loss rate of the ions to  
 96 these other sinks.

97  
 98 As we can see, the above equation does not explicitly depend on the size of the small ions nor can it  
 99 be directly used to describe the evolution of the size-dependent small ion size distribution. The time  
 100 evolution of small ions of certain size  $i$  are described by the charged general dynamics equations  
 101 (charged GDEs; Kulmala et al., 2012):

$$\frac{dN_i^{\pm}}{dt} = J_i + \chi N_i N_{d<i}^{\pm} - N_i^{\pm} CoagS_i - \alpha N_i^{\pm} N_{d<i}^{\mp} - \frac{GR}{\Delta d_i} N_i^{\pm} \quad (2)$$

102 Here,  $J_i$  is the formation rates of ions of size  $i$ . The second term on the right-hand side represents the  
 103 charging of neutral clusters by ions smaller than  $i$ , where  $\chi$  is the ion-cluster attachment coefficient.  
 104 The last term, where GR is the ion growth rate, describes the growth of ions  $i$  to larger sizes.  
 105 Considering Eq. 2, we can see that an increasing GR will shift the ion size distribution towards  
 106 larger diameters. CoagS is the highest for the smallest ions and if it increases, the concentrations of  
 107 smallest ions are decreasing the most, causing an apparent shift in the distribution towards larger  
 108 diameters. However, CoagS also affect the lifetime of small ions, so that with an increasing CoagS  
 109 the ions have less time to grow, reducing the concentration of larger small ions. If ion  
 110 concentrations are high, ion-ion recombination rate will be higher, which will also lead to shorter  
 111 small ion lifetime and possibly smaller concentrations of larger small ions. Through ion-cluster  
 112 attachment, the small ion size distribution depends on the size distribution of neutral clusters,  
 113 although this term is relatively small when compared to the coagulation loss and growth terms.

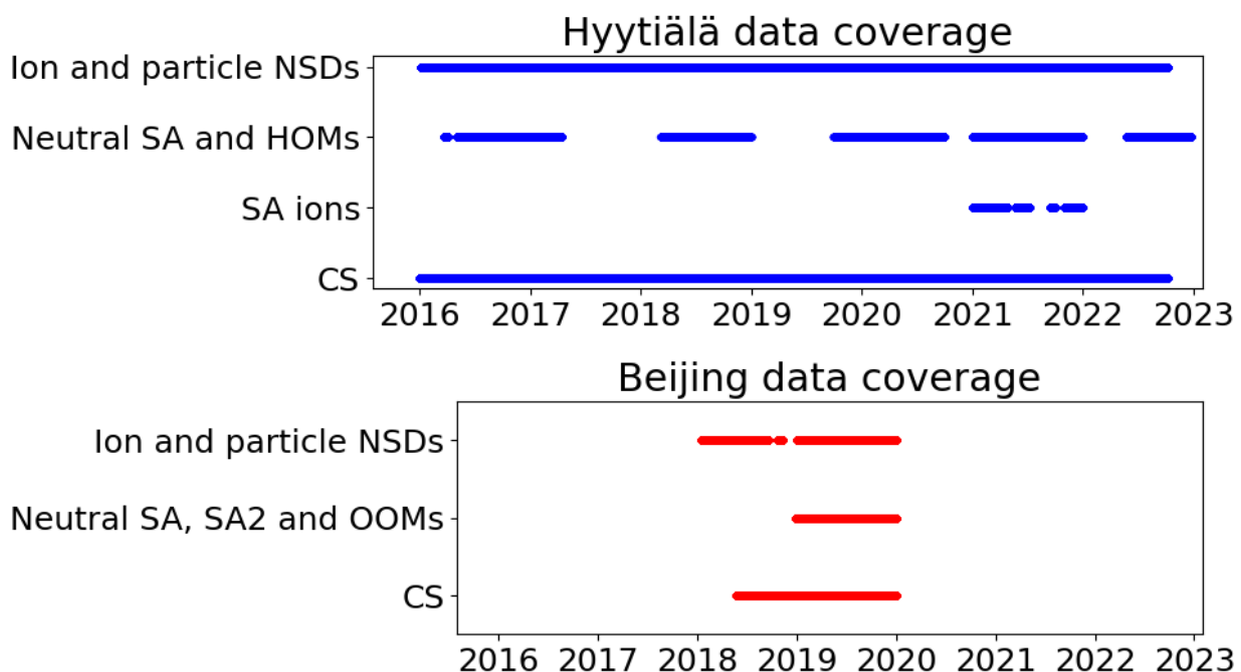
114 We assume that the role of direct transport of clusters on the changes in the small ion size  
 115 distribution is negligible due to their short lifetime of just a couple of minutes (Tamm et al.,  
 116 2006). Therefore, the observations are assumed to be very local. However, transport can indirectly  
 117 impact the size distribution of small ions i.e., through transport of trace gases and larger particles.  
 118 We also note that while meteorological conditions, such as temperature, strongly influence

119 processes such as HOM formation (e.g., Quéléver et al., 2019), we do not explicitly consider them  
120 in this study.

## 121 2.2 Measurement sites

122 Two different locations were considered in this study: SMEAR II measurement station in Hyytiälä,  
123 Finland (61°51' N, 24°17' E) and BUCT/AHL measurement station in Beijing, China (39°94' N,  
124 116°30' E). The former is a rural site surrounded by boreal forest while the latter is an urban site  
125 close to residential building and traffic roads. For more details on SMEAR II station, see Hari and  
126 Kulmala. (2005). For more details on BUCT/AHL site, see Liu et al. (2020). These two locations  
127 are included in the analysis due to their contrasting natures, providing an opportunity for insight  
128 into the variation of small ion size distribution and small ion dynamics in different environments.

## 129 2.3 Measurement and other data



130 **Fig.1:** Data coverage for the two sites, Hyytiälä, Finland, and Beijing, China, from which data were  
131 used in this study. NSD refers to number size distribution, while SA refers to sulfuric acid, SA2 to  
132 neutral sulfuric acid dimer, HOM to highly oxidized organic molecule and OOM to oxidized  
133 organic molecule.  
134

135  
136 Atmospheric ion and total particle number size distributions in Hyytiälä and Beijing were measured  
137 with Neutral cluster and Air Ion Spectrometer (NAIS; Manninen et al., 2009; Mirme and Mirme,  
138 2013). The NAIS measures both charged and total particle number size distributions in the ranges  
139 0.8-42 nm and 2.5-42 nm, respectively. Main focus of the analysis in this study is on the number  
140 size distributions of small ions (diameters below 2 nm). Ion concentrations between 2.0 and 2.3 nm  
141 were used to characterize the intensity of local clustering (Tuovinen et al., 2023) and new particle  
142 formation ranking data, characterizing the intensity of NPF, were also used. The NPF ranking was  
143 based on the total particle number concentration between 2.5 and 5 nm and determined according to  
144 the method presented by Aliaga et al. (2023).

145 The chemical composition of small ions typically differs between the polarities (Ehn et al., 2010;  
146 Zha et al., 2023). For example, Ehn et al. (2010) found that in Hyytiälä the daytime negative small  
147 ions consisted largely of sulfuric acid clusters, while positive small ions consisted of organic species  
148 such as alkyl pyridines and alkyl amines. Therefore, we cannot assume that the negative and  
149 positive small ion populations behave similarly with respect to i.e., increased sulfuric acid  
150 concentrations. Thus, both negative and positive polarity were separately considered.

151 All diameters used in study are electrical mobility diameters. We note that especially for the  
152 smallest of the ions the mobility diameter may not accurately describe the physical dimensions of  
153 the ion (see e.g., Ehn et al., 2011). Regardless, we refer to diameter rather than electrical mobility as  
154 we see it as more intuitively understandable parameter for the ion size.

155 From Hyytiälä, concentrations of neutral sulfuric acid and highly oxidized organic molecules  
156 (HOMs) were used to study the influence of cluster formation and growth on the small ion size  
157 distribution. These were measured with Chemical Ionization Atmospheric Pressure interface Time-  
158 Of-Flight (CI-APi-TOF) mass spectrometer (Jokinen et al., 2012). In addition, the signal counts of  
159 ionized sulfuric acid clusters measured with APi-TOF were used to give further insight into the  
160 composition of the small ions. The signal counts in the study are given as relative signals to the total  
161 measured ion current. From Beijing, neutral sulfuric acid, sulfuric acid dimer and total oxidized  
162 organic molecule (OOM) concentrations, which were measured with a nitrate based – long time-of-  
163 flight chemical ionization mass spectrometer (CIMS), were included in the analysis. We note that  
164 we use the term OOM instead of HOM for the organic molecules in Beijing based on previous  
165 results by Yan et al. (2021), suggesting that most of these measured organic molecules in Beijing do  
166 not meet the requirements for HOMs (see Bianchi et al., 2019).

167  
168 Condensation sinks (CS) for both sites were included in the analysis as proxies for the coagulation  
169 scavenging of the small ions. In Hyytiälä, the particle size distributions from which CS was derived  
170 from were measured with a twin Differential Mobility Particle Sizer (DMPS) system (Aalto et al.,  
171 2001). In Beijing, the particle size distributions for CS were measured with a particle size  
172 distribution (PSD) system (Liu et al., 2016).

173  
174 Data coverage for both sites is presented in Fig. 1.

## 175 **2.4 Determining the average small ion diameter**

176 From the small ion number size distributions, we determined the mean mobility diameter ( $d_{mean}$ ),  
177 and median mobility diameter ( $d_{median}$ ) of small ions. First, cubic interpolation was applied to the  
178 measured ion number size distributions. We note that nearest neighbor and linear interpolation  
179 methods were also tested, and the influence of the chosen method on the value of  $d_{mean}$  or  $d_{median}$  was  
180 found minor. The diameter range for the interpolation was from the lower detection limit to 2 nm  
181 with a step of 0.001 nm. Then,  $d_{peak}$  was determined by finding the diameter corresponding to the  
182 maximum concentration of small ions. Weighted mean and median were used to determine  $d_{mean}$  and  
183  $d_{median}$ , with the number concentrations of ions below 2 nm in diameter used as weights. The  
184 equation below was used to find weighted mean diameter:

$$185 \quad d_{mean} = \frac{\sum N_i d_i}{\sum N_i}, \quad (3)$$

186 where  $d_i$  is the diameter of ions of a certain size and  $N_i$  is their number concentration. The weighted  
187 median was determined by finding the diameter  $d_j$  satisfying

$$j = \min_k \left[ \sum N_i d_i > \frac{1}{2} \sum N_i d_i \right]. \quad (4)$$

188

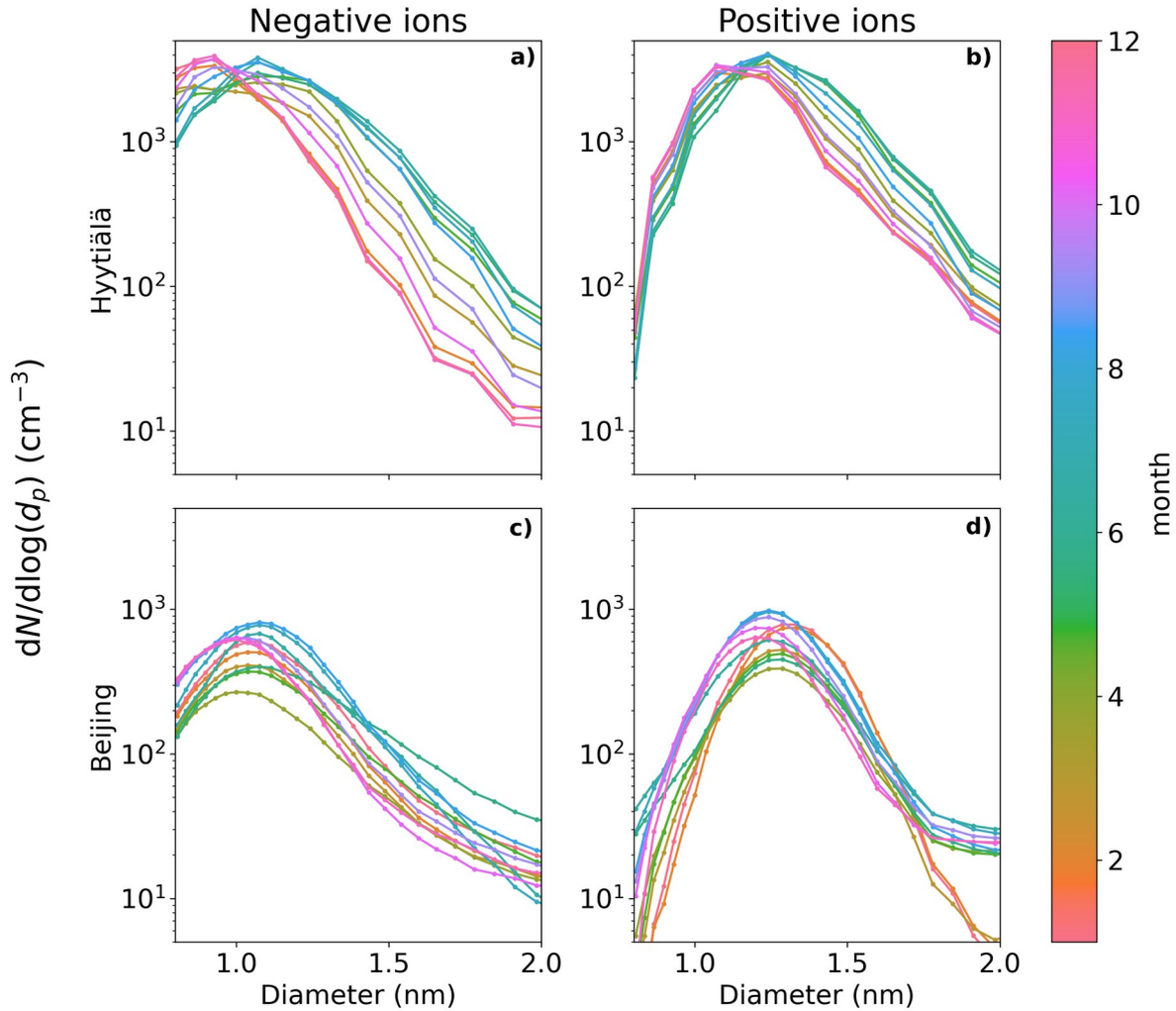
## 189 **3 Results**

### 190 **3.1 Seasonal variation of the small ion size distribution**

191 The Fig. 2a and b show the monthly median negative and positive ion distributions between 0.8 and  
 192 2 nm in Hyytiälä. Table 1 records the monthly mean and median diameters ( $d_{\text{mean}}$  and  $d_{\text{median}}$ ). Clear  
 193 month-to-month changes in the size distributions are observed, and these are more pronounced for  
 194 negative ions (Fig. 2a). During winter, the concentration of negative ions peaks already below 1.0  
 195 nm, while during summer the highest concentration is between 1.1 and 1.2 nm. In addition, the  
 196 concentrations of negative ions above 1.1 nm are increased from winter to summer. Close to 2 nm,  
 197 the ion concentration is almost one order of magnitude higher during summer. This behavior of the  
 198 size distribution is reflected in the values of  $d_{\text{mean}}$  and  $d_{\text{median}}$ , which are smallest during December  
 199 and January, with  $d_{\text{mean}} = 0.99$  nm and  $d_{\text{median}} = 0.95$  nm, and the largest during June and July, with  
 200  $d_{\text{mean}} = 1.15$  nm and  $d_{\text{median}} = 1.11$  nm.

201 Positive ion size distributions (Fig. 2b) behave similarly to the negative ones, however the changes  
 202 are less pronounced. Above 1.4 nm and up to 2.0 nm, the concentrations are roughly twice as high,  
 203 or less, during summer compared to winter. For positive small ions, the smallest value of  $d_{\text{mean}} =$   
 204 1.16 and  $d_{\text{median}} = 1.13$  nm (December and January) and the largest value of  $d_{\text{mean}} = 1.29$  nm and  
 205  $d_{\text{median}} = 1.27$  nm (June). The difference between the average diameters of negative and positive  
 206 small ions was around 0.15 nm, in line with previous studies (e.g., Hörrak et al., 2000).

207 The observed seasonal behavior of the size distributions in Hyytiälä follow expectations: during  
 208 spring and summer, the concentrations of low-volatility vapors are much higher due to increased  
 209 solar radiation and organic emissions (Sulo et al., 2021). Therefore, small ions should be able to  
 210 grow to larger diameters due to the uptake of these vapors. We will look further into how the small  
 211 ion size distribution varies with respect to low volatile vapor concentrations in Sect. 3.2.



212  
 213 **Fig. 2:** Median monthly sub-2 nm negative (a, c) and positive (b, d) ion size distributions for  
 214 Hyytiälä (a, b) and Beijing (c, d). The different months are marked by the different colors.  
 215

216 The Fig. 2c, d show the monthly median negative (Fig. 2c) and positive (Fig. 2d) ion distributions  
 217 between 0.8 and 2.0 nm in Beijing, while Table 2 records the monthly  $d_{\text{mean}}$  and  $d_{\text{median}}$ . As expected  
 218 due to the high CoagS (Eq. 1,2, and Fig. A1), the concentrations are lower than in Hyytiälä. For  
 219 negative ions (Fig. 2c), the concentrations during summer are higher than in other seasons below  
 220 1.6 nm and lower than in other seasons close to 2.0 nm. During spring, the concentrations of  
 221 negative ions below 1.4 nm are lower than in other seasons. The smallest value of negative  $d_{\text{mean}}$  and  
 222  $d_{\text{median}}$  are in November,  $d_{\text{mean}} = 1.04$  nm and  $d_{\text{median}} = 1.01$  nm. The largest values are during June,  
 223  $d_{\text{mean}} = 1.16$  nm and  $d_{\text{median}} = 1.12$  nm.

224 For positive small ions in Beijing (Fig. 2d), the concentrations of both the smallest and the largest  
 225 ions in the 0.8 to 2.0 nm range are both considerably lower from January to March compared to  
 226 later months. Otherwise, it is difficult to identify any clear seasonal patterns. The largest positive  
 227 average diameter is during February,  $d_{\text{mean}} = 1.32$  nm and  $d_{\text{median}} = 1.31$  nm, while the smallest values  
 228 are in November,  $d_{\text{mean}} = 1.22$  nm and  $d_{\text{median}} = 1.21$  nm.

229 Compared to Hyytiälä, the seasonal trends in Beijing are much less clear, which implies that the  
 230 factors controlling the small ion size distribution are less seasonal in Beijing than in Hyytiälä, where

231 the small ion size distribution show strong seasonal variation. However, we note that because there  
 232 are less data from Beijing compared to Hyytiälä, variation between years can have larger impact on  
 233 the results than in Hyytiälä.

234

235 **Table 1:** Mean and median monthly diameters (nm) of ions between 0.8 and 2.0 nm in Hyytiälä.

236 \*The highest concentration corresponds to the lowest detected diameter.

Month	Negative ions		Positive ions	
	$d_{\text{mean}}$	$d_{\text{median}}$	$d_{\text{mean}}$	$d_{\text{median}}$
1	0.99	0.95	1.16	1.13
2	1.00	0.96	1.17	1.15
3	1.05	1.02	1.2	1.18
4	1.08	1.06	1.21	1.20
5	1.12	1.10	1.27	1.25
6	1.15	1.11	1.29	1.27
7	1.15	1.11	1.28	1.26
8	1.13	1.10	1.25	1.23
9	1.11	1.08	1.22	1.20
10	1.05	1.02	1.19	1.17
11	1.01	0.98	1.17	1.14
12	0.99	0.95	1.16	1.13

237

238

239 **Table 2:** Mean and median monthly diameters (nm) of ions between 0.8 and 2.0 nm in Beijing.

240 \*The highest concentration corresponds to the lowest detected diameter.

Month	Negative ions		Positive ions	
	$d_{\text{mean}}$	$d_{\text{median}}$	$d_{\text{mean}}$	$d_{\text{median}}$
1	1.10	1.07	1.30	1.30
2	1.09	1.06	1.32	1.31
3	1.09	1.06	1.28	1.27
4	1.10	1.05	1.28	1.26
5	1.12	1.07	1.28	1.27
6	1.16	1.12	1.27	1.26
7	1.12	1.08	1.25	1.24
8	1.10	1.08	1.25	1.24
9	1.10	1.07	1.25	1.24
10	1.07	1.03	1.24	1.23

11	1.04	1.01	1.22	1.21
12	1.05	1.01	1.23	1.21

241

242

## 243 **3.2 Potential impact of low volatility vapors to small ion size distribution** 244 **in Hyytiälä**

### 245 **3.2.1 Highly oxidized organic molecules (HOMs)**

246 Fig. 3 shows the median ion number size distributions between 0.8 and 2.0 nm in Hyytiälä with  
247 respect to varying neutral highly oxidized organic molecule (HOM) concentration. HOM  
248 monomers, HOM dimers and total HOM are considered separately. Results for daytime (10:00-  
249 16:00) and evening (18:00-00:00) are both presented (Fig. 3a and 3b, respectively). The HOM  
250 concentrations are divided into percentiles.

251 A clear increase in the number of negative ions (Fig. 3a, i-iii, Fig. 3b, i-iii) above approx. 1.05 nm,  
252 and for positive ions (Fig. 3a, iv-vi, Fig. 3b, iv-vi) slightly larger than that, is seen with an  
253 increasing HOM concentration percentile for all the HOM categories. The difference is the largest  
254 for HOM monomers (i, iv) and HOM total (iii, vi), which is mainly dominated by the HOM  
255 monomers. The difference is also stronger for negative ions than positive ions, and is stronger  
256 during the evening (Fig. 3b) compared to daytime (Fig. 3a).

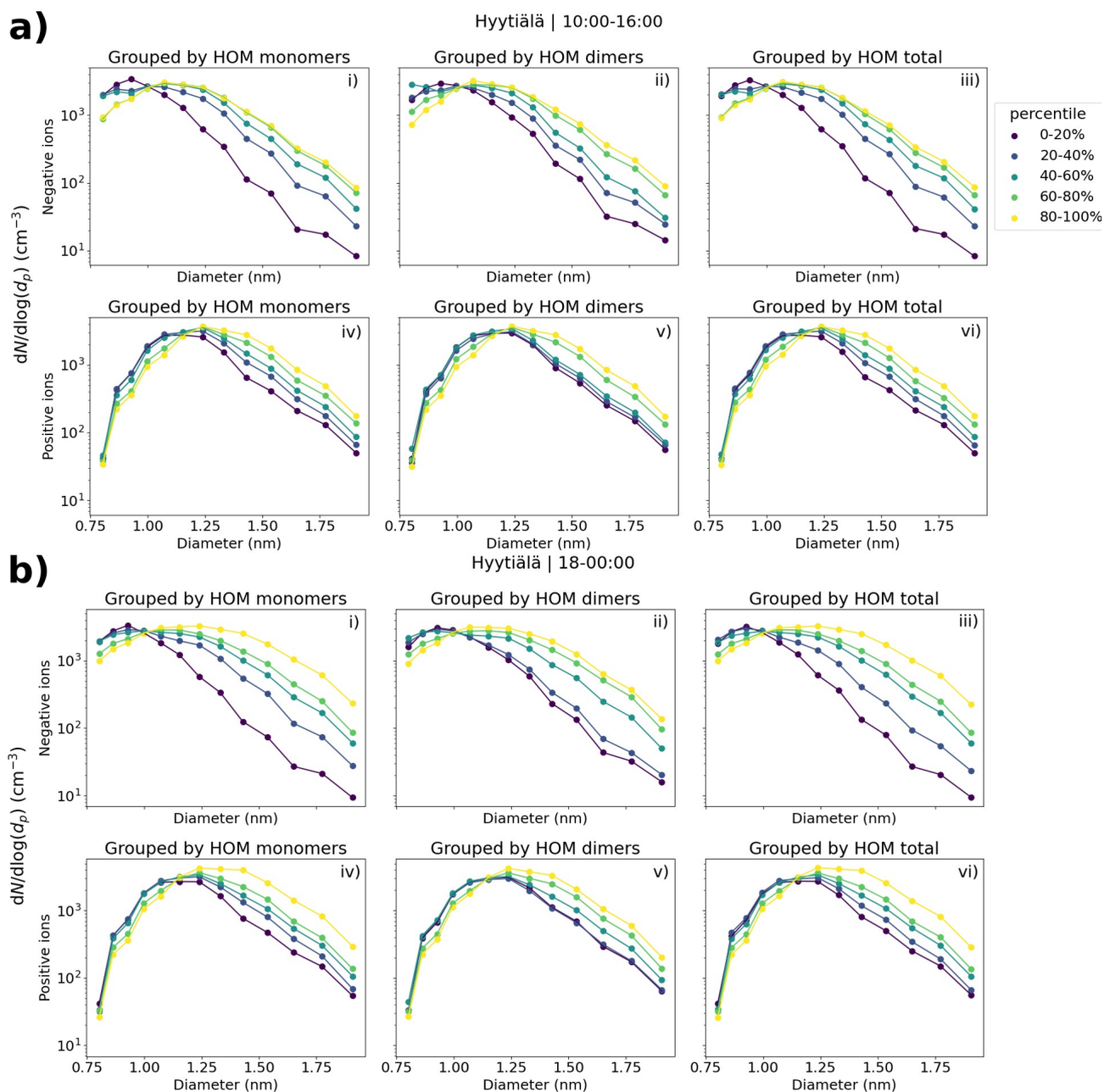
257 Comparing the negative ion size distributions between the HOM percentiles of 0-20% and 80-  
258 100%, we see that the difference in the concentrations increases with an increasing diameter, and  
259 that close to 2.0 nm this difference is approximately one order of magnitude during daytime (Fig.  
260 3a,i-iii) and a bit more during the evening (Fig. 3b, i-iii). Comparing the similar negative ion  
261 concentrations when HOM monomer concentration during the evening is in the 80-100% percentile  
262 compared to 0-20%, there's approx. a 0.5 nm shift in diameters (Fig. 3b, i), which is a big difference  
263 for the sub-2 nm ion population.

264 In line with the large differences in the small ion size distributions in Fig. 3 with respect to HOM  
265 concentration, a strong correlation between the small ion  $d_{\text{mean}}$  and the HOM concentrations was  
266 seen (Fig. 4). The Spearman correlation coefficients ( $r_s$ ) between  $d_{\text{mean}}$  and HOMs were 0.6 or  
267 above, for both daytime and evening. For daytime, the best correlation was between  $d_{\text{mean}}$  of positive  
268 ions and HOM monomer concentration,  $r_s = 0.74$  (Fig. 4e). During nighttime, the strongest  
269 correlation was between  $d_{\text{mean}}$  of negative ions and HOM monomer concentration,  $r_s = 0.73$  (Fig.  
270 4c).

271 The clear correlation between HOMs and the small ion size distribution in Hyytiälä suggests a  
272 strong impact of organic compounds to the small ion population. This interpretation, as opposed to  
273 the correlation being due to a correlation with another variable such sulfuric acid concentration, is  
274 supported by the observation of the correlation being stronger during evening when the  
275 concentrations of other precursors such as sulfuric acid are lower and organic ion cluster formation  
276 is known to take place in Hyytiälä (Mazon et al., 2016; Rose et al., 2018). We note that part of the  
277 increase in the diameters of the small ions when HOMs are abundant could be due to the large size  
278 of organic molecules when compared to sulfuric acid molecules. However, the clear increase in

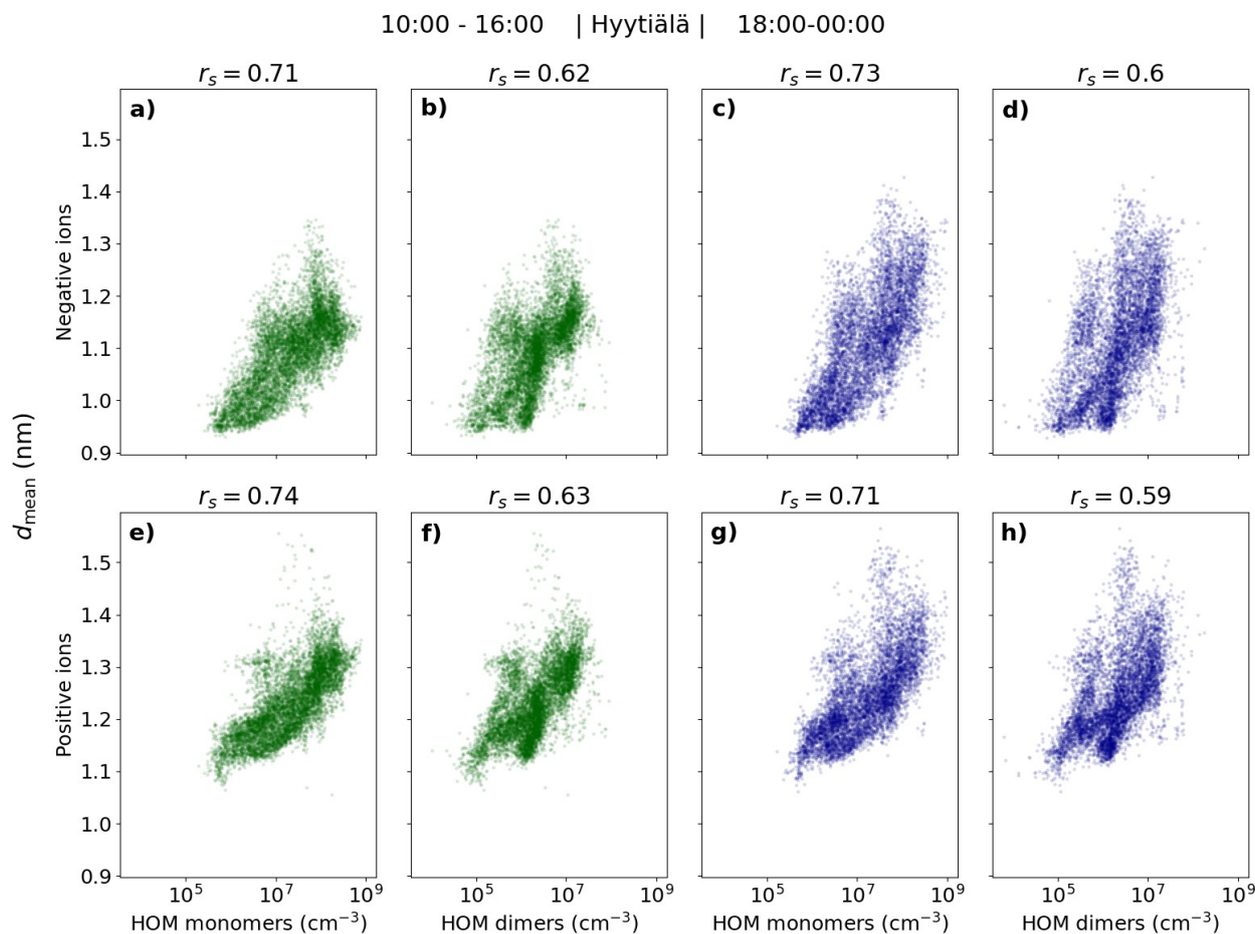
279 concentrations even close to 2.0 nm suggests that the impact is mainly due to the growth of small  
 280 ions by the uptake of organic vapors.

281 While the concentrations of larger small ions of both polarities increase with increasing HOM  
 282 percentile, the differences are larger for the negative ions (Fig. 3). This could be due to the uptake  
 283 of organics being more effective for the negatively charged ions. However, the equally strong  
 284 correlation between  $d_{\text{mean}}$  of positive small ions and HOM concentrations does not support this  
 285 interpretation. A possible explanation is the size difference between the negative and positive small  
 286 ions: due to the larger diameter of positive small ions, it might be that the impact of the growth to  
 287 the diameters of the positive ions is not as large. For a larger cluster (ion), more molecules are  
 288 needed to increase the diameter equally than for a smaller one.



289  
 290 **Fig. 3:** The median negative (i-iii) and positive (iv-vi) small ion (sub-2 nm) size distributions in  
 291 Hyytiälä, Finland grouped by the percentiles of neutral HOM monomer (i, iv), HOM dimer (ii, v)  
 292 and total HOM (iii, vi) concentrations. Both the evening (18:00-00:00) size distributions (b) and

293 daytime (10:00-16:00) size distributions (a) are shown. Daytime percentiles for HOM monomers,  
 294 dimers and total are 20%:  $4.30 \cdot 10^6$ ,  $5.93 \cdot 10^5$ , and  $5.10 \cdot 10^6$   $\text{cm}^{-3}$ ; 40%:  $1.34 \cdot 10^7$ ,  $1.45 \cdot 10^6$ , and  
 295  $1.50 \cdot 10^7$   $\text{cm}^{-3}$ ; 60%:  $4.00 \cdot 10^7$ ,  $2.44 \cdot 10^6$ , and  $4.18 \cdot 10^7$   $\text{cm}^{-3}$ ; 80%:  $9.70 \cdot 10^7$ ,  $7.62 \cdot 10^6$ , and  $1.05 \cdot 10^8$   $\text{cm}^{-3}$   
 296  $^3$ , respectively. Evening percentiles for HOM monomers, dimers and total are 20%:  $3.00 \cdot 10^6$ ,  
 297  $4.94 \cdot 10^5$ , and  $3.75 \cdot 10^6$   $\text{cm}^{-3}$ ; 40%:  $8.40 \cdot 10^6$ ,  $1.50 \cdot 10^6$ , and  $1.03 \cdot 10^7$   $\text{cm}^{-3}$ ; 60%:  $3.17 \cdot 10^7$ ,  $2.95 \cdot 10^6$ ,  
 298 and  $3.65 \cdot 10^7$   $\text{cm}^{-3}$ ; 80%:  $7.52 \cdot 10^7$ ,  $7.70 \cdot 10^6$ , and  $8.46 \cdot 10^7$   $\text{cm}^{-3}$ , respectively.



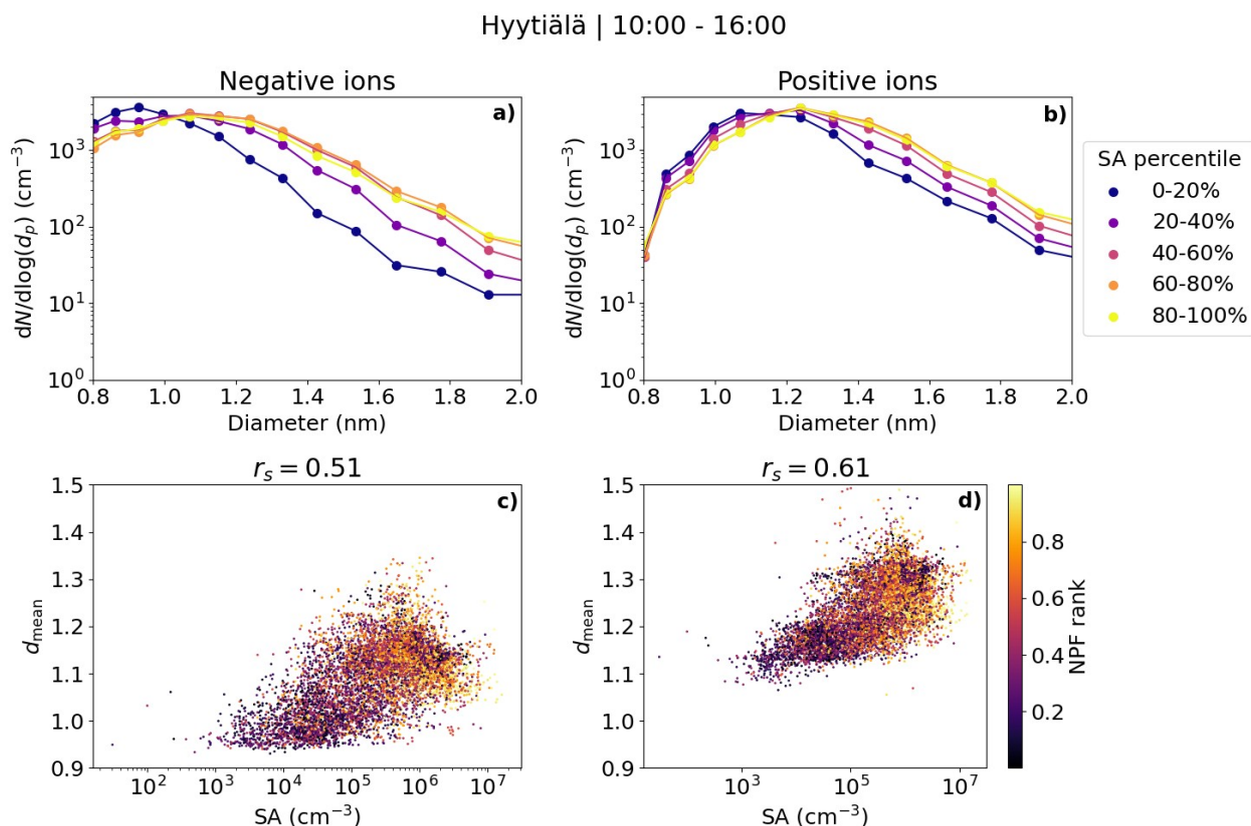
299  
 300 **Fig. 4:** Mean diameter ( $d_{\text{mean}}$ ) of negative (a-d) and positive (e-h) small (sub-2 nm) ions as a  
 301 function of HOM (monomer, dimer and total) concentration in Hyytiälä. The individual points are  
 302 hourly medians, and the daytime (10:00-16:00, marked in green) and evening (18:00-00:00, marked  
 303 in dark blue) are shown separately. Spearman correlation coefficients ( $r_s$ ) are shown.

304

### 305 3.2.2 Sulfuric acid

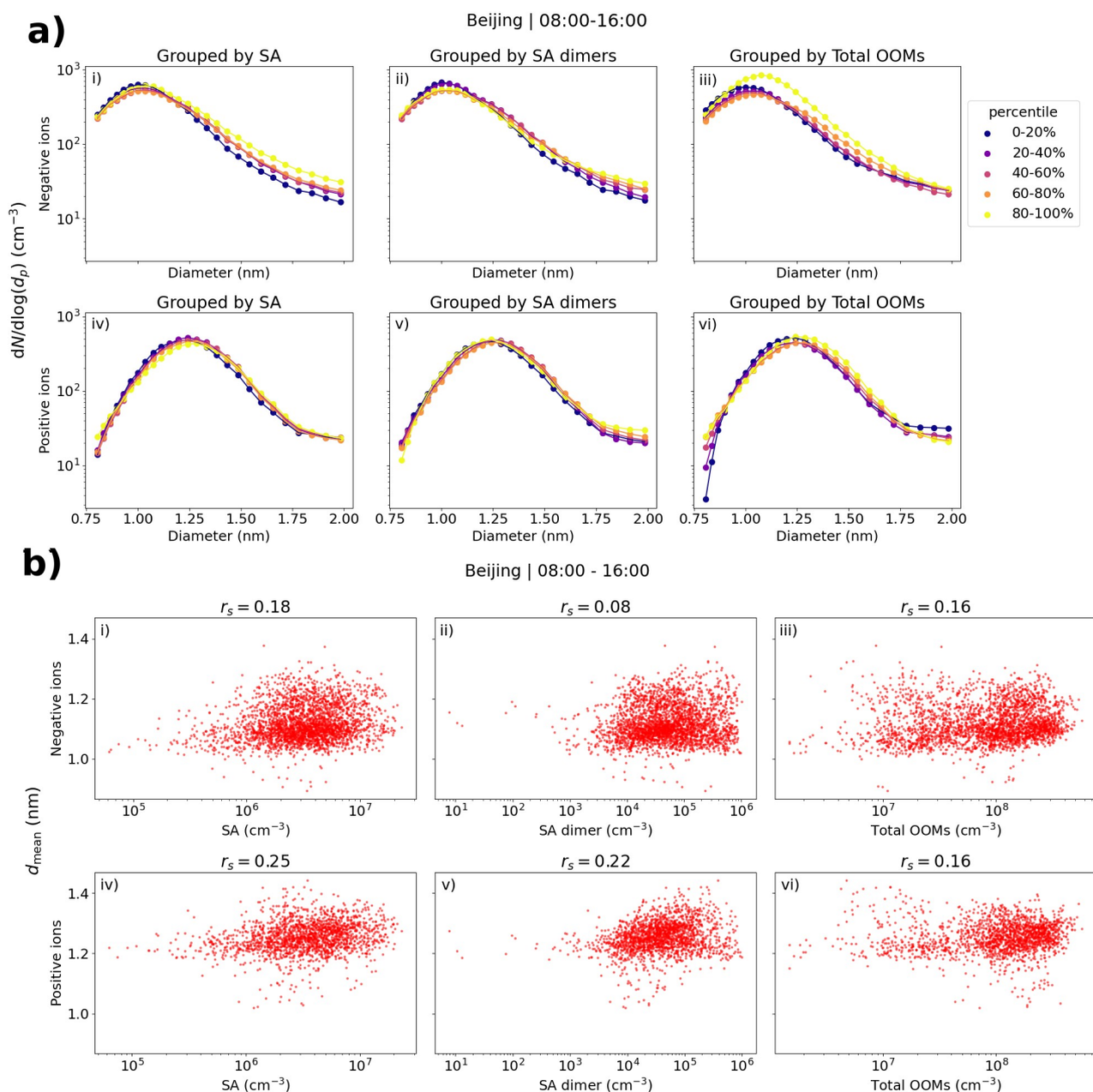
306 Fig. 5 shows the median daytime negative and positive small ion (0.8-2.0 nm) size distributions  
 307 grouped by percentiles of neutral sulfuric acid (SA) concentration (Fig. 5a and b). In addition, the  
 308 daytime hourly median  $d_{\text{mean}}$  values are shown as a function of the SA concentration (Fig. 5c and d).  
 309 We see a clear increase in the concentrations of negative (positive) small ions larger than  
 310 approximately 1.05 (1.1) nm when comparing SA concentrations in the lower percentiles to the  
 311 higher percentiles, until the behavior seems to stall so that the 60-80% and 80-100% percentiles  
 312 show similar size distributions. From previous studies, we know that while sulfuric acid is often  
 313 needed for the initial cluster formation, organic compounds tend to drive the cluster growth  
 314 (Kulmala et al., 2013). This could explain the small difference in the size distributions between  
 315 the 60-80% and 80-100% percentiles of the SA concentration.

316 A good positive correlation was seen between  $d_{\text{mean}}$  and SA concentration for both polarities,  $r_s =$   
 317 0.51 (0.61) for negative (positive) ions (Fig. 5c and d). The correlation is slightly weaker than what  
 318 was observed between  $d_{\text{mean}}$  and HOM concentrations. The majority of  $d_{\text{mean}}$  values above 1.1 nm  
 319 correspond to days with a high NPF ranking, while most values of  $d_{\text{mean}}$  below 1.1 nm correspond to  
 320 days with low NPF rank values below 0.5. Notably, for  $d_{\text{mean}}$  of negative small ions above approx.  
 321 1.1 nm, the values of  $d_{\text{mean}}$  do not seem to increase with an increasing SA concentration as clearly as  
 322 they do with an increasing HOM concentration (Fig. 4). As discussed above, organic compounds  
 323 might be needed to drive the growth of small ions further, and thus dependency of  $d_{\text{mean}}$  on SA is not  
 324 seen as clearly when  $d_{\text{mean}}$  is above 1.1 nm.



325  
 326 **Fig. 5:** The median number size distributions of small ions between 0.8 and 2.0 nm grouped by  
 327 percentiles of neutral sulfuric acid concentration (a) negative and b) positive ions) and scatter plots  
 328 and Spearman correlation coefficients ( $r_s$ ) of hourly mean diameter of small ions ( $d_{\text{mean}}$ ) and sulfuric  
 329 acid concentration (c) negative and d) positive ions) in Hyytiälä. In the scatter plot, the color  
 330 indicates the respective NPF rank of the day. Only daytime (10:00-16:00) values are included. The  
 331 percentile values for sulfuric acid are 20%:  $4.22 \cdot 10^4 \text{ cm}^{-3}$ , 40%:  $1.79 \cdot 10^5 \text{ cm}^{-3}$ , 60%:  $5.06 \cdot 10^5 \text{ cm}^{-3}$   
 332 and 80%:  $1.08 \cdot 10^6 \text{ cm}^{-3}$ .

333 **3.3 Relationship of small ion size distribution with low volatility vapors**  
 334 **in Beijing**



335 **Fig. 6:** (a) Small ion median daytime (08:00-16:00) number size distributions in Beijing, grouped  
 336 by percentiles of sulfuric acid (SA), SA dimer or total oxidized organic molecule (OOM)  
 337 concentrations. (b) Hourly daytime mean diameter ( $d_{\text{mean}}$ ) of small ions versus SA, SA dimer and  
 338 total OOM concentrations. Figures i-iii are for negative polarity and figures iv-vi for positive and  
 339 the figures with SA, SA dimer, and OOMs are in the respective order. Spearman correlation  
 340 coefficients ( $r_s$ ) are included. The percentile limits of SA, SA dimer and total OOM are 20%:  
 341 1.64·10<sup>6</sup>, 1.26·10<sup>4</sup>, and 2.19·10<sup>7</sup> cm<sup>-3</sup>; 40%: 2.53·10<sup>6</sup>, 3.11·10<sup>4</sup>, and 5.89·10<sup>7</sup> cm<sup>-3</sup>; 60%: 3.66·10<sup>6</sup>,  
 342 6.01·10<sup>4</sup> and 1.23·10<sup>8</sup> cm<sup>-3</sup>; and 80%: 5.17·10<sup>6</sup>, 1.32·10<sup>5</sup>, and 2.15·10<sup>8</sup> cm<sup>-3</sup>.

344 Fig. 6a shows the number size distributions of small ions grouped by percentiles of neutral sulfuric  
 345 acid, sulfuric acid dimer and total oxidized organic molecule (OOM) concentration in Beijing. We  
 346 can see that, especially compared to results already for Hyytiälä, the differences in the size

348 distributions with respect to different values of sulfuric acid or OOMs are small for both polarities.  
349 The concentration of negative ions below approx. 1.2 nm slightly decreases with increasing sulfuric  
350 acid (Fig. 6a, i) and sulfuric acid dimer (Fig. 6a, ii) concentration, while the concentrations above  
351 approx. 1.2 nm increase, indicating a weak but positive relationship between the sulfuric acid and  
352 the negative small ion growth. Close to 2.0 nm, where the increase is the highest, the concentration  
353 of negative ions is higher by around a factor of two when sulfuric acid concentration is in the 80-  
354 100% percentile compared to when it is in the 0-20% percentile (Fig. 6a, i). For both polarities, the  
355 concentrations below approx. 1.75 nm appear higher when the total OOM concentration is in the  
356 80-100% percentile compared to other times (Fig. 6a, iii and vi). However, the concentrations close  
357 to 2.0 nm are not simultaneously higher, indicating that despite the increased concentration of small  
358 ions, a larger number of them is not growing into intermediate ions.

359 Fig. 6b shows the scatter plots of  $d_{\text{mean}}$  and sulfuric acid, sulfuric acid dimer and total OOM  
360 concentrations. Weak positive correlation is seen, and the Spearman correlation coefficients ( $r_s$ ) are  
361 between 0.08 and 0.25. The differences in the values of  $d_{\text{mean}}$  are small. The relationship between the  
362 small ion size distribution or  $d_{\text{mean}}$  and low volatility vapor concentrations in Beijing appears weaker  
363 compared to Hyytiälä.

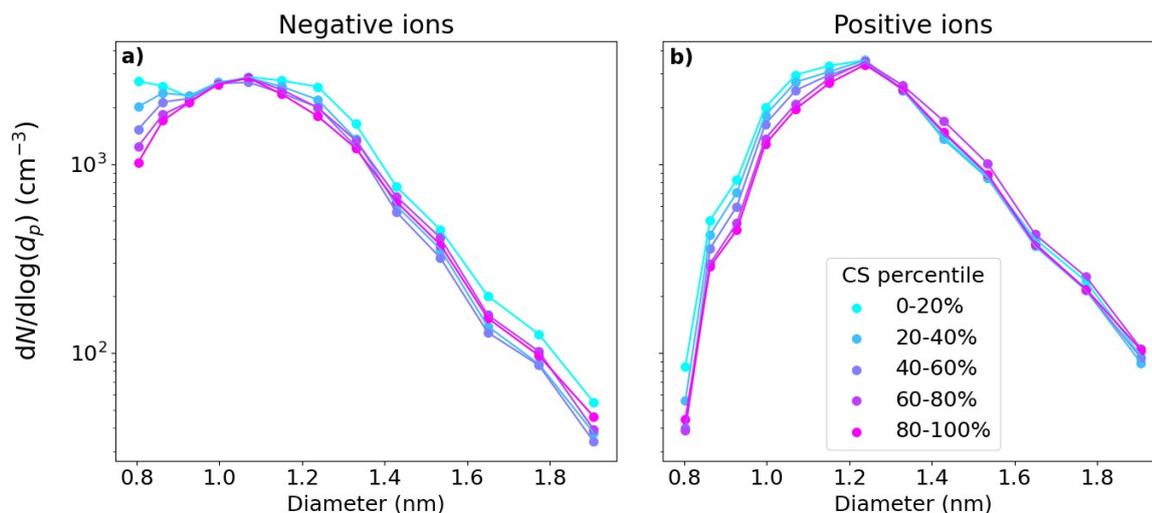
### 364 **3.4 Impact of coagulation scavenging on the small ion size distribution**

365 Fig. 7 shows the median small ion number size distributions in Hyytiälä and Beijing corresponding  
366 to the respective percentiles of condensation sink (CS). We see that the changes in the small ion size  
367 distribution with respect to changing CS are relatively small. In Hyytiälä (Fig. 7a, b), the  
368 concentration of small ions, especially that of the smallest in diameter, decreases slightly with  
369 increasing CS. The sink is relatively low in Hyytiälä, and therefore this result is not unexpected.

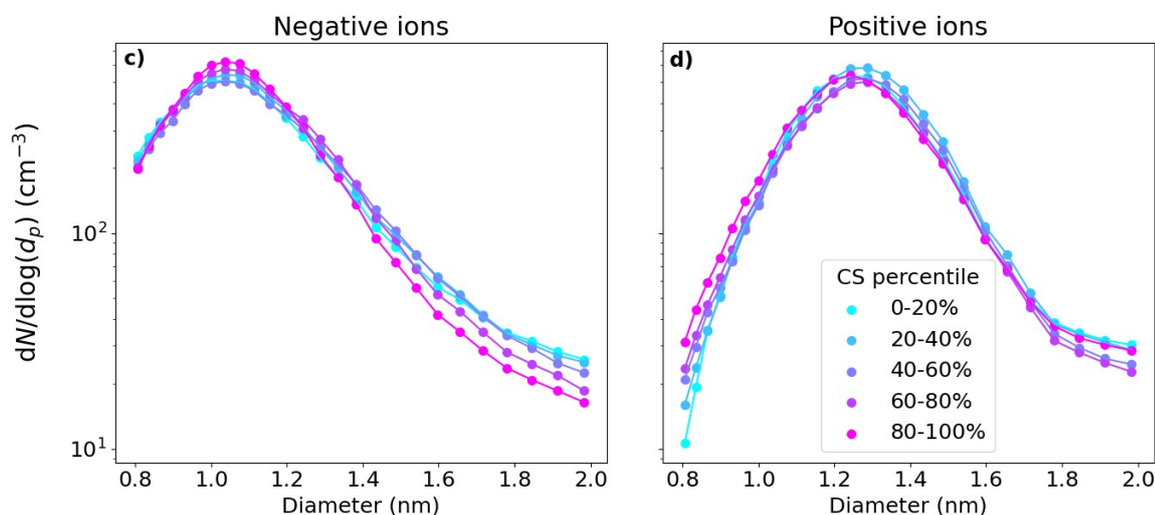
370 Based on Fig. A1, which shows the approximate values of the different terms in Eq. 2, we would  
371 expect CoagS to have a much stronger impact on the small ion dynamics in Beijing than in  
372 Hyytiälä. The impact should be most clear for the smallest sizes. Surprisingly, this was not  
373 observed. Fig. 7c shows that in Beijing, the negative small ion size distribution below approx. 1.3  
374 nm stays unchanged and the concentrations above decrease with an increasing CS. The larger  
375 positive small ion concentrations (Fig. 7d) also seem to slightly decrease with increasing sink, while  
376 the concentrations of the small ions close to 0.8 nm actually increase. Fig. A2a also shows that the  
377 total sub-2 nm concentration barely changes with changing CS. These results suggest that there  
378 could be a source of unknown nature for small ion cluster formation that is higher when CS is high,  
379 which would compensate for the increased coagulation scavenging of the ions.

380 Fig. A2d-f show the concentrations of the different low volatility vapors as a function of CS. We see  
381 that the concentration of sulfuric acid (Fig. A2d) and sulfuric acid dimers (Fig. A2e) decreases with  
382 increasing sink, as expected. However, the concentration of OOMs (Fig. A2f) increases with  
383 increasing sink. While this is purely speculation, if organic compounds are forming small ion  
384 clusters when the sink is higher, the weak apparent impact of CS on the small ion size distribution  
385 could be explained. Alternative potential explanation could be if there is a positive correlation  
386 between CS and the concentration of bases, which stabilize the small clusters. Regardless, the  
387 impact of CS on the statistics of the small ion size distribution in Beijing appears very small.

Hyytiälä | 10:00-16:00



Beijing | 08:00-16:00



388

389 **Fig. 7:** the median small ion size distributions grouped by the respective percentile of the  
 390 condensation sink (CS) values for Hyytiälä (a,b) and Beijing (c, d). The percentile value limits for  
 391 CS in Hyytiälä are  $1.3 \cdot 10^{-3} \text{ s}^{-1}$ ,  $2.3 \cdot 10^{-3} \text{ s}^{-1}$ ,  $3.5 \cdot 10^{-3} \text{ s}^{-1}$ , and  $5.5 \cdot 10^{-3} \text{ s}^{-1}$ . In Beijing, the percentile  
 392 value limits for CS are  $9.3 \cdot 10^{-3} \text{ s}^{-1}$ ,  $1.9 \cdot 10^{-2} \text{ s}^{-1}$ ,  $3.1 \cdot 10^{-2} \text{ s}^{-1}$ , and  $4.5 \cdot 10^{-2} \text{ s}^{-1}$ .

### 393 3.5 Correlation of small ion size distribution with sulfuric acid clusters 394 and NPF in Hyytiälä

395 Fig. 8a shows the median number size distribution of negative small ions grouped by percentiles of  
 396 the signals of SA ion clusters  $\text{HSO}_4^-$  (monomer),  $\text{H}_2\text{SO}_4 \cdot \text{HSO}_4^-$  (dimer) and  $(\text{H}_2\text{SO}_4)_2 \cdot \text{HSO}_4^-$  (trimer)  
 397 and their ratios in Hyytiälä. The median distributions are determined from daytime (10:00-16:00)  
 398 values. We observe a clear increase in the number of small ions with diameters above approx. 1.2  
 399 nm with an increased signal of SA ion monomers (Fig. 8a, i) and dimers (Fig. 8a, ii). The increase is  
 400 especially clear for dimers and the dimer to monomer ratio (Fig. 8a, iv), and the concentration of  
 401 small ions close 2.0 nm, where the differences are highest, is an order of magnitude higher when  
 402 dimer signal is in the 80-100% percentile compared to when the signal is in the 0-20% percentile.

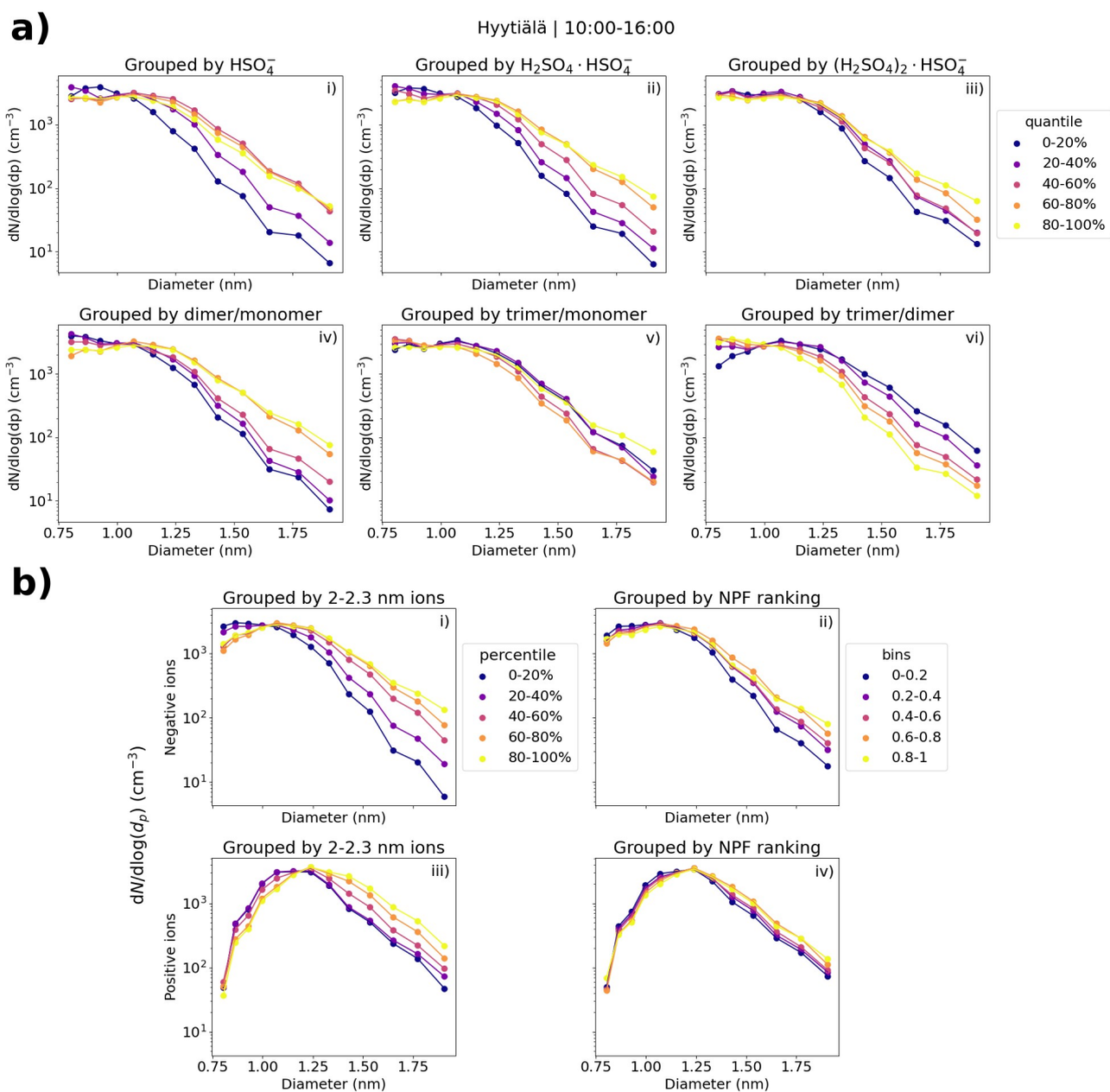
403 These results show that the dimer signal is a strong indicator for both the cluster formation and the  
404 growth of clusters to larger sizes in Hyytiälä.

405 We used the 2.0-2.3 nm ion concentrations and NPF ranking (Aliaga et al., 2023) as proxies for  
406 conditions that were favorable for cluster formation and growth. Fig. 8b shows the median daytime  
407 (10:00-16:00) size distributions for both polarities with respect to the percentiles of 2.0-2.3 nm ion  
408 concentrations and bins of NPF ranking in Hyytiälä. When the 2.0-2.3 nm ion concentration is  
409 higher, a clear increase in concentrations is seen above approx. 1.2 nm (Fig. 8b,i and iii). The  
410 difference in negative small ion concentrations close to 2.0 nm between 80-100% and 0-20% is over  
411 one order of magnitude (Fig. 8b, i). The small ion size distribution for both polarities shows this  
412 growth of small ions up to 2.0 nm, when local-scale intermediate ion formation is taking place.

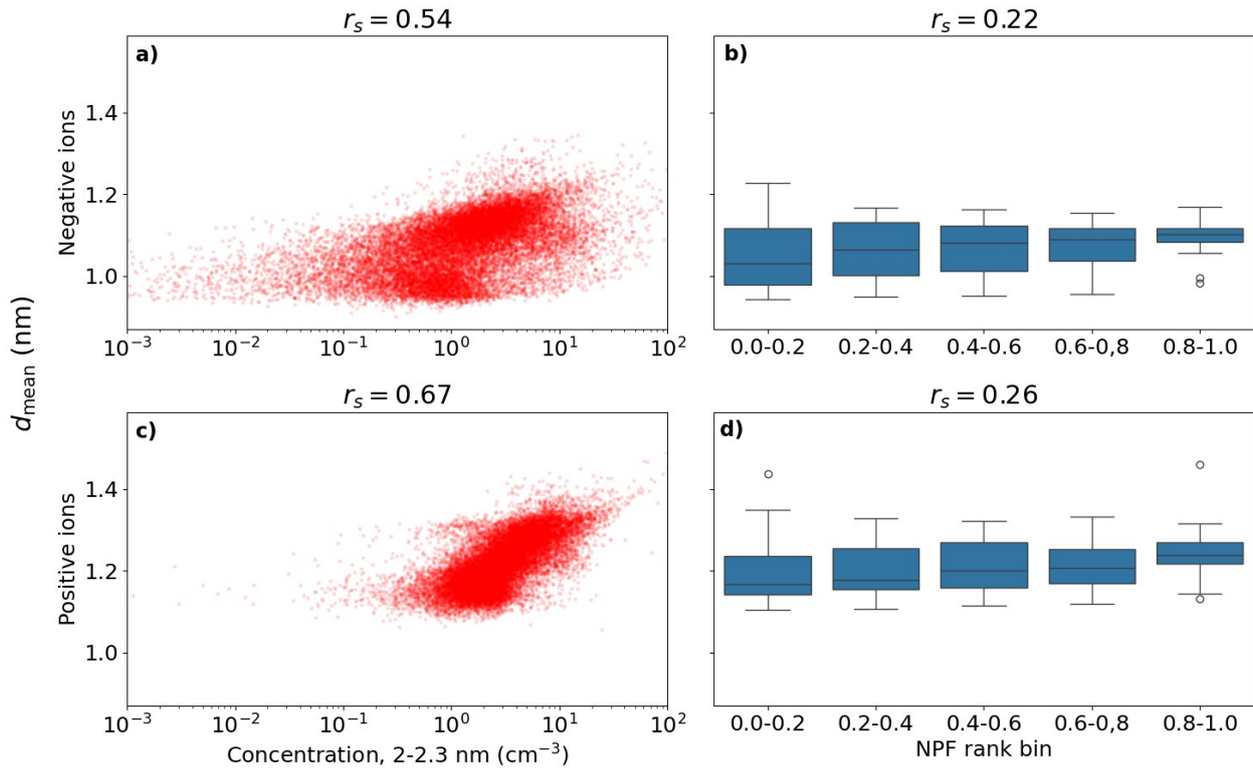
413 Similar observations can be made from the small ion size distributions with respect to the different  
414 NPF ranking values. However, the differences are smaller than with respect to 2.0-2.3 nm ions, and  
415 especially for positive small ions such differences are very small (Fig. 8b, iv). There is likely a  
416 combination of factors at play here. First of all, NPF ranking was determined for total particles  
417 between 2.5 and 5 nm and there might be differences stemming both from the ranking being less  
418 sensitive for local NPF, and for 2.0-2.3 nm ion concentrations being more sensitive for ion-induced  
419 clustering or NPF. In addition, differences between what is observed in the total particles versus  
420 ions can be caused by variation in the chemical compounds, which take up the available charges  
421 (Bianchi et al., 2017).

422 We note that the differences in the number size distribution of positive small ions are once again  
423 smaller than for negative small ions. Similarly to Sect. 3.2.1, we hypothesize that this is due to the  
424 size difference between the polarities.

425 Fig. 9 shows the scatter plot of hourly daytime negative (a) and positive (c) small ion  $d_{\text{mean}}$  and the  
426 concentration of 2.0-2.3 nm ions. As expected, a strong positive trend is seen between  $d_{\text{mean}}$  and 2.0-  
427 2.3 nm ion concentrations. The correlation coefficient is  $r_s = 0.54$  (0.67) for negative (positive) ions.  
428 Fig. 9 also shows the box plots of  $d_{\text{mean}}$  with NPF ranking, with negative small ions in Fig. 9b and  
429 positive in 9d. The median of  $d_{\text{mean}}$  increases with increasing NPF ranking, as expected. However,  
430 the variance for lower rankings is much higher, resulting in overall quite a low correlation between  
431  $d_{\text{mean}}$  and NPF ranking,  $r_s =$  for negative (positive small) ions.



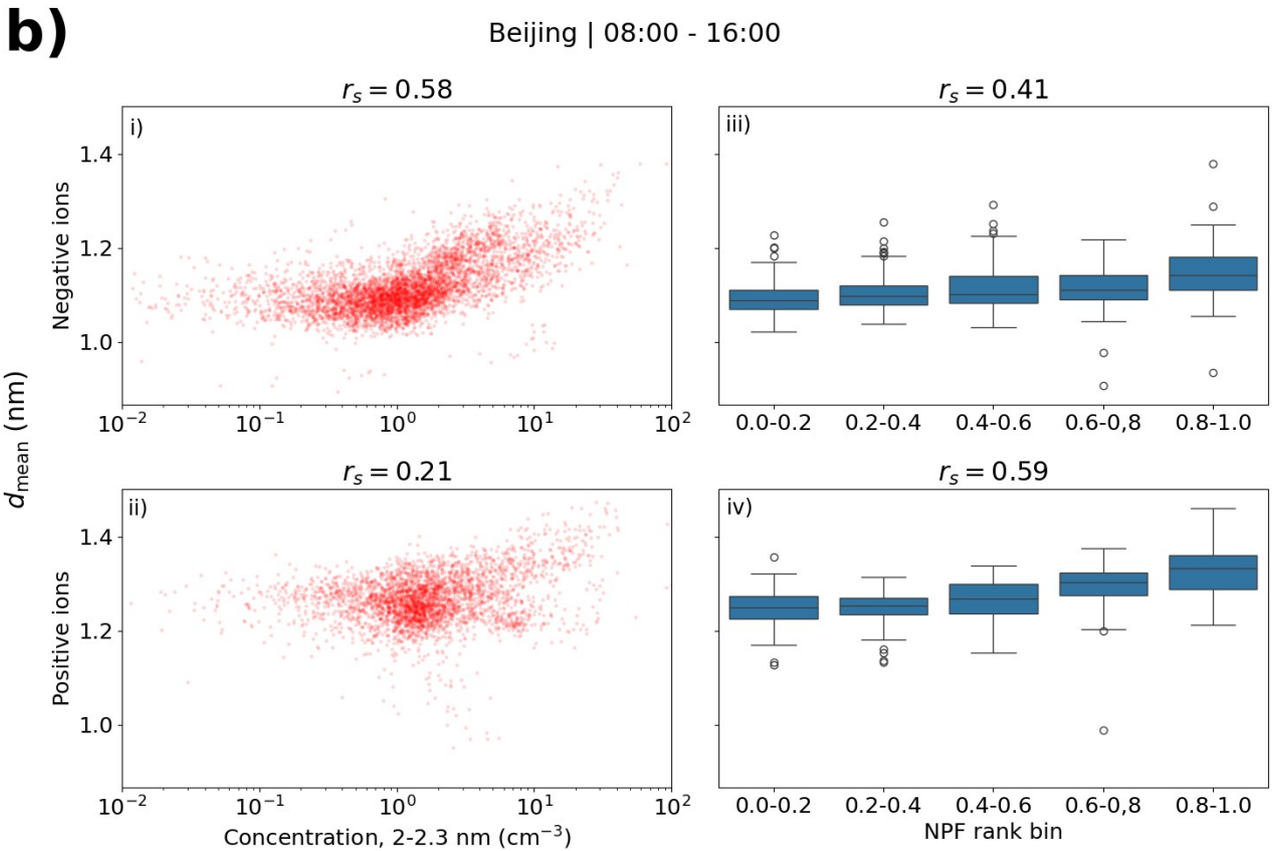
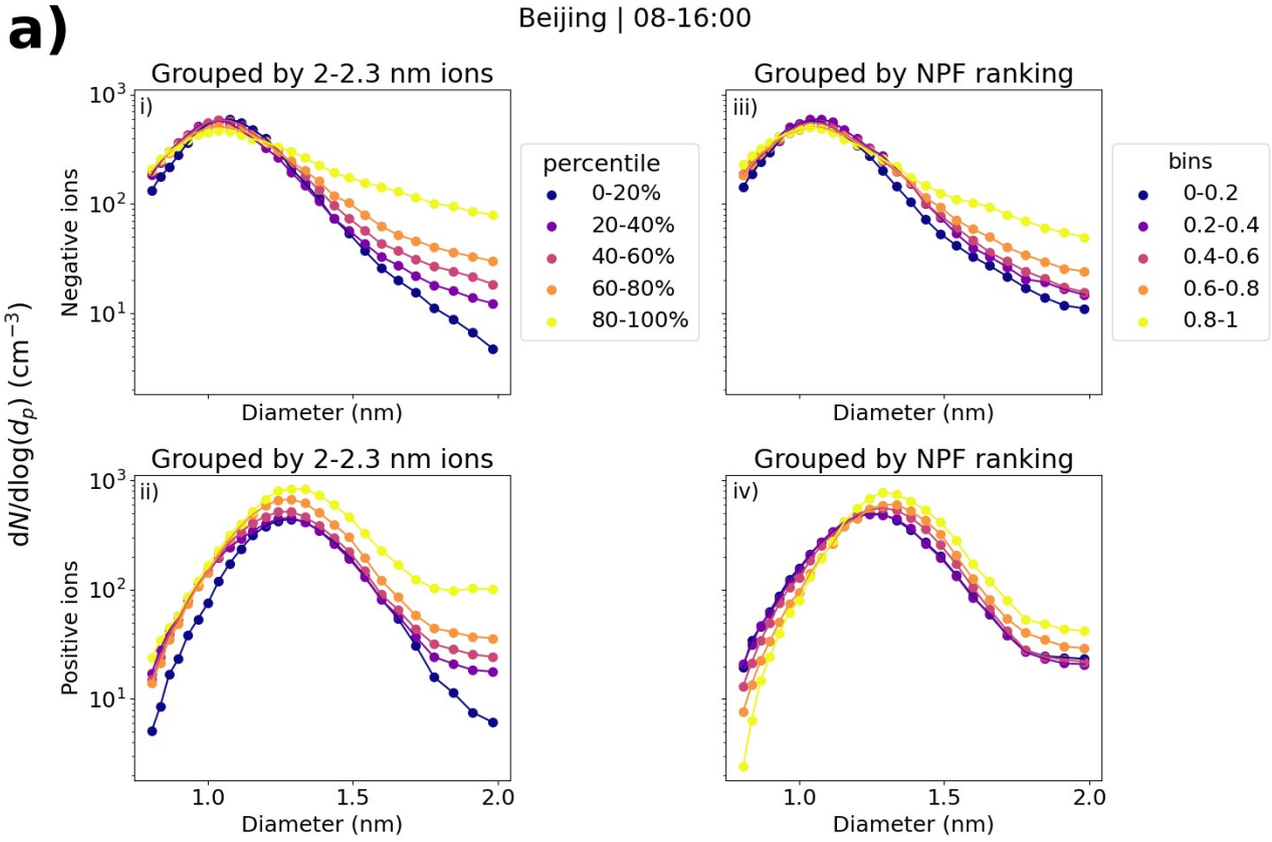
432  
 433 **Fig. 8:** (a) Hyytiälä daytime median negative small ion number size distributions grouped by  
 434 percentile of the signals of  $\text{HSO}_4^-$ ,  $\text{H}_2\text{SO}_4\cdot\text{HSO}_4^-$  or  $(\text{H}_2\text{SO}_4)_2\cdot\text{HSO}_4^-$  ions (figures i, ii, iii,  
 435 respectively) and their ratios (figures iv-vi). (b) Daytime median small ion size distributions for  
 436 both polarities grouped by the percentile of 2.0-2.3 nm ion concentrations (figures i, iii) of the  
 437 respective polarity or by NPF ranking (figures ii, iv). The percentile limits for negative (positive)  
 438 2.0-2.3 nm ion concentrations are 20%: 0.48 (1.75)  $\text{cm}^{-3}$ , 40%: 1.01 (2.34)  $\text{cm}^{-3}$ , 60%: 1.94 (3.49)  
 439  $\text{cm}^{-3}$ , and 80%: 3.39 (5.35)  $\text{cm}^{-3}$ .



440  
 441 **Fig. 9:** Hourly daytime negative (a,b) and positive (c, d) small ion diameter versus concentration of  
 442 2.0-2.3 nm ions of respective polarity (a, c) or NPF ranking (b, c) in Hyytiälä. Correlation  
 443 coefficients ( $r_s$ ) are also shown. The middle line of the box plots for  $d_{\text{mean}}$  and NPF rank are the  
 444 median values, while the boxes show the 25% and 75% percentiles and the lines the 10% and 90%  
 445 percentiles.

446 **3.6 Impact of NPF on small ion distribution in Beijing**

447



448

449 **Fig. 10:** (a) Median negative and positive small ion number size distributions in Beijing grouped by  
450 percentiles of 2.0-2.3 nm ion concentrations (b). Scatter plots of mean diameter and 2.0-2.3 nm ion  
451 concentrations (of respective polarity) or NPF ranking. Figures. i-ii are for negative polarity and iii-  
452 iv for positive. Values are for daytime (08:00-16:00). The percentile limits of 2.0-2.3 nm  
453 concentration for negative (positive) ions are 20%: 0.51 (0.85)  $\text{cm}^{-3}$ , 40%: 0.92 (1.27)  $\text{cm}^{-3}$ , 60%:  
454 1.33 (1.72)  $\text{cm}^{-3}$ , and 80%: 2.47 (3.22)  $\text{cm}^{-3}$ .

455

456 Fig. 10a shows the small ion size distributions with respect to the concentration of 2.0-2.3 nm ions  
457 or NPF ranking in Beijing. Negative ions are in Figures i-ii and positive in Figures iii-iv. For both  
458 polarities, clear differences are seen in the distributions depending on the percentile of the 2.0-2.3  
459 nm ion concentration. When the 2.0-2.3 nm ion concentrations are higher, the concentration of  
460 negative (positive) small ions above approx. 1.0 (1.3) nm is increased. The differences are largest  
461 close to 2.0 nm. Comparing the 0-20% and 80-100% percentiles, the difference in concentrations is  
462 around one order of magnitude when the diameter is approaching 2.0 nm, comparable to what was  
463 observed in Hyytiälä. Similar observations are seen with respect to NPF ranking, although to a  
464 lesser extent. For negative small ions (Fig. 10a, ii), the concentration at around 2.0 nm is four to  
465 five times higher when the NPF ranking is above 0.80 compared to when it is below 0.20. For  
466 positive ions, the concentration is less than two times higher (Fig. 10a, iv).

467

468 When looking at the small ion distributions in Beijing for different 2.0-2.3 nm ion concentrations or  
469 NPF ranking, unlike for low-volatility vapor concentrations, we are able to see the growth of small  
470 ions to intermediate ions in the size distribution. These results show that the growth of small ions to  
471 larger diameters in Beijing is not limited by the availability of sulfuric acid or oxidized organic  
472 vapors, unlike in Hyytiälä. In addition, based on our analysis it does not appear to be strongly  
473 limited by CS either. This is supported by the relatively weak correlation between CS and the NPF  
474 ranking or 2.0-2.3 nm ions (Fig. A2b,c). Therefore, we speculate that the small ion growth could be  
475 limited more by the availability of bases. However, due to the lack of long-term base concentration  
476 data, this question remains unanswered.

477

478 Fig. 10b shows the scatter plots of  $d_{\text{mean}}$  and 2.0-2.3 nm ion concentration and the box plots of  $d_{\text{mean}}$   
479 and NPF ranking. The correlation coefficients for negative ions are as expected,  $r_s = 0.58$  and  $r_s =$   
480  $0.41$  between  $d_{\text{mean}}$  and 2.0-2.3 nm ion concentration or NPF ranking, respectively. For positive ions,  
481 the correlation coefficient between  $d_{\text{mean}}$  and NPF ranking is  $r_s = 0.59$ , while it is only  $0.21$  between  
482  $d_{\text{mean}}$  and 2.0-2.3 nm ion concentration. From Fig. 10a we see that the concentrations of positive  
483 small ions below 1.0 nm also increase to some extent with increasing 2.0-2.3 nm ion concentration,  
484 which likely impacts the values of  $d_{\text{mean}}$ , resulting in a relatively poor overall correlation.

485

486 Notably, the differences in size distributions with respect to NPF ranking are clearer and the  
487 correlation between  $d_{\text{mean}}$  and the ranking is stronger in Beijing than in Hyytiälä for both polarities.  
488 One of the explaining factors could be the fact that intense NPF in Beijing is more common than in  
489 Hyytiälä (e.g., Dada et al., 2017; Deng et al., 2020), impacting the statistics of the ranking, and.  
490 Another possibility is that local clustering events, where ions or particles grow close to 2.0 nm but  
491 not much further, could be more common in Hyytiälä.

492

### 493 **3.7 Case studies**

494 Next, some case studies into the development of negative small ion size distributions, and other  
495 investigated variables, are presented for Hyytiälä and Beijing. These cases show that we are able to  
496 observe the cluster growth, driven by daytime NPF or evening clustering, from the ion number size  
497 distributions of individual days and not only from the statistics of the size distributions. Based on

498 the analysis presented in this study, the behavior of negative and positive small ion populations is  
499 mostly similar, and therefore, for simplicity, we have limited the analysis here to negative polarity.

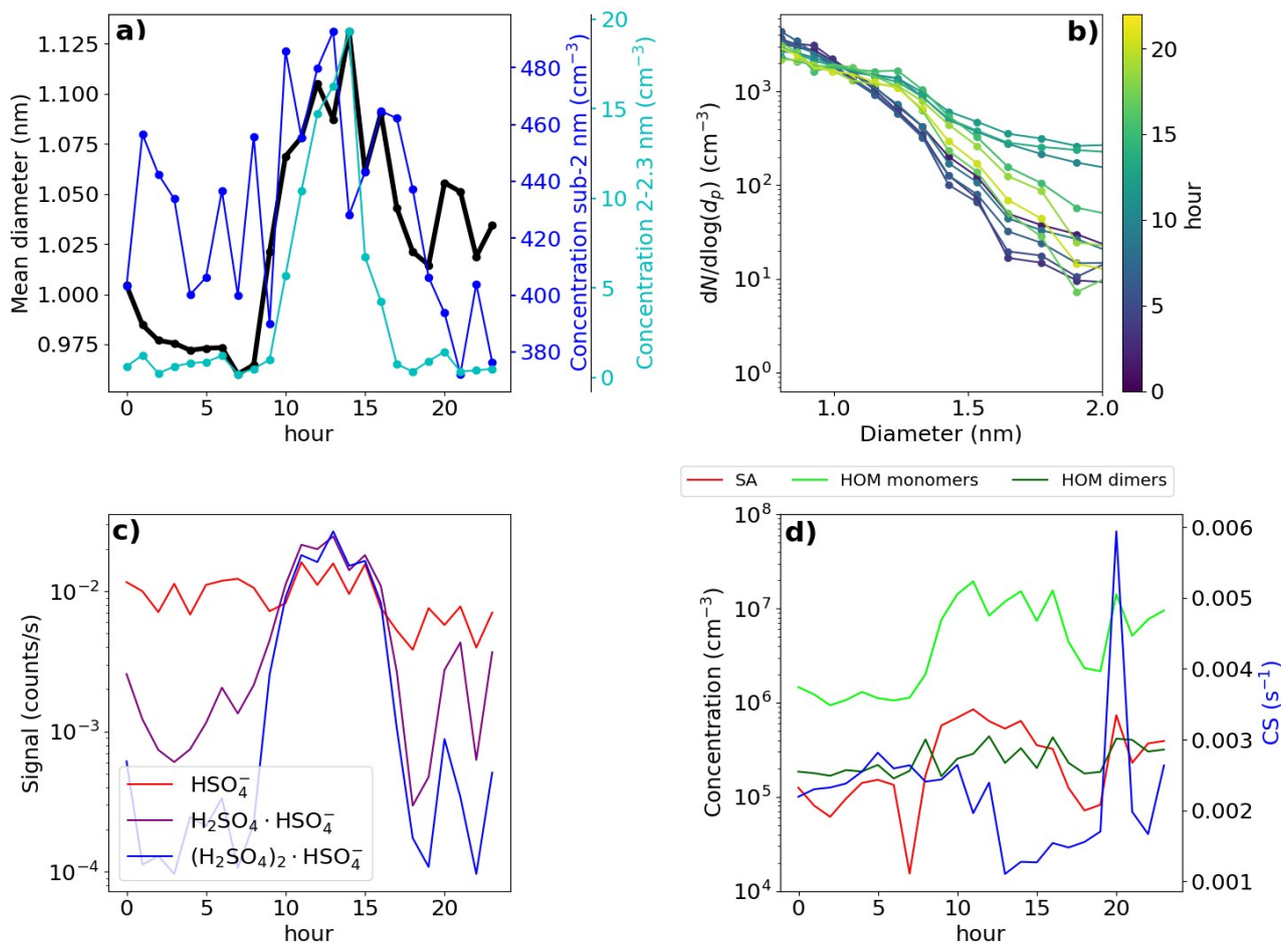
### 500 **3.7.1 Hyytiälä case 1 – an early spring day with NPF**

501 First of the investigated days was 10<sup>th</sup> of March, in 2021 and is presented in Fig. 11. During this  
502 day, a strong NPF event was observed with clear growth observed both in the total particle and ion  
503 number size distribution (see Fig. A3). In the morning, a strong increase in the SA ion dimer and  
504 trimer signals was detected after 07:00 (Fig. 11c), which occurred simultaneously with an increase  
505 in the concentration of neutral sulfuric acid. Shortly after, at around 08:00, neutral HOM monomer  
506 concentration started to increase (Fig. 11d). A strong increase in the concentration of 2.0-2.3 nm  
507 negative ions was observed after 09:00, indicating intense NPF on a local-scale (Fig. 11a).  
508 Approximately one hour before an increase in the concentration of 2.0-2.3 nm ions was first  
509 observed, the small ion  $d_{\text{mean}}$  started to increase from below 1.0 nm (Fig. 11a), showing that growth  
510 of clusters in the small ion population to larger sizes had begun. We can also see this from the  
511 negative ion number size distributions (0.8-2.0 nm; Fig. 11b): in the early hours of the day, the  
512 concentrations of the smallest ions are at their highest while the concentration of ions above approx.  
513 1.1 nm are at their lowest. Throughout the morning hours, we can see that the concentration of ions  
514 above approx. 1.1 nm increases and in the afternoon, around 14:00, the concentration of ions close  
515 to 2.0 nm is over a order of magnitude higher than during the night before. At around 14:00, the  
516 concentration of 2.0-2.3 nm ions and small ion  $d_{\text{mean}}$  also reach their peaks. At the same time, CS is  
517 at its lowest (Fig. 11d). Then, the concentrations of larger small ions, 2.0-2.3 nm and SA ion  
518 clusters starts to decrease, alongside with the concentration of HOM monomers.

519 We also took a look at the diameter specific concentrations in a smaller time frame (Fig. A6), which  
520 clearly shows how clear increase concentrations is observed for the diameters above 1.2 nm. A time  
521 delay between the increasing concentration of larger ions and smaller ions was seen, showing the  
522 growth of ions between 1.2 to 2.0 nm. Using the appearance time method (Lehtipalo et al., 2014),  
523 GR between 1.24 to 2.05 nm was estimated: GR = 0.40 nm/h. This value is somewhat lower than  
524 typical GRs reported in Hyytiälä (Hirsikko et al., 2005; Yli-Juuti et al., 2011), as expected due to  
525 the small size of the considered ions. Regardless, it shows that the growth of ions below 2.0 nm is  
526 non-negligible on this particular day. We note that this GR, or the ones presented for Hyytiälä Case  
527 2 and Beijing Case, is not a representative of the whole range of GRs for similar cases in the same  
528 location, and that there can be considerable variability.

529 This day clearly shows how the sulfuric acid and HOM driven particle formation is seen as the  
530 growth of the small ions to larger diameters. With the ion size distribution data, we have been able  
531 to get insight on when the cluster growth starts and how it progresses throughout this day.

2021-03-10



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538

**Fig. 11:** Data from Hyytiälä, 10<sup>th</sup> of March, 2021. (a) Hourly mean diameter of negative small ions (0.8-2.0 nm), total concentration of small ions, and concentration of 2.0-2.3 nm negative ions. (b) Two-hour median number size distribution of negative small ions. (c) Hourly signals of HSO<sub>4</sub><sup>-</sup>, H<sub>2</sub>SO<sub>4</sub>·HSO<sub>4</sub><sup>-</sup> and (H<sub>2</sub>SO<sub>4</sub>)<sub>2</sub>·HSO<sub>4</sub><sup>-</sup> ions. (d) Hourly median concentrations of neutral sulfuric acid (SA) and highly oxidized molecule (HOM) monomers and dimers, and the condensation sink (CS).

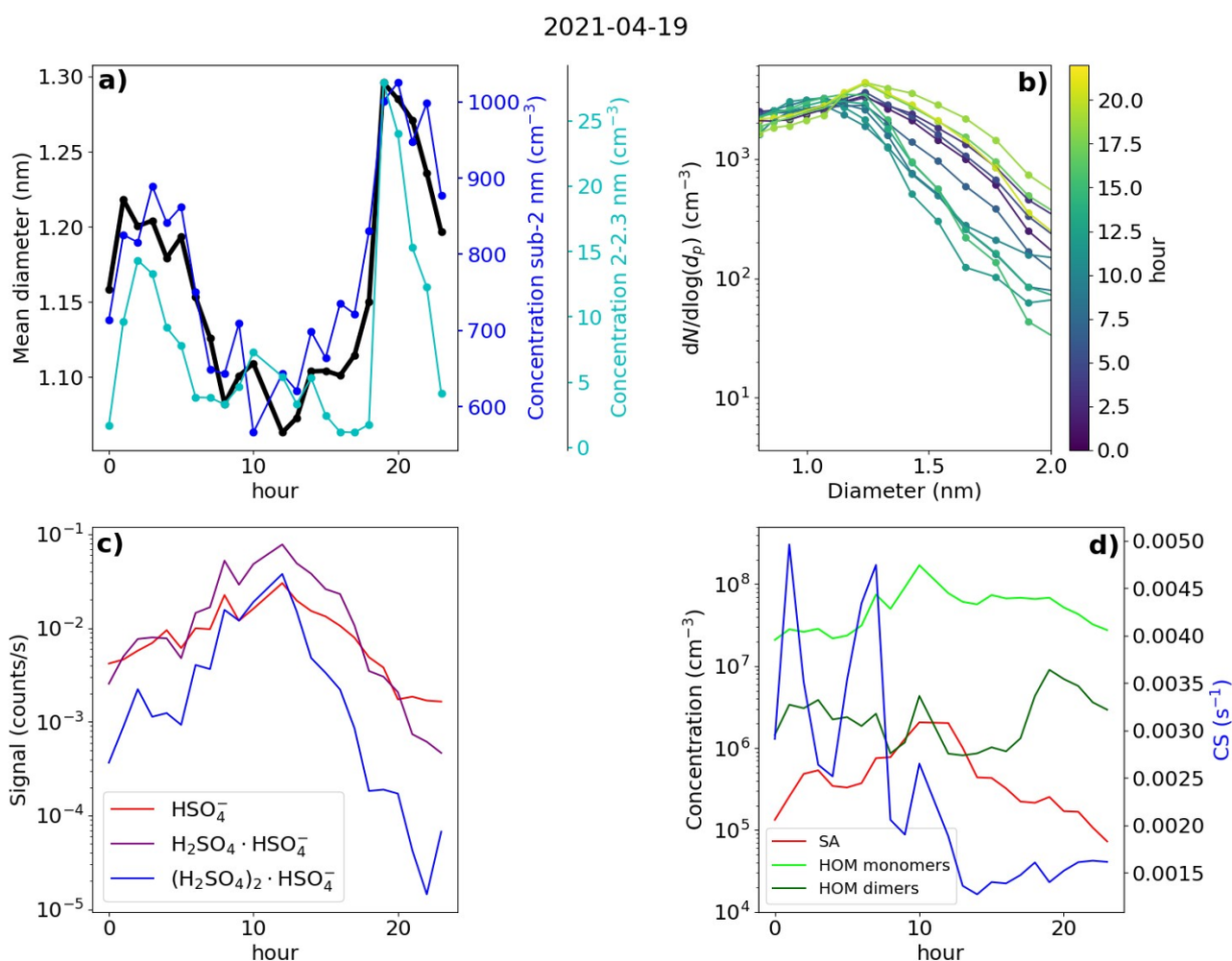
### 539 3.7.2 Hyytiälä Case 2 – a spring day with strong evening clustering

540 The second of the chosen days for Hyytiälä is 19<sup>th</sup> of April, 2021 (Fig. 12). The NPF ranking of this  
541 day was high, over 0.9, however the growth in the negative ion and total particle mode was  
542 discontinuous with the clearest growth observed above 5.0 nm, suggesting that the fraction of  
543 growing locally formed neutral clusters or ions was low (Fig. A4). However, strong evening ion  
544 cluster formation was observed on this day. Therefore, Case 2 illustrates both the contribution of  
545 organic vapors to initiate the growth of larger particles and the evening ion cluster formation  
546 attributable to HOM dimers (Mazon et al., 2016).

547 Starting from the early hours of the day, the signals of SA ions and neutral SA concentration  
548 increase (Fig. 12c and 12d), reaching their maxima around 13:00 in the early afternoon. Compared  
549 to Hyytiälä Case 1, the signal from trimers is lower in relation to the signal from monomer and  
550 dimer. At the same time, CS decreases (Fig. 12d). From the negative ion number size distributions  
551 (Fig. 12b), we see that the concentration of negative ions below approx. 1.2 nm increases and the  
552 concentration of small ions above approx. 1.2 nm strongly decreases starting from the early hours of

553 the day until afternoon. This is reflected in the value of  $d_{\text{mean}}$ , which decreases from over 1.2 nm to  
 554 below 1.1 nm (Fig. 12a). The concentration of 2.0-2.3 nm negative ions decreases until 08:00 in the  
 555 morning, after which it increases briefly before decreasing again (Fig. 12a). The small ion total  
 556 concentration also strongly decreases from over 800  $\text{cm}^{-3}$  to 600  $\text{cm}^{-3}$  (Fig. 12a). Unlike in Case 1,  
 557 on this day, the growth of small ions during daytime is negligible and an increased fraction of the  
 558 available charge is taken up by small, below 1.2 nm ions, many of which are likely composed of  
 559 sulfuric acid monomers or dimers. This explains the behavior of the ion size distributions,  $d_{\text{mean}}$  and  
 560 the total small ion concentration.

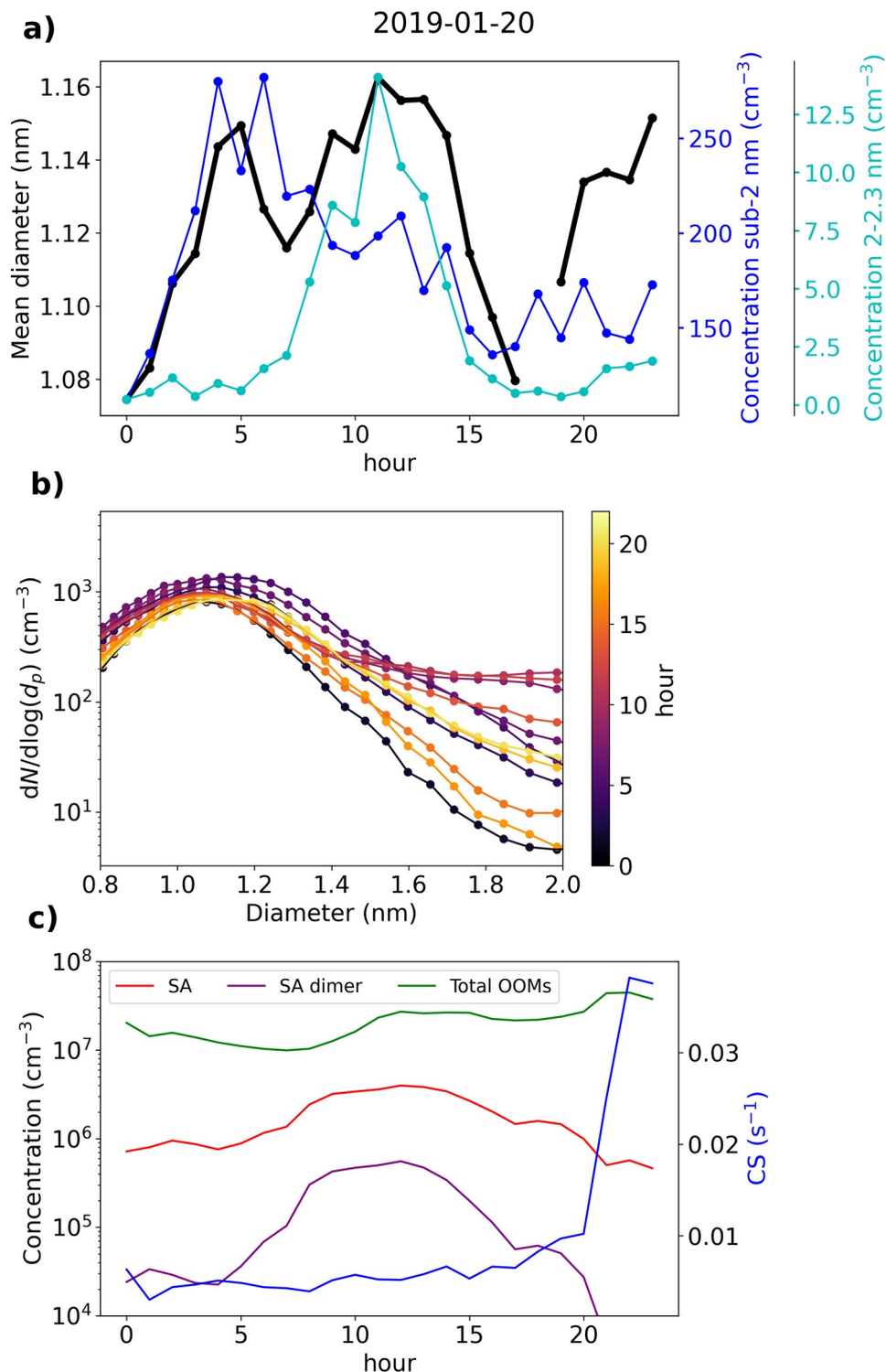
561 After 14:00 in the afternoon, the concentration of neutral HOM dimers starts to increase, and  
 562 reaches a peak at around 19:00 (Fig. 12d). Compared to Case 1, the HOM dimer concentration is  
 563 over one order of magnitude higher. Notably, at the same time as the HOM dimer concentration  
 564 starts increase, clear growth of total particles above 5.0 nm is observed (Fig. A4). Concentration of  
 565 small ions larger than approx. 1.2 nm (Fig. 12b) and 2.0-2.3 nm ion concentration (Fig. 12a)  
 566 strongly increase. Small ion  $d_{\text{mean}}$  increases from approx. 1.1 nm to 1.3 nm, while the total negative  
 567 small ion concentration increases from around 600  $\text{cm}^{-3}$  to 1000  $\text{cm}^{-3}$ . The negative ion GR between  
 568 1.43 to 2.05 nm was estimated to be 1.28 nm/h (Fig. A6), which is over twice as high as the GR  
 569 estimated for Case 1, likely due to the high concentration of lower volatility HOMs driving the  
 570 small ion growth during this particular evening.



571  
 572 **Fig. 12:** Data from Hyytiälä, 19<sup>th</sup> of April, 2021. (a) Hourly mean diameter of negative small ions

573 (0.8-2.0 nm), total concentration of small ions, and concentration of 2.0-2.3 nm negative ions. (b)  
 574 Two-hour median number size distribution of negative small ions. (c) Hourly signals of  $\text{HSO}_4^-$ ,  
 575  $\text{H}_2\text{SO}_4\cdot\text{HSO}_4^-$  and  $(\text{H}_2\text{SO}_4)_2\cdot\text{HSO}_4^-$  ions. (d) Hourly median concentrations of neutral sulfuric acid  
 576 (SA) and highly oxidized molecule (HOM) monomers and dimers, and the condensation sink (CS).

577 **3.7.3 Beijing Case – a day with intense NPF**



578

579 **Fig. 13:** Data from Beijing, 20<sup>th</sup> of January, 2019. (a) Hourly mean diameter of negative small ions  
580 (0.8-2.0 nm), total concentration of small ions, and concentration of 2.0-2.3 nm negative ions. (b)  
581 Two-hour median number size distribution of negative small ions. (c) Hourly median concentrations  
582 of neutral sulfuric acid (SA), SA dimer, and total oxidized organic molecules (OOMs), and  
583 condensation sink (CS).

584 Fig. 13 presents data from Beijing on 20<sup>th</sup> of January, 2019. This day was characterized by an  
585 intense NPF event, observed both in the ion and the total particle size distribution (Fig. A5). We see  
586 that from 00:00 until 05:00 in the morning, the negative small ion concentrations seem to increase,  
587 which is apparent for the whole sub-2 nm size range (Fig. 13a and 13b). Simultaneously,  $d_{\text{mean}}$  of the  
588 small ions increases (Fig. 13a). The concentration of 2.0-2.3 nm negative ions stays low (Fig. 13a),  
589 indicating that there is no significant growth of small ions to intermediate ions. CS is similar  
590 throughout the night and the early morning. Based on the data presented here, we hypothesize that  
591 the increase in small ion concentration is attributed to a larger fraction of the ions being detected by  
592 the instrument as  $d_{\text{mean}}$  increases.

593 After 05:00 in the morning, an increase in neutral sulfuric and sulfuric acid dimer concentration is  
594 observed (Fig. 13c). Simultaneously, the concentration of 2.0-2.3 nm ions increases sharply,  
595 indicating the formation of intermediate ions. Two changes in the small ion size distribution are  
596 shown: first, the concentration of small ions below approx. 1.5 nm ions, decreases and second, the  
597 concentration of small ions above that increases. Increasing growth of small ions to larger sizes  
598 causes a shift in their size distribution. Notably, no growth in the surface plots (Fig. A5) is observed  
599 yet, likely due to locality of or insufficient intensity of the ion formation. After 12:00, the  
600 concentrations of small ions larger than approx. 1.5 nm start to decrease, as does the concentration  
601 of 2.0-2.3 nm ions. While the growth of ions and particles at larger diameters continues, the  
602 intensity of the cluster growth decreases.

603 In this case, the negative small ion GR was estimated to be 0.24 nm/h from 1.72 to 2.06 nm (Fig.  
604 A7), which is lower than the values determined for the two Hyytiälä cases and is on the lower range  
605 of values of particle GRs for Beijing (Deng et al., 2020). Another noteworthy observation can be  
606 made from the diameter specific concentrations (Fig. A7): as already seen from the size  
607 distributions and more clearly here, the concentrations of ions up to around 1.5 nm decrease, while  
608 the concentrations above increase at the same time. This implies that the ions, which actually start  
609 to grow to larger sizes are close to 1.5 nm in diameter, though at such a low GR their survival  
610 probability to larger sizes is likely very low (Kulmala et al., 2017).

611 In Sect. 3.3, we saw how in Beijing there does not seem to be correlation between the small ion  
612 number size distribution and the concentration of sulfuric acid. On this day, the increased  
613 concentrations of sulfuric acid occurred approx. simultaneously with the observed small ion growth.  
614 Previous studies have shown the importance of sulfuric acid in particle formation in Beijing (Yao et  
615 al., 2018; Cai et al., 2021; Yan et al., 2021). As such, it seems likely that the growing small ions  
616 seen on this day are composed of sulfuric acid. However, while sulfuric acid forms these growing  
617 clusters, their growth also requires other ingredients.

## 618 **4 Conclusions**

619 We studied the seasonality of small ion number size distribution and the relationship of the small  
620 ion size distribution with low-volatility organic vapors, sulfuric acid, coagulation sink (CoagS) and  
621 NPF in a rural boreal forest location of Hyytiälä, Finland and an urban megacity location of Beijing,  
622 China. Both analysis of long time series of data and daily case studies were carried out. We found a  
623 clear seasonality of the small ion size distribution in Hyytiälä, where the small ions of both  
624 polarities were the smallest in size during winter and the largest during late spring and summer. In  
625 Beijing, while there were month-to-month variations in the size distribution, but no clear seasonal  
626 pattern was identified.

627 We found that in Hyytiälä the small ion size distribution strongly varied with respect to the  
628 concentration of organic, especially highly oxidized organic (HOM) monomer, compounds and that  
629 the concentration of small ions above approx. 1.2 nm increased strongly with increasing HOM  
630 monomer concentration. This was observed more strongly for negative polarity and during the  
631 evening, which was found to be connected to the evening ion cluster formation driven by organics  
632 in Hyytiälä. The small ion size distribution also showed clear increase in the size of the small ions  
633 in Hyytiälä with respect to neutral sulfuric acid and ionized sulfuric acid dimers, associated with  
634 daytime cluster formation and growth. In contrast, there was no clear relationship between the  
635 concentration of either organic vapor or sulfuric acid and the size of the small ions in Beijing. The  
636 reason for this remains to be identified, but we hypothesize that the concentration of bases is the  
637 limiting factor determining if growth of small ions is seen in Beijing.

638 We found that the small ion size distribution in either location did not change strongly with  
639 changing CoagS. In Hyytiälä, small decrease in the concentration of especially the smallest ions  
640 was seen, as expected. However, despite our expectations, this was not observed in Beijing. The  
641 reason for this remains to be answered in future research.

642 When the concentration of ions in the range 2.0-2.3 nm increased, indicating the occurrence of local  
643 NPF, we observed clear signs of growth in the small ion size distribution. This was seen in both  
644 locations, even in Beijing, where no clear association of small ion size with organic vapor or  
645 sulfuric acid was found. To a lesser extent, an increase in the small ion size was also seen with  
646 respect to NPF rank, a parameter, which characterizes the intensity of NPF. These results support  
647 the conclusion that in Beijing the concentration of sulfuric acid or organic vapor does not determine  
648 whether small ions grow to intermediate ions.

649 Overall, we have shown in a novel way how the atmospheric cluster formation and growth  
650 processes impact the number size distribution of small ions. The sub-2 nm size range is integral for  
651 understanding of the first steps of new particle formation and the activation of the clusters to grow  
652 into particles. Our results can be applied in research into the dynamics of charged clusters and how  
653 they grow from clusters to particles.

## 654 **Author contributions**

655 ST analyzed the data and wrote the manuscript. JL was responsible for the ion measurements in  
656 Hyytiälä. CL and NS were responsible for the measurements of low-volatility vapors and ion

657 clusters. YL was responsible for the measurements in Beijing. MK and VMK conceptualized the  
658 study. All authors contributed to reviewing and editing the manuscript.

## 659 Code and data availability

660 The Hyytiälä DMPS dataset can be accessed through the SmartSMEAR data portal at  
661 <https://smear.avaa.csc.fi/>. Other data and the codes used to produce the figures in this paper are  
662 available upon request from the authors.e

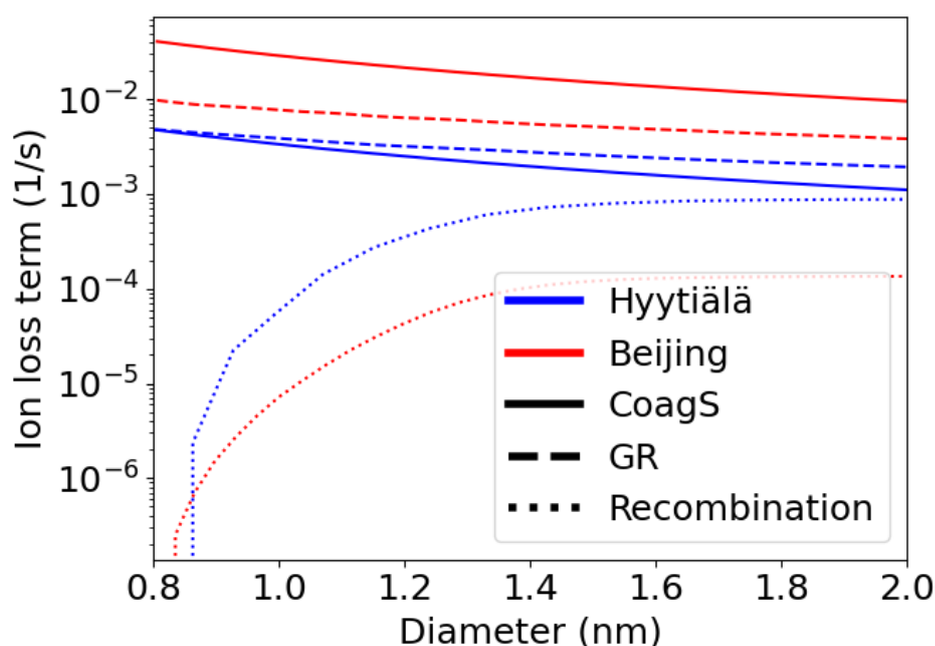
## 663 Competing interests

664 At least one of the (co-)authors is a member of the editorial board of Aerosol Research. Authors  
665 have no other competing interests to declare.

## 666 Acknowledgments

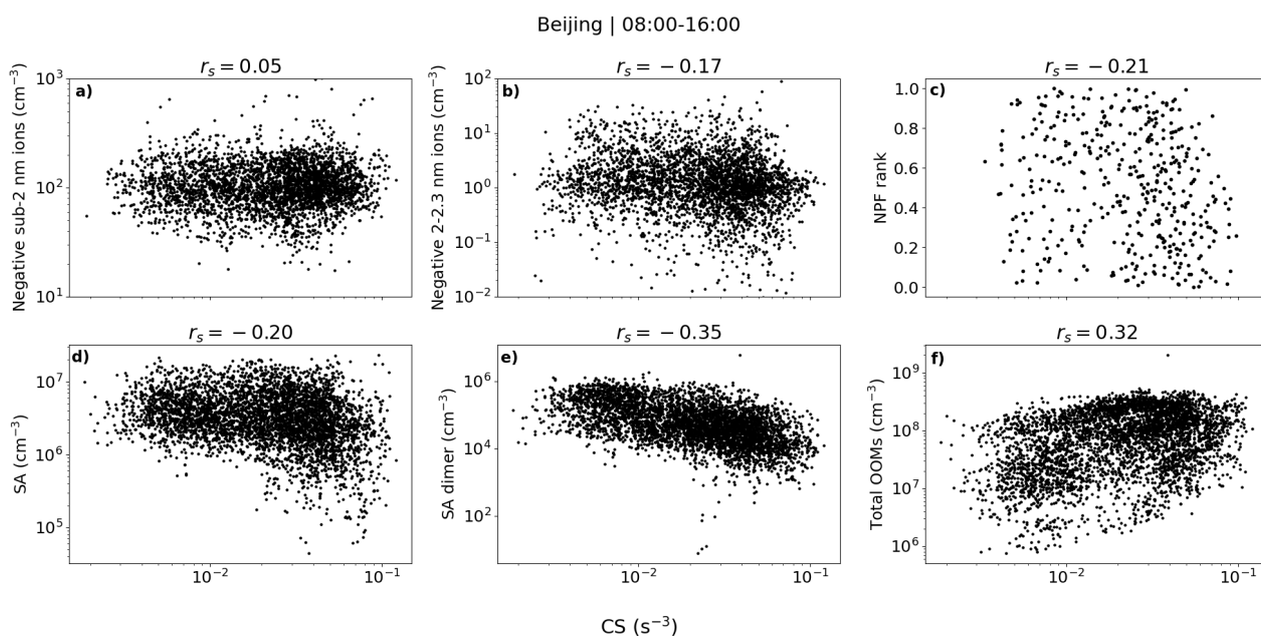
667 This work has been supported by the ACCC Flagship funded by the Academy of Finland grant nos.  
668 337549 (UH) and 337552 (FMI), and the “Gigacity” project funded by the Jenny and Antti Wihuri  
669 Foundation. We acknowledge the SMEAR II and AHL/BUCT technical and scientific staff.

## 670 Appendix



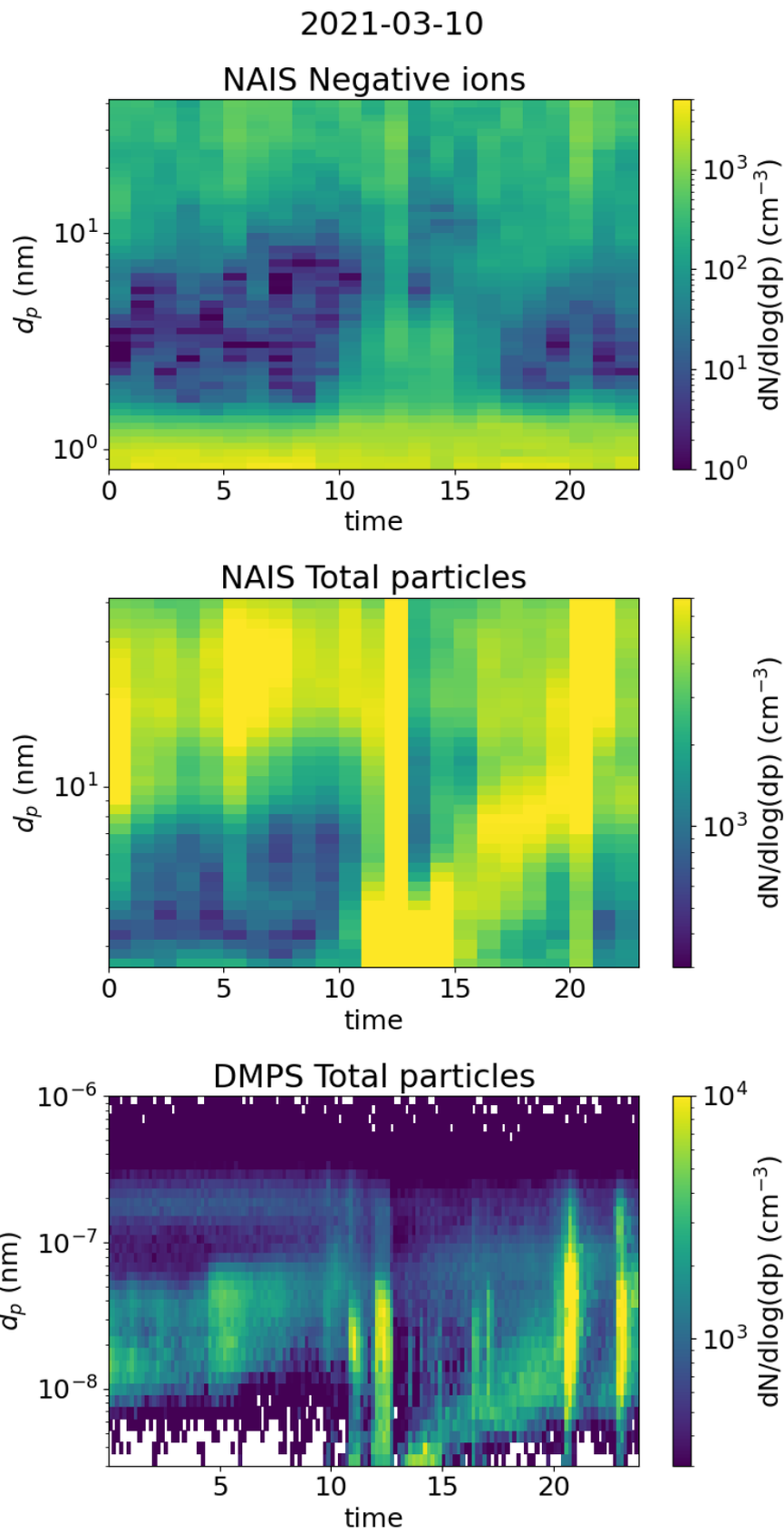
671

672 **Fig. A1:** A rough approximation of the different loss terms of negative ions as a function of ion  
673 diameter based on Eq. 2 in both Hyytiälä and Beijing. The concentration of positive ions were based  
674 on the median number size distributions. GR was assumed to equal 1 nm/h and was assumed to be  
675 constant with diameter. CoagS was scaled based on the median CS, and assuming that the charge  
676 enhancement of the sink was by a factor of 2. The median CS were  $CS=2.9 \cdot 10^{-3} \text{ s}^{-1}$  and  $2.5 \cdot 10^{-2} \text{ s}^{-1}$ ,  
677 for Hyytiälä and Beijing, respectively.



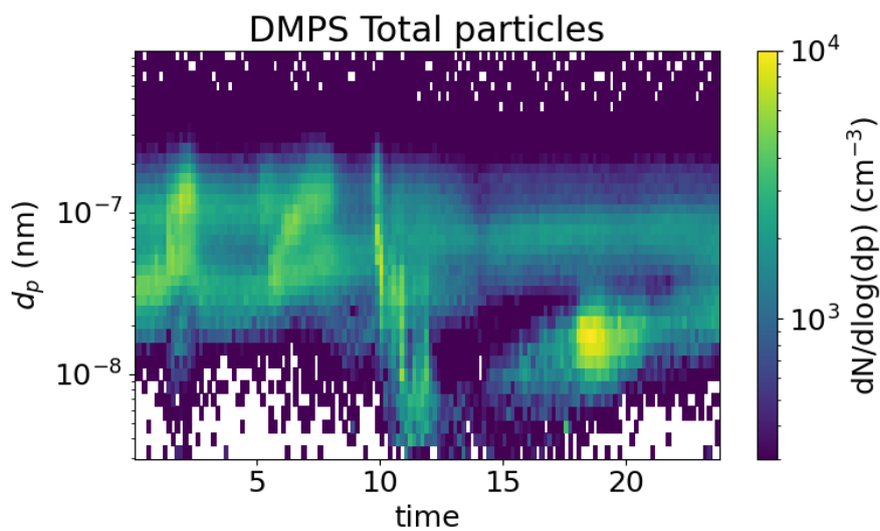
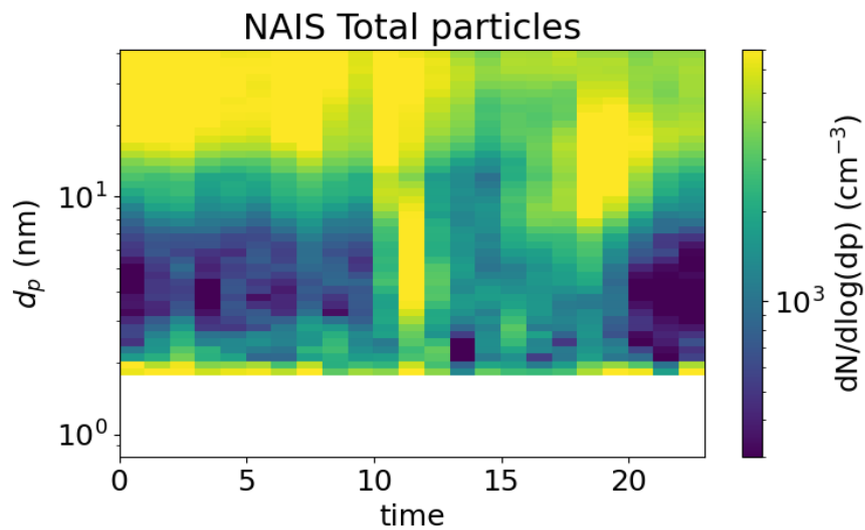
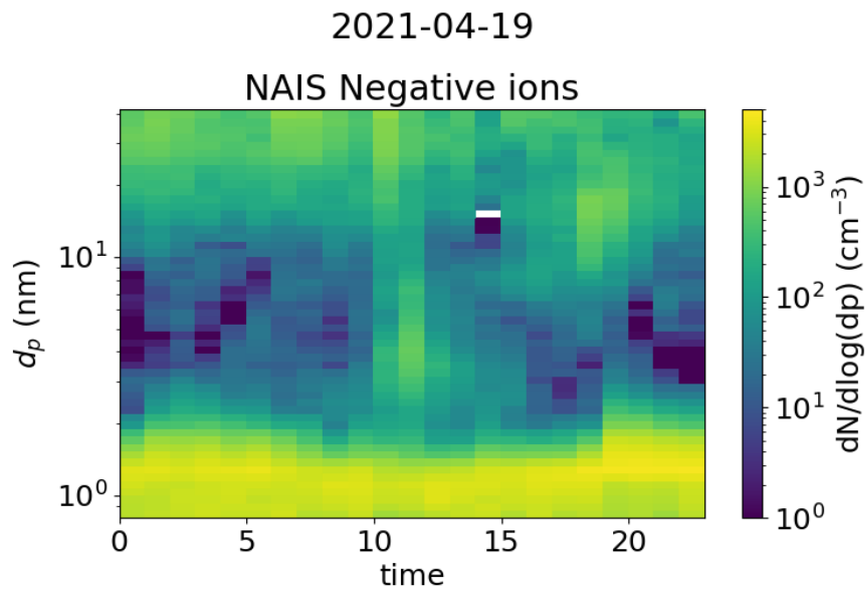
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679 **Fig. A2:** The concentrations of negative sub-2 nm (a) and 2.0-2.3 nm (b) ions, NPF ranking values  
 680 (c), and the concentrations of neutral sulfuric acid (SA, d), SA dimer (e), and total oxidized organic  
 681 molecules (OOMs, f) with respect to condensation sink (CS) in Beijing. The values are hourly  
 682 medians, except for c), where NPF ranking is a daily parameter and CS is the daytime median. The  
 683 Spearman correlation coefficients ( $r_s$ ) are also shown.



684

685 **Fig. A3:** Surface plots of negative ion number size distribution and total particle number size  
 686 distribution measured by NAIS and DMPS in Hyytiälä on 10<sup>th</sup> of March, 2021.

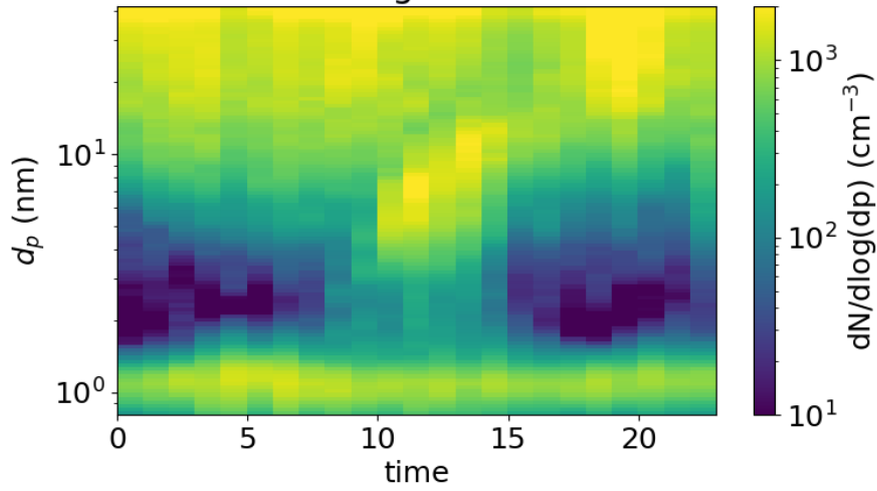


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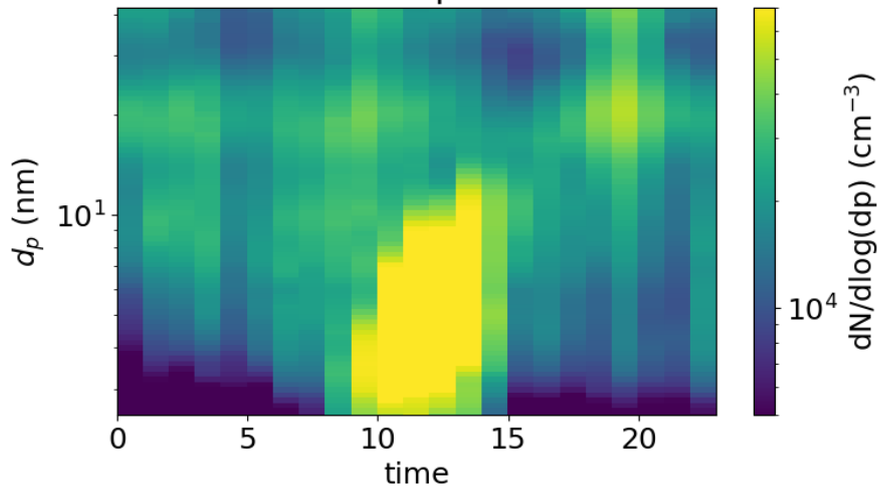
688 **Fig. A4:** Surface plots of negative ion number size distribution and total particle number size  
 689 distribution measured by NAIS and DMPS in Hyytiälä on 19<sup>th</sup> of April, 2021.

2019-01-20

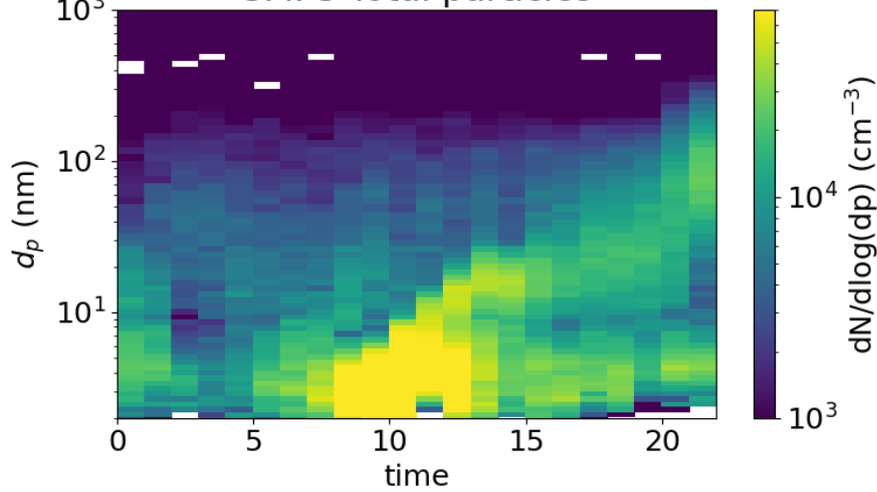
NAIS Negative ions



NAIS Total particles

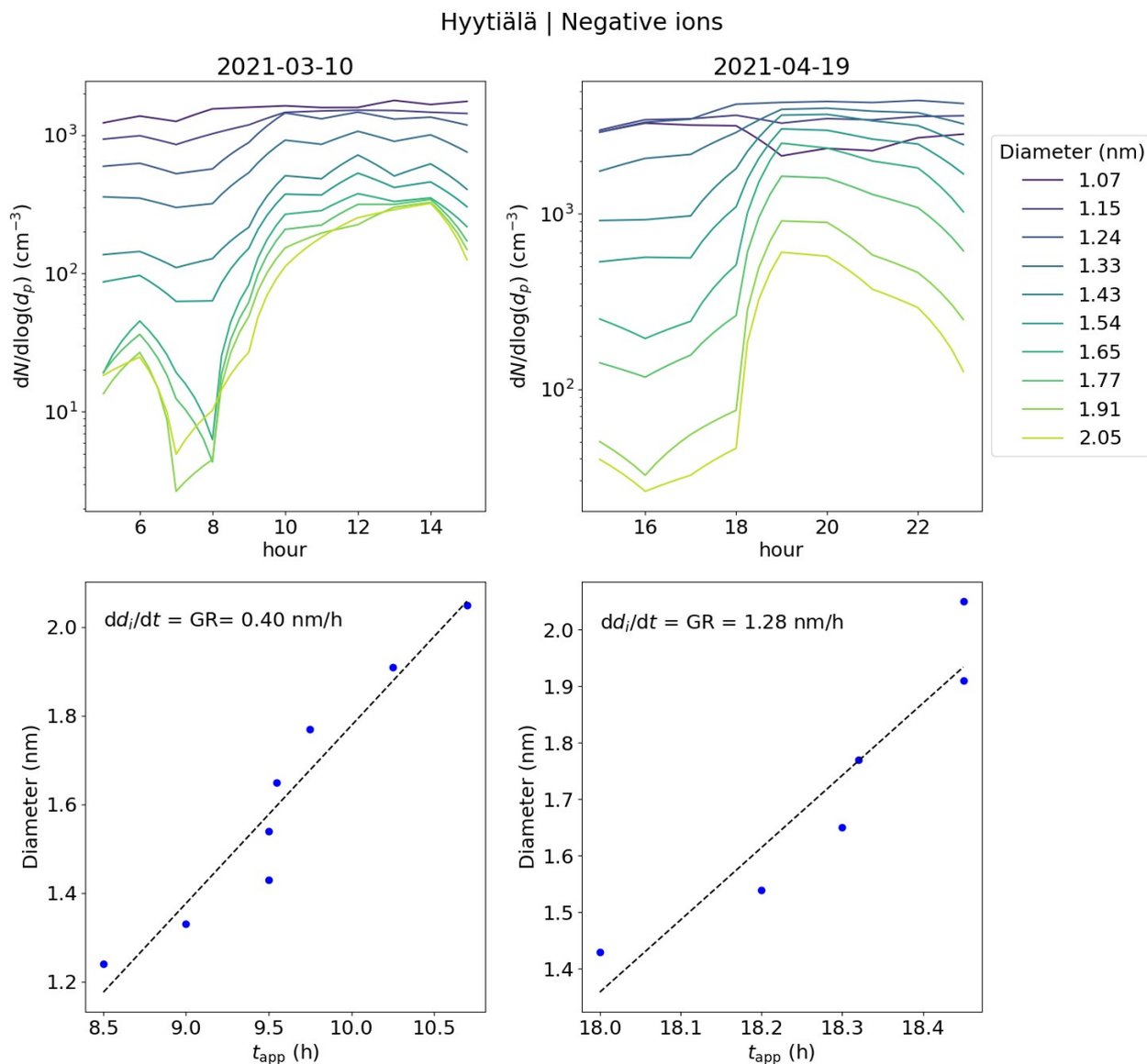


SMPS Total particles



691 **Fig. A5:** Surface plots of negative ion number size distribution and total particle number size  
 692 distribution measured by NAIS and SMPS (see Liu et al., 2016 for more information) in Beijing on  
 693 20<sup>th</sup> of January, 2019.

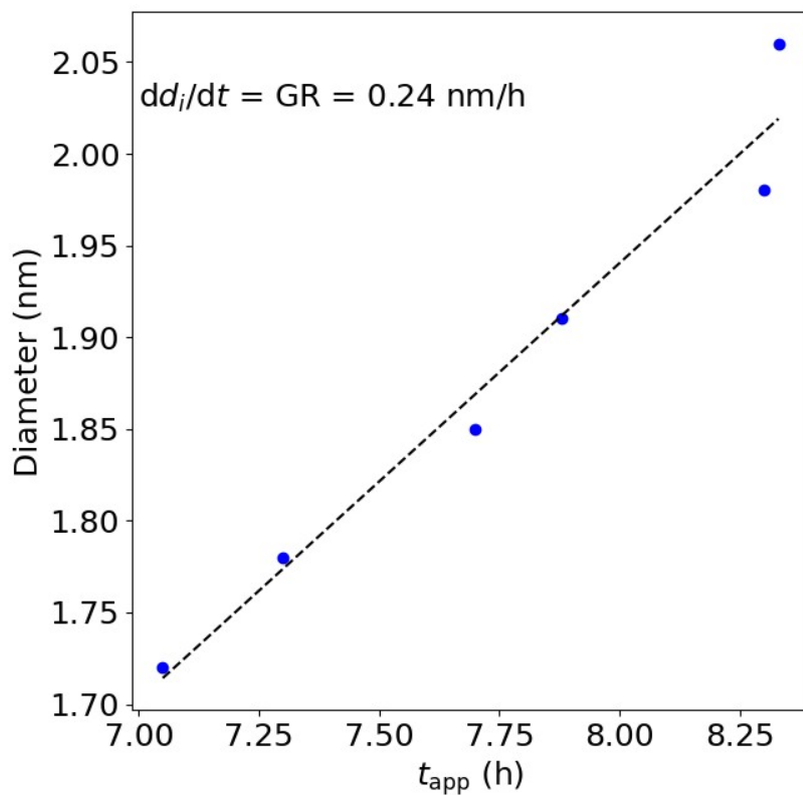
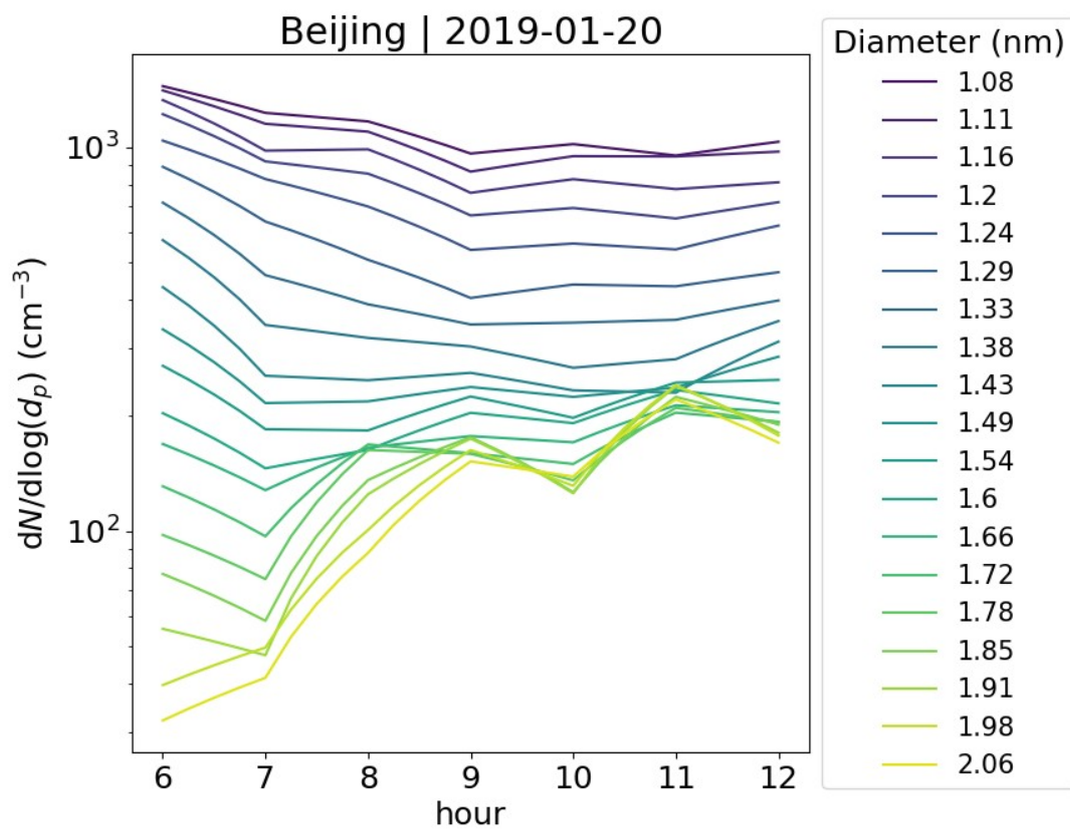
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695

696 **Fig. A6:** The upper panels show concentrations of ions of a certain diameter with the hour of the  
 697 day on 10<sup>th</sup> of March, 2021 and 19<sup>th</sup> or April, 2021 in Hyytiälä, Finland. The different colors of the  
 698 line indicate the respective ion diameter ( $d_i$ ). The bottom panels show the appearance time, defined  
 699 as the time that the concentration reaches 50% of its maximum, and the respective  $d_i$ . The ion  
 700 growth rate (GR) derived from these values as a slope of linear regression is shown. For 10<sup>th</sup> of  
 701 March, the GR was determined from 1.24 to 2.05 nm and for 19<sup>th</sup> of April from 1.43 to 2.05 nm.

702



703

704 **Fig. A7:** The upper panel shows the concentrations of ions of a certain diameter with the hour of the  
 705 day on 20<sup>th</sup> of January, 2019 Beijing, China. The different colors of the line indicate the respective

706 ion diameter ( $d_i$ ). The bottom panel shows the appearance time, defined as the time that the  
707 concentration reaches 50% of its maximum, and the respective  $d_i$ . The ion growth rate (GR)  
708 derived from these values as a slope of linear regression is shown. The GR was determined from  
709 1.72 to 2.06 nm.

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