

1 **Analysis of insoluble particles in hailstones in China**

2 Haifan Zhang¹, Xiangyu Lin¹, Qinghong Zhang^{1*}, Kai Bi^{2*}, Chan-Pang Ng¹, Yangze Ren¹, Huiwen Xue¹, Li Chen³, Zhuolin
3 Chang⁴

4 ¹Department of Atmospheric and Oceanic Sciences, School of Physics, Peking University, Beijing 100871, China

5 ²Field Experiment Base of Cloud and Precipitation Research in North China, China Meteorological Administration, Beijing
6 101200, China

7 ³Electron Microscopy Laboratory, Peking University, Beijing 100871, China

8 ⁴Key Laboratory for Meteorological Disaster Monitoring and Early Warning and Risk Management of Characteristic
9 Agriculture in Arid Regions, China Meteorological Administration, Yinchuan 750002, China

10 *Corresponding author: Qinghong Zhang (qzhang@pku.edu.cn); Kai Bi (bikai_picard@vip.sina.com)

11

Abstract. Insoluble particles influence weather and climate by heterogeneous freezing process. Current weather and climate models face considerable uncertainties in freezing processes simulation due to limited information regarding species and number concentration of heterogeneous ice-nucleating particles, particularly insoluble particles. Here, for the first time, size distribution and species of insoluble particles are analyzed in 30 shells of 12 hailstones collected from China, using scanning electron microscopy and energy dispersive X-ray spectrometry. A total of 289,461 insoluble particles were detected and identified into 3 species: organics, dust, and bioprotein, utilizing machine learning methods. The size distribution of insoluble particles of each species varies greatly among the different hailstones but little in their shells. Further, classic size distribution of organics and dust followed logarithmic normal distributions, which could potentially be adapted in future weather and climate models, despite the existence of uncertainties. ~~Our findings highlight the need for atmospheric chemistry to be considered in the simulation of ice freezing process. Our finding suggests the aerosol species and number concentration variance in different storms should be considered in model simulation of the ice freezing process.~~

1 Introduction

Insoluble particles, acting as main heterogeneous ice-nucleating particles in the atmosphere (Lamb and Verlinde, 2011), influence precipitation formation and radiative forcing (Hoose and Möhler, 2012; DeMott et al., 2015), and further impact weather and climate (Vergara-Temprado et al., 2018). Temperature and vapor supersaturation are used to calculate the number concentration of ice crystal particles in microphysics parameterization rather than considering the physical properties of ice-nucleating particles in weather and climate models (DeMott et al., 2010). Few models used the freezing parameterization, which establishes a direct connection between the number concentration of ice-nucleating particles and the number concentration of ice crystals. The absence of description regarding the number concentration of ice-nucleating particles in models can result in an incorrect estimation of ice crystals and lead to significant bias in radiative simulations (Vergara-Temprado et al., 2018).

An improved description of the number concentrations of ice-nucleating particles is needed (DeMott et al., 2010), while obstructed by a lack of complete microphysical observation in clouds about ice-nucleating particles. There are two ways to sample ice-nucleating particles: The first involves an airborne instrument, named continuous flow thermal gradient diffusion chamber (Rogers et al., 2001; Prenni et al., 2009; DeMott et al., 2010). The second is done in the laboratory, where scientists conduct freezing experiments (Hoose and Möhler, 2012). In most cases, it is necessary for an aircraft to collect air parcels for measurement of the physical properties of ice-nucleating particles in the air. However, former field projects sampled air parcels in anvils of convective clouds, cirrus and winter mixed-phase stratiform clouds. No flight report or article has reported that they sampled air parcels through cores in deep convection. This phenomenon is consistent with consideration for flight security. Thus, current observation is insufficient for describing the whole convective cloud, especially the deep convection in severe

42 storms. Absence about microphysical observations of ice-nucleating particles within severe storms leads to uncertainty in
43 understanding cold cloud process.

44 Hailstones, as a product of deep convective clouds, serves as a carrier of information within these clouds. Recently,
45 analysis revealed large diversity in number concentration of soluble ions among hailstones from different hailstorms (Li et al.,
46 2018). Further, the detection of soluble ions along with isotopic analysis of a huge hailstone revealed an up-and-down hailstone
47 growth trajectory, which demonstrated that the different shells were formed at different heights (Li et al., 2020). These studies
48 have proved that aerosol information in convective cloud may be recorded in soluble particles within hailstones (Li et al., 2018,
49 2020). Similarly, insoluble particles in hailstones can also record aerosol information in severe storms.

50 Former studies showed that species and number concentration of insoluble particles in hailstones (Vali, 1968; Rosinski,
51 1966; Michaud et al., 2014) would influence heterogeneous nucleation process (Hoose and Möhler, 2012) and further hailstone
52 formation (Knight, 1981). Information on the species of insoluble particles can determine the freezing temperature when these
53 particles participate in the initiation of ice crystal formation and subsequently impact hailstone embryo growth. Biological
54 particles in hailstones, such as pollen and bacteria, are more efficient ice-nucleating particles than dust within the ice nucleation
55 region of storm clouds (Michaud et al., 2014). They can raise the freezing threshold temperature above -15°C , while dust
56 particles are activated to form ice crystals at temperatures below -15°C (Michaud et al., 2014). In addition to species, number
57 concentration of insoluble particles can also influence the hailstone formation. When more dust particles were considered, a
58 model simulation resulted in larger number concentration of ice crystals, smaller graupels (one type of hailstone embryos) size,
59 and suppression of the hailstone growth (Chen et al., 2019). Nonetheless, previous studies involving analysis of insoluble
60 particles in hailstones mainly focused on substances analysis or total number concentration statistics. A size distribution of
61 insoluble particles in hailstones with species information, which is beneficial for completing microphysical observation in
62 severe storms, has not been given so far.

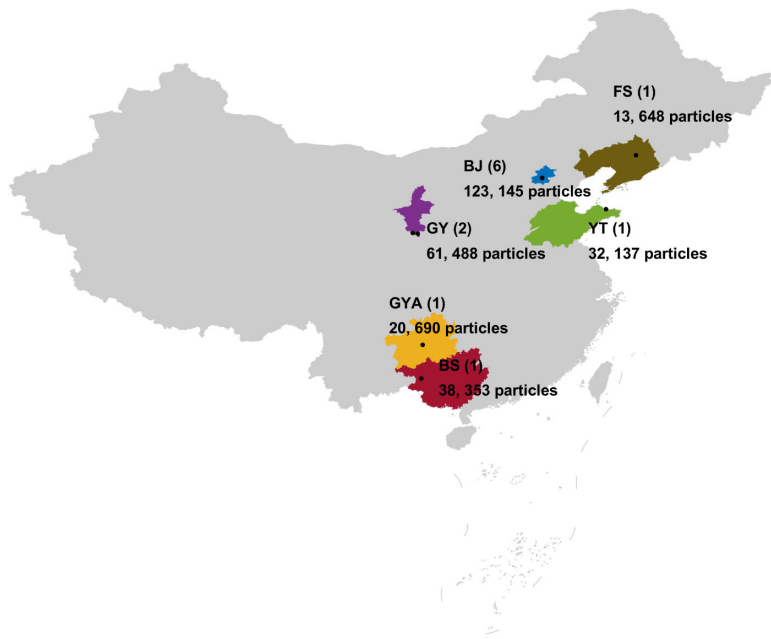
63 This study analyzed insoluble particles in hailstones collected from 8 hailstorms that occurred in China between 2016
64 and 2021. The identification of insoluble particles in hailstones was conducted using Scanning Electron Microscopy (SEM)
65 and Energy Dispersive X-ray spectrometry (EDX). The insoluble particles were identified into three species using Self-
66 Organized Maps (SOMs) and the random forest method. The variation in size distribution of insoluble particles in embryos
67 and different shells was explored. Based on the size distributions, logarithmic normal distributions were fitted to describe the
68 concentration of organics and dust in deep convection.

69 **2 Methods**

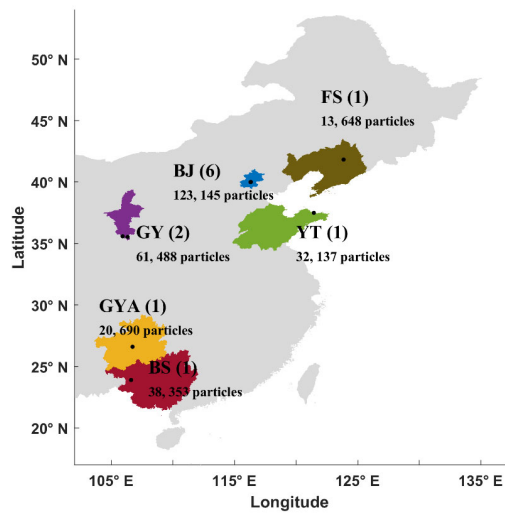
70 **2.1 Sample information and experimental design**

71 Hailstones were collected from eight hailstorms that occurred in six provinces of China during warm seasons from 2016

72 to 2021 (Table 1, Fig. 1). Volunteers stored the hailstones in clean containers, including plastic bags, glass containers, and
73 tinfoil, either during or immediately after the hail events. All hailstone samples were transported to a laboratory at Peking
74 University in Beijing and kept at temperatures ranging from -18 °C to -4 °C. The hailstones were then transferred into vacuum-
75 sealed plastic pockets and preserved in a freezer, maintaining an internal temperature ranging from -29°C to -23°C, until they
76 underwent further processing and analysis.



77



78
 79 **Fig. 1: Geographical distribution of collected hailstones. The collecting locations of hailstones are indicated by black dots. Provinces**
 80 **of China from which the hailstones were collected are represented by six different colors. The number of hailstones we analyzed was**
 81 **indicated in parentheses. Abbreviations (corresponding to Table 1): BJ - BeiJing; BS - BaiSe; FS - FuShun; GY - GuYuan; GYA -**
 82 **GuiYAng; YT - YanTai.**

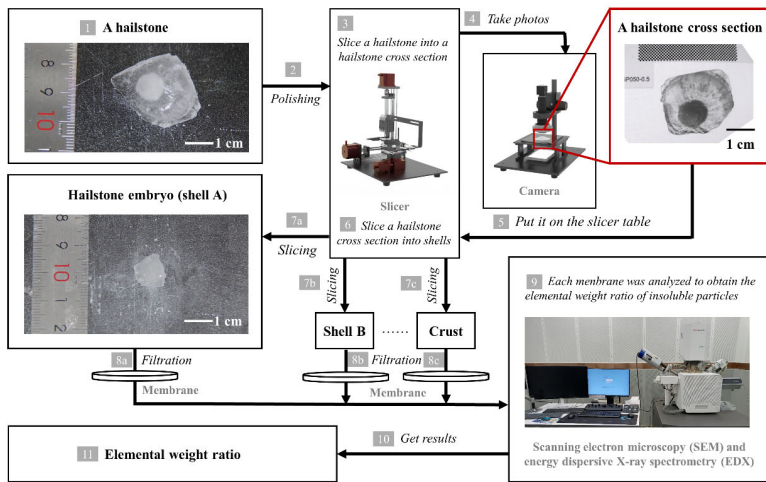
83 **Table 1: Information about collected hailstones.**

Date & Beijing Local Time ^a	Latitude & Longitude ^b	Total column	Freezing level	Location & Sample abbreviation ^c	Samples ^f	Diameter ^g (mm)	Particle number ^h
		water vapor ^e (kg / m ²)	height – orography altitude ^d (m)				
19 June 2018, 18:30	41.82° N, 123.85° E	26.35 ^[18]	3241.66 ^[18]	FuShun (FS)	1	13.80	13,648
10 June 2016, 15:00	40.00° N, 116.32° E	36.86 ^[14]	3780.52 ^[14]	BeiJing (BJ1)	1	—	35,291
30 June 2021, 20:18	39.95° N, 116.30° E	31.84 ^[20]	3852.76 ^[20]	BeiJing (BJ2)	5	25.38	14,865
				BeiJing (BJ3)		24.11	20,233
				BeiJing (BJ4)		16.30	20,350
				BeiJing (BJ5)		14.86	14,350
				BeiJing (BJ6)		22.80	18,056
01 Oct 2021, 14:02	37.49° N, 121.44° E	32.81 ^[13]	3642.42 ^[13]	YanTai (YT)	1	45.00	32,137
25 Aug 2020, 18:00	35.53° N, 106.32° E	17.83 ^[17]	422.58 ^[17]	GuYuan (GY1)	1	15.00	29,341
26 Aug 2020, 16:00	35.58° N, 105.93° E	17.01 ^[15]	835.04 ^[15]	GuYuan (GY2)	1	18.50	32,107
14 Apr 2016, 20:00	26.60° N, 106.72° E	31.62 ^[19]	2147.58 ^[19]	GuiYAng (GYA)	1	26.20	20,690
09 May 2016, 18:51	23.90° N, 106.60° E	47.45 ^[18]	4572.70 ^[18]	BaiSe (BS)	1	—	38,353

84

^a Date and Beijing local time of hailstorms occurrences. Hailstones were collected within 30 min during hail.^b Latitude and longitude where the hailstone were collected.^c Total column water vapor values (Beijing local time of ERA5 reanalysis data in square brackets (Hersbach et al., 2018)).^d Depth between freezing level and orography (Beijing local time of ERA5 reanalysis data in square brackets(Hersbach et al., 2018)).^e Location and sample abbreviations.^f Numbers of hailstones used in the experiments.^g Diameter of hailstone (— means no record).^h Insoluble particle number in hailstones.

85 Insoluble particles were extracted in the experiments (Fig. 2). The surface of each hailstone was polished to remove any
 86 attached grass or soil. Subsequently, the hailstones were sliced into cross-sections along the major axis, corresponding to the
 87 size of the hailstone embryo. The cross-section were further sliced into shells using heated Fe-Cr alloy wire at an air
 88 temperature below -8°C . The shells within a hailstone were distinguished based on their natural transparency or opacity.
 89 However, hailstones with a major axis < 7 mm could not be sliced due to the mass loss resulting from heating using our
 90 experimental apparatus.



91
 92 **Fig. 2:** Schematic diagram illustrating the experimental framework. [1-2] The surface of each hailstone was polished to remove any
 93 attached grass or soil. [3] Subsequently, the hailstones were sliced into cross-sections along the major axis, corresponding to the size
 94 of the hailstone embryo. [4-7] After photographing the hailstone cross-sections, they were further subdivided into shells using heated
 95 Fe-Cr alloy wire at an air temperature below -8°C . The shells were distinguished based on their natural transparency or opacity. [8]
 96 The solution of melting shell samples was then passed through a filter membrane to isolate the insoluble particles. [9] Each shell
 97 sample underwent analysis using scanning electron microscopy and energy-dispersive X-ray spectrometry to determine the
 98 elemental weight ratios of the insoluble particles within approximately 4 hours. [11] Finally, the elemental weight ratio information
 99 of hailstones was obtained.

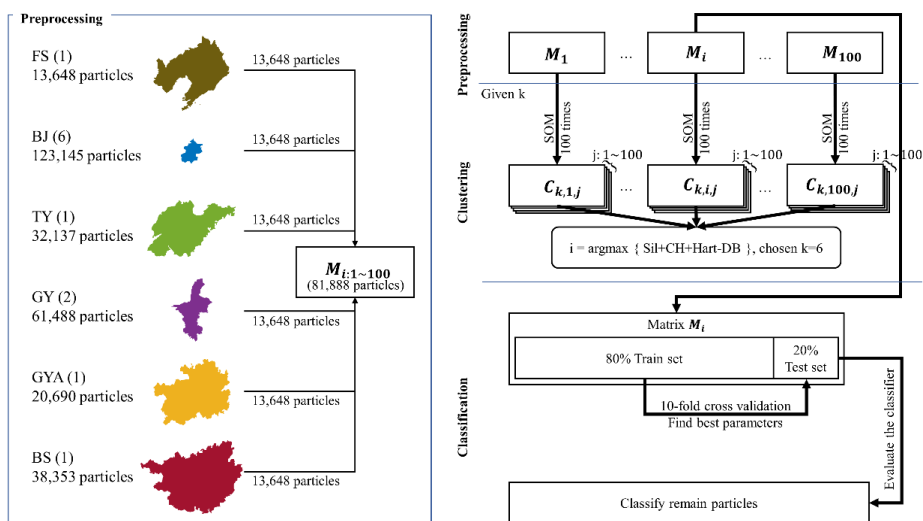
100
 101 The shells were sequentially labeled with capital letters in alphabetical order, starting from the embryo (designated as
 102 shell A) and progressing toward the crust. After the ice shells melting into a solution, the solution was filtered through a
 103 membrane (VSWP01300, Merck KGaA, Germany) with a pore size of 30 nm. The 1 mL (a total of 5 mL) of distilled water
 104 underwent five passes through the filter membrane to ensure maximum retention of insoluble particles on the membrane.

105 Subsequently, the filter membrane was dried under an air temperature of approximately 40°C to satisfy the dry-environment
 106 requirements of SEM.

107 The number of insoluble particles in each shell was determined using scanning electron microscopy (SEM), with a focus
 108 on particles larger than 0.16 μm. The length along the major axis of the particles was measured using Aztec software (Aztec
 109 software, Oxford Instruments plc, UK) on SEM images. The software was able to randomly capture electron microscopy
 110 photos of the membrane (Aztec User Manual). No particle will be counted repeatedly. Energy-dispersive X-ray spectrometry
 111 (EDX) was utilized to determine the elemental weight ratios of the particles. Only elements with an atomic number greater
 112 than 4 could be detected due to the X-ray input window being made of beryllium. Each shell sample was analyzed within
 113 approximately 4 hours by SEM and EDX. The scanning mode of SEM was set in a random order to reduce errors caused by
 114 bias in the detection area.

115 2.2 Clustering and classification

116 The number of insoluble particles was measured using Aztec on SEM images, but the species could not be determined
 117 directly and were identified by machine learning method. The criteria of species classification were established by the SOMs
 118 method to determine the species of unclassified particles. These labeled particles were then regarded as training set in random
 119 forest classifier. Details are presented in Fig. 3.



120
 121 **Fig. 3:** Schematic diagram illustrating the methodological framework used for particle identification in this study. A total of 100
 122 matrices M_i , with i ranging from 1 to 100, were utilized in self-organized maps clustering analyses, each containing 81,888
 123 unidentified particles with 19 elemental features (N, Na, Mg, Al, Si, P, S, Cl, K, Ca, Ti, Cr, Mn, Fe, Ni, Cu, Br, Ba, and Pb). The
 124 centroid matrix $C_{k,i,j}$ represents the clustering results obtained through the self-organized maps method with a given cluster

125 number k . The self-organized maps operation with the same k was repeated 100 times to ensure result robustness, where j
126 denotes the number of repetitions ranging from 1 to 100. Four indices, Silhouette index (Sil), Calinski–Harabasz index (CH),
127 modified Hartigan index (Hart), and Davies–Bouldin index (DB), were employed to determine the optimal parameters k , i , and j .
128 The matrix \mathbf{M}_i containing identified 81,888 particles was randomly divided into a training set (80 %) and a test set (20 %) for
129 random forest classification. The 10-fold cross-validation was utilized to determine the best tree. Abbreviations (corresponding to
130 Table 1): BJ - BeiJing; BS - BaiSe; FS - FuShun; GY - GuYuan; GYA - GuiYAng; YT - YanTai.

131

132 With reference to the studies of Ault et al. (2012) and Kirpes et al. (2018) and considering the results of elemental weight
133 ratios determined by EDX analysis, 19 elements (N, Na, Mg, Al, Si, P, S, Cl, K, Ca, Ti, Cr, Mn, Fe, Ni, Cu, Br, Ba, and Pb)
134 were selected to confirm the species of particles. C and O were not taken in account when clustering or classifying particles as
135 the membrane filters were made from cellulose acetate and cellulose nitrate, which contain C, H, N, and O. We could not detect
136 H because the ray-input window was made of beryllium. All particles showed high contents of C and O but different contents
137 of N, so N was retained as a feature of classification.

138 Species of aerosol particles vary with sampling location (Tao et al., 2017). Therefore, when establishing the matrices
139 of elemental weight ratios for clustering, equal amounts of data were randomly extracted from the sample data from each
140 province to ensure the inclusion of a consistent proportion of samples from each region in the training process. A hailstone FS
141 collected from Fushun City, Liaoning Province was shown to contain 13,648 insoluble particles, which was the smallest among
142 all samples from six provinces (Fig. 1). With random sampling of 13,648 particles from each province, the matrix used in
143 clustering analyses included 81,888 particles. This operation was repeated 100 times to obtain 100 matrices \mathbf{M}_i with i
144 ranging from 1 to 100.

145 Each matrix \mathbf{M}_i was clustered using the SOMs method. SOMs belong to the category of competitive learning algorithms
146 and are a type of artificial neural network (Kohonen, 1990). A basic SOMs network consists of an input layer, weight vectors,
147 and an output layer. Each neuron in the output layer possesses a set of weight vectors, which represent the topological structure
148 of the neurons in the output layer, associated with the inputs. SOMs are commonly used as dimensionality reduction algorithms,
149 enabling the representation of high-dimensional data in a lower-dimensional structure while preserving the original topology.
150 When SOMs are trained on unlabeled data for clustering purpose, it proves highly beneficial in clustering unlabeled and high-
151 dimensional inputs into visualized two-dimensional outputs.

152 We utilized the SOMs code from MATLAB's deep learning toolbox. The input of SOMs is \mathbf{M}_i . At begin, the neural
153 network in the output layer was initialized as 1-D dimension with k neurons. The number of neurons in the output layer
154 matches k ranging from 2 to 10. The operation of SOMs with the same initialized k neurons and input matrix \mathbf{M}_i was
155 repeated 100 times to ensure result robustness. The clustering result was stored in matrix $\mathbf{C}_{k,i,j}$, which corresponded to the

156 given k centroids in \mathbf{M}_i with j^{th} SOMs operation. Each $\mathbf{C}_{k,i,j}$ matrix consists of k rows and 19 columns (corresponding
157 to the number of elemental features). Four indices, namely, the Silhouette index (Rousseeuw, 1987), the Calinski–Harabasz
158 index (Calinski and Harabasz, 1974), the modified Hartigan index (Sibson and Hartigan, 1976), and the Davies–Bouldin index
159 (Davies and Bouldin, 1979), were selected as evaluation indicators to determine the parameters k , i and j . The Silhouette
160 index, Davies–Bouldin index, and Calinski–Harabasz index assess the similarity between a particle and others within the same
161 cluster, as well as the dissimilarity across different clusters for a given k . Hartigan index evaluates whether it is worthy to
162 increase the k . Notably, Hartigan index has undergone modifications that preserve its statistical meaning while conserving
163 computational resources.

164 Hartigan index (Sibson and Hartigan, 1976) is defined as:

$$165 \quad H(k) = (N - k - 1) \left[\frac{err(k)}{err(k+1)} - 1 \right], k = 2 \sim 10 \quad (1)$$

$$166 \quad err(k) = \sum_{g=1}^k \sum_{x_g \in C_g} (x_g - C_g)^2 \quad (2)$$

167

168 k : the number of clusters.

169 C : the centroid of all data

170 N : the number of observations in data

171 C_g : the centroid of cluster g

172 x_g : the observation of cluster g

173 x_n : the observation of data

174 The calculation of $H(k)$ requires clustering for values of k ranging from 2 to 11 in order to obtain $H(2)$, $H(3)$, ...,
175 $H(10)$. Clustering particles into 11 clusters would require performing an additional 10,000 iterations of the SOMs, with 100
176 iterations of extracting \mathbf{M}_i and 100 iterations of SOMs for each \mathbf{M}_i . Additionally, we observed that the SOMs did not perform
177 well in the Silhouette index (Sil), the Calinski–Harabasz index (CH), and the Davies–Bouldin index (DB) when $k = 2$. As a
178 result, we introduced modifications to the Hartigan index.

$$179 \quad Hart(k) = [N - (k - 1) - 1] \left[\frac{err(k-1)}{err(k)} - 1 \right], k = 2 \sim 10 \quad (3)$$

$$180 \quad err(k) = \sum_{g=1}^k \sum_{x \in C_g} (x - C_g)^2, k \geq 2 \quad (4)$$

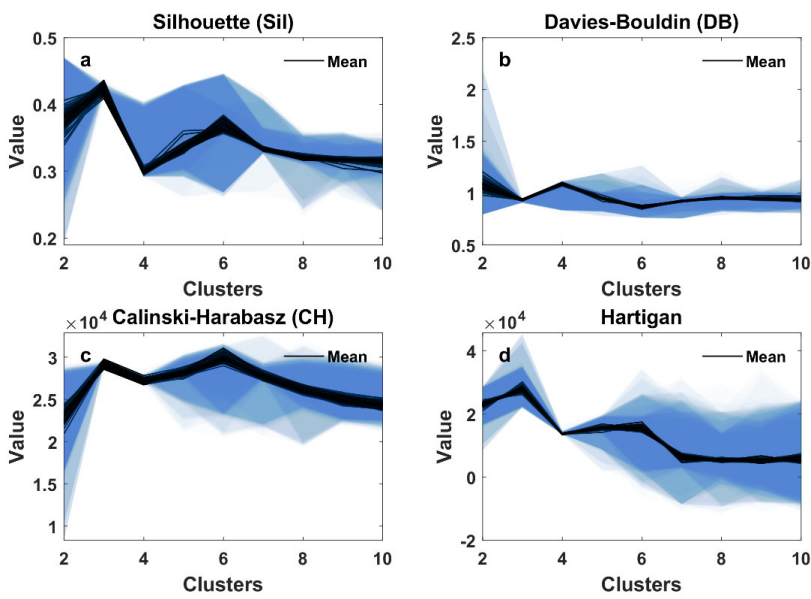
181

182 When $k = 1$, it indicates that all particles are belong to one cluster.

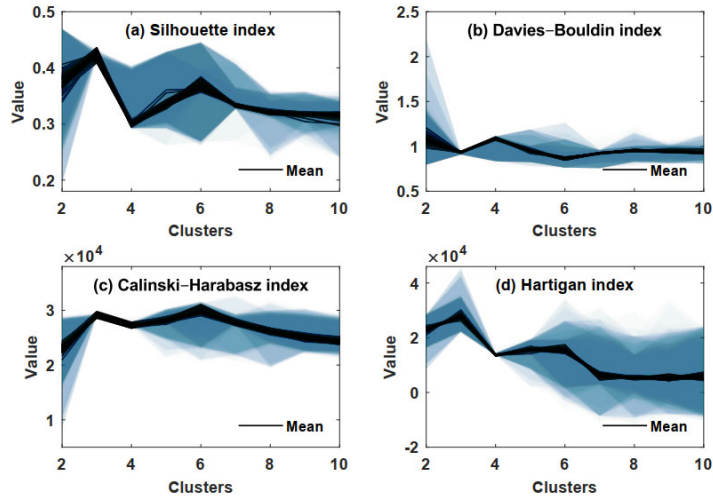
$$183 \quad err(1) = \sum_{n=1}^N (x_n - C)^2 \quad (5)$$

184 In clustering with a specific value of k , our objective is to have particles tightly grouped together in feature space while
 185 ensuring that the centroids exhibit a significant dispersion compared to $k - 1$. A higher value of $Hart(k)$ for a given k
 186 indicates improved clustering performance. The best k , i and j was chosen by combining the evaluation of the four indices
 187 (Fig. 4). We applied max normalization to rescale the four indices, $Sil(k)$, $CH(k)$, $DB(k)$, and $Hart(k)$. Subsequently, the
 188 best combination of k , i and j was determined, resulting in $\{Sil(k, i, j) + CH(k, i, j) + Hart(k, i, j) - DB(k, i, j)\}$
 189 reaching its maximum.

Formatted: Indent: First line: 2 ch



191



192

193 Fig. 4: Evaluation of self-organized maps clustering results. The clustering results of self-organized maps were evaluated using (a)
 194 Silhouette index, (b) Davies–Bouldin index, (c) Calinski–Harabasz index, and (d) Hartigan index. The self-organized maps operation
 195 was repeated 100 times to ensure result robustness. The solid lines and shading represent the average and spread of 100 repetitions,
 196 respectively.

197

198 The centroid matrix $C_{k,i,j}$ with best k , i and j was treated as a training set for random forest classification. The chosen
 199 centroid matrix $C_{k,i,j}$ with the top four elements is shown in Fig. 5 with $k = 6$. The first species with low elemental weight
 200 ratio except C and O contents was considered to be organics. The second species with high Fe content and low Cr content was
 201 introduced by the material of the slicer used in the experiment. The third species had a high Al content representing oxides or
 202 carbonates of aluminum. The fourth and fifth species were mineral silicates. So that, the third, fourth, and fifth species were
 203 referred to as “dust”. The last species with high N content was protein-containing biological aerosol.

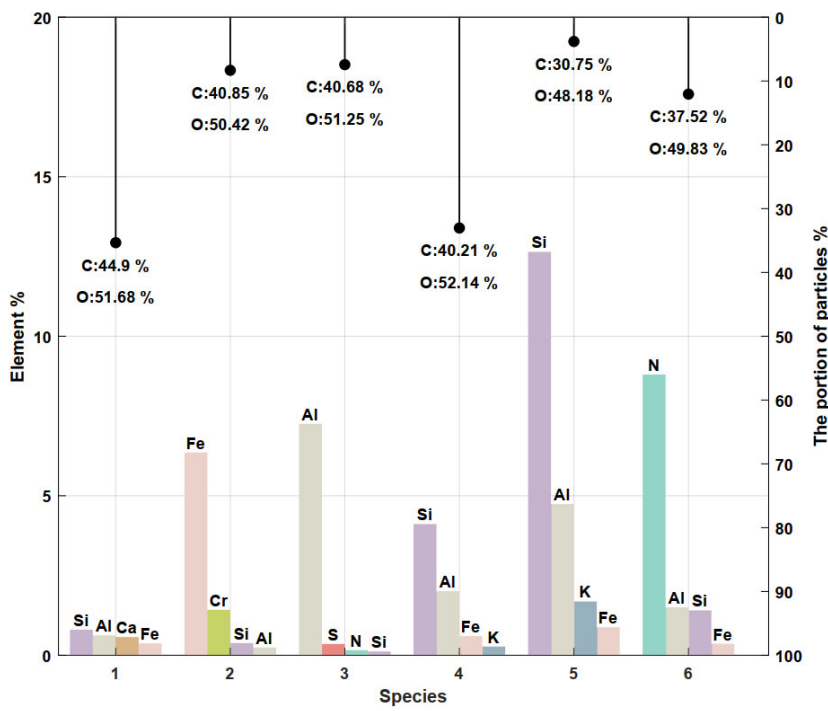
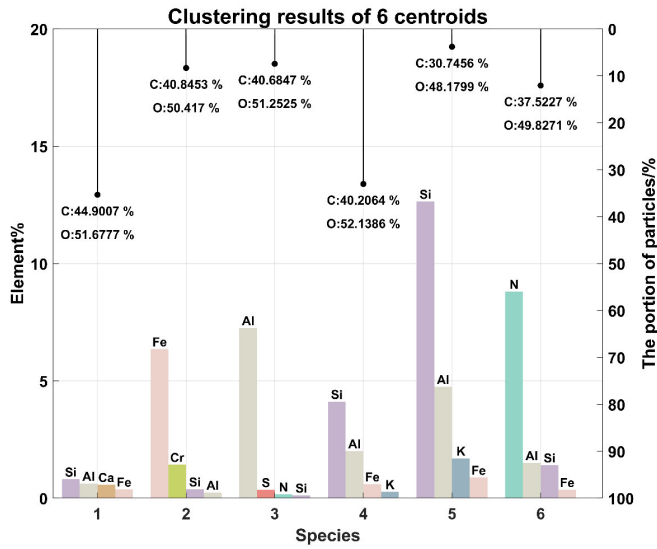


Fig. 5: Centroids of clustering with six clusters from self-organized maps results and each species portion. Colored bars show the

207 top four elements of each species. The stem bars show the portion of each species. The average contents of C and O of each species
 208 are marked at the end of the stem bars.

209
 210 The random forest method was applied in classifying insoluble particles, which involves randomly growing 100
 211 classification trees. The training set consisted of 80 % of M_1 and 10-fold stratified cross-validation was applied during the
 212 training process to find the best tree among the 100 random trees. The remaining 20 % particles of M_1 was used as the test set
 213 to evaluate the best tree. The confusion matrix of classification results are shown in Fig. 6. All remaining insoluble particles
 214 were classified by this tree. Finally, we identified three species: organics, dust, and bioprotein aerosols.

		Insoluble Particle Classification						Recall	
True type	1	937	7					99.3%	0.7%
	2	5	6076	1	28	5		99.4%	0.6%
	3		2	786			2	99.5%	0.5%
	4		41		4577	19		98.7%	1.3%
	5	1	11		7	2967		99.4%	0.6%
	6			2			904	99.8%	0.2%
		99.92%	99.39%	99.96%	99.42%	99.74%	99.98%		
Precision		99.4%	99.0%	99.6%	99.2%	99.2%	99.8%		
		0.6%	1.0%	0.4%	0.8%	0.8%	0.2%		
		1	2	3	4	5	6		
		Predicted type							

215

		Recall							
	1	4515	16		38			98.8%	1.2%
	2	15	2947	1	12			99.1%	0.9%
	3	1		833			2	99.6%	0.4%
	4	26	8	1	6100	9		99.3%	0.7%
	5			2	13	985		98.5%	1.5%
	6						854	100.0%	
		99.41%	99.68%	99.96%	99.35%	99.85%	99.99%		
Precision		99.1%	99.2%	99.5%	99.0%	99.1%	99.8%		
		0.9%	0.8%	0.5%	1.0%	0.9%	0.2%		
		1	2	3	4	5	6		
		Predicted species							

Fig. 6: Confusion matrix of the best random forest classifier tree. The numbers on the diagonal are accurately predicted insoluble particles. Numbers in bold indicate the accuracy of prediction of each type.

2.3 Conversion of insoluble particle number concentration

Particle number was converted to a number concentration per cubic centimeter volume water (hereinafter referred to as number concentration) using the following formula:

$$n_{liquid} \cdot V_{liquid} = N_{liquid} = N_{diluted} = n_{diluted} \cdot V_{diluted} \quad (6)$$

The number of insoluble particles in the melted shell solution (N_{liquid}) can be calculated by multiplying their number concentration (n_{liquid}) with the volume of the shell solution (V_{liquid}). Part of the solution was not used up in the experiments and was kept as a backup. Therefore, the shell solution was diluted in some experiments and part of the solution was consumed in the experiments. As in the melting solution, the number of insoluble particles in the diluted solution ($N_{diluted}$) can be calculated by multiplying their number concentration ($n_{diluted}$) with the volume of the diluted solution ($V_{diluted}$). The total particle number in the melted shell (N_{liquid}) remains unchanged during the dilution process ($N_{diluted}$).

$$n_{diluted} = n_{used} = \frac{N_{used}}{V_{used}} \quad (7)$$

231 The number concentration of the diluted solution ($n_{diluted}$) is equal to that of the consuming part (n_{used}). Assuming the
 232 rinsing operation ensures all insoluble particles in the shell were on the membrane, the number of insoluble particles in the
 233 consumed solution (N_{used}) is equal to the number of insoluble particles counted on the membrane (N_{filter}).

234 We use SEM to capture electron microscopy images of the membrane. Assuming a uniform distribution of insoluble
 235 particles on the filter membrane, a software randomly capture electron microscopy photos of the membrane and count the
 236 visible insoluble particles in those images. The relationship between total number of visible insoluble particles counted in the
 237 images (N_{count}) and N_{filter} is:

$$238 \quad \frac{S_{images}}{S_{filter}} = \frac{N_{count}}{N_{filter}} \quad (8)$$

239 That is, N_{filter} is determined by multiplying N_{count} by the ratio of the areas between the entire filter membrane (S_{filter})
 240 and the electron microscopy images (S_{images}). These three formulas Eq. (6-8) were reduced to Eq. (9):

$$241 \quad n_{liquid} = \frac{1}{V_{liquid}} \cdot \frac{S_{filter}}{S_{images}} \cdot \frac{V_{diluted}}{V_{used}} \cdot N_{count} \quad (9)$$

242 Here, S_{filter} , S_{images} , N_{count} , $V_{diluted}$, and V_{used} can be measured. The liquid volume (V_{liquid}) was determined as the
 243 average of readings obtained by two experimenters from the test tube. Take the logarithm on both sides:

$$244 \quad \ln n_{liquid} = -\ln V_{liquid} + \ln S_{filter} - \ln S_{images} + \ln V_{diluted} - \ln V_{used} + \ln N_{count} \quad (10)$$

245 Based on Eq. (10), a tiny change in n_{liquid} can be represented as dn_{liquid} :

$$246 \quad dn_{liquid} = n_{liquid} \cdot \left(-\frac{dV_{liquid}}{V_{liquid}} + \frac{dV_{diluted}}{V_{diluted}} - \frac{dV_{used}}{V_{used}} + \frac{dN_{count}}{N_{count}} \right) \quad (11)$$

247 As,

$$248 \quad dS_{filter} = dS_{images} = 0 \quad (12)$$

249 The uncertainty (Δ) of n_{liquid} comes from the measurement error of the experimental instruments, following below
 250 (Taylor, 1997):

$$251 \quad \Delta = n_{liquid} \cdot \sqrt{\left(\frac{dV_{liquid}}{V_{liquid}}\right)^2 + \left(\frac{dV_{diluted}}{V_{diluted}}\right)^2 + \left(\frac{dV_{used}}{V_{used}}\right)^2 + \left(\frac{dN_{count}}{N_{count}}\right)^2} \quad (13)$$

252 Here, the accuracy of the test tube is 0.1 mL. The term dV represents the greatest reading error caused by human and
 253 was set to 0.05 mL. The quantity $\frac{dN_{count}}{N_{count}}$ corresponds to the uncertainty associated with size of insoluble particles and the
 254 scan settings.

$$255 \quad \frac{dN_{count}}{N_{count}} = \frac{dPs}{Ps} = \frac{3}{6,340,608} \quad (14)$$

256 The term dPs represents the minimum number of pixels that can be detected in an image. Ps denotes the total number
 257 of pixels in the micrograph.

258 2.4 Curves fitting

259 We aggregated insoluble particles into 0.2- μm intervals (0.2 μm bin interval in Fig. 7 and Fig. 10, and 2 μm bin interval
260 in Fig. 8 and Fig. 9) to fit the logarithmic normal distribution:

$$261 \quad n(\ln D) = \frac{N}{\sqrt{2\pi} \ln \sigma_g} \cdot \exp\left[-\frac{(\ln D - \ln D_g)^2}{2 \ln^2 \sigma_g}\right] \quad (15)$$

262 N denotes the total number concentration of particles. Both $n(\ln D)$ and $n(D)$ represent the size distributions of
263 particles, where D is the diameter of insoluble particles. $n(\ln D)$ and $n(D)$ can be converted to each other by D .

$$264 \quad n(D) = \frac{1}{D} \cdot n(\ln D) \quad (16)$$

265

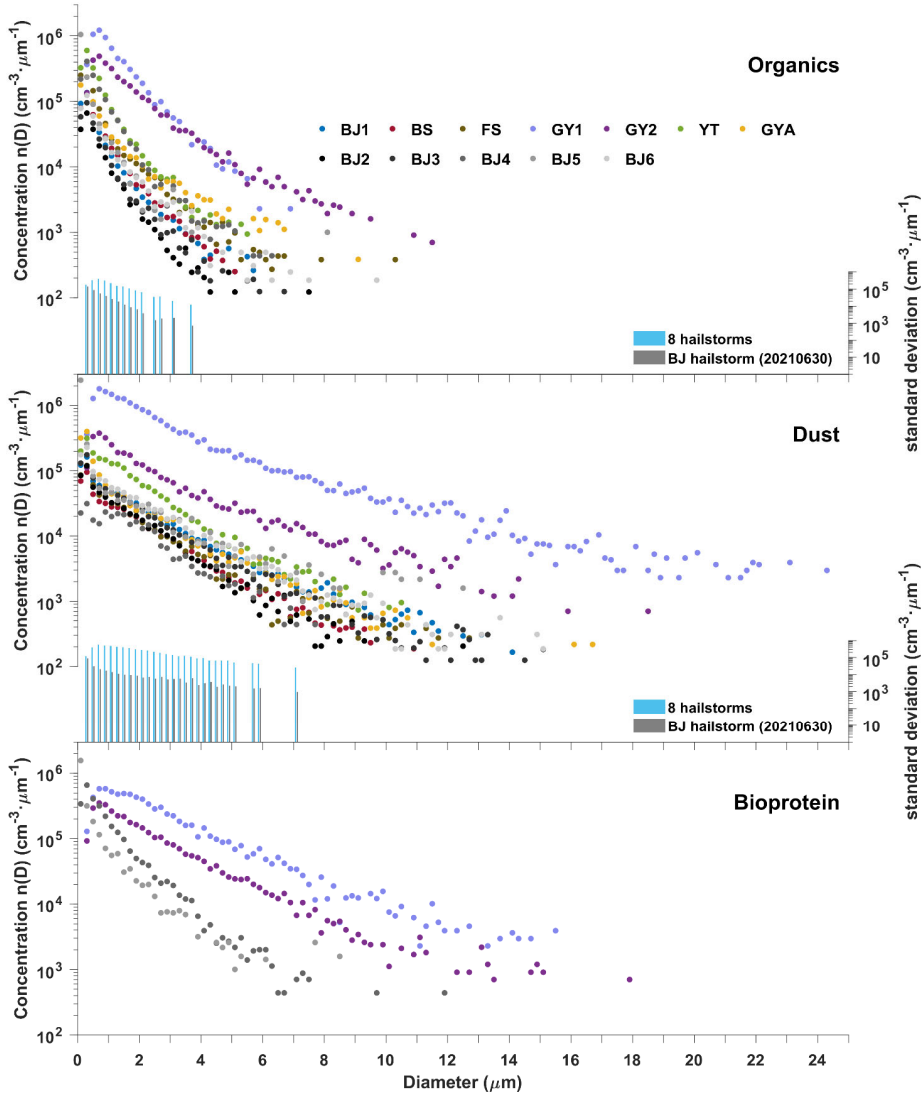
266 When the N_{count} in an interval equals 1, the number concentration will exhibit a flat tail due to the conversion to obtain
267 n_{liquid} . The fitting data were selected with intervals equals to 0.2 μm . The least squares method was applied to determine the
268 fitting parameters and R^2 was used to estimate the goodness of fit. The two centroids of fitting parameters of organics and dust
269 were determined by K-means method.

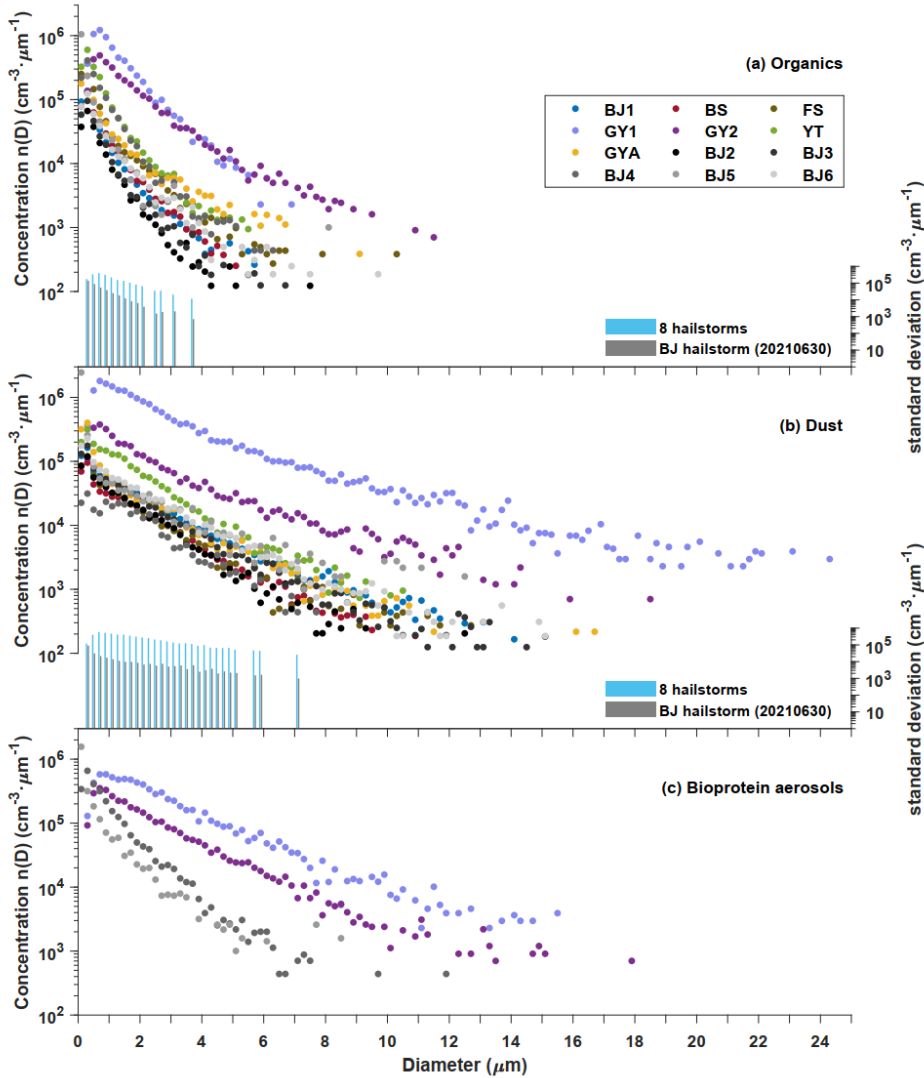
270 3 Results

271 A total of 289,461 insoluble particles were detected from 30 shells of 12 hailstones using SEM. The identification of
272 insoluble particles employed SOMs for clustering and random forest for classification. Four indices were utilized to determine
273 the appropriate parameters in clustering. The clustering results ($C_{k,l}$) were divided into a training and a testing set for
274 classification. The confusion matrix of the best classifier showed an accuracy, precision, and recall of 99.7 %, 99.4-3 %, and
275 99.25 %, respectively. All particles were classified as organics, dust, and bioprotein aerosols (i.e., the fraction of biological
276 aerosols with protein content).

277 3.1 Sample similarity

278 Five of the 12 hailstones (BJ2–BJ6) were from the same hailstorm that occurred in Beijing on June 30, 2021. The insoluble
279 particles present in BJ2–BJ6 showed similarity in the size distribution of organics, dust, and bioprotein aerosols, while those
280 from 8 hailstones (BJ1, BJ2, BS, FS, GY1, GY2, YT and GYA) exhibited a wider dispersion (Fig. 7). The results were similar
281 to those of Li et al., who reported that the number concentrations of water-soluble ions varied among hailstorm events but
282 showed similarity in the same storm (Li et al., 2018). These analyses suggested that insoluble particles in the hailstorm may
283 come from local natural or anthropogenic emissions (e.g., soil dust, aerosols from biomass and fossil fuel combustion, products
284 of the conversion of gaseous precursors), which is also suggested by the results on water-soluble ions (Beal et al., 2022). The
285 updraft within the hailstorm is likely to bring insoluble particles from local surfaces or boundary layers into deep convective
286 clouds, as hailstorms are among the most severe storms with strong updrafts (Battaglia et al., 2022).





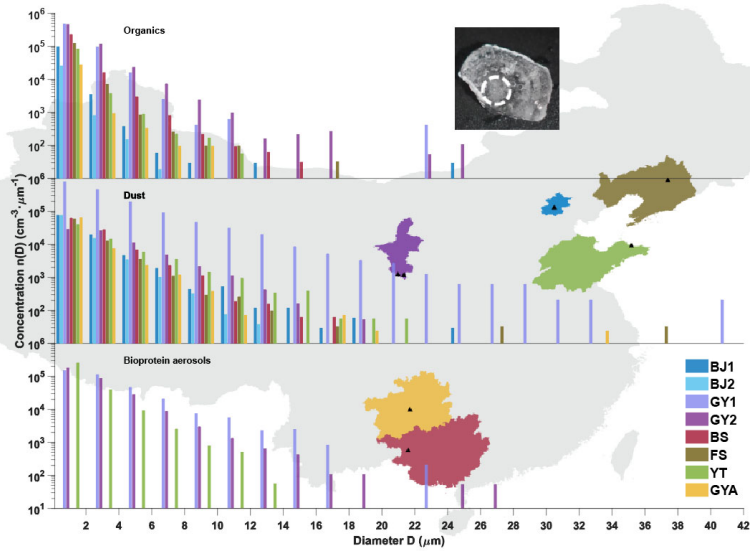
288
 289 Fig. 7: Size distribution of (a) organics, (b) dust, and (c) bioprotein aerosols of insoluble particles in 12 hailstones. The colored dots
 290 represent data from 7 hailstones BJ1, BS, FS, GY1, GY2, YT and GYA which were from different hailstorms. The black and gray
 291 dots correspond to data from hailstones (BJ2 to BJ6) that were from the same hailstorm occurring in Beijing on June 30, 2021. The
 292 blue and gray bars indicate the standard deviation of number concentration of insoluble particles from 8 hailstones (BJ1, BJ2, BS,
 293 FS, GY1, GY2, YT and GYA) from 8 cases and 5 hailstones (BJ2 to BJ6) from one case, respectively. Abbreviations (corresponding
 294 to Table 1): BJ - BeiJing; BS - BaiSe; FS - FuShun; GY - GuYuan; GYA - GuiYAng; YT - YanTai.

Formatted: Centered

295 **3.2 Size distribution in embryos**

296 All hailstone embryos analyzed in this study are graupel particles, which grows from the initial ice particles through
297 accretion of supercooled droplets (Knight, 1981). These initial ice particles are formed through nucleation of insoluble particles
298 where heterogeneous nucleation take place (Lamb and Verlinde, 2011). In other words, insoluble particles in graupels influence
299 the formation of ice crystals and subsequently affect the formation of hailstone embryos.

800



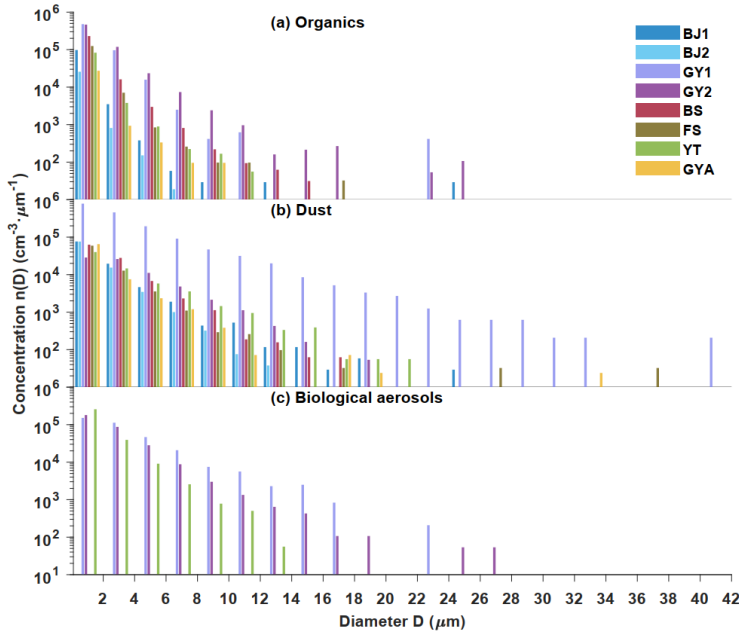


Fig. 8: Size distribution of (a) organics, (b) dust, and (c) bioprotein aerosols –insoluble particles in hailstone embryos. Colors represent different hailstones. Different colors represent the provinces in China from which the hailstones were collected. Black triangles indicate the locations of hailstone sample collection. The white dashed circle highlights part of the hailstone embryo.

Abbreviations (corresponding to Table 1): BJ - BeiJing; BS - BaiSe; FS - FuShun; GY - GuYuan; GYA - GuiYAng; YT - YanTai.

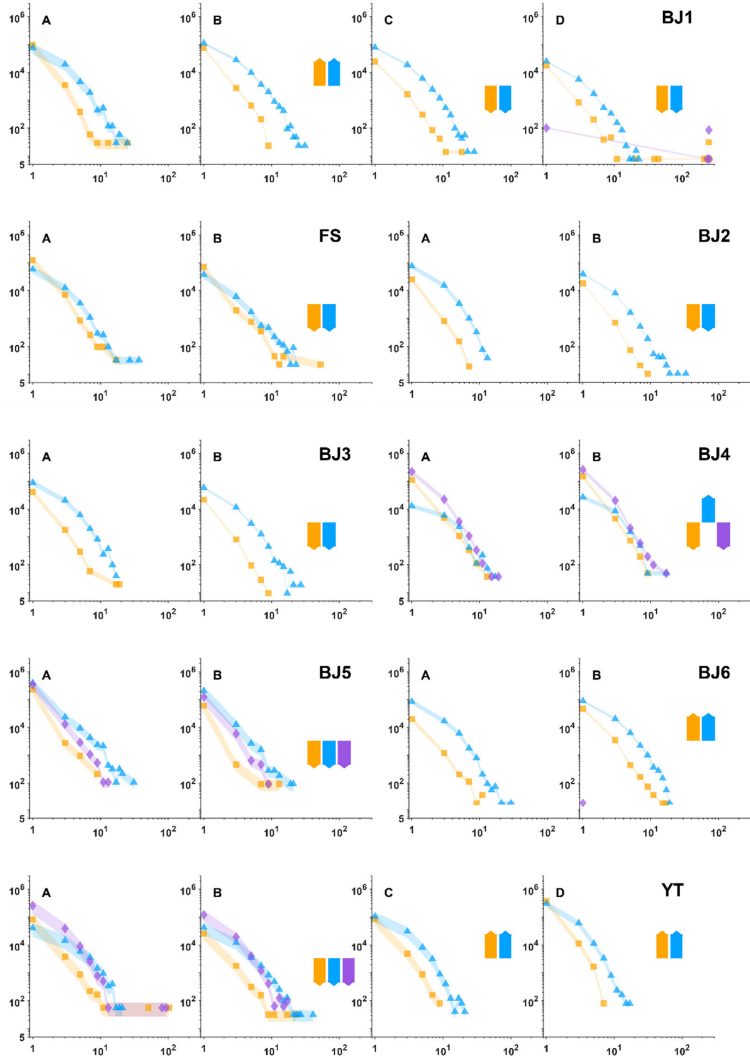
The variations in number concentrations of dust and bioprotein insoluble particles indicate that particle number concentrations decrease exponentially with particle diameter, with markble variation observed among hailstorms (Fig. 8). BJ2 was selected to represent five hailstones from the same storm to simplify comparison. The size distribution distinguishes organics from dust and bioprotein aerosols. The number concentrations of organics from all samples decrease with particle diameter less than 8 μm , while those of GY1 and GY2 fluctuate starting at diameters of 8 μm and 12 μm , respectively. Compared to other hailstones, GY1 and GY2 were collected in remote areas, where is fields of rural areas dedicated to growing crops near the south of the Gobi Desert. Therefore, GY1 and GY2 have a coarse mode of organics with particle diameters larger than 12 μm , possibly might due to the emission of spring-wheat straw burning and unrestricted diesel engine vehicles. The transport of coal combustion in surrounding cities may also contribute to the coarse mode organics. Among all cases, there is a significant variance in the size distribution of both organics and dust. The number concentration of organics from a hailstone embryo varied from 1 to 390 times, compared to those at the same particle diameter in hailstone embryos from

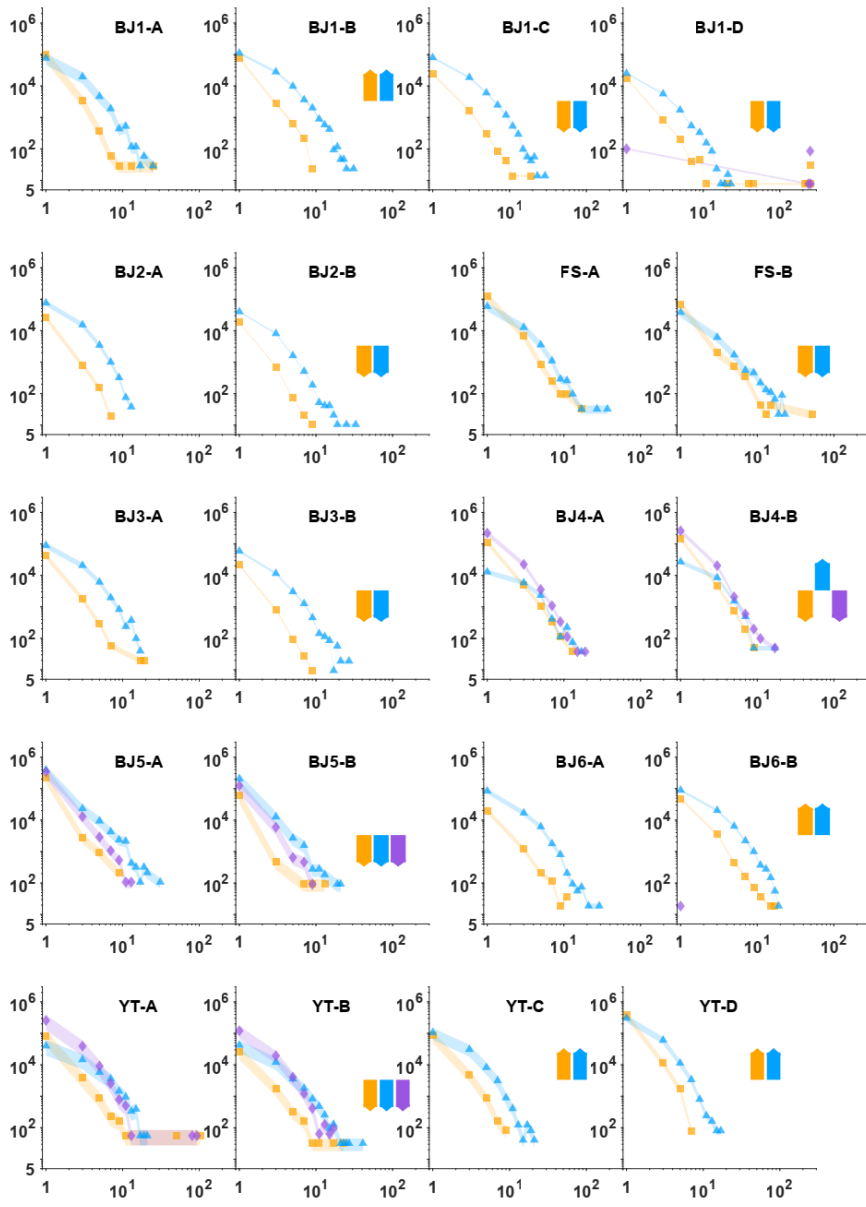
318 different cases. The number concentration of dust from a hailstone embryo varied from 1 to 527 times, compared to those at
319 the same particle diameter in hailstone embryos from different cases. The number concentrations of dust from BJ1, BJ2, and
320 GY1 are at least 3 times higher than organics in particles of the same diameter in the range of 2–24 μm .

321 Moreover, dust showed a wider size distribution than organics and bioproteins among all samples. Dust from GY1 had a
322 higher number concentration and larger maximum size (42 μm) compared to other hailstone embryos. Hailstone samples with
323 high insoluble particle content, i.e., GY1 and GY2, showed significantly lower total column water vapor values and smaller
324 depth between freezing level height and orography within one hour before hailstorm occurrence, compared to other hailstones
325 (Table 1). The competition of condensation and relative shorter updraft pathway might be responsible for the high number
326 concentrations of organics, dust, and bioproteins in GY1 and GY2, as compared with other hailstones. Bioprotein aerosols,
327 with high freezing efficiency, may have formed initial ice particles in GY1, GY2, and YT, while dust or organics formed initial
328 ice particle in hailstorms in the other five cases. All hailstone embryos contained organics and dust, but not all hailstone
329 embryos contained a significant amount of bioprotein aerosols. Due to limited comprehension of the transportation and
330 transformation processes of biological materials, it is challenging that to establish a definitive relationship between biological
331 protein particles and biological aerosols (Fröhlich-Nowoisky et al., 2016).

332 3.3 Size distribution in shells

333 Size distribution of each species varied little in characteristics between outer shells with the embryos (Fig. 9). In a four-
334 shell hailstone, the number concentrations of insoluble particles exhibited V-shaped distributions (BS and YT) or inverse V-
335 shaped distributions (BJ1) from embryo to crust. Five of nine two-shell hailstones showed higher number concentrations of
336 dust in crusts than embryos, while seven of them showed higher number concentrations of organics in embryos than crusts.
337 Moreover, the quantification of differences in number concentration varied little among shells. The 90.5 % points showed that
338 differences in number concentration of the same kind particles in a shell compared to the previous shell at the same diameter
339 was within twice (294 data points in Fig. 9). This observation is attributed to the fact that the growth of hailstones beyond the
340 embryo stage relies on the accretion of supercooled water rather than ice crystals (Lamb and Verlinde, 2011). Consequently,
341 the hailstone recorded not only insoluble particles during the embryo formation, but also insoluble particle in the hailstone
342 growth zone throughout the hailstorm. As a result, the size distribution of particles within the entire hailstones may represent
343 the distribution of insoluble particles in deep convection regions where the hailstones went through.

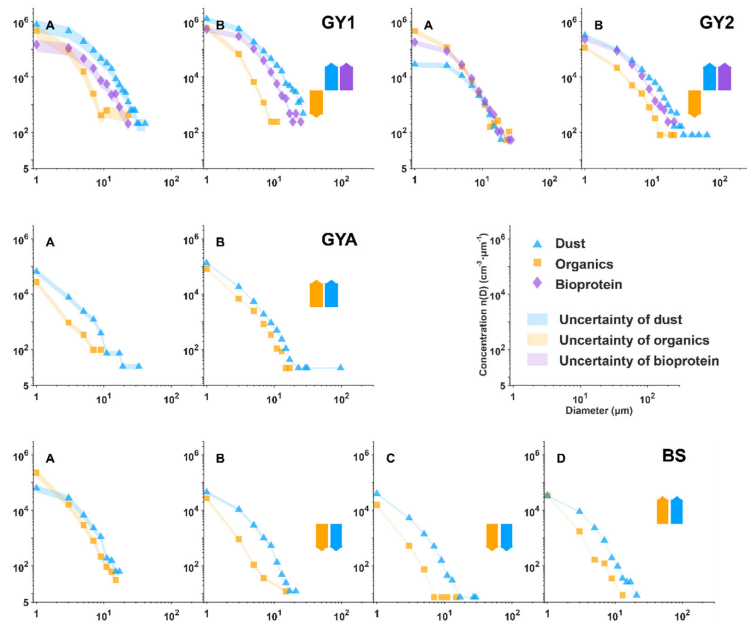




345
 346 Fig. 9: Size distribution of insoluble particles within the natural shells of 12 hailstones is represented. Blue triangles, orange squares,
 347 and purple diamonds are used to indicate dust, organics, and bioprotein aerosols, respectively. The natural shells are denoted
 348 alphabetically with capital letters (shell A refers to embryos, and shell B/D refers to the crust of hailstones). The arrow direction
 349 illustrates the tendency of particle number concentration in each layer compared to the previous shell. Shading is employed to

350 indicate uncertainty. Detailed calculations are provided in the supplementary information. Abbreviations (corresponding to Table
351 1): BJ - BeiJing; BS - BaiSe; FS - FuShun; GY - GuYuan; GYA - GuiYAng; YT - YanTai.

352



353

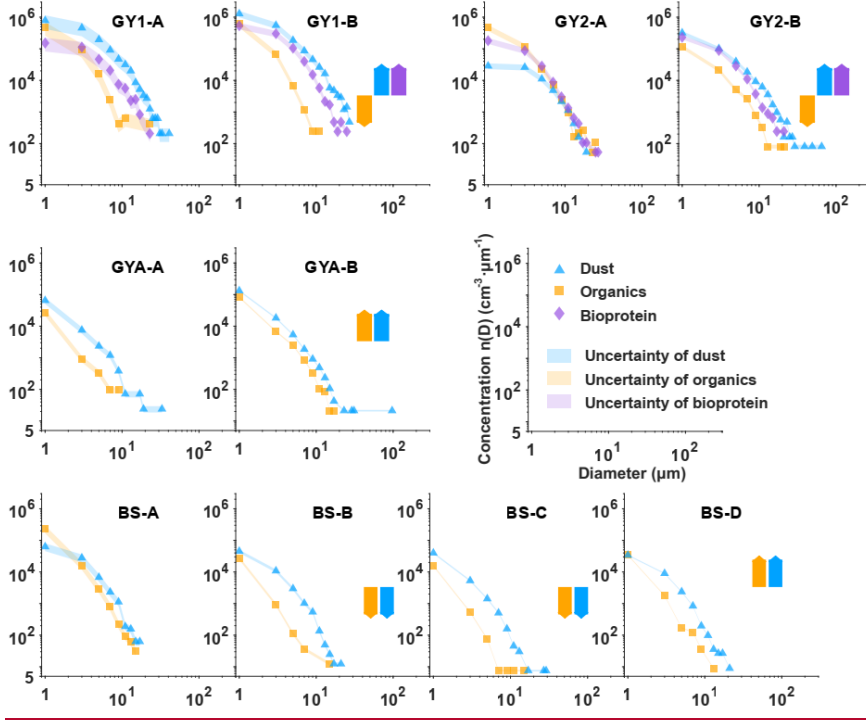


Fig. 9_b is a continuation of Fig. 9a continued.

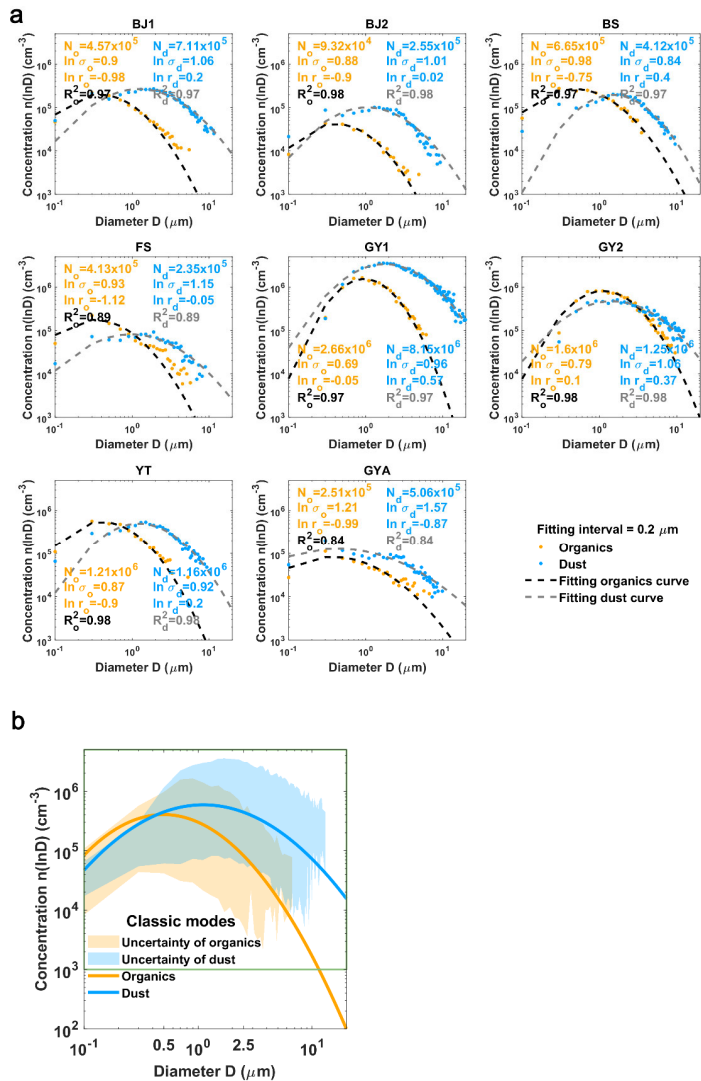
3.4 Logarithmic normal distribution of dust and organics

The size distributions of dust and organics in the whole hailstone can be described by a logarithmic normal distribution (Fig. 10a) (Lamb and Verlinde, 2011):

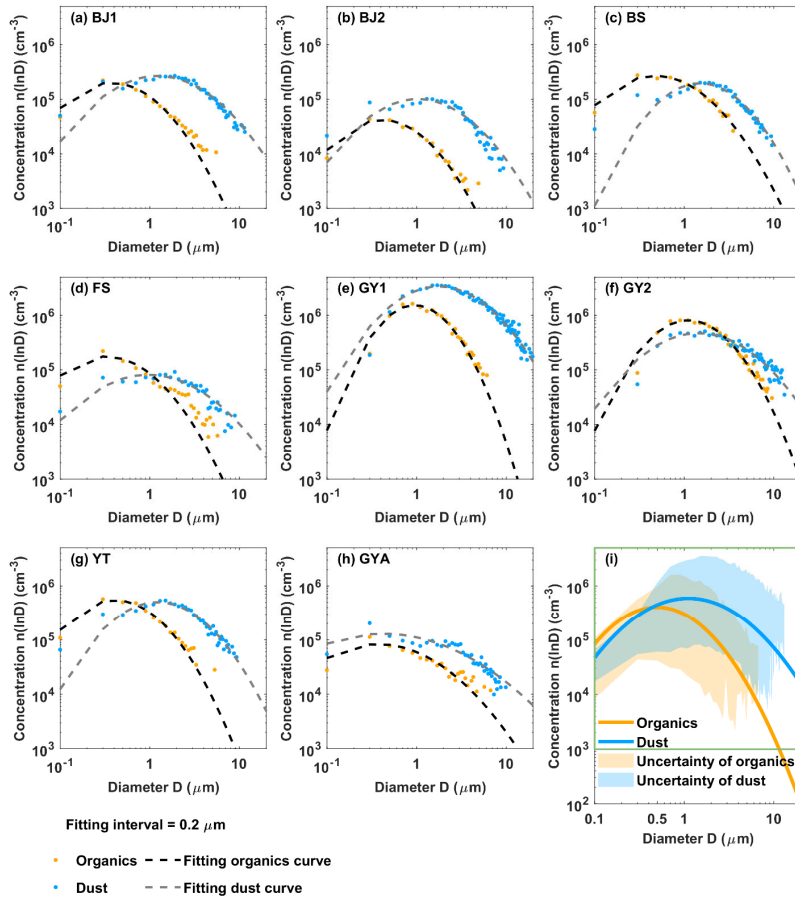
$$n(\ln D) = \frac{N}{\sqrt{2\pi} \ln \sigma_g} \cdot \exp \left[-\frac{(\ln D - \ln D_g)^2}{2 \ln^2 \sigma_g} \right], (D > 0.2 \mu\text{m}) \quad (17)$$

Where $n(\ln D)$ is the number concentration of insoluble particles per cubic centimeter volume water ranging from $\ln D - \frac{1}{2} \ln \sigma_g$ to $\ln D + \frac{1}{2} \ln \sigma_g$. Here, D represents the diameter of particles (in micrometers), $\ln D_g$ is the geometric mean diameter, and $\ln \sigma_g$ is the geometric standard deviation (Lamb and Verlinde, 2011). The number of bioprotein aerosols was below the limit of detection in some samples, so that, only the curves of organics and dust were fitted. The fitting parameters of the same species were aggregated in parameter space, and were suspected to be related to the physical properties of each species, requiring further studies for confirmation. Moreover, the fitting parameters of organics and dust particles were clustered into two centroids (Fig. 10b) by the K-means method, which indicated that organics and dust have two classic modes

368 (classic mode of organics: $\ln D_o = -0.70$, $\ln \sigma_o = 0.91$, and $N_o = 9.19 \times 10^5 \text{ cm}^{-3}$; classic mode of dust: $\ln D_d = 0.11$, $\ln \sigma_d$
 369 $= 1.07$, and $N_d = 1.59 \times 10^6 \text{ cm}^{-3}$). That is, insoluble organics in hailstones are usually smaller in diameter and present in lower
 370 amounts than dust. Regardless of fine or coarse particles ($D < 0.5 \mu\text{m}$ in diameter were not considered in reference to DeMott
 371 et al. (DeMott et al., 2010)), the number concentration of dust was up to 2 orders of magnitude higher than the number
 372 concentration of organics. These observations indicated that dust accounted for the major portion of particles in eight
 373 hailstorms (no considering about bioprotein), which was consistent with the observations of embryos described above.



374



375
 376
 377 Fig. 10: Fitting size distribution functions of organics and dust contained in the whole hailstone. (a)-(h) Fitting parameters of
 378 logarithmic normal distributions of BJ1, BJ2, BS, FS, GY1, GY2, YT, GYA. (bi) Classic modes of dust and organics (interval of data
 379 is 0.2 μm and fitting curves painted with interval of 0.02 μm). The fitting parameters for subfigures (a)-(h) are listed in Table 2. The
 380 fitting range of (a)-(h) is shown with a green rectangle in (i). The centroid of the organics fitting parameter (orange line) is $\ln \sigma_o =$
 381 0.91 , $\ln D_o = -0.70$, and $N_o = 9.19 \times 10^5 \text{ cm}^{-3}$. The centroid of the dust fitting parameter (blue line) is $\ln \sigma_d = 1.07$, $\ln D_d = 0.11$,
 382 and $N_d = 1.59 \times 10^6 \text{ cm}^{-3}$. Shading showed uncertainty of organics and dust. Abbreviations (corresponding to Table 1): BJ - BeiJing;
 383 BS - BaiSe; FS - FuShun; GY - GuYuan; GYA - GuiYAng; YT - YanTai.

385 **Table 2: The fitting parameters of dust and organics size distribution in Fig. 10 (a)-(h).**

Sample	N_o (cm^{-3})	$\ln D_o$	$\ln \sigma_o$	R_o^2	N_d (cm^{-3})	$\ln D_d$	$\ln \sigma_d$	R_d^2
BJ1	4.57×10^5	-0.98	0.90	0.97	7.11×10^5	0.20	1.06	0.93
BJ2	9.32×10^4	-0.90	0.88	0.98	2.55×10^5	0.02	1.01	0.89
BS	6.65×10^5	-0.75	0.98	0.97	4.12×10^5	0.40	0.84	0.91
FS	4.13×10^5	-1.12	0.93	0.89	2.35×10^5	-0.05	1.15	0.87
GY1	2.66×10^6	-0.05	0.69	0.97	8.15×10^6	0.57	0.96	0.98
GY2	1.60×10^6	0.10	0.79	0.98	1.25×10^6	0.37	1.06	0.95
YT	1.21×10^6	-0.90	0.87	0.98	1.16×10^6	0.20	0.92	0.94
GYA	2.51×10^5	-0.99	1.21	0.84	5.06×10^5	-0.87	1.57	0.79

Formatted: Line spacing: 1.5 lines, Tab stops: 40 ch, Left

Formatted Table

386

387 4 Conclusions

388 This was the first study to simultaneously analyze both the number concentrations and species (including organics, dust
389 and bioproteins) of insoluble particles in hailstones. The findings from this analysis offer valuable insights into particle
390 observations within severe storms. Understanding the number concentration and composition of these insoluble particles is
391 crucial, as they play a significant role as ice-nucleating particles during the heterogeneous nucleation process in deep
392 convection.

393 The size distribution of insoluble particles in hailstones from the same hailstorm showed less variation than those from
394 different hailstorms. One possible reason is that updrafts of hailstorms brought insoluble particles from local surfaces or
395 boundary layers into deep convective clouds. Moreover, almost all insoluble particles in hailstone embryos analyzed in this
396 study showed an exponential size distribution, which was consistent with the effects of gravity. The number concentrations of
397 organics and dust from different hailstone embryos differed up to 389 times and 526 times at the same diameter, respectively.
398 The changes in particle concentration may lead to at least one-order-of-magnitude variance in ice-nucleating particle (DeMott
399 et al., 2010). Additionally, size distribution of insoluble particles varied in shells up to 27 times, which was much smaller than
400 differences with different hailstorms.

401 Two logarithmic normal distribution models were applied to fit the size distribution of organics and dust within hailstones,
402 providing a description of insoluble particles in the deep convection during hailstone formation. The analysis of the two classic
403 size distribution modes of insoluble particles indicated a significant presence of dust, without considering bioprotein.
404 Furthermore, a positive correlation exists between the number concentrations of insoluble particles and that of ice-nucleating
405 particles in hailstones, specifically for corresponding species (Ren et al., 2023, submitted, figure not shown). A further

406 measurement of ice-nucleating particles by drop-freezing experiments will establish the relationship between insoluble
407 particles and immersion ice-nucleating particles. Combination of these results with future experiments to determine the number
408 concentrations and species of particles from local observations will establish the relationship between surface observation and
409 ice-nucleating particles in deep convective clouds, which will lead to improvement of the parameterization of ice-nucleating
410 particles in both weather and climate models.

411 Nonetheless, two kinds of classic size distribution modes of organics and dust in hailstones were performed, but a more
412 robust classic mode required a larger number of samples. In future, for climate or weather models, the classic mode can be
413 assumed as the mean state to describe the characteristics of insoluble particles in supercooling water. In addition, this study
414 did not attempt to parameterize bioprotein aerosols, because there was a great uncertainty in quantification due to poor
415 understanding of biological processes (Fröhlich-Nowoisky et al., 2016). Further collaborative studies are required to gain a
416 better understanding of biological processes to establish the classic bioprotein mode.

417 **Code availability**

418 Self-organized maps algorithm is functions on MATLAB

419 <https://ww2.mathworks.cn/help/deeplearning/ref/selforgmap.html>

420 Random forest algorithm is functions on MATLAB

421 https://ww2.mathworks.cn/help/stats/treebagger.html?searchHighlight=TreeBagger&s_tid=srchtitle_TreeBagger_1

422 The 10-fold stratified cross-validation algorithm is functions on MATLAB

423 https://ww2.mathworks.cn/help/stats/cvpartition.html?searchHighlight=cvpartition&s_tid=srchtitle_cvpartition_1

424 Identification algorithms are coded on MATLAB and will be made available on request.

425 **Data availability**

426 Data will be made available on request.

427 **Author contributions**

428 Haifan Zhang wrote the original draft under the concept presented by Qinghong Zhang. Haifan Zhang, Xiangyu Lin and
429 Chan-Pang Ng participated in preprocess and reservation of hailstones from volunteers. Haifan Zhang and Xiangyu Lin sliced
430 hailstones using machine manufactured by Kai Bi and performed the experiments on analyzing element weight ratio of
431 insoluble particles with help of Li Chen. Kai Bi also provided hailstones BJ2 ~ BJ6. Machine learning on identification of
432 particles is operated by Haifan Zhang. Yangze Ren and Huiwen Xue compared ice nucleation particles from drop-freezing

433 experiments with our data. Zhuolin Chang provided hailstones GY1 and GY2. All authors discussed and contributed to the
434 final manuscript. Qinghong Zhang directed this project.

435 **Competing interests**

436 The authors declare no competing interests.

437

438 **Acknowledgments**

439 This study was supported by the National Natural Science Foundation of China (Grant Nos. 42030607 and 41930968),
440 the Innovation Project of the China Meteorological Administration (Grant No. CXFZ2021J038) and the Key R & D projects
441 in Ningxia Hui Autonomous Region (2022BEG02010). The authors thank Cai Yao from the Meteorological Bureau of
442 Guangxi, China in collecting hailstones BS in Guangxi. The authors thank volunteers in collecting hailstones. The authors
443 thank Prof. Jiwen Fan from Pacific Northwest National Laboratory of the United States for discussions.

444 **References:**

445 Aztec User Manual:

446 <https://utw10193.utweb.utexas.edu/InstrumentManuals/Oxford%20EDS%20AZtec%20User%20Manual.pdf>, last access:
447 22 August 2023.

448 Battaglia, A., Mroz, K., and Cecil, D.: Satellite hail detection, in: *Precipitation Science*, Elsevier, 257–286,

449 <https://doi.org/10.1016/B978-0-12-822973-6.00006-8>, 2022.

450 Beal, A., Martins, J. A., Rudke, A. P., de Almeida, D. S., da Silva, I., Sobrinho, O. M., de Fátima Andrade, M., Tarley, C. R.

451 T., and Martins, L. D.: Chemical characterization of PM_{2.5} from region highly impacted by hailstorms in South America,

452 *Environ. Sci. Pollut. Res.*, 29, 5840–5851, <https://doi.org/10.1007/s11356-021-15952-6>, 2022.

453 Calinski, T. and Harabasz, J.: A dendrite method for cluster analysis, *Commun. Stat. - Theory Methods*, 3, 1–27,

454 <https://doi.org/10.1080/03610927408827101>, 1974.

455 Chen, Q., Yin, Y., Jiang, H., Chu, Z., Xue, L., Shi, R., Zhang, X., and Chen, J.: The Roles of Mineral Dust as Cloud

456 Condensation Nuclei and Ice Nuclei During the Evolution of a Hail Storm, *J. Geophys. Res. Atmos.*, 124, 14262–14284,

457 <https://doi.org/10.1029/2019JD031403>, 2019.

458 Davies, D. L. and Bouldin, D. W.: A Cluster Separation Measure, *IEEE Trans. Pattern Anal. Mach. Intell.*, PAMI-1, 224–

459 227, <https://doi.org/10.1109/TPAMI.1979.4766909>, 1979.

460 DeMott, P. J., Prenni, A. J., Liu, X., Kreidenweis, S. M., Petters, M. D., Twohy, C. H., Richardson, M. S., Eidhammer, T.,

461 and Rogers, D. C.: Predicting global atmospheric ice nuclei distributions and their impacts on climate, *Proc. Natl. Acad.*

462 *Sci.*, 107, 11217–11222, <https://doi.org/10.1073/pnas.0910818107>, 2010.

463 DeMott, P. J., Prenni, A. J., McMeeking, G. R., Sullivan, R. C., Petters, M. D., Tobo, Y., Niemand, M., Möhler, O., Snider,

464 J. R., Wang, Z., and Kreidenweis, S. M.: Integrating laboratory and field data to quantify the immersion freezing ice

465 nucleation activity of mineral dust particles, *Atmos. Chem. Phys.*, 15, 393–409, <https://doi.org/10.5194/acp-15-393-2015>,

466 2015.

467 Fröhlich-Nowoisky, J., Kampf, C. J., Weber, B., Huffman, J. A., Pöhlker, C., Andreae, M. O., Lang-Yona, N., Burrows, S.

468 M., Gunthe, S. S., Elbert, W., Su, H., Hoor, P., Thines, E., Hoffmann, T., Després, V. R., and Pöschl, U.: Bioaerosols in

469 the Earth system: Climate, health, and ecosystem interactions, *Atmos. Res.*, 182, 346–376,

470 <https://doi.org/10.1016/j.atmosres.2016.07.018>, 2016.

471 Hersbach, H., Bell, B., Berrisford, P., Biavati, G., Horányi, A., Muñoz Sabater, J., Nicolas, J., Peubey, C., Radu, R., Rozum,

472 I., Schepers, D., Simmons, A., Soci, C., Dee, D., and Thépaut, J.-N.: ERA5 hourly data on single levels from 1959 to

473 present, Copernicus Clim. Chang. Serv. Clim. Data Store (CDS).[data set],

474 <https://doi.org/https://doi.org/10.24381/cds.adbb2d47>, 2018.

475 Hoose, C. and Möhler, O.: Heterogeneous ice nucleation on atmospheric aerosols: a review of results from laboratory
476 experiments, *Atmos. Chem. Phys.*, 12, 9817–9854, <https://doi.org/10.5194/acp-12-9817-2012>, 2012.

477 Knight, N. C.: The Climatology of Hailstone Embryos, *J. Appl. Meteorol.*, 20, 750–755, <https://doi.org/10.1175/1520->
478 0450(1981)020<0750:TCOHE>2.0.CO;2, 1981.

479 Kohonen, T.: The self-organizing map, *Proc. IEEE*, 78, 1464–1480, <https://doi.org/10.1109/5.58325>, 1990.

480 Lamb, D. and Verlinde, J.: *Physics and Chemistry of Clouds*, First edi., Cambridge University Press, Cambridge,
481 <https://doi.org/10.1017/CBO9780511976377>, 2011.

482 Li, X., Zhang, Q., Zhu, T., Li, Z., Lin, J., and Zou, T.: Water-soluble ions in hailstones in northern and southwestern China,
483 *Sci. Bull.*, 63, 1177–1179, <https://doi.org/10.1016/j.scib.2018.07.021>, 2018.

484 Li, X., Zhang, Q., Zhou, L., and An, Y.: Chemical composition of a hailstone: evidence for tracking hailstone trajectory in
485 deep convection, *Sci. Bull.*, 65, 1337–1339, <https://doi.org/10.1016/j.scib.2020.04.034>, 2020.

486 Michaud, A. B., Dore, J. E., Leslie, D., Lyons, W. B., Sands, D. C., and Priscu, J. C.: Biological ice nucleation initiates
487 hailstone formation, *J. Geophys. Res. Atmos.*, 119, 12,186–12,197, <https://doi.org/10.1002/2014JD022004>, 2014.

488 Prenni, A. J., Demott, P. J., Rogers, D. C., Kreidenweis, S. M., Mcfarquhar, G. M., Zhang, G., and Poellot, M. R.: Ice nuclei
489 characteristics from M-PACE and their relation to ice formation in clouds, *Tellus B*, 61, 436–448,
490 <https://doi.org/10.1111/j.1600-0889.2009.00415.x>, 2009.

491 Rogers, D. C., DeMott, P. J., Kreidenweis, S. M., and Chen, Y.: A Continuous-Flow Diffusion Chamber for Airborne
492 Measurements of Ice Nuclei, *J. Atmos. Ocean. Technol.*, 18, 725–741, <https://doi.org/10.1175/1520->
493 0426(2001)018<0725:ACFDCF>2.0.CO;2, 2001.

494 Rosinski, J.: Solid Water-Insoluble Particles in Hailstones and Their Geophysical Significance, *J. Appl. Meteorol.*, 5, 481–
495 492, [https://doi.org/10.1175/1520-0450\(1966\)005<0481:SWIPIH>2.0.CO;2](https://doi.org/10.1175/1520-0450(1966)005<0481:SWIPIH>2.0.CO;2), 1966.

496 Rousseeuw, P. J.: Silhouettes: A graphical aid to the interpretation and validation of cluster analysis, *J. Comput. Appl.*
497 *Math.*, 20, 53–65, [https://doi.org/10.1016/0377-0427\(87\)90125-7](https://doi.org/10.1016/0377-0427(87)90125-7), 1987.

498 Sibson, R. and Hartigan, J. A.: Clustering Algorithms., *Appl. Stat.*, 25, 70, <https://doi.org/10.2307/2346526>, 1976.

499 Tao, J., Zhang, L., Cao, J., and Zhang, R.: A review of current knowledge concerning PM2.5 chemical composition, aerosol
500 optical properties and their relationships across China, *Atmos. Chem. Phys.*, 17, 9485–9518, <https://doi.org/10.5194/acp->
501 17-9485-2017, 2017.

502 Taylor, J. R.: *An Introduction to Error Analysis*, Second edi., University Science Books, 330 pp., 93570275X, 1997.

503 Vali, G.: *Ice Nucleation Relevant to Formation of Hail*, Ph. D. thesis, McGill University, 122 pp., 1968.

504 Vergara-Temprado, J., Miltenberger, A. K., Furtado, K., Grosvenor, D. P., Shipway, B. J., Hill, A. A., Wilkinson, J. M.,
505 Field, P. R., Murray, B. J., and Carslaw, K. S.: Strong control of Southern Ocean cloud reflectivity by ice-nucleating

506 particles, Proc. Natl. Acad. Sci., 115, 2687–2692, <https://doi.org/10.1073/pnas.1721627115>, 2018.

507

