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Fractal Dimensions and Mixing Structures of Soot Particles during Atmospheric Processing

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1 Abstract

Soot particles strongly absorb sunlight and hence act as a short-lived warming agent. 2 Atmospheric aging of soot particles changes their morphology and mixing state and 3 4 consequently alter their optical properties. Here we collected soot particles at tunnel, urban, mountaintop, and background sites in Northern China and analyzed their 5 mixing structures and morphology using transmission electron microscopy. Soot 6 particles were further classified into three types: bare-like, partly coated, and 7 embedded. Bare-like soot particles were dominant at the tunnel site, while most soot 8 particles were partly coated or embedded type at other sites. Fractal dimensions (D_f) 9 of different types of soot particles ranged from 1.80 to 2.16 and were ordered as: 10 bare-like < partly coated < embedded. Moreover, their average D_f changed from 1.8 11 12 to 2.0 from the tunnel to the background site. We conclude that the D_f can characterize the shape of soot aggregates reasonably well and its variation reflects 13 soot aging processes. Compared with the reported D_f of soot particles, we found that 14 $D_f = 1.8$ used in previous optical models primarily represents freshly emitted soot 15 16 aggregates, rather than the ambient ones.

18 **1 Introduction**

Soot particles, also known as black carbon (BC) or elemental carbon (EC), are 19 fractal-like aggregates produced from the incomplete combustion of biomass and 20 21 fossil fuels. Soot particles strongly absorb sunlight and heat the air, altering the radiative forcing of the atmosphere and affecting global and regional climate.¹⁻⁴ 22 During transport and aging, fresh soot particles mix with organic and inorganic 23 aerosols, changing their morphology and compactness, which leads to changes of 24 their optical properties and radiative forcing.^{5, 6} Jacobson⁷ proposed that the sulfate 25 coating on soot particles can enhance optical absorption by ~2 through treating the 26 mixture of soot and sulfate as a core-shell model. However, Cappa et al.⁸ observed 27 that the absorption enhancement of aged soot particles in Sacramento was 6% on 28 average at 532 nm by in-situ measurements. Different conclusions about the optical 29 absorption of soot particles should be attributed to their complicated shapes and 30 various mixing states in the atmosphere.^{9, 10} Due to the lack of quantification on the 31 variation of shapes and mixing structures of soot particles, the debate on optical 32 33 properties of soot particles still continues.

Some experimental methods such as combination of single-particle soot 34 photometer (SP2), three-wavelength photoacoustic soot spectrometer (PASS-3), and 35 Aerodyne soot particle-aerosol mass spectrometer (SP-AMS) were used to well 36 characterize physicochemical properties of soot and measured their optical 37 properties.¹¹⁻¹⁴ However, these measurements could not provide accurate morphology 38 of soot aggregates for the modeling study. Many numerical optical models such as the 39 Rayleigh-Debye-Gans (RDG) approximation¹⁵, T-Matrix^{16, 17}, and Discrete Dipole 40 Approximation $(DDA)^{18}$ can be used to calculate the optical properties of soot 41 aggregates.^{10, 19-21} Except RDG, other numerical models require the morphology of 42 soot aggregates, which can be generated numerically using fractal dimension (D_f) . 43 Among the available algorithms to generate fractal aggregates, the tunable method²² is 44 preferred due to its capability of generating aggregates of a prescribed D_{f} , which is the 45 most important morphological parameter of fractal aggregates. Adachi et al.²³ used 46

47 electron tomography in transmission electron microscopy (TEM) to calculate the D_f of individual soot particles. The method requires a sophisticated system of TEM coupled 48 with tomography, which is not commonly available. Xiong and Friedlander²⁴ 49 calculated the D_f of individual soot particles by drawing circles around the primary 50 particles and then determining the size and position of the primary particles in the 51 TEM image using scaling laws.^{25, 26} The method is inefficient in obtaining the D_f of 52 hundreds of soot particles, because it requires 10-30 minutes for each soot aggregate. 53 54 Later, an approach for image characterization of soot aggregates was proposed by Brasil et al.²⁷ and Oh and Sorensen.²⁸ The method can conveniently derive various 55 parameters of individual soot particles in the scaling law and obtain a D_f to represent 56 their ensemble morphology. Recently, China et al.²⁹ successfully used this method to 57 calculate D_f of soot particles freshly emitted by wildfire. However, there are only 58 quite few available reports about the D_f of ambient soot particles, whose D_f values are 59 very important to understand their optical properties in different environments. 60

In this study, we report a detailed analysis of a large number of soot particles 61 62 collected at tunnel, urban, mountaintop, and background sites in polluted air in the North China Plain (NCP). At each site, soot particles are classified into three types 63 based on their mixing states and morphology and then their corresponding D_f values 64 are calculated and compared systematically for the first time. We use a method 65 combining TEM analysis and numerical calculation to obtain a D_f to represent the 66 ensemble morphology of soot aggregates. At last, we discuss their morphological 67 and mixing properties and the implication of these properties on aging. 68

69

70 2 Materials and Methods

71 **2.1 Aerosol Sampling**

The NCP was covered by the regional haze layer during the sampling period, so we defined our samples from the continental polluted air. Aerosol samples were collected at four sampling sites in NCP: a tunnel site, an urban site, a mountaintop site, and a background site (Figure S1), where the relative humidities (RHs) were at 76 about 52%, 16%, 64%, and 56%, respectively. The RHs at the four sampling sites were lower than 65% during the sampling period, indicating that the hazes were 77 mainly dry. The Kaiyuan tunnel site is a busy highway that enters Jinan City. The 78 urban site in Jinan City is a typical downtown site with strong vehicle and residential 79 emissions. The Mountain Tai (at 1534 m above sea level) is the highest mountain in 80 NCP. The aerosol particles collected at the mountaintop site reflect regional transport 81 of aerosol particles in NCP.³⁰ The background Changdao Island in the Bohai Sea is a 82 downwind site of Shandong province and the Jing-Jin-Ji area (i.e., Beijing, Tianjin, 83 and Hebei province) during winter (Figure S1). Aerosol samples were 84 simultaneously collected at the urban, mountaintop, and background sites during 85 13-23 December 2014. At the tunnel site, aerosol samples were collected on 8 86 November 2016. A total of 779 soot particles from 31 samples were analyzed to 87 determine their size and elemental composition using TEM/EDS. We note that the 88 distribution of aerosol particles on TEM grids was not uniform. Therefore, we chose 89 three to four areas from the center and periphery of each grid to ensure that the 90 91 analyzed particles were representative. Once the internally mixed soot particles are under the strong electron beam (Figure S2), they can easily damage the sulfates and 92 nitrates but do not change morphology of soot aggregates. This microscopic analysis 93 is explained in the Supporting Information. 94

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- 96

2.2 Morphology Analysis of Soot Particles

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Fractal dimension of soot particles can be characterized using the scaling law²⁵:

$$N = k_g \left(\frac{2R_g}{d_p}\right)^{D_f} \tag{1}$$

where N is the total number of monomers in each aggregate, R_g is the radius of 99 gyration of the soot aggregate, d_p is the monomer diameter, k_g is the fractal prefactor, 100 and D_f is the mass fractal dimension. Note that the D_f in this study is the mass fractal 101 102 dimension of soot aggregates that excludes the coating. In this study, the D_f and k_g 103 are estimated from a power law fit of a scatter plot of N vs the values of $2R_{g}/d_{p}$.

N can also be scaled with the aggregate projected area in the following 104

105 power-law relationship:

106

$$N = k_a \left(\frac{A_a}{A_p}\right)^{\alpha} \tag{2}$$

107
$$\delta = \frac{2a}{l} \tag{3}$$

108 where A_a is the projected area of the soot aggregate, A_p is the mean projected area of 109 the monomer, k_a is a constant, and α is an empirical projected area exponent. The 110 exact values of k_a and α depend on the overlap parameter (δ),²⁸ which can be 111 calculated using equation (3) with a being the monomer radius and l the lattice 112 spacing in TEM images. The number of monomers N can then be calculated using 113 equation (2).²⁸

114 The parameters d_p and R_g are also required to determine D_f . While d_p can be 115 obtained directly from analysis of TEM images, estimation of the actual radius of 116 gyration (R_g) is complicated. Here we used the following simple correlation

117
$$L_{\rm max}/(2R_{\rm e}) = 1.50 \pm 0.05$$
 (4)

to calculate R_g^{27} , where L_{max} is the maximum length of the soot aggregate obtained from TEM images.

120

121 **3 Results and Discussion**

122 3.1 Morphology and Mixing State of Soot Particles

Fresh soot particles are normally chain-like aggregates. Once soot particles mix 123 with other aerosol components in the air, the aging process can rearrange the 124 structure of the inner soot aggregates.³¹ Based on their morphology and the visual 125 estimation of coating on soot particles in TEM images, we classified them into three 126 types: bare-like, partly coated, and embedded. Bare-like soot particles in TEM 127 images display clear monomers without any visible coating on their surface (Figure 128 1a-1/1b-1/1c-1). Partly coated soot particles mean that individual soot particles are 129 partly coated by other aerosol components (Figure 1a-2/1b-2/1c-2). Embedded soot 130 particles refer to individual soot particles that are heavily coated or are entirely 131

embedded within other aerosol components (Figure 1a-3/1b-3/1c-3). Figure 1 shows 132 the three types of soot particles collected at the urban, mountaintop, and background 133 sampling sites. Bare-like soot particles are dominant in the tunnel samples, because 134 vehicles emit large amounts of fresh soot particles (Figure S3). Similar results have 135 been found near freeways.³² Based on their different mixing structures, embedded 136 soot particles are normally considered as more aged than the partly coated soot 137 particles.³³ We also calculated the area ratios of coating/soot core for internally 138 mixed soot particles. Figure S4 shows partly coated soot particles mostly have ratios 139 smaller than one, indicating smaller coatings. In contrast, embedded soot particles 140 have lager ratios, some of which are more than 20 times larger. These results are 141 consistent with our classification. 142

Figure 2 shows the fractions of three types of soot particles in different 143 atmospheric environments. The result shows that the bare-like soot particles 144 accounted for 64% of all particles in tunnel air but only 1~25% in urban polluted air 145 (Figure 2). Wang et al.³⁴ also found a fairly low fraction (31.2%) of externally mixed 146 147 soot particles in urban Xi'an City of China through a single-particle soot photometer (SP2). As a result, the polluted air likely accelerated the transformation from 148 bare-like into partly coated or embedded soot particles.³³ It should be noted that 149 bare-like soot particles accounted for 25% at the urban site and 21% at the 150 151 background site, but embedded soot particles significantly increased from 12% to 39% (Figure 2). These results indicate that the polluted air masses from the Jing-Jin-Ji 152 area and Shandong province (Figure S1) brought a large number of aged soot 153 particles into the downwind background air. In addition, the fraction of embedded 154 soot particles at the mountaintop site is largest at 55% and bare-like soot is lowest at 155 1% among the three sampling sites. *China et al.*³⁵ found that most soot particles from 156 North America became internally mixed at the summit caldera of the Pico Volcano. 157 Therefore, soot particles that are emitted mostly at ground level but transported into 158 the upper atmospheric layers could undergo intense aging processes during their 159 160 transports. The RH is a critical factor to enhance heterogeneous reactions of acidic gases on particle surface because secondary aerosols can deliquesce at about 60-80% 161

RH and form liquid phases.³⁶ During the sampling period, there was higher RH around 64% in the upper air than the 16-55% on the ground. Indeed, many embedded soot particles on the mountaintop left a water rim around sulfate coating after drying on the substrate (Figure 1b-3), which indicates that secondary aerosol components existed as the liquid phase in the air.³³ We conclude that soot particles likely underwent more complicated ageing processes due to the higher RH of the upper layers than at the polluted ground sites.

169

170 **3.2 Quantifying the Shapes of Soot Particles**

It is widely acknowledged that D_f of soot particles reflects their combustion 171 conditions and aging processes.²³ Compact soot particles often have larger D_f than 172 lacy aggregates.³⁷ Here we calculated the D_f of soot particles collected at the four 173 sampling sites (Figure 3). D_f of bare-like soot particles at different sampling sites was 174 very close, at ~1.82 (Figure 3a, 3b, 3d). Bare-like soot particles have the lowest D_f 175 followed by partly coated and embedded soot particles (Figure 3), suggesting that 176 177 bare-like soot particles were more lacy compared to the partly coated and embedded types. D_f of partly coated soot particles tends to be ~1.87, smaller than the 1.90~2.16 178 of embedded soot particles (Figure 3). Similarly, China et al.²⁹ found the same 179 properties (i.e., bare-like < partly coated < embedded) of D_f of the three types of soot 180 particles emitted by wildfires. Peng et al.³⁸ also found that the morphology of soot 181 particles was modified heavily during aging processes. For the background soot 182 particles, D_f ranges between 1.83 and 2.16, with a medium of 2.00 (Figure 3d). In 183 contrast, D_f of the urban soot particles has lower values, between 1.83 and 1.90. We 184 multiplied the number fraction of each type of soot by their corresponding D_f to 185 calculate the statistical weighting of D_f . The statistical weighting of D_f values of the 186 urban, mountaintop, and background site are 1.87, 1.90, and 1.97, respectively, which 187 have an average value of 1.91. 188

The convexity (*CV*), roundness (*RN*), and D_f of the three types of soot particles at the four sampling sites are listed in Table S1. The *CV* and *RN* distributions of the three types of soot particle at the same sampling site (Figure S5) clearly prove their

192 D_f changes (Figure 3). The *CV* and *RN* of bare-like soot particles are smallest 193 followed by those of partly coated and embedded soot particles at the four sampling 194 sites. We therefore conclude that larger *CV* and larger *RN* represent more 195 compactness for aged soot particles, consistent with the study of *China et al.*²⁹

We found that D_f of fresh soot particles retained a consistent value (~1.82) at 196 different sampling sites in the polluted air (Figure 3), although fresh soot particles 197 display slightly different D_f due to their different sources and combustion 198 conditions.³⁹ Many researchers obtained D_f of soot particles of the primary sources, 199 such as D_f from biomass burning at 1.67-1.83⁴⁰, D_f from vehicle emissions 200 1.52-1.94³², and D_f from diesel at 1.6-1.9⁴¹. D_f of soot particles becomes larger when 201 soot aggregates are coated by other components during atmospheric processes.⁴² 202 This indicates that soot particles likely collapse during the coating processes. In 203 addition, the wide range of D_f of soot particles in the background air is somewhat 204 expected because they originate from multiple sources, such as industries, residential 205 heating, and transportation,³⁰ and have undergone different atmospheric aging 206 durations and processes.²³ In addition, the D_f of embedded soot particles at 207 1.90~2.16 in this study are much lower than 2.3~2.6 reported by some previous 208 studies.^{21, 23, 43} Adachi et al.^{21, 23} used cube-counting method to calculate D_f of soot 209 particles. In most cases the images of fractal aggregates cannot be decomposed at all 210 scales into an integer number of square boxes using this method,⁴⁴ which may lead 211 to a larger D_f . In the study by *Bambha at al.*,⁴³ the smaller monomer diameter may 212 cause lesser structural compaction.⁴⁵ Besides, the coating material condensed at a 213 low humidity often causes no restructuring, whereas the coating liquefies at a higher 214 humidity and restructuring occurs promptly.⁴⁶ In a word, these differences could be 215 attributed to data processing methods, aging environments, and soot aggregate 216 properties. 217

Using the scaling law method, the previous studies reported D_f at 1.52~1.94 for soot particles at road side³² and $D_f > 2$ for soot particles at a remote marine troposphere site.³⁵ Here we systematically studied D_f of ambient soot particles collected at three representative polluted sites. These data are crucial to assess the accurate shape of soot particles in the dry continental air.

223

224 4 Atmospheric Implications

Previous studies reported that the fractal dimensions (D_f) of fresh soot particles 225 from vehicles, biomass, diesel, and wildfire emissions are around 1.73³², 1.75⁴⁰, 226 1.75^{41} , and 1.89^{29} , respectively, which are close to $1.80 \sim 1.83$ (D_f) of the bare-like 227 soot particles obtained in this study (Figure 3). The reason is that fresh soot particles 228 are generally formed via a cluster-dilute aggregation mechanism in a small-scale 229 burning regime.^{39,47} These fresh soot particles are hydrophobic before being affected 230 by secondary aerosols and condensable vapors in the atmosphere.⁴⁸ Therefore, fresh 231 soot particles can hardly collapse and their structures remain largely unchanged. In 232 contrast, once soot particles interact with secondary organic and inorganic aerosols 233 and water vapor during long-range transport, they became more compact as 234 evidenced by the larger D_f in mountaintop and background air (Figure 3). TEM 235 images further show that the morphology of soot particles not only became more 236 compact from vehicular emission to background air (Figure 1), but also possibly 237 underwent reconstruction under the influence of water vapor.^{5, 49} Therefore, these 238 hygroscopic secondary aerosols heavily caused morphological changes of soot 239 240 particles in the atmosphere.

In this study, the D_f of soot particles were found to vary from 1.80 to 2.16 241 (Figure 3d) for different mixing structures, which indicate that the mixing structure 242 of soot particles can represent their aging degree.^{31, 33} At present, many studies set D_f 243 as ~1.8 to simulate the complex structure of soot particles and to further calculate 244 their optical properties.^{10, 50-52} However, some studies have suggested that the highly 245 compact soot particles have substantially different optical properties from the lacy 246 ones.^{20, 53, 54} In particular, the mass-specific scattering cross sections (MSC) of soot 247 particles follow the order: $D_f = 2.1 > D_f = 1.78 > D_f = 1.4$.²⁰ Therefore, it is essential 248 to select suitable D_f values to construct accurate optical models of soot particles. Our 249 results show that the statistical weighting of D_f of soot samples collected at the urban, 250

mountaintop, and background site has an average value at 1.91, suggesting that D_f = 1.91 could be more representative for ambient soot particles in continental polluted air. In particular, D_f = 1.91 can well represent soot particles in dry (RH<65%), winter polluted air in North China. Further studies are required to quantify the D_f of soot particles in different atmospheric environments, such as in humid, troposphere, and strongly photochemical air, because they all can accelerate soot aging in the atmosphere.^{5, 34}

258

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264

265 Associated Contents

Supporting Information Available: specific microscopic analysis, related
 geometric parameters of soot particles, and some supplementary tables and figures.

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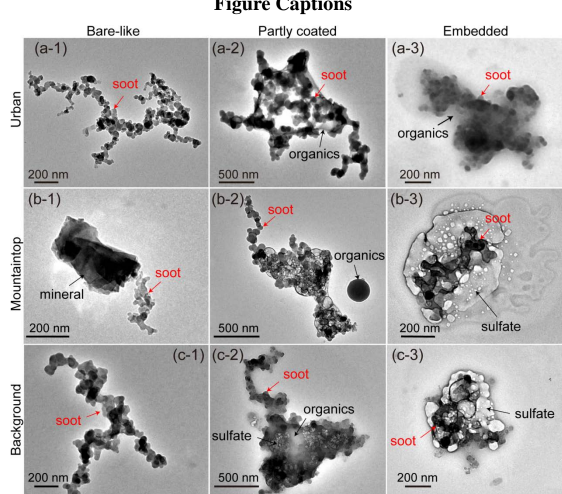




Figure Captions



433 Figure 1. TEM images of individual soot particles collected at the urban site (a-1/2/3), the 434 mountaintop site (b-1/2/3), and the background site (c-1/2/3). Soot particles are classified into 435 three types: bare-like (a/b/c-1), partly coated (a/b/c-2), and embedded (a/b/c-3).

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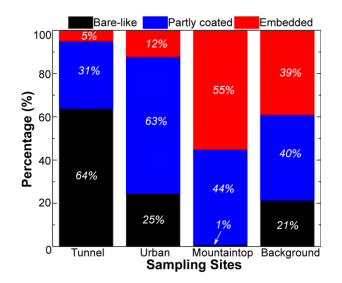


Figure 2. The percentages of bare-like, partly coated, and embedded soot particles collected at
four sampling sites. 147, 216, 295, and 121 soot particles were analyzed in the samples collected
at the tunnel, urban, mountaintop, and background sites, respectively.

441

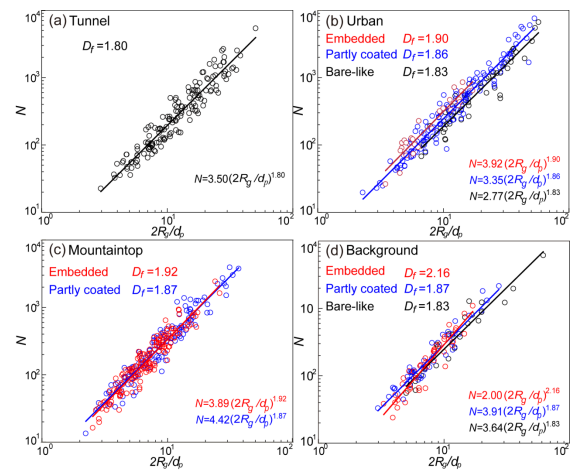


Figure 3. The fractal dimensions of different types of soot collected at the tunnel (a), urban (b),
mountaintop (c), and background (b) site. For each site, the lines and circles represent bare-like

- 445 (black), partly coated (blue), and embedded (red) soot particles.
- 446

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450	
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