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# **Physicochemical and temporal characteristics of individual atmospheric aerosol particles in urban Seoul during KORUS-AQ campaign: insights from single-particle analysis**

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**Abstract.** Single-particle analysis was conducted to characterize atmospheric aerosol particles collected at Olympic Park in Seoul, South Korea, as a part of the Korea–United States Air Quality (KORUS-AQ) campaign which was carried out during May–June 2016. The KORUS-AQ campaign aimed to understand the temporal and spatial characteristics of atmospheric pollution on the Korean Peninsula through an international cooperative field study. A total of 8004 individual particles from 52 samples collected between 23 May–5 June 2016 were investigated using a quantitative electron probe X-ray microanalysis (low-Z particle EPMA), resulting in the identification of seven major particle types. These included genuine and reacted mineral dust, sea-spray aerosols, secondary aerosol particles, heavy-metal-containing particles, combustion particles, Fe-rich particles, and others (particles of biogenic and humic-like substances – HULIS). Distinctly different relative abundances of individual particle types were observed during five characteristic atmospheric situations, namely (a) a mild haze event influenced by local emissions and air mass stagnation; (b) a typical haze event affected by northwestern air masses with a high proportion of sulfate-containing particles; (c) a haze event with a combined influence of northwestern air masses and local emissions; (d) a clean period with low particulate matter concentrations and a blocking pattern; and (e) an event with an enhanced level of heavy-metal-containing particles, with Zn, Mn, Ba, Cu, and Pb being the major species identified. Zn-containing particles were mostly released from local sources such as vehicle exhausts and waste incinerations, while Mn-, Ba-, and Cu-containing particles were attributed to metal alloy plants or mining. The results suggest that the morphology and chemical compositions of atmospheric aerosol particles in urban areas vary depending on their size, sources, and reaction or aging status and are affected by both local emissions and long-range air masses.

# <span id="page-0-0"></span>**1 Introduction**

Atmospheric aerosols, originating from various anthropogenic and natural sources, have significant impacts on climate change and human health (IPCC, 2021). Anthropogenic emissions greatly influence the composition and behavior of airborne particulate matter (PM) (Kim et al., 2018b; Chowdhury et al., 2018; Nault et al., 2018). On the Korean Peninsula, anthropogenic pollutants primarily come from local emissions and long-range-transported air masses (Cho et al., 2021; Park et al., 2020; Choi et al., 2021; Kumar et al., 2021; Nault et al., 2018). Studies have observed changes in the characteristics of aerosols composed of organic and inorganic compounds influenced by different air mass flows. Secondary organic aerosols (SOAs) are particularly affected by local emissions, while inorganic particles can be influenced by either local emissions or long-range-transported pollutants (Nault et al., 2018; Kim et al., 2018b, 2020; Park et al., 2018; Chen et al., 2017). Local emissions, including biomass burning, cooking, and traffic exhaust, primarily influence the formation of SOAs in urban areas (Nault et al., 2018; Park et al., 2018; Kim et al., 2018b). On the other hand, transboundary transport of pollutants is significantly affected by comprehensive climatic conditions and can lead to air pollution episodes dominated by inorganic components, including sulfate, nitrate, and ammonium (Kumar et al., 2021; Lee et al., 2019a; Choi et al., 2019).

East Asia has seen a significant decline in air quality over the past few decades due to increased emissions of gaseous and particulate pollutants as a result of rapid industrial and economic growth. The Korean Peninsula, surrounded by China, Japan, and Russia, exhibits complex aerosol characteristics influenced by a combination of local emissions, surrounding seas, and transboundary long-range-transported air masses (Pochanart et al., 2004; Crawford et al., 2020; Peterson et al., 2019; Ramachandran et al., 2020; Kim et al., 2018a). To further investigate factors affecting air pollution on the Korean Peninsula, an international cooperative field study, the Korea–United States Air Quality (KORUS-AQ) campaign, was conducted during May–June 2016 (Crawford et al., 2020). Through this campaign, the temporal and spatial characteristics of various gaseous and particulate pollutants on the Korean Peninsula were successfully elucidated, making it an important study in the field of atmospheric science (Crawford et al., 2020). On the Korean Peninsula, ammonium was found to be the most sensitive factor affecting PM<sub>2.5</sub> exposure, followed by  $NO_x$ ,  $SO_2$ , organic carbon (OC), and black carbon (BC) (Choi et al., 2019). The presence of anthropogenic ammonium on the Korean Peninsula leads to the formation of ammonium sulfate (AS) and ammonium nitrate (AN) particles (Kim et al., 2021, 2020). Regarding the composition of atmospheric  $PM<sub>1</sub>$  in Seoul, the most populated metropolitan area in South Korea, OC content was found to be the highest, followed by sulfate, nitrate, ammonium, and BC (Kim et al., 2018b).

While previous studies have effectively examined the impact of anthropogenic emissions on the formation of submicron particles during the KORUS-AQ campaign, research on supermicron particles remains limited. Aerosol particles in the supermicron fraction, which mainly originate from natural sources like mineral dust and sea-spray aerosols (SSAs), make up a significant proportion of the total aerosol mass (Andreae and Rosenfeld, 2008; Seinfeld and Pandis, 2006). Airborne mineral dust particles in East Asia, directly emitted from arid regions of Mongolia and northern China, can undergo physicochemical changes during long-range transportation, for example, through atmospheric reactions with anthropogenic  $NO<sub>x</sub>$  and  $SO<sub>2</sub>$ , resulting in the formation of nitrates and sulfates. This leads to alterations in chemical compositions, morphology, size, and radiative forcing capabilities (Sullivan et al., 2007; Yu et al., 2020; Geng et al., 2014; Heim et al., 2020; Sobanska et al., 2012). The investigation of the characteristics of supermicron particles, including their particle–particle variability, formation dynamics, and atmospheric fate, is important to gain a comprehensive understanding of the behavior and impact of atmospheric aerosols of natural and anthropogenic origin on air quality and climate change.

This study utilized a quantitative electron probe X-ray microanalysis (EPMA) technique based on scanning electron microscopy coupled with X-ray spectrometry, so-called low-Z particle EPMA, to examine the physicochemical characteristics of individual aerosol particles collected at Olympic Park in Seoul, South Korea, during the KORUS-AQ campaign. Low-Z particle EPMA is a powerful single-particle analytical technique for providing information on unique features of individual aerosol particles, including morphology, elemental compositions, and particle–particle variability (Geng et al., 2009, 2011; Li et al., 2017; Wu et al., 2019). Differences in these features are attributed to particle sources, formation mechanisms, and atmospheric fate (Wu et al., 2019; Song et al., 2022). This article consists of two parts: (1) an examination of the differences in physicochemical characteristics based on particle species and (2) an analysis of the temporal variations of individual aerosol particles during the KORUS-AQ campaign. The characterization of individual particles, combined with other studies on atmospheric aerosols during the KORUS-AQ period, provides valuable insights into the unique features of urban atmospheric particles.

# **2 Experiments**

#### 2.1 Sampling

Ambient aerosol particles were collected at Olympic Park (37.52◦ N, 127.12◦ E) in Seoul, the capital of South Korea (Fig. S1 in the Supplement). The Seoul metropolitan area (SMA), with high population density, numerous local emissions, and transboundary long-range transport, provides a suitable location for investigating the complex characteristics of atmospheric aerosols (Kim et al., 2018b, 2020). A threestage cascade Dekati  $PM_{10}$  Impactor (Dekati Ltd.) with an aerodynamic cut-off size of 10, 2.5, and  $1.0 \,\mu m$  for stages 1–3 at a 10 L min−<sup>1</sup> flow rate, respectively, was used to collect aerosol particles on Al foils. Each sample set was analyzed for particles collected at stages 2 and 3, corresponding to PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub>, respectively. A total of 52 sets of samples were collected in the morning and afternoon (09:00–10:00 and 15:00–16:00 KST) from 23 May– 5 June 2016. The sampling duration for each stage was controlled to obtain an optimum number of particles without overloading on the Al foils, such as 10–30 min for stage 2 and 5–15 min for stage 3. The 72 h backward air mass trajectories were generated using the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model for different receptor heights of 250, 500, and 1000 m a.g.l. The HYSPLIT model is available on the NOAA's Air Resources Laboratory's website (Stein et al., 2015; Rolph et al., 2017; [http://www.arl.noaa.gov/ready/hysplit4.html,](http://www.arl.noaa.gov/ready/hysplit4.html) last access: 11 January 2024).

# 2.2 Determination of individual particle types by low-*Z* particle EPMA

The physicochemical characteristics of individual aerosol particles were examined using a scanning electron microscope (SEM) (JEOL JSM-6390) equipped with an Oxford Link SATW ultrathin window energy-dispersive X-ray (EDX) detector under vacuum conditions. The resolution of the detector was 133 eV for Mn-K $\alpha$  X-rays, and X-ray spectra were recorded using INCA Oxford software (Oxford Instruments Analytical Ltd, INCA suite version 4.09). Routine measurement was conducted using an accelerating voltage of 10 kV and beam current of 0.5 nA, while 20 kV and 0.25 nA were used to confirm heavy-metal elements of specific particles. To obtain sufficient X-ray counts for quantitative analysis, a typical measurement time of 15 s was chosen. The net X-ray intensities for the elements were obtained using a non-linear least-squares fitting of the collected spectra using the AXIL program (Vekemans et al., 1994). The elemental concentrations of the individual particles were determined from their X-ray intensities using a Monte Carlo simulation combined with reverse successive approximations (Ro et al., 2001, 2002). The chemical species of individual aerosol particles were determined based on their size, morphology, and elemental composition. Measurements under vacuum conditions may result in the evaporation of volatile organic components in individual aerosol particles, but these effects are negligible for ambient supermicron aerosols given their general chemical compositions.

# **3 Results and discussion**

# 3.1 Characteristics and abundances of individual particle types

Individual particles were classified into 13 species based on their morphology and elemental composition and further categorized into seven major groups based on their sources and/or formation mechanism. These groups are (1) secondary aerosol particles including SOAs and secondary organic and inorganic aerosols (SOIAs); (2) genuine and aged/reacted mineral dust; (3) reacted SSAs; (4) combustion particles; (5) Fe-rich particles; (6) heavy-metal-containing particles; and (7) others, including particles of biogenic and humiclike substances (HULIS). More information on the classification can be found in the Supplement (Sect. A and Table S1). While the internal mixing state of individual aerosol particles can offer valuable insights into the sources and formation mechanisms of ambient aerosols (Adachi and Buseck, 2013; Li et al., 2021; Zhang et al., 2022), this study primarily focuses on the overall physicochemical characteristics and relative abundances of the ambient aerosols due to the extensive number of particles investigated.

#### 3.1.1 Secondary aerosol particles (SOAs and SOIAs)

Secondary aerosol particles, including SOAs and SOIAs, account for 5.6% and 29.3% in the PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub> fractions, respectively. These particles, likely formed through gas-to-particle conversion, photochemical processes, and the condensation of semi-volatile organic compounds (Hallquist et al., 2009; Kim et al., 2018a), are significantly more abundant in the fine  $PM_{1-2.5}$  fraction than in the  $PM_{2.5-10}$  fraction. The morphology, X-ray spectra, and elemental compositions of typical secondary aerosol particles are presented in Fig. 1. SOA particles appear as dark droplets in their secondary electron image (SEI) and are primarily composed of C and  $O$  ( $> 90\%$  in low-Z particle EPMA analysis) (Fig. 1a). The spread-droplet-like morphology of SOA particles collected on the hydrophilic Al foil suggests that they are likely low viscous and water soluble. In contrast, SOIAs, which are mixtures of SOA and inorganic constituents such as  $NH_4^+$ , NO<sub>3</sub><sup>-</sup>, and/or SO<sub>4</sub><sup>2-</sup>, exhibit C, N, O, and S in their X-ray spectra and are apparently susceptible to damage by electron beams (Fig. 1b, c). The morphology of SOIA particles varies depending on the organic and inorganic contents. Those with high inorganic content appear as bright, crystalline shapes surrounded by a water-soluble footprint (Fig. 1b), while those with a high organic content resemble dark droplets (Fig. 1c). An inset marked with  $*$  in Fig. 1c shows a SOIA particle that appears as a core–shell structure with a SOIA core surrounded by a dark droplet shade mainly containing C and O. The differences in the crystalline morphology of SOIAs indicate that the heterogeneous nucleation and/or crystallization of particles can vary depending on the chemical species present (Wu et al., 2020). Furthermore, the significant water-soluble footprint surrounding SOA and SOIA particles indicates that aqueous-phase chemistry is a crucial process in the formation of secondary aerosol particles in the urban area of Seoul. Previous studies have reported that SOA particles in South Korea are primarily influenced by local emissions, while the sources of inorganic components are highly relevant to both local emissions and transboundary long-range-transported air masses (Nault et al., 2018; Kim et al., 2018b; Choi et al., 2019).

# 3.1.2 Mineral dust particles

Genuine and reacted mineral dust particles are the most abundant particle types among the seven major ones in this study, accounting for  $73.2\%$  and  $44.5\%$  in the PM<sub>2.5−10</sub> and PM1−2.<sup>5</sup> fractions, respectively. These mineral dust par-



**Figure 1.** Morphology, X-ray spectra, and elemental compositions of (a) SOA, (b) SOIA (high inorganic), (c) SOIA (high organic), (d) reacted aluminosilicate, and (e) reacted carbonate particles.

ticles are irregularly shaped and appear bright in SEI, mainly consisting of crustal elements such as Al, Si, Ca, Mg, K, and others. The observed chemical species of mineral dust particles include aluminosilicates (such as feldspar, muscovite, montmorillonite, illite, kaolinite, talc, and pyrophyllite), quartz  $(SiO<sub>2</sub>)$ , carbonates (calcite – CaCO<sub>3</sub>; dolomite – CaMg $(CO_3)_2$ ), magnesite (MgCO<sub>3</sub>), TiO<sub>2</sub>, and their reacted/aged ones. Genuine mineral dust particles, tending to be larger in size, are significantly more abundant in the  $PM_{2.5-10}$  fraction than in the  $PM_{1-2.5}$  fraction, whereas the proportion of reacted minerals is slightly higher in the PM<sub>1−2.5</sub> fraction (71.0 %) compared to the PM<sub>2.5−10</sub> fraction (66.2%) due to the larger specific surface area of  $PM_{1-2.5}$ particles, making them more prone to chemical reactions in the air.

The reactivity of mineral dust particles varies depending on their chemical species and size, as shown in Table 1. In the PM2.5−<sup>10</sup> fractions, particles are highly associated with nitrate compared to sulfate  $(46.2\% \text{ vs. } 30.0\%)$ , while the abundance of sulfate is comparatively higher than nitrate for PM<sub>1−2.5</sub> particles (34.3 % vs. 20.0 %), indicating that sulfate formation occurs more frequently in smaller particles. The proportion of reacted particles is significantly higher in carbonate particles than in aluminosilicates (93.9 % vs. 56.2 %), indicating that carbonate mineral dust has a higher reactivity than aluminosilicates. Reacted aluminosilicate particles appear bright and irregular, being surrounded by watersoluble moieties (Fig. 1d), implying that the chemical reaction mostly occurred on the surface, while reacted carbonate species show dark, lumpy, core–shell shapes (Fig. 1e), indicating that the reaction readily occurred from the surface to the internal part. Further analysis reveals that the carbonate particles tend to react with sulfate, while aluminosilicates were more likely to interact with nitrate (Table 1). The different abundances of sulfate and nitrate in the reacted mineral particles not only depend on the particle species and size, but also on the source, transport pathway, and formation process (Geng et al., 2011, 2014; Sullivan et al., 2007). These findings suggest that (a) carbonate minerals are more sensitive to changes in atmospheric conditions than aluminosilicates, and (b) carbonate minerals react with sulfate before nitrate due to the prevailing neutralization by sulfate (Takahashi et al., 2014; Matsuki et al., 2005; Seinfeld and Pandis., 2006; Sullivan et al., 2017).

# 3.1.3 Sea-spray aerosols (SSAs)

Nascent SSAs are rich in characteristic elements such as Na, Mg, and Cl, as indicated by their X-ray spectra. They are released into the atmosphere from the sea surface through film drops and jet drops caused by bubble bursting (Eom et al., 2016; Cochran et al., 2017). Freshly emitted SSAs are a mixture of inorganic Na, Mg, and Cl and organic compounds such as fatty acids, amino acids, and liposaccharides, which are closely related to the biological activity of micro-organisms in the marine environment (Eom et al., 2016; Cochran et al., 2017). Once released into the atmosphere, these nascent SSAs tend to react with various acidic species such as sulfuric, nitric, and organic acids to form reacted/aged SSAs. All SSAs for both  $PM_{2.5-10}$  and  $PM_{1-2.5}$ fractions were found to be in the reacted form (Table 1), despite their short transport distances (∼ 50–100 km until they reach the sampling site from the Yellow Sea), suggesting that they are susceptible to atmospheric reactions (Laskin et al., 2003; Gupta et al., 2015; Li et al., 2017; Chen et al., 2020). As shown in Table 1, the reacted SSAs accounted for 12.5 % and 15.8 % in the PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub> fractions, respectively, in which the nitrate-containing SSAs were more abundant than the sulfate-containing ones in the  $PM_{2.5-10}$  fraction

| Type               |                  | Genuine  | Reacted<br>(containing elements) |          |             | % of reacted<br>particles |           | Total    |
|--------------------|------------------|----------|----------------------------------|----------|-------------|---------------------------|-----------|----------|
|                    |                  |          | N                                | S        | <b>Both</b> |                           |           |          |
| $PM_{2.5-10}$      |                  |          |                                  |          |             |                           |           |          |
| Mineral dust       | Aluminosilicates | $23.2\%$ | 17.9%                            | $4.9\%$  | 5.8%        | 55.2%                     | 66.3%     | $73.2\%$ |
|                    | Carbonates       | $1.5\%$  | $8.5\%$                          | $6.9\%$  | $4.5\%$     | $92.9\%$                  | (overall) |          |
| Sea-spray aerosols |                  | $0\%$    | $4.5\%$                          | $2.9\%$  | 5.1%        | $100.0\%$                 |           | $12.5\%$ |
| $PM_{1-2.5}$       |                  |          |                                  |          |             |                           |           |          |
| Mineral dust       | Aluminosilicates | $12.1\%$ | $5.0\%$                          | $7.7\%$  | $3.4\%$     | 57.2%                     | $70.9\%$  | $44.4\%$ |
|                    | Carbonates       | $0.8\%$  | $3.6\%$                          | $10.2\%$ | $1.6\%$     | $95.0\%$                  | (overall) |          |
| Sea-spray aerosols |                  | $0\%$    | $4.4\%$                          | $9.4\%$  | $2.0\%$     | $100.0\%$                 |           | $15.8\%$ |

**Table 1.** Relative abundances of genuine and reacted mineral dust and SSA particles.

 $(9.6\% \text{ vs. } 8.0\%)$ , while those containing sulfates were more abundant in the PM<sub>1−2.5</sub> fraction (11.4 vs. 6.4 %), indicating that sulfate formation occurs more in smaller SSA particles. The higher abundance of SSAs containing both nitrates and sulfates in the larger size fraction may be attributed to the availability of sufficient anions to accumulate acidic cations, which is associated with a decrease in acidity as particle size increases (Angle et al., 2021).

#### 3.1.4 Combustion particles

The combustion particles include soot agglomerates, tar balls, fly ash, and char particles, accounting for 1.3 % and 2.8 % in the PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub> fractions, respectively. Most elemental carbon (EC) particles, such as soot agglomerates, tar balls, and char particles, have similar elemental compositions, but they can be differentiated based on their unique morphology (Fig. 2 and Table S1).

Soot agglomerates are remnants of incomplete combustion and are formed through the vaporization–condensation mechanism (Bond et al., 2004; Chen et al., 2006). Based on their morphology and elemental compositions, soot agglomerates can be classified into two types: fresh and aged. The fresh soot agglomerates appear bright and have a characteristic chain-like structure with fractal geometry, as shown in the right-side SEI of Fig. 2a. The complex geometry of the soot agglomerates provides an active area for the deposition of gaseous or particulate species. The morphology of aged soot agglomerates shown in Fig. 2a is more compact than that of the fresh ones. The aging of the soot agglomerates is attributed to several mechanisms such as oxidation, absorption, or condensation of gaseous species and coagulation with other particles. This aging process can cause the soot agglomerates to shrink and restructure into a more compact shape, as shown in Fig. 2a (Bond et al., 2004; Zhang et al., 2008; Chen et al., 2006).

Tar ball particles are composed of organic oligomers and are a representative particle type from smoldering combustions such as biomass burning or biofuel combustion (Adachi et al., 2019; Girotto et al., 2018; Psfai et al., 2004). The spherical shape of the tar ball particles (Fig. 2b) results from post-physical and chemical transformation of the organic matter. The formation of the tar balls can vary depending on factors such as oligomerization of organics, condensation, photochemical processes, water loss, and temperature changes, leading to different internal structures (Tth et al., 2018; Adachi et al., 2019).

Char particles are incomplete combustion residues of liquid or solid carbonaceous fuel materials, appearing compact and irregular in shape on the SEI, as shown in Fig. 2c (Chen et al., 2006).

Fly ash particles, as shown in Fig. 2d, have a similar elemental composition to aluminosilicate mineral particles but with a distinct bright spherical shape on the SEI. These particles were rarely found in both size fractions, accounting for 0.08 % and 0.42 % in PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub> fractions, respectively. The spherical morphology of fly ash particles is attributed to their formation mechanism, which involves rapid cooling after being released from high-temperature combustion at industrial plants (Geng et al., 2011).

#### 3.1.5 Heavy-metal-containing particles (HMs)

Particles containing heavy-metal elements (HMs), such as Zn, Pb, Cu, Mn, Ba, Zr, Sr, Cd, As, Cr, V, Ni, Sn, and Co, are of particular concern due to their adverse impact on human health. In this study, a significant number of HMs were observed, accounting for 2.7% and 4.4% in the  $PM_{2.5-10}$ and  $PM_{1-2.5}$  fractions, respectively. Among the 14 types of HMs observed, Zn, Pb, Ba, Cu, and Mn were frequently encountered (Fig. 3). HMs can be released from both anthropogenic and natural sources, with thermal power plants,



Figure 2. Morphology, X-ray spectra, and elemental compositions of (a) aged soot aggregates, (b) tar balls, (c) char, (d) fly ash, (e), (f) Zn-HMs, and (g) Mn/Ti-containing particles (Mn/Ti-HMs).



**Figure 3.** Heavy metals observed in HMs for (a) PM<sub>2.5−10</sub> and (b)  $PM_{1-2.5}$  fractions.

vehicle exhaust, battery manufacture, and the metallurgical industry being some of the most common anthropogenic sources (Tian et al., 2015; Xu et al., 2004). Tracing the sources of HMs during the KORUS-AQ campaign can be done based on coexisting elements, morphologies, and relative abundances.

As shown in Fig. 3, the most abundant type of HMs observed in this study was Zn-containing particles (Zn-HMs), accounting for 32.4 % and 57.9 % of the total HMs in PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub>, respectively. Zn-HMs can be emitted from various anthropogenic sources such as waste incinera-

tion, vehicle emissions, rubber tire wear, and coal combustion (Hopke et al., 1991; Chow et al., 2004; Hjortenkrans et al., 2007). In the sampling site, which is an urban area with heavy traffic, Zn-HMs may be attributed to vehicle emissions such as rubber tire and brake pad wear. Two major identified types of Zn-HMs were C–Zn–Cl and C–Zn–Cl + (N or S) (Fig. 2e, f), which made up 54.4 % and 29.8 % of the total Zn-HMs, respectively. A significant proportion (84.9 %) of Zn-HMs was observed to contain Cl, likely due to incomplete atmospheric reactions of  $ZnCl<sub>2</sub>$ .  $ZnCl<sub>2</sub>$  can easily undergo aqueous-phase chemical reactions in the atmosphere due to its hygroscopic nature. The presence of N or S on the X-ray spectra and dark-droplet morphology on the SEI of the Zn-HMs indicate that the particles had undergone atmospheric reactions with  $NO_x / SO_x$  (Moffet et al., 2008). The temporal variations of Zn-HMs will be discussed in Sect. 3.2.

A total of 20 Pb-containing particles (Pb-HMs) were observed in this study, in the forms of mixtures with SSAs (8 particles), Pb–Cl–other heavy metals (6 particles), mineral dust (4 particles), and Pb–As (2 particles). They were likely emitted from vehicle exhaust and coal-fired power plants (Lee et al., 2019b). Among the 39 Mn-containing particles (Mn-HMs) observed, 24 particles were associated with mineral dust, coexisting mainly with Al, Si, Ca, and Mg; 6 particles with Fe; 2 particles with SSAs; 4 particles with Mg, Cl, and S; and 3 particles with F. They might originate from natural soil or anthropogenic sources such as ore-crushing plants, ferroalloy plants, and similar facilities (Moreno et al., 2011). The morphology and elemental composition of an Mn-HM are shown in Fig. 2g. Among the total 33 Cu-containing particles (Cu-HMs), 17 particles were mixed with Fe, followed



**Figure 4.** Relative abundances of various particle types in the (a)  $PM_{2.5-10}$  and (b)  $PM_{1-2.5}$  fractions.

by the mineral dust form (10 particles) and minor forms such as Cu–C–S and Cu–C–N–O (6 particles). Major sources of atmospheric Cu include non-ferrous metal plants, mining, and smelting complexes (Choi et al., 2013; Eichler et al., 2014). Among the total of 30 Ba-containing particles (Ba-HMs), 17 particles were mixed with Fe, followed by mineral dust (5 particles),  $BaSO<sub>4</sub>$  (3 particles), and other minor forms (5 particles). Ba-HMs could be released from natural sources in the form of barite (BaSO<sub>4</sub>) and witherite (BaCO<sub>3</sub>) and anthropogenic sources such as ore crushing plants, mining, refining, and manufacture of barium products (Choudhury et al., 2009; Beddows et al., 2004). The observation that Mn- , Ba-, and Cu-HMs appear abundantly as a mixture of Fe or mineral dust in this study suggests a possibility that their major source might be ferroalloy plants, mining, or ore crushing plants.

# 3.1.6 Fe-rich, biogenic, and HULIS particles

Fe-rich particles, which have an irregular shape and appear bright on the SEI, usually contain more than 20 % Fe in elemental concentration. These particles account for 1.7 % and 2.2 % in the PM<sub>2.5−10</sub> and PM<sub>1−2.5</sub> fractions, respectively, and likely originate from steel production, metallurgical industries, and the abrasion of brake linings (Geng et al., 2011).

Biogenic particles, primarily originating from natural sources (Martin et al., 2010), are relatively more abundant in the PM<sub>2.5−10</sub> fraction (2.83 %) than the PM<sub>1−2.5</sub> fraction  $(0.81\%)$ . They can be identified by their unique morphologies and the presence of minor elements such as Na, Mg, N, K, P, S, and/or Cl (Ro et al., 2002; Geng et al., 2011). In this study, most of the observed biogenic particles were attributed to trichomes, plant fragments, pollen, or spores, as their sizes were generally larger than 2 µm (Matthias-Maser et al., 2000; Coz et al., 2010). Typical examples of biogenic particles are displayed in Fig. S2a–c, corresponding to fungal spores, micro-organisms, and trichomes or leaf fragments, respectively.

The HULIS particles, consisting mainly of water-insoluble organic carbon (WISOC), are characterized by high C and O content and unique morphology. There are 17 out of 8004 particles, only accounting for 0.2 %. They might be released from soil, wetland, and sewage treatment plants.

# 3.2 Temporal chemical composition variations of individual aerosol particles during the KORUS-AQ campaign

Based on differences in relative abundances of individual particle types, ambient PM concentrations (Fig. S3), and



**Figure 5.** Typical secondary electron images of PM1−2.5 aerosol particles collected on (a) 23 May (afternoon) and (b) 25 May (morning) (RM: reacted mineral dust).

backward air mass trajectories (Fig. S4), the sampling period (23 May–5 June) of the KORUS-AQ campaign was divided into five characteristic atmospheric situations as follows: Period I (23 May), an SOA-dominant period influenced by local emissions and air mass stagnation; Period II (25–28 May), an SOIA-rich haze episode with the influence of long-rangetransported air-masses; Period III (20–31 May), haze events with the combined influence of long-range-transported airmasses and local emissions; Period IV (1–3 June), a clean air period; and Period V (4–5 June), a period dominantly influenced by local emissions. The relative abundances of individual particle types are shown in Fig. 4. There are significant differences over the sampling period, especially in the  $PM_{1-2.5}$  fractions.

# 3.2.1 Period I (23 May)

On 23 May, the first day of the sampling period, individual aerosol particles showed clear distinctions in morphology and elemental compositions, particularly for secondary aerosol particles. As shown in Fig. 5a, most secondary aerosol particles in the  $PM_{1-2.5}$  fraction, including SOA and SOIA, had dark-droplet morphology, indicating that their major chemical species are organic carbon. Figure 6 highlights a significant increase in the ratio of SOA to secondary aerosol particles. The SOA / secondary aerosol particle ratio of the  $PM_{1-2.5}$  fraction was notably higher (55.2 %) in the 23 May sample compared to the average for the overall samples during the campaign  $(22.0\%)$ , emphasizing the enhanced contribution of organic carbon to secondary aerosol particle formation. Figure 7 shows that the proportion of combustion particles increased by 1.8 and 1.6 times compared to the overall average in the  $PM_{2.5-10}$  and PM<sub>1−2.5</sub> fractions, respectively. A slightly elevated PM concentration on 23 May (Fig. S3) suggests mild air pollution on that day. Our findings align with other bulk studies that confirmed an increased proportion of organic carbon in PM<sup>1</sup> aerosols during 17–23 May (Kim et al., 2018a, b). Stagnant

conditions under a persistent anticyclone prevented the transport of pollutants from other regions, suggesting a dominant influence of local emissions during this period (Kim et al., 2018b; Peterson et al., 2019; Heim et al., 2020). The formation of SOA in South Korea, particularly in urban areas, was reported to be predominantly influenced by local emissions (Nault et al., 2018). Consequently, the rise in the proportion of organic carbon during Period I can be attributed to the augmented contribution of local emissions to the formation of secondary aerosol particles due to air mass stagnation (Peterson et al., 2019; Kim et al., 2018a, b; Crawford et al., 2021). The enhanced level of combustion particles suggests the contribution of local emissions, while it could be associated with additional influences from Siberian wildfires between 20 and 23 May, as reported by previous studies (Song et al., 2022; Peterson et al., 2019). Overall, the data from 23 May indicate a clear influence of local emissions on aerosol particle composition and concentration.

# 3.2.2 Period II (25–28 May)

After the rainfall on 24 May, the morphology, elemental composition, and relative abundance of individual aerosol particles during 25–29 May (Period II) differed significantly compared to those observed in Period I. In terms of particle morphology, Fig. 5b shows that SOIA particles on 25 May exhibited a bright crystalline morphology, suggesting that these particles are primarily composed of inorganic components such as sulfate, nitrate, and ammonium, as described in Sect. 3.2.1. These bright crystalline SOIA particles mostly contain high sulfur contents, as shown in Fig. 1b, suggesting that their major composition is likely ammonium sulfate (AS) (Wu et al., 2019). Ammonium-rich conditions in East Asia facilitate the existence of secondary particles in AS or AN forms (Kim et al., 2020, 2021). The ratio of SOIA particles to total particles increased dramatically during Period II, as shown in Fig. 6. In the  $PM_{1-2.5}$  fraction, the proportion of SOIA particles out of the total particles increased signif-



**Figure 6.** Relative abundances of (a) secondary aerosols, (b) reacted mineral dust, and (c) reacted SSA particles during the KORUS-AQ campaign.

icantly to 61.5 % on 25 May, compared to the overall average of 24 %, and remained high at 46.3 % during Period II. Additionally, the reacted/aged mineral dust and SSA particles containing sulfate were dominant during Period II both in the PM1−2.<sup>5</sup> and PM2.5−<sup>10</sup> fractions. The drastic increase in ambient PM concentration during this period (Fig. S3) is indicative of an air pollution (haze) episode. In contrast to Period I, which is considered to be influenced mainly by local emissions due to air mass stagnation, the drastic increase in sulfate composition of secondary aerosols, reacted mineral dust, and reacted SSA particles during Period II seems to be driven by other external factors than local emissions. As shown in Fig. S4b, air mass transportation from northeastern China at low altitudes (250 m a.g.l) was observed during this haze episode, which contrasts with Period I. Mild southwesterly winds  $(< 5 \text{ m s}^{-1})$  facilitate the transport of pollutants from China to the study region (Peterson et al., 2019; Nault et al., 2018; Heim et al., 2020; Choi et al., 2019). An elevated level of secondary inorganic constituents, including sulfate, nitrate, and ammonium, was also reported during this period (Kim et al., 2018a, b; Song et al., 2022). The humid conditions (RH  $> 60\%$ ) sustained during this period provided a favorable environment for the formation of secondary particles (SIA and SOA) (Peterson et al., 2019). Also, sulfate-

containing mineral dust and SSA particles were abundantly observed because the air masses were buffered rapidly with sulfate when they passed through urban and industrial areas during long-range transportation (Yu et al., 2020). Overall, the data from Period II suggest a significant influence of long-range-transported air masses from northeastern China on the composition and concentration of aerosol particles in the study region.

#### 3.2.3 Period III (29–31 May)

The relative abundances of individual particles during 29– 31 May (Period III) differed from those of 25–28 May (Period II), despite consistently high PM concentrations during the period (Fig. S3) and air mass flow from northeastern China (Fig. S4c, d). During Period III, the proportion of SOIA particles decreased to 14.4 % compared to 46.3 % in Period II (Fig. 6). Concurrently, the proportions of reacted mineral dust and SSA particles increased with a noticeable increase in nitrate-containing ones (Fig. 6). The increase in nitrate-containing particles in the urban area suggests a strong influence of local emissions (Yan et al., 2015). Changes in the relative abundances of individual HM particles were also noticeable (Fig. 7). The proportion of Zn-



**Figure 7.** Relative abundances of (a) combustion particles and (b) heavy-metal-containing particles during the KORUS-AQ campaign.

HMs increased rapidly from 0.8 % during Period II to 2.8 % during Period III, suggesting an elevated influence of local emissions, given that the major sources of Zn-HMs are local emissions (Sect. 3.1.5). The proportion of other HMs also somewhat increased to 2.8 % compared to the overall average of 1.9 %. The changes in the relative abundances of individual particles observed in this study are fairly different from other bulk studies in which air pollution episodes with consistently high inorganic contents were observed during 25–31 May (Kim et al., 2018a, b). Similar to Period II, weak westerly winds facilitated the transport of pollutants during Period III; however, the formation of secondary particles appears to be relatively reduced due to ∼ 20 % lower RH compared to Period II (Peterson et al., 2019). Additionally, as shown in Fig. S4b, c, during Period III, the travel distance and residence time within the Korean Peninsula were longer relative to Period II, suggesting an increased mixing of transported and local pollutants. Based on the decrease in the proportion of secondary aerosol particles formed mainly through gas–particle conversion and the increase in the reacted forms of the primary aerosol particles, including mineral dust and SSAs, it is plausible that aggregation or mixing between individual particles intensified during Period III. Overall, the changes in particle abundances during Period III indicate a complex interplay of local emissions and long-range transport. The decrease in secondary aerosol particles and the increase in reacted primary aerosol particles, along with the elevated proportions of Zn-HMs and other HMs, suggest intensified mixing of pollutants from various sources during this period.

# 3.2.4 Period IV (1–3 June)

After a series of air pollution episodes, ambient PM concentrations decreased drastically from 1 June (Fig. S3). The relative abundance of individual particle types during 1–3 June manifested distinct differences compared to Periods I-III. During Period IV, the proportion of secondary aerosol particles decreased to 12.9 %, compared to an overall average of 29.2 %, while the proportion of genuine mineral dust particles increased from an overall average of 12.9 % to 18.3 % (Fig. 4). An increase in nitrate-containing reacted mineral dust and SSA particles was observed in both size fractions (Fig. 6). Zn-HMs from local emissions were also frequently encountered during this period (Fig. 7). Moreover, an increase in tar balls and char particles was observed (Fig. 7). Increases in nitrate-containing particles, Zn-HMs, and combustion particles suggest an intensified influence of local emissions. Additionally, it was reported that a blocking pattern influenced by high atmospheric pressure was observed over East Asia during this period, which minimized the transportation of pollutants from other Asian mainland areas (Heim et al., 2020; Peterson et al., 2019). This blocking pattern could have contributed to the increased influence of local emissions and the observed rise in nitrate-containing particles, Zn-HMs, and combustion particles during Period IV. Overall, the data from Period IV suggest that local emissions played a dominant role in shaping the aerosol composition during this period, with limited influence from long-rangetransported air masses. The decrease in secondary aerosol particles and the increase in genuine mineral dust particles and locally emitted pollutants such as Zn-HMs and combustion particles further support this conclusion.

#### 3.2.5 Period V (4–5 June)

Ambient PM concentrations slightly increased during Period V compared to Period IV (Fig. S3). There was a noticeable increase in Zn-HMs during this period, with the proportion of Zn-HM increasing from an average of  $2.6\%$  to  $6.6\%$ . This increase was particularly noticeable during the afternoon hours (Fig. 7) and might be related to heightened weekend traffic, as the sampling area is a park surrounded by thoroughfares (Fig. S1). In addition, the proportion of secondary aerosol particles increased drastically to 28.8 % compared to 12.9 % in Period IV, and the proportion of combustion particles somewhat increased to 3.6 % compared to an average of 2.6 %. Air mass trajectories shown in Fig. S4f suggest an intensified influence from inland South Korea during Period V, resulting in the increased levels of particles from local emissions and secondary aerosol particles. When the air mass travels a short distance ( $\sim$  250 km d<sup>-1</sup>), urban areas could be primarily influenced by local emissions (Lee et al., 2019a). Continued blocking patterns from Period IV effectively exclude pollutant transport from outside areas but occasionally lead to stagnant conditions (Peterson et al., 2019). Overall, the changes in particle abundances during Period V indicate an intensified influence of local emissions and secondary aerosol particles, likely due to weekend traffic and stagnant conditions. The increase in Zn-HMs and combustion particles further supports the impact of local anthropogenic emissions on air quality during this period. Efforts to manage and control local emission sources, including vehicle emissions, waste incineration, and fossil fuel combustions, could play a crucial role in improving air quality in the urban area.

# **4 Conclusions**

Individual aerosol particles collected at Olympic Park, Seoul, South Korea, during the KORUS-AQ campaign were analyzed using low-Z particle EPMA. A total of 8004 particles from 52 samples were examined to identify their chemical species, particle–particle variability, sources, and atmospheric fate. The major constituents in the  $PM_{2.5-10}$  and PM<sub>1−2.5</sub> fractions were mineral dust, SSAs, and secondary aerosol particles. However, the relative abundance of individual particle types varied depending on changes in air mass flow and differences in emission sources. The reacted mineral dust and reacted SSA particles containing nitrate were abundant in the  $PM_{2.5-10}$  fraction, whereas sulfatecontaining ones were relatively higher in the  $PM_{1-2.5}$  fraction. Of particular interest, heavy metals were found to account for a relatively high proportion of particles both in the PM<sub>2.5−10</sub> (2.65 %) and PM<sub>1−2.5</sub> (4.42 %) fractions, with Zn, Pb, Ba, Mn, and Cu being the major species. Zn and Pb are mainly released from sources such as waste incineration, vehicle exhaust, and coal-fired power plants, while Mn, Ba, and Cu are primarily released from mining and metal alloy industries.

The relative abundances of secondary aerosol particles varied significantly during the sampling period, reflecting changes in air mass stagnation and emission sources. During the haze episodes, sulfate-containing particles, including SOIA, mineral dust, and SSAs, were predominant, and the proportion of SOA particles increased as local influence intensified. During the clean period of 1–3 June, nitratecontaining particles were abundantly observed, indicating a high contribution of  $NO<sub>x</sub>$  emissions from local sources. Zn-HMs from local sources such as vehicle emissions and waste incineration were noticeably observed during 4–5 June when the air mass stagnated over the Korean Peninsula.

The temporal variations in the abundance and physicochemical characteristics of individual aerosol particles provide valuable insights into the behavior and emission sources of atmospheric urban aerosols. The changes in the composition of organic and inorganic components resulted in distinct morphological and crystalline structures of secondary aerosol particles, influencing properties such as hygroscopic behavior and radiative forcing. The relative abundance of HMs, particularly those containing Zn, effectively reveals the impact of local emissions such as vehicle emissions and waste incineration. The highly hygroscopic nature of the observed Zn-HMs suggests a potential threat to human health, as they are prone to absorbing or reacting with other organic and inorganic components in the atmosphere. The observed changes in the abundance of particles from typical combustion events and secondary aerosol particles emphasize the need to manage local emission sources to maintain air quality. The complexity of aerosol particle behavior highlights the importance of a comprehensive understanding of the interplay between local emissions, long-range transport, and meteorological conditions to develop effective air pollution mitigation strategies.

**Code and data availability.** The data set is available upon request from Chul-Un Ro (curo@inha.ac.kr).

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**Author contributions.** RCU and GH designed and supervised the entire experimental program, provided guidance on the quantitative analysis of individual particles, and reviewed the manuscript. HJY conducted the single-particle analysis, analyzed the data, and wrote the manuscript. LW reviewed and provided feedback on the manuscript. All authors read and approved the final paper.

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

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