Characterization of chemical and physical changes in atmospheric aerosols during fog processing at Baengnyeong Island, South Korea

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HIGHLIGHTS

- Particle composition and size measurements were used to explore fog processing.
- Organic scavenging efficiency $\eta$ (7–34%) was lower in less-oxidized particles.
- Mass decreased after 3 fog events and increased, notably in organics, in 2 events.
- The aerosol carbon oxidation state decreased by the end all fog episodes.

ABSTRACT

In situ interactions between aerosols and atmospheric fog droplets were explored using aerosol physical and chemical measurements and fog samples taken at the Baengnyeong Island Intensive Air Quality Monitoring Station, South Korea during June 16th to July 21st, 2014. To investigate wet scavenging by fog, aerosols were characterized by aerosol mass spectrometer before, during, and after five fog events. Total non-refractory PM\textsubscript{1} mass decreased by an average of 22.8% with the onset of fog. The scavenging efficiency ($\eta$) of organics was 7–34%, where lower $\eta$ was associated with lower organic oxidation and hygroscopicity. The $\eta$ was 16–68% for NH\textsubscript{4}\textsuperscript{+}, -1–35% for SO\textsubscript{4}\textsuperscript{2–}, and 17–86% for NO\textsubscript{3}–, consistent with previous studies and the relatively high NO\textsubscript{3}– hygroscopicity. The variation in scavenging was size dependent for both organics and NO\textsubscript{3}–. Particles with lower NO\textsubscript{3}– $\eta$ on 7/18 were smaller (mode = 346 nm), while those with higher NO\textsubscript{3}– $\eta$ were larger (593 nm). Differences in mass were also explored before and after fog events. Three episodes showed overall decreases in all the components after the fog event, while two episodes showed overall increases, most notably in organics. Both organic mass and SO\textsubscript{2}−/H\textsubscript{2}SO\textsubscript{4}− ratio were used to indicate the prevalence of hydroxymethanesulfonate (HMS), an aqueous secondary organic aerosol (aqSOA) formation marker, which increased after the 7/18 fog episode. The aerosol carbon oxidation state decreased toward the end of all fog episodes suggesting possible molecular fragmentation and loss of highly oxidized functional groups. On 7/18, the organic content developed lower O:C and higher H:C moving “up” a slope of ~1 in a van Krevelen diagram over the course of the fog episode, indicating carboxylic acid or hydroxycarbonyl loss. This study explores relatively rare time-resolved aerosol fog processing measurements. The results confirm previously established relationships between scavenging and particle size, demonstrate overall increases and decreases in particle mass after fog processing based on dominant local chemistry and wet deposition, and show predominant decreases in organic oxidation after fog processing, thereby, contributing to our understanding of particle processing and organic cycling in the atmosphere.

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https://doi.org/10.1016/j.atmosenv.2022.119091

Received 25 August 2021; Received in revised form 1 March 2022; Accepted 27 March 2022

Available online 28 March 2022

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1. Introduction

Fog and cloud processing can influence the fate of atmospheric particulate species, which play important roles in climate and radiative balance (Ghan et al., 2007), air quality, human health (Pope et al., 2002), nutrient cycling (Benedict et al., 2013), and other important biogeochemical processes. Fog processing involves both physical and chemical mechanisms including scavenging and removal of atmospheric particles by droplets, which can serve to reduce particle and gas concentrations in the atmosphere (Collett et al., 2001), hygroscopic particle growth (Tang and Munkelwitz, 1993), aqueous processing such as aqueous oxidation of both gas phase precursors and dissolved semi- and low-volatile compounds (Ervens et al., 2011; Harris et al., 2014; Kaul et al., 2011; Shen et al., 2012), and droplet re-evaporation, which releases both particulate and gaseous species (Ervens, 2015). Fog and cloud droplets also play a vital role in the hydrological cycle and influence atmospheric radiative transfer (Pruppacher and Klett, 2010). Fogs comprise of a complex multiphase system combining gases, particles, and water droplets.

Fog scavenging of particles includes impaction and nucleation mechanisms. Impaction scavenging involves the inclusion of interstitial particles into fog droplets via Brownian diffusion, inertial impaction, and phoretic processes, while nucleation scavenging involves the absorption of water by particles to become activated fog droplets in a supersaturated environment (van Pinxteren et al., 2016). Numerous studies have found preferential scavenging of species with higher hygroscopicity including NO$_3^-$ (Gilardoni et al., 2014; Varutbangkul et al., 2006) and more-oxidized organic compounds (Decesari et al., 2005; Noone et al., 1992). Our understanding of changes in particle chemical composition, especially the organic fraction, due to fog processing continues to develop and will benefit from additional ambient observations (Herckes et al., 2013).

Scavenging is also a function of particle size both in total and for individual species featuring various size-scavenging relationships. For example, Gilardoni et al. (2014) found a size-segregated scavenging efficiency ($\eta$) following sigmoidal curves for both NO$_3^-$ and organic aerosols with $\eta$ decreasing to near zero with decreasing vacuum aerodynamic diameter (d$_{va}$) below 100 nm and increasing to ~1 at d$_{va}$ values of above 700 nm. Other studies have also found size dependencies even though they were not resolved by particle composition (Noone et al., 1992; Sellegri et al., 2003). Overall, few studies have investigated compositionally resolved fog scavenging as a function of particle size.

However, scavenging of particles by fog droplets when fog event first impacts the study site is only one of the many processes that affect particle mass and composition during fog processing. For instance, aqueous reactions can both produce and consume gaseous and particulate species, and subsequent fog droplet evaporation may release some of these species in the form of physicochemically altered particles. Such aqueous secondary aerosol formation can be indicated by increases in particle size within the 0.4–1 μm range (Hering and Friedlander, 1982; John et al., 1990) in contrast to gas-phase particle formation, which predominantly forms particles of <0.2 μm (Ervens et al., 2011). Aqueous secondary organic aerosol (aqSOA) formation can be indicated by the detection of methane sulfonic acid (MSA), which is produced by the oxidation of dimethyl sulfide (DMS) emitted from oceanic organisms (Phinney et al., 2006; von Glasow and Crutzen, 2004; Zorn et al., 2008).

Fig. 1. Location of the Baengnyeong Island sampling site in South Korea. Reproduced with permission from Lee et al. (2015).
In addition, oxalic acid is a typical marker of aqSOA formation by the oxidation of glyoxal and other gas-phase precursors (Carlton et al., 2007; Graban et al., 2004). Another aqSOA formation tracer, hydroxy-methanesulfonate (HMS) is an important tracer in ambient fog and cloud processing, HMS is produced by the formaldehyde–HSO₅⁻ (or SO₂⁻) reaction in aqueous-phase (Gilariondi et al., 2014; Munger et al., 1986; Whiteaker and Prather, 2003).

To better understand particle chemical composition and the interactions between aerosols and atmospheric droplets, fog sampling and aerosol physical and chemical measurements were conducted at the Baengnyeong Island Intensive Air Quality Monitoring Station, South Korea, during June–July 2014. The chemical composition and physical properties of aerosol particles were characterized before, during, and after fog events using a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne) and a scanning mobility particle sizer (SMPS, TSI). The collected fog/cloud samples were also characterized and are presented separately (Boris et al., 2016). This work will examine changes in particle chemical composition and microphysics due to fog and cloud processing in the boundary layer.

2. Methods

2.1. Sampling site

The interstitial particles in the fog and ambient particles were measured at the Baengnyeong Island Intensive Air Quality Monitoring Station, South Korea (37.97°N, 124.63°E; 150 m above mean sea level (AMSL); Fig. 1) during June 17th to July 21st, 2014. This site, which houses a variety of instruments and is used by the National Institute of Environmental Research (NIER) for routine air quality monitoring, is located on a relatively small and low-lying island at the northwesternmost boundary of South Korea and features frequent fog events and well-studied atmospheric chemistry. Baengnyeong Island is influenced by both photochemically aged and somewhat less-oxidized transported industrial emissions, biomass burning and dust from the Pearl River Delta (generally more aged and richer in SO₄²⁻), Korean Peninsula and East sea (typically containing more abundant organics), and East China Sea (Boris et al., 2016; Choi et al., 2016; Kang et al., 2018, 2020; Lee et al., 2015).

2.2. Instrumentation and data analysis

The particle chemical composition was characterized before, during, and after fog events using a HR-ToF-AMS in the V-mode with an averaging time of 5 min. Ambient air was drawn from an inlet ~100 m above sea level at a total flow rate of 3 LPM through a URG cyclone (Dμ = 2.5 μm) and dried to < 40% RH using Perma Pure dryers (MD-110-24) before being focused into the AMS interior via an aerodynamic lens (aerodynamic diameters = 60–600 nm; Liu et al., 2007). Within the AMS, particles are vaporized at 600 °C, ionized at 70 eV, and the resulting fragmented molecules are separated by mass in a ToF-MS and counted to produce total mass spectra (DeCarlo et al., 2006). In this study, total organics, SO₄²⁻, NO₃⁻, NH₄⁺, and Cl⁻ were quantified using the standard fragmentation tables and adjustments for composition-dependent collection efficiency (Allan et al., 2004; Middlebrook et al., 2012) provided in SQUIRREL (v1.56C) and PIKA (v1.15C) AMS data analysis software (DeCarlo et al., 2006; Supeer, 2009). Although fragmentation limits our ability to determine organic speciation, some speciation was explored (discussed in Section 3.5) and elemental analysis was used to investigate changes in organic aerosol oxidation state (including O/C, H/C, and OM/OC) during fog processing. Ammonium nitrate particles (300 nm, ~350 particles cm⁻³) were used for NO₃⁻ ionization efficiency (IE) calibrations, and default relative IE values were used for other species.

Positive Matrix Factorization (PMF) analysis was performed on the AMS data using the Igor Pro-based PMF Evaluation Toolkit (PET; v3.05). PMF has been described in detail elsewhere (Paatero and Tapper, 1994; Paatero, 1997; Ulbrich et al., 2009). Ulbrich et al. (2009) and others have previously applied PMF to AMS data using PET, which deconvolutes spectral patterns, or ‘factors’ (such as hydrocarbon-like organic aerosol, biomass burning aerosol, and oxidized organic aerosol, or OOA) from the measured mass spectra and constructs a representative spectrum and timeline for each factor (e.g., Jimenez et al., 2009; Ulbrich et al., 2009; Sun et al., 2011; Zhang et al., 2011; Saarikoski et al., 2012; Lee et al., 2015). Descriptive statistics for the PMF analysis (i.e., FPEAK and Q values) were provided in Fig. S1.

Aerodynamic properties were measured using a scanning mobility particle sizer (SMPS, model 3034, TSI Inc., USA) with a particle size scanning range of 10.4–469.8 nm and scanning time of 5 min as described in Hogrefe et al. (2006). The SMPS was preceded by a krypton-85 (Kr) neutralizer (model 3077, TSI Inc., USA). The fog liquid water content (LWC) was measured by a Gerber particle sizer (SMPS, TSI) with a particle size scanning range of 10.4–469.8 nm and scanning time of 5 min as described in Hogrefe et al. (2006). The SMPS was preceded by a krypton-85 (Kr) neutralizer (model 3077, TSI Inc., USA).

The fog liquid water content (LWC) was measured by a Gerber particulate volume monitor (PVM-100, GSI Inc., USA; Gerber, 1991) and the fog event was determined as periods in which the LWC exceeds 30 mg m⁻³ (Boris et al., 2016). The World Meteorological Organization (WMO) defines fog by thickness (or intensity) according to the restriction of horizontal visibility. The visibilities of dense, moderate, and light fogs are less than 1 km, 1 to less than 5 km, and 5 to less than 11 km,
3. Results and discussion

3.1. Liquid water content and chemical composition of fog samples

Fog events (LWC >30 mg m⁻³) impacted the monitoring station on 6/30, 7/1, 7/2, 7/3, 7/18, 7/19, and 7/20 of 2014. Of these seven events, the data processing herein focuses on five, namely episodes on 6/30, 7/1, 7/2, 7/3, 7/18, 7/19, and 7/20 (Table 1). The 7/3 and 7/19 episode was excluded from the analysis because the LWC exceeded 30 mg m⁻³ only intermittently. The compositions of these fog water samples are reported in depth in Boris et al. (2016) and they are summarized here. Boris et al. (2016) collected and analyzed fog samples, which were larger than 16 μm, and small droplets in the range of 4–16 μm. Overall, the dominant components of the fog water were NH₄⁺ (mean = 2220 μM), NO₃⁻ (1260 μM), SO₄²⁻ (730 μM), Na⁺ (551 μM), Cl⁻ (253 μM), and the fog water pH was 3.48–5.00 (3.94). Marine sources as indicated by Ca²⁺ and other species from sea-salt and biomass burning as suggested by K⁺ in conjunction with nitrophenol, possible levoglucosan enhancement (Weber et al., 2007), and fire detected upwind (via MODIS and meteorological analysis) contributed considerably to particulate mass on some dates (Boris et al., 2016).

Acetic, formic, oxalic, succinic, maleic, and other organic acids are often formed from aqueous reactions and contribute significantly to the fog water organic fraction, indicating the influence of aqueous (in addition to dry) photochemical oxidation during long-range transport (Boris et al., 2016).

3.2. Time series of non-refractory PM₁ mass and chemical composition

Fig. 2 shows the variations in non-refractory PM₁ (NR-PM₁) component mass during the five fog episodes explored in this study. The total average NR-PM₁ (NR-PM₁ = Organics, SO₄²⁻, NO₃⁻ and NH₄⁺) concentration with standard deviation was 6.30 ± 4.26 μg m⁻³ during on June 17th to July 21st, 2014. SO₄²⁻ and organics were the predominant aerosol species at 1–11 μg m⁻³ (48.6 ± 12.1%) and 1–7 μg m⁻³ (23.0 ± 12.7%), respectively. NO₃⁻ and NH₄⁺ were lower at 0.2–4 μg m⁻³ (10.5 ± 7.42%) and 0.3–4 μg m⁻³ (17.8 ± 3.57%), respectively. NH₄⁺ was higher than NO₃⁻, which implies that the NH₄⁺ can also combine with SO₄²⁻. These total particle mass and percent composition values are consistent with previous measurements at this site (Lee et al., 2015) and are similar to those of other marine and anthropogenically influenced coastal sites (Choi et al., 2017; Rivellini et al., 2017; Schulze et al., 2018).

Positive matrix factorization (PMF) analysis of earlier data from Baengnyeong Island indicated the presence of hydrocarbon-like organic aerosol (HOA) and two oxidized organic aerosol (OOA) factors, namely, more oxidized (MO-OOA) and less oxidized (LO-OOA) OOA (Lee et al., 2015). In this study, O:C averaged 0.71 ± 0.11 (range: 0.68–0.74) indicating that the organic aerosol was predominantly within the low-volatility oxygenated organic aerosol (LV-OOA) range as defined in previous studies (Aiken et al., 2008; Canagaratna et al., 2015; Kroll et al., 2009; Ng et al., 2010). The relationships between organic oxidation (and thus, hygroscopicity) and scavenging are discussed in Section 3.3.

3.3. Fog aerosol scavenging efficiency and aerosol changes after fog processing

Total NR-PM₁ averaged 4.86 ± 4.36 μg m⁻³ during fog episodes and 6.30 ± 4.26 μg m⁻³ during non-fog periods. Particle concentrations generally decreased (average decrease = 22.8%) with the onset of the fog episodes (see Fig. 2), which is consistent with previously measured scavenging behavior by Collett et al. (2001) and Gigardoni et al. (2014), who measured 60% reduction in NR-PM₁ during fog events at a background site.
The fog scavenging efficiency ($\eta$) is defined as the fraction or percentage of a given particle component X remaining in the atmosphere immediately after the onset of the fog event (Eqn 1) (Noone et al., 1992).

\[
\eta = 1 - \frac{[X]_{\text{interstitial}}}{[X]_{\text{before fog}}}
\]  

(Eq 1)

where $[X]_{\text{interstitial}}$ refers to the concentration of unscavenged particles suspended between fog droplets. We used 30-min average data for the fog scavenging efficiency calculations. $[X]_{\text{before fog}}$ was used “before period” datasets in Table 1, and $[X]_{\text{interstitial}}$ data was used 30-min averaged data right after of “before fog period” in Table 1. Fig. 3 shows the $\eta$ of each particle constituent during each of the fog episodes. NO$_3^-$ is scavenged most efficiently. Note that the very slight increase in SO$_4^{2-}$ on 7/20 is well within the instrument error. However, this lack of decrease in particulate SO$_4^{2-}$ concentration is atypical of the fog process, and it may indicate the arrival of a new and higher SO$_4^{2-}$ air mass with the fog event. In addition, by checking the wind direction of the 7/20 case, the air masses came from the sea southwest of the monitoring site before the fog event, but during the fog event, the air masses came from inland of the island by the southeast wind (Table 1). The significant differences in $\eta$ between the components during some of the fog events suggest external mixing and differences in hygroscopicity and or size-dependent composition within the particle population, assuming that the fog events were not accompanied by transported air masses that differed significantly in composition from the pre-fog air masses. For instance, on 6/30 and 7/2, the $\eta$ patterns of NH$_4^+$ and NO$_3^-$ are similar, but somewhat (not statistically) different from the scavenging efficiencies shared by organics and SO$_4^{2-}$. This may indicate externally mixed NH$_4$NO$_3$ and Org (organics)-SO$_4$ (or Org-SO$_4$-NH$_4$) particles, of which NH$_4$NO$_3$ scavenged more efficiently.

The Org $\eta$ values measured herein range from 7% to 34% and include two values of <20%, which are slightly lower than the Org $\eta$ values measured elsewhere. For instance, $\eta$ was measured at 20–60% in the Po Valley, Italy (Gilardoni et al., 2014), 33–90% in Angiola, California, and 41% in Fresno, California (Collett et al., 2008). These instances of relatively low organic aerosol scavenging efficiency may have arisen from lower organic hygroscopicity, which can be a characteristic of less-oxidized organic particles (Ge et al., 2012; Lambe et al., 2011; Massoli et al., 2010). Indeed, during the 7/20 episode, which featured particularly low organic scavenging, the particles had relatively high H:C (1.42) and low O:C (0.47) values as compared to the other fog events. The $\eta$ for ammonium measured 16–68%, while that for SO$_4^{2-}$ ranged from –1 (indicating a slight increase in mass) to 35%. The NO$_3^-$ $\eta$ values measured herein were 17–86% (average = 48%). The higher values are consistent with NO$_3^-$ scavenging observed in the Po Valley, Italy, where...
it averaged 71% (Gilardoni et al., 2014). For both Org and NO$_3^-$, the variation in scavenging may be size dependent. Gilardoni et al. (2014) found significant decreases in Org and NO$_3^-$ scavenging efficiencies with decreasing size of particles <400 nm and <300 nm, respectively. Indeed, the particle sizes had a mode of 346 nm on 7/18 (Fig. S4) when the lowest NO$_3^-$ $\eta$ values was observed, and a larger mode of 593 nm during the 7/1 event when NO$_3^-$ $\eta$ was the highest.

However, scavenging, which reveals the net uptake of particles when the fog event first impacts the study site, is not the only process that affects the ultimate particle mass and composition during fog processing. Aqueous reactions can both produce and consume gaseous and particulate species, and subsequent fog droplet evaporation will release some of these species in physicochemically altered aerosol particles. To explore such processing, and assuming negligible change in aerosol properties due to air mass transport during the given events, Fig. 4 shows the overall mass reduction efficiency (in which, like $\eta$, negative numbers denote increases in mass) for Org, NO$_3^-$, SO$_4^{2-}$, and NH$_4^+$ during each of the fog episodes. Note that, while three episodes feature overall decreases in the masses of the chemical components after the conclusion of the fog event, two episodes (namely 7/2 and 7/18) show overall increases in the component masses after the event, despite the initial scavenging (Fig. 4). Organics increase most notably, which is consistent with the increases in SOA due to fog processing found in India (Kaul et al., 2011) and California (Ge et al., 2012). The mass of NO$_3^-$ also increases with organics after fog processing on both 7/2 and 7/18. Farmer et al. (2010) reported that the NO$_3^-$/NO$_2^-$ ratio was increased with the organonitrate. The NO$_3^-$/NO$_2^-$ ratio was increased after fog event on 7/2 case (Table S1), but not 7/18. The increase of organics and NO$_3^-$ on 7/2 case it possibly affected by organonitrate formation. The reduction efficiency of 7/18 case may have been overestimated due to the low concentration (Fig. 4). This finding is consistent with numerous observations showing that aqueous reactions in atmospheric droplets can form secondary organonitrate-containing particles (Ervens et al., 2011; Zhao et al., 2013).

3.4. Hydroxymethanesulfonate indicates aqueous-phase secondary organic aerosol formation

The aqueous-phase reaction can produce many kinds of compounds, such as MSA, oxalic acid, and HMS by a variety of chemical reactions (Carlton et al., 2007; Crahan et al., 2004; Munger et al., 1986; Phinney et al., 2006; Zorn et al., 2009). HMS, a tracer for one route of aqSOA formation, was analyzed to further explore the possible aqSOA formation found in the previous section. HMS formation rate is 1.6 nM h$^{-1}$ to 0.1 M h$^{-1}$ at pH 3 to pH 7, respectively (Munger et al., 1986), and the fog water pH was 3.48–5.00 (3.94) during the fog event (Boris et al., 2016). HMS fragments form AMS mass spectra dominated by fragments such as CHO$^+$, CH$_2$O$^+$, SO$_3^+$, and SO$_2^+$, which are common to a variety of ambient aerosol types. No characteristic HMS ions were found in AMS spectra (Ge et al., 2012). However, HMS does not produce SO$_3^+$, HSO$_3^+$, or H$_2$SO$_4^+$ ions under AMS ionization, while ammonium sulfate does. Thus, we can use the changes in SO$_3^+/SO_2^+$, SO$_3^+/HSO$_3^+$, SO$_2^+/H_2$SO$_4^+$, SO$_2^+/SO_3^+$, SO$_2^+/HSO$_3^+$, and SO$_2^+/H_2$SO$_4^+$ ratios to explore possible HMS contributions to the organic aerosol in question (Ge et al., 2012), where higher ratios indicate higher amounts of HMS in the ambient aerosol. Of these ratios, SO$_3^+/H_2$SO$_4^+$ and SO$_2^+/H_2$SO$_4^+$ fluctuate most dramatically over time (and between fog and non-fog periods; Fig. S5). SO$_2^+/H_2$SO$_4^+$ is used herein (Fig. 5). Both the organic mass and SO$_2^+/H_2$SO$_4^+$ ratio increased after the 7/18 fog episode even though the latter did not to a significant extent. This SO$_2^+/H_2$SO$_4^+$ increase may arise from a decrease in inorganic SO$_4^{2-}$ (as indicated by the decrease in the mass of total SO$_4^{2-}$) and/or real increase in the HMS contribution, but may also be changed within error. Nonetheless, the change in this ratio suggests that the aqueous-phase SOA contributions increased in importance relative to inorganic SO$_4^{2-}$ during this episode. SO$_2^+/H_2$SO$_4^+$ did not increase along with organic mass on 7/2, but NO$_3^-$ did. This concurrent increase in organics and NO$_3^-$ may indicate the formation of organonitrates, which can contribute to the SOA mass increases in atmospheric droplets.

**Fig. 5.** SO$_3^+/H_2$SO$_4^+$ ratios before and after the fog episodes.

**Fig. 6.** Change in carbon oxidation state between just before and after the fog event.
3.5. Changes in aerosol oxidation state after fog events

As shown in Fig. 6, the aerosol carbon oxidation state was lower after the fog event than before the fog onset for all fog episodes. In addition, we separated OA into 3-factors as HOA, LO-OOA, and MO-OOA (Figs. S2 and S3). The HOA is similar to Primary OA (POA), LO-OOA and MO-OOA correspond to SOA (Zhang et al., 2011). The MO-OOA decreased after fog in all fog episodes, but relatively fresh LO-OOA and HOA increased (Table 2). The HOA increased in all cases except 7/2, and LO-OOA increased in 6/30, 7/2, and 7/18 cases. The carbon oxidation state of after fog event in 7/20 case was the lowest of five episodes (Fig. 6). In addition, the highest HOA and lowest SOA (LO-OOA + MO-OOA) ratios were observed after fog compared to other cases. This phenomenon suggests that the processing of SOA precursors forms less-oxidized SOA and/or causes the fragmentation and loss of oxidized functional groups, which can occur during aqueous photooxidation (Schurman et al., 2018). Such changes in oxidation are often enabled by the uptake of high-order unsaturated VOCs such as light aromatics (e.g., toluene and benzene) and biogenic VOCs (e.g., isoprene), which are subsequently oxidized via aqueous reactions (McNeill, 2015). The resulting semi-volatile SOA can then be released into the particle phase via droplet evaporation (Ervens et al., 2011). In addition, OA increases its hygroscopicity as it is oxidized (Massoli et al., 2010). Massoli et al. (2010) reported that the LV-OOA had higher hygroscopicity than the semi-volatility OOA (SV-OOA) in laboratory SOA experiments. Xu et al. (2017) reported the water-solubility of various OA factors calculated by the PMF. The HOA had the lowest water-solubility, followed by the LO-OOA and the MO-OOA, where oxidation proceeds (Xu et al., 2017). The increase in hygroscopicity of OA creates conditions for increased absorption and growth in fog droplets during fog events (Ming and Russell, 2004). The OA absorbed in fog droplets can be easier to scavenger by wet deposition in the atmosphere, which may increase the proportion of relatively fresh OA by scavenged OA.

During the 7/18 event, which featured the most notable change in carbon oxidation state and overall decreases in all particle components, the organic content developed lower O:C and higher H:C over the course of the fog episode indicating a decrease in oxidation (Fig. 7a). This oxidation process clearly falls along a slope of $1$ (see linear regression for the fog period in dark green), which is indicative of carboxylic acid or hydroxycarbonyl loss (when moving “up” the line toward higher H:C and lower O:C; Heald et al., 2010). In addition, during fog events, H:C and the fraction of HOA was increased and then decreased, while O:C and the fraction of SOA decreased and only O:C and LO-OOA increased (Fig. 7). This behavior is consistent with particles produced from drying aerosolized cloud water during laboratory photooxidation (Schurman et al., 2018). Interestingly, although not as relevant to the discussion of fog aerosol processing herein, the particles follow approximately the same slope toward higher O:C and lower H:C after the fog event as they are re-oxidized in the atmosphere, which is likely by adding carboxylic acid or hydroxycarbonyl groups (Fig. 7b).

3.6. Changes in aerosol physical properties during fog

The particle size distribution and size-resolved chemical compositions before (02:30–03:00 h) and after (08:00–08:30 h) the 6/30 fog event were compared (Fig. 8 and Fig. S6). Unfortunately, the analysis...
range of the SMPS was 10.4–469.8 nm, which could not be precisely compared with that of the HR-ToF-AMS, but it was used for understanding the particle distribution below 500 nm (Fig. 8a). The mode diameter before the fog event was 111.4 nm (3566 particles cm⁻³), and the mode diameter increased to 138.2 nm (5831 particles cm⁻³) after the fog event. The increase in the number of particles was observed to be below the 300 nm size. The all size-resolved chemical composition of NR-PM₁ were decreased within the 300–1000 nm (Fig. 8b–e) particle range, probably because of wet deposition. The concentrations of Org, SO₄²⁻, and NH₄⁺ were increased below 300 nm. The concentration of NO₃⁻ did not increase below 300 nm after the fog event. During the fog event, gaseous compounds (e.g., VOCs, H₂SO₄, and NH₃) react to the condensation mode particles (Ge et al., 2012), and secondary aerosol, such as (NH₄)₂SO₄ and NH₄HSO₄, is formed by aqueous phase reactions in the fog droplet. This supports the increase in Org, NO₃⁻, SO₄²⁻, and number concentrations below 300 nm (Fig. 8). In addition, the correlation between SO₄²⁻ and NH₄⁺ after the fog event was observed to be 0.82 (correlation coefficient), while the correlation between NO₃⁻ and NH₄⁺ was 0.2, suggesting internal mixing of SO₄²⁻ and NH₄⁺ in the particle.

4. Conclusions

The aerosol physical and chemical measurements were conducted at
Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgements

This work was supported by grants from the National Research Foundation of Korea (Grant No. NRF-2019R1A1A1007947). The additional data analysis was supported by the National Institute of Environmental Research (NIER-2020-04-02) and the extended experiment was conducted by Korea Basic Science Institute (National research Facilities and Equipment Center) grant funded by the Ministry of Education (2019R1A6C1020041).

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.atmosenv.2022.119091.

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