

Source-Resolved Volatility and Oxidation State Decoupling in Wintertime Organic Aerosols in Seoul

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Abstract.

Organic aerosols (OA) are key components of wintertime urban haze, but the relationship between their oxidation state and volatility—critical for understanding aerosol evolution and improving model predictions—remains poorly constrained. While oxidation–volatility decoupling has been observed in laboratory studies, field-based evidence under real-world conditions is scarce, particularly during severe haze episodes. This study presents a field-based investigation of OA sources and their volatility characteristics in Seoul during a winter haze period, using a thermodenuder coupled with a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS).

Positive matrix factorization resolved six OA factors: hydrocarbon-like OA, cooking, biomass burning, nitrogen-containing OA (NOA), less-oxidized oxygenated OA (LO-OOA), and more-oxidized OOA (MO-OOA). Despite having the highest oxygen-to-carbon ratio (~ 1.15), MO-OOA exhibited unexpectedly high volatility, indicating a decoupling between oxidation state and volatility. We attribute this to fragmentation-driven aging and autoxidation under stagnant conditions with limited OH exposure. In contrast, LO-OOA showed lower volatility and more typical oxidative behavior.

Additionally, NOA—a rarely resolved factor in wintertime field studies—was prominent during cold, humid, and stagnant conditions and exhibited chemical and volatility features similar to biomass burning OA, suggesting a shared combustion origin and meteorological sensitivity.

These findings provide one of the few field-based demonstrations of oxidation–volatility decoupling in ambient OA and highlight how source-specific properties and meteorology influence OA evolution. The results underscore the need to refine OA representation in chemical transport models, especially under haze conditions.

Keywords: Organic aerosol volatility, HR-ToF-AMS, Thermodenuder, elemental ratios, aging, fragmentation

30 1 Introduction

31 Atmospheric aerosols affect both human health and the environment by reducing visibility (Ghim et al., 2005; Zhao
32 et al., 2013) and contributing to cardiovascular and respiratory diseases (Hamanaka et al., 2018; Manisalidis et al.,
33 2020). In addition, aerosols play a significant role in climate change by scattering or absorbing solar radiation and
34 modifying cloud properties (IPCC AR6). Among the various aerosol components—including sulfate, nitrate,
35 ammonium, chloride, crustal materials, and water—organic aerosols (OA) are particularly important to characterize,
36 as they account for 20–90% of submicron particulate matter (Zhang et al., 2007). Identifying OA sources and
37 understanding their behavior are critical for effective air quality management; however, this is particularly
38 challenging due to the vast diversity and dynamic nature of OA compounds, which originate from both natural and
39 anthropogenic sources. Unlike inorganic aerosols, organic aerosols (OAs) evolve continuously through complex
40 atmospheric reactions, influenced by emission sources, meteorological conditions, and aerosol properties (Jimenez
41 et al., 2009; Hallquist et al., 2009; Robinson et al., 2007; Donahue et al., 2006; Ng et al., 2010; Cappa and Jimenez,
42 2010).

43 Volatility is a key parameter for characterizing organic aerosol (OA) properties, as it governs gas-to-particle
44 partitioning behavior and directly influences particle formation yields (Sinha et al., 2023). The classification of OA
45 species based on their volatility—from extremely low-volatility (ELVOC) to semi-volatile (SVOC) and
46 intermediate-volatility (IVOC) compounds—is central to the conceptual framework of secondary OA (SOA)
47 formation and growth (Donahue et al., 2006). It also affects atmospheric lifetimes and human exposure by
48 determining how long aerosols remain suspended in the atmosphere (Glasius and Goldstein, 2016). Therefore,
49 accurately capturing OA volatility is essential for improving predictions of OA concentrations and their
50 environmental and health impacts. However, chemical transport models often significantly underestimate OA mass
51 compared to observations (Matsui et al., 2009; Jiang et al., 2012; Li et al., 2017), largely due to incomplete
52 precursor inventories and simplified treatment of processes affecting OA volatility. For instance, aging—through
53 oxidation reactions such as functionalization and fragmentation—can significantly alter volatility by changing OA
54 chemical structure (Robinson et al., 2007; Zhao et al., 2016). Early volatility studies primarily utilized thermal
55 denuders (TD) coupled with various detection instruments to investigate the thermal properties of bulk OA
56 (Huffman et al., 2008). The subsequent coupling of TD with the Aerosol Mass Spectrometer allowed for
57 component-resolved volatility measurements, providing critical, quantitative insight into the properties of OA
58 factors (e.g., SV-OOA vs. LV-OOA) across different regions (Paciga et al., 2016; Cappa and Jimenez, 2010). These
59 component-resolved volatility data are often used to constrain the Volatility Basis Set (VBS)—the current state-of-

the-art framework for modeling OA partitioning and evolution (Donahue et al., 2006). However, a limitation in many field studies is that the TD-AMS thermogram data are rarely translated into quantitative VBS distributions for individual OA factors, which limits their direct use in chemical transport models. Furthermore, the volatility of OOA during extreme haze conditions, where the expected inverse correlation between oxidation (O:C) and volatility can break down (Jimenez et al., 2009), remains poorly characterized, particularly in East Asia's highly polluted winter environments. A recent study in Korea further highlighted the importance of accounting for such processes when interpreting OA volatility under ambient conditions (Kang et al., 2022). Given its central role in OA formation, reaction, and atmospheric persistence, volatility analysis is critical for bridging the gap between measurements and model performance.

Traditionally, due to the complexity and variability of OA, the oxygen-to-carbon (O:C) ratio has been used as a proxy for estimating volatility. In general, higher O:C values indicate greater oxidation and lower volatility (Jimenez et al., 2009). Accordingly, many field studies classify oxygenated OA (OOA) into semi-volatile OOA (SV-OOA) and low-volatility OOA (LV-OOA) based on their O:C ratios (Ng et al., 2010; Huang et al., 2010; Mohr et al., 2012). However, this relationship is not always straightforward. Fragmentation during oxidation can increase both O:C and volatility simultaneously, disrupting the expected inverse correlation (Jimenez et al., 2009). In laboratory experiments, yields of highly oxidized SOA have been observed to decrease due to fragmentation (Xu et al., 2014; Grieshop et al., 2009). These findings suggest that while O:C can offer useful insights, it is insufficient alone to represent OA volatility. Direct volatility measurements, especially when paired with chemical composition data, are necessary to improve our understanding of OA sources and aging processes.

In this study, we investigate the sources and volatility characteristics of OA in Seoul during winter. Wintertime OA presents additional challenges due to its high complexity. During winter, emissions from combustion sources such as biomass burning and residential heating significantly increase, contributing large amounts of primary OA (Kim et al., 2017). Meanwhile, low ambient temperatures and reduced photochemical activity affect the formation and evolution of secondary OA (SOA). Frequent haze events further complicate the aerosol properties by extending aging times and increasing particle loadings. These overlapping sources and atmospheric conditions make winter OA particularly difficult to characterize and predict. Despite Seoul's significance for air quality management, comprehensive studies on OA volatility during winter remain limited. To address these goals, we conducted real-time, high-resolution measurements using a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS) coupled with a thermodenuder (TD). The objectives of this study are to: (1) improve the understanding of

89 wintertime OA in Seoul, (2) characterize the volatility of OA associated with different sources, and (3) explore the
90 relationship between OA volatility and chemical composition.

91 **2 Experimental methods**

92 **2.1 Sampling Site and Measurement Period**

93 We conducted continuous real-time measurements in Seoul, South Korea, from 28 November to 28 December 2019.
94 The sampling site was located in the northeastern part of the city (37.60° N, 127.05° E), approximately 7 km from
95 the city center, surrounded by major roadways and mixed commercial–residential land use. Air samples were
96 collected at an elevation of approximately 60 meters above sea level, on the fifth floor of a building. A detailed site
97 description has been reported previously for winter Seoul (Kim et al., 2017). During this period, the average
98 ambient temperature was 1.76 ± 4.3 °C, and the average relative humidity (RH) was $56.9 \pm 17.5\%$, based on data
99 from the Korea Meteorological Administration (<http://www.kma.go.kr>).

100 **2.2 Instrumentation and Measurements**

101 The physico-chemical properties of non-refractory PM₁ (NR-PM₁) species—including sulfate, nitrate, ammonium,
102 chloride, and organics—were measured using an Aerodyne high-resolution time-of-flight aerosol mass
103 spectrometer (HR-ToF-AMS) (DeCarlo et al., 2006). PM₁ mass in this study is taken as NR-PM₁ (from AMS) +
104 black carbon (BC; measured by MAAP), which is appropriate for winter Seoul where refractory PM₁ (metal/sea-
105 salt/crustal) is minor and dust events were excluded (e.g., Kim et al., 2017; Nault et al., 2018; Kang et al., 2022;
106 Jeon et al., 2023). Data were acquired at 2.5-minute intervals, alternating between V and W modes. The V mode
107 provides higher sensitivity but lower resolution, suitable for mass quantification, whereas the W mode offers higher
108 mass resolution but lower sensitivity, used here for OA source apportionment. Simultaneously, black carbon (BC)
109 concentrations were measured at 1-minute intervals using a multi-angle absorption photometer (MAAP; Thermo
110 Fisher Scientific, Waltham, MA, USA). Total PM₁ mass was calculated as the sum of NR-PM₁ and BC.

111 Hourly trace gas concentrations (CO, O₃, NO₂, SO₂) were obtained from the Gireum air quality monitoring station
112 (37.61° N, 127.03° E), managed by the Seoul Research Institute of Public Health and Environment. Meteorological

113 data (temperature, RH, wind speed/direction) were collected from the nearby Jungreung site (37.61° N, 127.00° E).
 114 All data are reported in Korea Standard Time (UTC+9).

115 To examine aerosol volatility, a thermodenuder (TD; Envalytix LLC) was installed upstream of the HR-ToF-AMS.
 116 Details are provided in Supplementary Section S1 Kang et al. (2022). Briefly, ambient flow alternated every 5
 117 minutes between a TD line and a bypass line at 1.1 L min⁻¹. Residence time in the TD line was ~6.3 s. The TD
 118 setup included a 50 cm heating section followed by an adsorption unit. Heated particles were stripped of volatile
 119 species, while the downstream carbon-packed section prevented recondensation. TD temperature cycled through
 120 12 steps (30 to 200 °C), with each step lasting 10 min (total cycle = 120 min). AMS V and W modes were alternated
 121 during the same cycle. The heater was pre-adjusted to the next temperature while the bypass was active.
 122

123 **2.3 Data Analysis**

124

125 **2.3.1 Data analysis and OA Source Apportionment**

126 HR-AMS data were processed using SQUIRREL v1.65B and PIKA v1.25B. Mass concentrations of non-refractory
 127 PM₁ (NR-PM₁) species were derived from V-mode data, while high-resolution mass spectra (HRMS) and the
 128 elemental composition of organic aerosols (OA) were obtained from W-mode data. NR-PM₁ quantification
 129 followed established AMS protocols (Ulbrich et al., 2009; Zhang et al., 2011). Both the bypass and TD streams
 130 were processed using a time-resolved, composition-dependent collection efficiency CE(t) following Middlebrook
 131 et al. (2012). TD heating can modify particle water and phase state/mixing and thereby influence CE beyond
 132 composition (Huffman et al., 2009), but prior TD-AMS studies indicate that such effects are modest and largely
 133 multiplicative, which do not distort thermogram shapes or T₅₀ ordering (Faulhaber et al., 2009; Cappa & Jimenez,
 134 2010). In our data, the CE(t) statistics for the two lines were similar (campaign-average CE: TD = 0.55 ± 0.08;
 135 bypass = 0.53 ± 0.04; Δ = 0.02 ≈ 3.7%, below the combined uncertainty ≈ 0.09). We therefore report volatility
 136 metrics with these line-specific CE(t) corrections applied and interpret potential residual CE effects as minor. For
 137 organics, elemental ratios (O:C, H:C, and OM/OC) were calculated using the Improved-Ambient (IA) method
 138 (Canagaratna et al., 2015). Positive Matrix Factorization (PMF) was applied to the HRMS of organics using the
 139 PMF2 algorithm (v4.2, robust mode) (Paatero and Tapper, 1994). The HRMS and corresponding error matrices
 140 from PIKA were analyzed using the PMF Evaluation Tool v2.05 (Ulbrich et al., 2009). Data pretreatment followed
 141 established protocols (Ulbrich et al., 2009; Zhang et al., 2011).

A six-factor solution ($f_{\text{Peak}} = 0$; $Q/Q_{\text{expected}} = 3.56$) was selected as optimal (Fig. S1). The resolved OA sources included hydrocarbon-like OA (HOA; 14%; O:C = 0.13), cooking-related OA (COA; 21%; O:C = 0.18), nitrogen-enriched OA (NOA; 2%; O:C = 0.22), biomass-burning OA (BBOA; 13%; O:C = 0.25), less-oxidized oxygenated OA (LO-OOA; 30%; O:C = 0.68), and more-oxidized oxygenated OA (MO-OOA; 20%; O:C = 1.15) (Figs. S2 and S3). Alternative five- and seven-factor solutions were also evaluated. In the five-factor solution, the biomass burning source was not clearly resolved and appeared to be distributed across multiple factors. In the seven-factor solution, BBOA was further split into two separate factors without clear distinction or added interpretive value, making the six-factor solution the most physically meaningful and interpretable (Figs. S4 and S5). [To ensure the statistical robustness of this solution, we calculated uncertainties for each PMF factor using the bootstrap method \(100 iterations\) with the PET toolkit \(v2.05\) \(EPA, 2014; Xu et al., 2018; Srivastava et al., 2021\) \(Table S2 and Fig. S13\).](#)

2.3.2 Thermogram and Volatility Estimation

The chemical composition dependent mass fraction remaining (MFR) was derived at each TD temperature by dividing the corrected mass concentration of the TD line [p] by the average of the adjacent bypass lines [p-1] and [p+1]. Thermograms were corrected for particle loss, estimated using reference substances like NaCl, which exhibit minimal evaporation (Huffman et al., 2009; Saha et al., 2014; Kang et al., 2022). OA factor concentrations at each TD temperature were derived via multivariate linear regression between post-TD HRMS and ambient OA factor HRMS profiles as described in Zhou et al., 2017.

Volatility distributions were modeled using the thermodenuder mass transfer model from Riipinen et al. (2010) and Karnezi et al. (2014), implemented in Igor Pro 9 (Kang et al., 2022). OA mass was distributed into eight logarithmic saturation concentration bins (C^* : 1000 to $0.0001 \mu\text{g m}^{-3}$). Modeled MFRs were fit to observations using Igor's "FuncFit" function, repeated 1,000 times per OA factor to determine best-fit results. The model assumes no thermal decomposition and includes adjustable parameters: mass accommodation coefficient (α_m) and enthalpy of vaporization (ΔH_{exp}), randomly sampled within literature-based ranges (Table S1).

3 Results and discussion

169 3.1 Overview of PM₁ Composition and OA Sources

170 We conducted continuous measurements from 28 November to 28 December 2019, characterizing a winter period
171 with a mean PM₁ concentration of $27.8 \pm 15.3 \mu\text{g m}^{-3}$. This concentration is characterized as moderate; it closely
172 matches historical winter PM₁ means in Seoul (Kim et al., 2017) and implies an equivalent PM_{2.5} concentration is
173 about $34.8 \mu\text{g m}^{-3}$ (using a Korea-specific PM₁/PM_{2.5} ≈ 0.8 (Kwon et al., 2023), which is near the national 24-h PM_{2.5}
174 standard ($35 \mu\text{g m}^{-3}$) (AirKorea). The full co-evolution of PM₁, gaseous pollutants, and meteorological conditions
175 is provided in Fig. S6, showing an average ambient temperature of $1.76 \pm 4.3^\circ\text{C}$ and average relative humidity (RH)
176 of $56.9 \pm 17.5\%$ during the study.

177 Figure 1 summarizes the overall non-refractory submicron aerosol (NR-PM₁) composition and the identified OA
178 factors. Organics (41%) and nitrate (30%) were the most abundant chemical components of PM₁, followed by
179 ammonium (12%), sulfate (10%), BC (5%), and chloride (3%) (Fig. 1a). Among the organic aerosols, six OA
180 factors were identified during the winter of 2019: hydrocarbon-like OA (HOA; 14%; O:C = 0.13), cooking-related
181 OA (COA; 21%; O:C = 0.18), nitrogen-enriched OA (NOA; 2%; O:C = 0.22), biomass burning OA (BBOA; 13%;
182 O:C = 0.25), and two types of secondary organic aerosols—less-oxidized oxygenated OA (LO-OOA; 30%; O:C =
183 0.68) and more-oxidized oxygenated OA (MO-OOA; 20%; O:C = 1.15) (Fig. 1e and Fig. S2). These compositions
184 are consistent with previous wintertime observations in Kim et al. (2017), with the exception of [newly resolved](#)
185 [NOA source. In the following sections, we describe each OA factor in the order of secondary OA \(SOA\), primary](#)
186 [OA \(POA\) and, and finally introduce NOA, which—while related to combustion POA—emerged as a distinct,](#)
187 [nitrogen-rich factor under the winter conditions of this study.](#)

188 PM₁ mass concentrations varied widely, ranging from 4.61 to $91.4 \mu\text{g m}^{-3}$, largely due to two severe haze episodes
189 that occurred between December 7–12 and December 22–26 (Fig. 1). During these episodes, average
190 concentrations increased significantly, driven primarily by elevated levels of nitrate and organic aerosols—
191 particularly MO-OOA and NOA (Fig. 1f,g). Back-trajectory clustering shows frequent short-range recirculation
192 over the Seoul Metropolitan Area during haze (Cluster 1; Fig. S8), and the time series indicates persistently low
193 surface wind speeds during these periods (1.73 ± 0.89 vs. 2.34 ± 1.18 (clean)) (Fig. S6). [These patterns indicate](#)
194 [stagnation-driven accumulation of local emissions, consistent with the simultaneous increase of MO-OOA and](#)
195 [NOA that are examined in detail in subsequent sections. together pointing to stagnation-driven accumulation of](#)
196 [local emissions; the concurrent increases in MO-OOA and NOA are therefore consistent with enhanced in-city](#)

~~formation under stagnant conditions.~~ Such haze episodes, characterized by local emission buildup and secondary aerosol production, are a typical wintertime feature, as also reported in Kim et al. (2017).

3.1.2.1 Secondary organic aerosols (SOA)

In this study, two OOA factors—more-oxidized OOA (MO-OOA) and less-oxidized OOA (LO-OOA)—were identified, together accounting for approximately half of the total organic aerosol (OA) mass. This fraction is notably higher than that reported in previous wintertime urban studies (Kim et al., 2017; Zhang et al., 2007). Both OOAs exhibited characteristic mass spectral features, including prominent peaks at m/z 44 (CO_2^+) and m/z 43 ($\text{C}_2\text{H}_3\text{O}^+$), which are widely recognized as markers of oxygenated organics (Fig. S2e, S3f). The oxygen-to-carbon (O:C) ratios for MO-OOA and LO-OOA were 1.15 and 0.68, respectively, indicating both factors are highly oxidized relative to the primary OA factors (HOA, COA, BBOA) and that MO-OOA is substantially more oxidized than LO-OOA. ~~highly oxidized chemical compositions.~~ The O:C ratio of MO-OOA was especially elevated, exceeding those reported in previous Seoul campaigns—0.68 in winter 2015 (Kim et al., 2017), 0.99 in spring 2019 (Kim et al., 2020), and 0.78 in fall 2019 (Jeon et al., 2023)—while the LO-OOA ratio was within a similar range.

~~MO-OOA showed strong correlations with secondary inorganic species such as nitrate ($r = 0.90$), ammonium ($r = 0.92$), and sulfate ($r = 0.81$), consistent with its formation through regional and local photochemical aging processes (Fig. S3). In contrast, LO-OOA exhibited only modest correlations with sulfate, nitrate, and ammonium ($r = 0.50$, 0.51 , and 0.42 , respectively). This weaker coupling indicates that LO-OOA represents a less aged oxygenated OA component (fresh SOA), distinguishable from the more aged, highly processed MO-OOA which tracks closely with secondary inorganic species. Regarding potential primary influence, LO-OOA does not exhibit a pronounced m/z 60 (levoglucosan) signal (Figs. S2 and 9). While the levoglucosan marker (f_{60}) is known to diminish with atmospheric aging and can become weak or undetectable downwind (Hennigan et al., 2010; Cubison et al., 2011), the absence of a distinct peak combined with the separation from inorganic salts suggests that LO-OOA is best characterized as freshly formed secondary organic aerosol likely originating from the rapid oxidation of local anthropogenic precursors, rather than a primary emission source.~~

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photochemical aging processes (Fig. S3). In contrast, LO-OOA exhibited only modest correlations with sulfate, nitrate, and ammonium ($r = 0.50, 0.51$, and 0.42 , respectively), indicating that suggesting LO-OOA represents a less aged oxygenated OA component with possible influence from semi-primary precursors, rather than a factor strongly tied to secondary inorganic aerosol formation. additional contributions from semi-primary sources not closely linked to inorganic secondary formation (e.g., cooking, traffic, biomass burning). LO-OOA does not exhibit a pronounced m/z 60 (levoglucosan) signal (Fig. S2); however, the levoglucosan marker (f_{60}) is known to diminish with atmospheric aging and can become weak or undetectable downwind (Hennigan et al., 2010; Cubison et al., 2011). Taken together, the weaker coupling with secondary inorganics and the absence of a strong m/z 60 peak indicate that LO-OOA is a mixture of aged secondary organics and aged semi-primary urban emissions, while a contribution from aged biomass burning influence cannot be ruled out.

3.1.3 Primary organic aerosols (POA)

3.1.2 Primary organic aerosols (POA)

Three primary organic aerosol (POA) factors were identified in this study: hydrocarbon-like OA (HOA), cooking-related OA (COA), and biomass burning OA (BBOA). These three components exhibited mass spectral and temporal characteristics consistent with previous observations in Seoul and other urban environments. HOA was characterized by dominant alkyl fragment ions ($C_nH_{2n+1}^+$ and $C_nH_{2n-1}^+$; Fig. S2a) and a low O:C ratio (0.13), consistent with traffic-related emissions (0.05–0.25) (Canagaratna et al., 2015). It showed strong correlations with vehicle-related ions $C_3H_7^+$ ($r = 0.79$) and $C_4H_9^+$ ($r = 0.86$) (Kim et al., 2017; Canagaratna et al., 2004; Zhang et al., 2005), and exhibited a distinct morning rush hour peak (06:00–08:00), followed by a decrease likely driven by boundary layer expansion (Fig. S3a).

COA, accounting for 21% of OA, showed higher contributions from oxygenated ions than HOA, with tracer peaks at m/z 55, 84 and 98 (Fig. S2b) consistent with cooking emissions (Sun et al., 2011). COA showed an enhanced signal at m/z 55 relative to m/z 57, with a 55/57 ratio of 3.11, substantially larger than that of HOA (1.10). This elevated ratio is consistent with previously reported AMS COA spectra in urban environments (e.g., Allan et al., 2010; Mohr et al., 2012; Sun et al., 2011), supporting our factor assignment. It correlated strongly with cooking-

related ions such as $\text{C}_3\text{H}_3\text{O}^+$ ($r = 0.94$), $\text{C}_5\text{H}_8\text{O}^+$ ($r = 0.96$), and $\text{C}_6\text{H}_{10}\text{O}^+$ ($r = 0.98$) (Fig. S3h), and displayed prominent peaks during lunch and dinner hours, reflecting typical cooking activity patterns.

BBOA was identified based on characteristic ions at m/z 60 ($\text{C}_2\text{H}_4\text{O}_2^+$) and 73 ($\text{C}_3\text{H}_5\text{O}^+$), both of which are associated with levoglucosan—a well-established tracer for biomass burning (Simoneit et al., 2002). Its relatively high f_{60} and low f_{44} values (Fig. S8a9) indicate that the BBOA observed in this study was relatively fresh and had not undergone extensive atmospheric aging (Cubison et al., 2011). Furthermore, BBOA exhibited moderate correlations with NOA in both diurnal profiles and time series (Fig. 2), particularly with nitrogen-containing ions such as $\text{C}_2\text{H}_4\text{N}^+$ ($r = 0.67$) and $\text{C}_2\text{H}_6\text{N}^+$ ($r = 0.56$) (Fig. 2 and S3), which are also dominant peaks in the NOA mass spectrum. This overlap suggests a potential shared emission source or co-emission scenario, consistent with the co-emission of both organic aerosols and reduced nitrogen-containing compounds. Regarding source location, several pathways can influence Seoul's biomass burning signature. First, urban/peri-urban small-scale burning (e.g., solid-fuel use in select households, restaurant charcoal use, and intermittent waste burning) has been reported and can enhance BBOA locally (Kim et al., 2017). Second, nearby agricultural-residue burning in surrounding provinces occurs seasonally and can episodically impact the metropolitan area (Han et al., 2022). Third, regional transport from upwind regions (e.g., northeastern China/North Korea) can bring biomass burning influenced air masses under northerly/northwesterly flow (Lamb et al., 2018; Nault et al., 2018). In this dataset, the nighttime and early-morning enhancements, the BBOA–NOA co-variation, and trajectory clusters showing regional recirculation indicate a predominantly local/near-source contribution during the study period (Yoo et al., 2024), with episodic non-local influences remaining possible.

3.1.43 Nitrogen-containing organic aerosol (NOA)

A distinct nitrogen-containing organic aerosol (NOA) factor was resolved in this study, whereas earlier wintertime AMS–PMF analyses in Seoul did not isolate such a component. Unlike previous wintertime aerosol studies in Seoul, this study successfully resolved a nitrogen-containing organic aerosol (NOA) factor by applying positive matrix factorization (PMF) to high-resolution AMS data. NOA contributed approximately 2% of the total organic aerosol (OA) mass comparable to urban observations in

Guangzhou (3%; Chen et al., 2021), Pasadena (5%; Hayes et al., 2013), and New York (5.8%; Sun et al., 2011).

Detection of particulate NOA using real time measurement has been challenging due to its low concentration and high volatility. Although Baek et al. (2022) identified nitrogen-containing species in Seoul via year-round filter-based molecular analysis, PMF-based resolution of NOA in real time has not been previously reported. The successful identification in this study is likely attributable to favorable winter meteorological conditions—specifically low temperatures (-0.24°C) and persistently high relative humidity ($\sim 57\%$) compared to the 2017 winter season (Kim et al., 2017)—that enhanced gas-to-particle partitioning of semi-volatile amines, thereby enabling their detection (Fig. S2). NOA concentrations frequently exceeded $1\text{ }\mu\text{g m}^{-3}$ when RH surpassed 60% (Fig. 2), supporting the importance of RH-driven partitioning and the subsequent formation of low-volatility aminium salts (Rovelli et al., 2017). Although extremely low temperatures may inhibit NOA formation due to the transition of aerosol particles into solid phase (Ge et al., 2011; Srivastava et al., 2022), the combination of consistently cold and humid conditions during the measurement period likely promoted the partitioning of semi-volatile amines into the particle phase.

In addition, episodic haze events further elevated NOA levels, increasing its contribution to OA from 1% during clean periods to as much as 3% (Fig. 1f-h). These high-concentration events likely improved the signal-to-noise ratio, facilitating PMF resolution. Back-trajectory analysis linked these events to regional recirculation patterns (Cluster 1, Fig. S7), suggesting a predominantly local origin—consistent with the short atmospheric lifetimes and high reactivity of most amines (Ge et al., 2011; Nielsen et al., 2012; Hanson et al., 2014).

The NOA factor exhibited the highest nitrogen-to-carbon (N:C) ratio (0.22) and the lowest oxygen-to-carbon (O:C) ratio (0.19) among all POA factors (Fig. S2), indicating a chemically reduced, nitrogen-rich composition. The factor represents semi-volatile, reduced nitrogen species that originate from primary urban combustion sources but whose observed mass in the particle phase is enhanced by rapid secondary partitioning and salt formation (Ge et al., 2011; You et al., 2014). The NOA mass spectrum was dominated by amine-related fragments including m/z 30 (CH_4N^+), 44 ($\text{C}_2\text{H}_6\text{N}^+$), 58 ($\text{C}_3\text{H}_8\text{N}^+$), and 86 ($\text{C}_5\text{H}_{12}\text{N}^+$) (Fig. 3a). The spectral signature of the factor is defined by the characteristic dominance of the m/z 44 fragment, which typically serves as the primary marker for dimethylamine (DMA)-related species, closely followed by m/z 58 (trimethylamine, TMA) and m/z 30 (methylamine, MA). This profile is in strong agreement with NOA factors resolved via PMF in other polluted

environments. For instance, the dominance of m/z 44 and m/z 30 aligns with amine factors reported in New York City (Sun et al., 2011) and Pasadena, California (Hayes et al., 2013). This DMA-dominated signature is also consistent with seasonal characterization of organic nitrogen in Beijing (Xu et al., 2017) and Po Valley, Italy (Saarikoski et al., 2012), reinforcing the common chemical signature of reduced organic nitrogen across diverse urban and regional environments. Furthermore, the presence of non-negligible signals at m/z 58 and m/z 86 supports the contribution of slightly larger alkylamines, a pattern that aligns well with established AMS laboratory reference spectra for these reduced nitrogen compounds (Ge et al., 2011; Silva et al., 2008).

These amines are commonly emitted during the combustion of nitrogen-rich biomass and proteinaceous materials and are frequently associated with biomass-burning emissions (Ge et al., 2011). Previous molecular analyses in Seoul also indicate DMA, MA, and TMA as the dominant amine species in December (Baek et al., 2022). While other amines such as triethylamine (TEA), diethylamine (DEA), and ethylamine (EA) may contribute via industrial/solvent pathways (e.g., chemical manufacturing, petrochemical corridors, wastewater treatment), our HR-AMS spectra are dominated by small alkylamine fragments (m/z 30, 44, 58, 86) and the diurnal behavior covaries with combustion markers ([below Fig. 2](#)), indicating a primarily combustion-linked influence. Nevertheless, recent urban measurements and sector-based analyses show that industrial activities can contribute measurable amines in cities (Tiszenkel et al., 2024; Zheng et al., 2015; Mao et al., 2018; Shen et al., 2017; Yao et al., 2016). Accordingly, a minor NOA contribution from solvent/industrial amines cannot be excluded. [NOA exhibited a nighttime-early-morning enhancement \(Fig. 2a\), similar to BBOA, indicating that both factors are influenced by wintertime combustion and residential heating, which are known sources of small alkylamines and amides \(You et al., 2014; Yao et al., 2016\). Strong correlations of NOA with \$\text{CH}_4^+ \text{N}^+\$ \(\$r = 0.95\$ \) and \$\text{C}_2\text{H}_6\text{N}^+\$ \(\$r = 0.91\$ \) \(Fig. 2\) further support the presence of reduced-nitrogen species associated with these combustion activities. However, the time series of NOA and BBOA are not strongly correlated \(Fig. 2 and Fig. S7\). This contrast reflects their differing behaviors: BBOA follows a relatively regular daily emission pattern, whereas NOA appears predominantly during stagnant haze periods \(Fig. 1\) when cold, humid, and low-wind conditions allow semi-volatile amines to partition to the particle phase and form low-volatility aminium salts. Thus, NOA in wintertime Seoul likely reflects a combination of shared primary combustion influences and enhanced secondary processing of amine-containing precursors under meteorological conditions that favor partitioning and accumulation. Supporting this, NOA exhibited a diurnal pattern similar to that of BBOA, with both peaking at night and in the early morning \(Fig. 2a\), suggesting shared sources or formation mechanisms. Biomass burning under cold, oxygen-limited conditions is known to emit various amines and amides \(You et al., 2014; Yao et al., 2016\), which may contribute directly to](#)

NOA or serve as precursors for its secondary formation. Strong correlations with CH_3N^+ ($r = 0.95$) and $\text{C}_2\text{H}_5\text{N}^+$ ($r = 0.91$) (Fig. 2) further support the presence of reduced nitrogen compounds, typically associated with residential fuel combustion and wintertime heating. However, the time series of NOA and BBOA were not well correlated (Fig. 2 and S7), likely because NOA episodes preferentially occurred during haze periods under stagnant conditions (Fig. 1), whereas BBOA emissions tend to follow a more regular, daily emission pattern. Under cold, humid, and stagnant conditions, these semi-volatile amines can readily partition into the particle phase and form low-volatility aminium salts, enhancing the observed NOA signal. Taken together, these results suggest that NOA during wintertime in Seoul is strongly influenced by a combination of combustion-related primary emissions and subsequent atmospheric processing of amine-containing species, facilitated by seasonally favorable conditions.

Detection of particulate NOA using real time measurement has been challenging due to its low concentration and high volatility. Although Baek et al. (2022) identified nitrogen-containing species in Seoul via year-round filter-based molecular analysis, PMF-based resolution of NOA in real time has not been previously reported. The successful identification in this study is likely attributable to favorable winter meteorological conditions—specifically low temperatures (-0.24°C) and persistently high relative humidity ($\sim 57\%$) compared to the 2017 winter season (Kim et al., 2017)—that enhanced gas-to-particle partitioning of semi-volatile amines, thereby enabling their detection (Fig. S2). NOA concentrations frequently exceeded $1\ \mu\text{g m}^{-3}$ when RH surpassed 60% (Fig. 2), supporting the importance of RH-driven partitioning and the subsequent formation of low-volatility aminium salts (Rovelli et al., 2017). Although extremely low temperatures may inhibit NOA formation due to the transition of aerosol particles into solid phase (Ge et al., 2011; Srivastava et al., 2022), the combination of consistently cold and humid conditions during the measurement period likely promoted the partitioning of semi-volatile amines into the particle phase. In addition, episodic haze events further elevated NOA levels, increasing its contribution to OA from 1% during clean periods to as much as 3% (Fig. 1f–h). These high-concentration events likely improved the signal-to-noise ratio, facilitating PMF resolution. Back-trajectory clustering indicates that NOA-enhanced events were dominated by short-range recirculation (Cluster 1; Fig. S7), consistent with the short atmospheric lifetimes and high reactivity of alkylamines (Nielsen et al., 2012; Hanson et al., 2014). Overall, the factor reflects semi-volatile, reduced-nitrogen species originating from primary urban combustion sources, with their observed particle-phase mass amplified by rapid secondary partitioning and salt formation under seasonally favorable conditions.

NOA contributed approximately 2 % of total OA, comparable to urban contributions reported in Guangzhou (3 %; Chen et al., 2021), Pasadena (5 %; Hayes et al., 2013), and New York (5.8 %; Sun et al., 2011). These similarities suggest that the NOA factor observed in Seoul reflects a broader class of urban wintertime reduced-nitrogen aerosols rather than a site-specific anomaly. In most urban environments, the detectability of NOA appears to depend strongly on the interplay between emission strength, stagnation, and humidity—which together govern the particle-phase partitioning of volatile amines.

~~atmospheric processing of amine-containing species, facilitated by seasonally favorable conditions.~~

~~3.1.2 Secondary organic aerosols (SOA)~~

~~In this study, two OOA factors—more oxidized OOA (MO-OOA) and less oxidized OOA (LO-OOA)—were identified, together accounting for approximately half of the total organic aerosol (OA) mass. This fraction is notably higher than that reported in previous wintertime urban studies (Kim et al., 2017; Zhang et al., 2007). Both OOAs exhibited characteristic mass spectral features, including prominent peaks at m/z 44 (CO_2^+) and m/z 43 ($\text{C}_2\text{H}_5\text{O}^+$), which are widely recognized as markers of oxygenated organics (Fig. S2e, S3f). The oxygen-to-carbon (O:C) ratios for MO-OOA and LO-OOA were 1.15 and 0.68, respectively, indicating highly oxidized chemical compositions. The O:C ratio of MO-OOA was especially elevated, exceeding those reported in previous Seoul campaigns—0.68 in winter 2015 (Kim et al., 2017), 0.99 in spring 2019 (Kim et al., 2020), and 0.78 in fall 2019 (Jeon et al., 2023)—while the LO-OOA ratio was within a similar range.~~

~~MO-OOA showed strong correlations with secondary inorganic species such as nitrate ($r = 0.90$), ammonium ($r = 0.92$), and sulfate ($r = 0.81$), consistent with its formation through regional and local photochemical aging processes (Fig. S3). In contrast, LO-OOA exhibited only modest correlations with sulfate, nitrate, and ammonium ($r = 0.50$, 0.51 , and 0.42 , respectively, suggesting additional contributions from semi-primary sources not closely linked to inorganic secondary formation (e.g., cooking, traffic, biomass burning). LO-OOA does not exhibit a pronounced m/z 60 (levoglucosan) signal (Fig. S2); however, the levoglucosan marker (f_{60}) is known to diminish with atmospheric aging and can become weak or undetectable downwind (Hennigan et al., 2010; Cubison et al., 2011). Taken together, the weaker coupling with secondary inorganics and the absence of a strong m/z 60 peak indicate that LO-OOA is a mixture of aged secondary organics and semi-primary urban emissions, while a contribution from aged biomass burning influence cannot be ruled out.~~

3.1.3 Primary organic aerosols (POA)

Three primary organic aerosol (POA) factors were identified in this study: hydrocarbon-like OA (HOA), cooking-related OA (COA), and biomass burning OA (BBOA). These three components exhibited mass spectral and temporal characteristics consistent with previous observations in Seoul and other urban environments. HOA was characterized by dominant alkyl fragment ions ($C_nH_{2n-1}^+$ and $C_nH_{2n-1}^+$; Fig. S2a) and a low O:C ratio (0.13), consistent with traffic-related emissions (0.05–0.25) (Canagaratna et al., 2015). It showed strong correlations with vehicle-related ions $C_3H_7^+$ ($r=0.79$) and $C_4H_9^+$ ($r=0.86$) (Kim et al., 2017; Canagaratna et al., 2004; Zhang et al., 2005), and exhibited a distinct morning rush hour peak (06:00–08:00), followed by a decrease likely driven by boundary layer expansion (Fig. S3a).

COA, accounting for 21% of OA, showed higher contributions from oxygenated ions than HOA, with tracer peaks at m/z 55, 84 and 98 (Fig. S2b) consistent with cooking emissions (Sun et al., 2011). It correlated strongly with cooking-related ions such as $C_3H_5O^+$ ($r=0.94$), $C_3H_7O^+$ ($r=0.96$), and $C_4H_9O^+$ ($r=0.98$) (Fig. S3h), and displayed prominent peaks during lunch and dinner hours, reflecting typical cooking activity patterns.

BBOA was identified based on characteristic ions at m/z 60 ($C_2H_4O_2^+$) and 73 ($C_3H_6O^+$), both of which are associated with levoglucosan—a well-established tracer for biomass burning (Simoneit et al., 2002). Its relatively high f_{OC} and low f_{tr} values (Fig. S8a) indicate that the BBOA observed in this study was relatively fresh and had not undergone extensive atmospheric aging (Cubison et al., 2011). Furthermore, BBOA exhibited moderate correlations with NOA in both diurnal profiles and time series (Fig. 2), particularly with nitrogen-containing ions such as $C_2H_4N^+$ ($r=0.67$) and $C_2H_6N^+$ ($r=0.56$) (Fig. 2 and S3), which are also dominant peaks in the NOA mass spectrum. This overlap suggests a potential shared emission source or co-emission scenario, consistent with the co-emission of both organic aerosols and reduced nitrogen-containing compounds. Regarding source location, several pathways can influence Seoul's biomass burning signature. First, urban/peri-urban small-scale burning (e.g., solid fuel use in select households, restaurant charcoal use, and intermittent waste burning) has been reported and can enhance BBOA locally (Kim et al., 2017). Second, nearby agricultural residue burning in surrounding provinces occurs seasonally and can episodically impact the metropolitan area (Han et al., 2022). Third, regional transport from upwind regions (e.g., northeastern China/North Korea) can bring biomass burning influenced air masses under northerly/northwesterly flow (Lamb et al., 2018; Nault et al., 2018). In this dataset, the nighttime and early morning enhancements, the BBOA–NOA co-variation, and trajectory clusters showing

~~regional recirculation indicate a predominantly local/near source contribution during the study period (Yoo et al., 2024), with episodic non-local influences remaining possible.~~

3.2 Volatility of Non-Refractory Species

Figure 4 presents thermograms of non-refractory (NR) species measured by HR-ToF-AMS. The mass fraction remaining (MFR) after thermodenuder (TD) treatment follows the typical volatility trend reported in previous studies (Xu et al., 2016; Kang et al., 2022; Jeon et al., 2023; Huffman et al., 2009): nitrate was the most volatile, followed by chloride, ammonium, organics, and sulfate. Nitrate showed the steepest decline with increasing temperature, with a T_{50} of ~ 67 °C—substantially higher than that of pure ammonium nitrate (~ 37 °C; Huffman et al., 2009). At 200 °C, $\sim 2\%$ of the initial nitrate signal remained (Fig. 4). Since pure ammonium nitrate fully evaporates well below this temperature (Huffman et al., 2009), this small residual fraction likely represents the least volatile portion of organic nitrates. Nitrate showed the steepest decline with temperature, with a T_{50} of ~ 67 °C—higher than pure ammonium nitrate (~ 37 °C; Huffman et al., 2009), suggesting contributions from less volatile species like organonitrates or metal nitrates (Feng et al., 2023). This small residual mass is consistent with previous observations of thermally refractory or semi-refractory nitrate species—including organonitrates with low effective saturation concentrations and nitrate associated with transition metal ions—which are incompletely vaporized in AMS instruments (Farmer et al., 2010; Xu et al., 2015; Canagaratna et al., 2007). As AMS cannot directly distinguish between inorganic and organic nitrate at high TD temperatures, this residual fraction reflects low volatility nitrate-containing material rather than ammonium nitrate itself. Nearly complete evaporation occurred by 200 °C ($\sim 2\%$ remaining). Compared to previously reported fall conditions ($T_{50} \sim 73$ °C, incomplete evaporation), winter nitrate appeared more volatile, indicating relatively fewer non-volatile nitrate forms (e.g., Kang et al., 2022; Jeon et al., 2023). Sulfate exhibited the highest thermal stability among the measured species. The thermogram showed a relatively stable mass fraction (MFR > 0.8) up to ~ 130 °C, followed by a sharp decline at temperatures above 140 °C (Fig. 4). This profile is consistent with the typical volatilization behavior of ammonium sulfate in TD-AMS, which requires higher temperatures to evaporate compared to nitrate or organics (Huffman et al., 2009). At 200 °C, approximately 25% of the sulfate mass remained. This residual suggests the presence of a sulfate fraction with lower volatility than pure ammonium sulfate, likely associated with organosulfates or low-volatility mixtures, whereas refractory metal sulfates are not efficiently detected by the AMS (Canagaratna et al., 2007). Sulfate was the least volatile ($T_{50} \sim 170$ °C), consistent with ammonium sulfate (Scott and Cattell, 1979). A subtle slope change near 140 °C likely reflects ammonium sulfate

449 ~~morphology or /phase state changes and/or interactions between organosulfate and inorganic mixingsulfate,~~
450 ~~rather than contributions from metallic (refractory) sulfates, which are not efficiently detected by AMS (Huffman~~
451 ~~et al., 2009; Canagaratna et al., 2007). Approximately 25% of the initial sulfate signal remained at 200 °C (Fig.~~
452 ~~4), suggesting the presence of low volatility sulfate containing material or mixed organic inorganic components,~~
453 ~~although the AMS alone cannot resolve their specific chemical identity. About 25% remained at 200 °C,~~
454 ~~indicating possible contributions from metallic or organic sulfates.~~ Ammonium showed intermediate volatility,
455 with T_{50} between nitrate and sulfate. Its slightly lower winter T_{50} suggests stronger nitrate association. Residual
456 ammonium at 200 °C was consistent (~4%) in previously reported spring/fall measurements (Kang et al., 2022;
457 Jeon et al., 2023). ~~Chloride volatility was broadly consistent with prior AMS studies, with T_{50} values comparable~~
458 ~~across seasons (e.g., Xu et al., 2016; Jeon et al., 2023). The near-complete evaporation observed in winter (~4%~~
459 ~~residual at 200 °C, Fig. 4) indicates that the chloride measured here was dominated by volatile inorganic chloride,~~
460 ~~specifically ammonium chloride (NH₄Cl), which fully evaporates at relatively low temperatures (Huffman et al.,~~
461 ~~2009). By contrast, metal chlorides (e.g., NaCl, KCl) are refractory and far less volatile; they are also poorly~~
462 ~~detected by the AMS (Canagaratna et al., 2007). The lower residual in winter compared to fall (~10%) therefore~~
463 ~~suggests that wintertime chloride consisted almost exclusively of pure ammonium chloride, whereas the fall~~
464 ~~samples may have contained a minor fraction of less volatile or refractory chloride species. “Chloride volatility~~
465 ~~was broadly consistent with prior AMS studies, with T_{50} values comparable across seasons (e.g., Xu et al., 2016;~~
466 ~~Jeon et al., 2023). The near-complete evaporation observed in winter (~4% residual at 200 °C, Fig. 4) suggests~~
467 ~~that the chloride measured here was dominated by volatile inorganic chloride, such as ammonium chloride~~
468 ~~(NH₄Cl), which fully evaporates at relatively low temperatures (Huffman et al., 2009; Middlebrook et al., 2012).~~
469 ~~By contrast, metal chlorides (e.g., NaCl, KCl) are far less volatile and are poorly detected by AMS due to their~~
470 ~~refractory nature (Canagaratna et al., 2007). The lower winter residual compared to fall (~10%) therefore likely~~
471 ~~reflects a shift toward more volatile chloride sources—such as residential heating or road salt related NH₄Cl~~
472 ~~formation—rather than contributions from refractory metal chlorides. Chloride volatility was also comparable~~
473 ~~across seasons in prior studies in terms of T_{50} , but exhibited more complete evaporation in winter (~4% residual~~
474 ~~vs. ~10% in fall), possibly reflecting a shift in source to more volatile forms like road salt during wintertime.~~
475 ~~Organics exhibited moderate volatility (T_{50} ~120 °C), and their thermogram showed a gradual, continuous~~
476 ~~decrease in mass fraction with increasing TD temperature. This smooth profile reflects the presence of a broad~~
477 ~~distribution of organic compounds spanning SVOC to LVOC ranges, in contrast to inorganic species such as~~
478 ~~nitrate or ammonium chloride, which often show more abrupt losses at characteristic temperatures (Huffman et~~
479 ~~al., 2009; Xu et al., 2016). This behavior is consistent with previous TD-AMS observations in Seoul during~~

spring and fall (Kang et al., 2022; Jeon et al., 2023). Organics exhibited moderate volatility ($T_{50} \sim 120^\circ\text{C}$), and their thermogram showed a gradual, continuous decrease in mass fraction with increasing TD temperature. This smooth profile reflects the presence of a broad distribution of organic compounds spanning SVOC to LVOC ranges, in contrast to inorganic species such as nitrate or ammonium chloride, which often show more abrupt losses at characteristic temperatures (Huffman et al., 2009; Xu et al., 2016). This behavior is consistent with previous TD-AMS observations in Seoul during spring and fall (Kang et al., 2022; Jeon et al., 2023). OA exhibited moderate volatility ($T_{50} \sim 120^\circ\text{C}$), consistent with the presence of a wide variety of compounds with differing volatilities. This trend aligns with previously reported spring and fall observations in Seoul, Korea (Kang et al., 2022; Jeon et al., 2023).

3.2.1 Volatility Profiles of Organic sources

Figure 5 presents the volatility distributions of six OA sources within the volatility basis set (VBS) framework. Volatility is expressed as the effective saturation concentration (C^* , $\mu\text{g m}^{-3}$), where higher C^* values correspond to higher volatility. Following Donahue et al. (2009), C^* values are categorized into four bins: extremely low-volatility organic compounds (ELVOCs, $\log C^* < -4.5$), low-volatility organic compounds (LVOCs, $-4.5 < \log C^* < -0.5$), semi-volatile organic compounds (SVOCs, $-0.5 < \log C^* < 2.5$), and intermediate-volatility organic compounds (IVOCs, $2.5 < \log C^* < 6.5$).

Among the primary OA (POA) sources, hydrocarbon-like OA (HOA) exhibited the highest volatility, with mass predominantly distributed in the SVOC and IVOC ranges, consistent with its chemically reduced nature ($\text{O:C} = 0.13$) and direct combustion origin. Mass fraction remaining (MFR) results (Fig. S9) further support this, showing rapid mass loss at lower temperatures. Biomass burning OA (BBOA) and nitrogen-containing OA (NOA) also showed high volatility, peaking in the SVOC–IVOC range ($\log C^* = 1\text{--}3$), but displayed slightly higher O:C ratios (0.25 and 0.19, respectively). This modest enhancement in O:C reflects their source composition—biomass combustion produces partially oxygenated organic species (e.g., levoglucosan, phenols), and NOA contains nitrogen-bearing functional groups—rather than enhanced atmospheric oxidation. Among the primary OA (POA) sources, hydrocarbon-like OA (HOA) exhibited the highest volatility, with mass predominantly distributed in the SVOC and IVOC ranges. This is consistent with its low oxidation state ($\text{O:C} = 0.35$) and primary emission characteristics. Mass fraction remaining (MFR) results (Fig. S9) further support this, showing rapid mass loss at

lower temperatures. Biomass burning OA (BBOA) and nitrogen-containing OA (NOA) also showed high volatility, peaking in the SVOC–IVOC range ($\log C^* = 1–3$), and had lower O:C ratios of 0.25 and 0.19, respectively. Their slightly more oxidized nature relative to HOA, despite a similar volatility range, may reflect emissions occurring under nighttime or cooler conditions, which promote condensation of otherwise volatile species. The similar volatility distributions and MFR profiles of BBOA and NOA (Fig. S9) further support the possibility of a shared emission source or formation pathway (Section 3.1.1). Cooking-related OA (COA) showed a more moderate volatility profile, with mass more evenly distributed across the LVOC and SVOC bins. This behavior differs from that of BBOA, which is slightly more oxidized yet more volatile. This apparent decoupling between oxidation state and volatility is a characteristic feature of COA reported in previous volatility studies (Paciga et al., 2016; Kang et al., 2022). These studies attribute the lower volatility of COA to its abundance of high-molecular-weight fatty acids (e.g., oleic, palmitic, and stearic acids) and glycerides (Mohr et al., 2009; He et al., 2010). Unlike the smaller, fragmented molecules typical of biomass burning, these lipid-like compounds possess high molar masses that suppress volatility, even though their long alkyl chains result in low O:C ratios.

Cooking-related OA (COA) showed a more moderate volatility profile, with mass more evenly distributed across the LVOC and SVOC bins. This pattern reflects its diverse cooking sources and variable emission profiles as previously reported (Kang et al., 2022).

For secondary OA (SOA), less-oxidized oxygenated OA (LO-OOA) exhibited the lowest volatility, with substantial mass in the LVOC and ELVOC bins ($C^* \approx 10^{-3}$ – 10^{-4}). This is in agreement with previous findings in Seoul during spring (Kang et al., 2022). In contrast, more-oxidized OOA (MO-OOA), despite its higher oxidation state (O:C = 1.15), displayed greater volatility, with a peak at $C^* \approx 10^1$. This discrepancy likely reflects differences in formation and aging processes, as discussed further in Section 3.3.

Overall, the volatility characteristics across OA factors suggest that oxidation state alone does not fully explain volatility. Rather, volatility is shaped by a combination of emission source, emission timing, temperature, and atmospheric processing. These findings highlight the importance of integrating both chemical and physical characterization to better understand OA formation and aging across seasons.

3.3 Aging effect on volatility from 2D VBS

Generally, the oxygen-to-carbon (O:C) ratio of organic aerosols (OA) is inversely related to their volatility. As O:C increases through aging, the effective saturation concentration (C^*) typically decreases, resulting in lower volatility

(Donahue et al., 2006; Jimenez et al., 2009). This relationship arises because oxidative functionalization introduces polar groups (e.g., hydroxyl, carboxyl) that increase molecular weight and enhance intermolecular hydrogen bonding, thereby reducing the effective saturation concentration (C^*) and promoting particle-phase retention (Jimenez et al., 2009; Kroll and Seinfeld, 2008; Donahue et al., 2011). ~~This common relationship arises because the addition of oxygen-containing functional groups (e.g., hydroxyl, carboxyl, carbonyl), which increases molecular weight and enhances intermolecular interactions such as hydrogen bonding, thereby reducing vapor pressure (Jimenez et al., 2009; Kroll and Seinfeld, 2008). Moreover, oxidative aging often leads to oligomerization or functionalization, promoting particle-phase retention and reducing the effective saturation concentration (C^*) (Donahue et al., 2011; Robinson et al., 2007).~~ However, in this study, the most oxidized OA factor—MO-OOA, with a high O:C ratio of 1.15—exhibited unexpectedly high volatility. Its volatility distribution was skewed toward SVOCs and IVOCs (Fig. 5), and its rapid mass loss in MFR thermograms (Fig. S9) further indicated low thermal stability. This observation appears to contradict the usual inverse O:C–volatility relationship; however, under winter haze conditions—with suppressed O_3 /low OH, particle-phase autoxidation and fragmentation can yield higher-O:C yet more volatile products, with enhanced condensation on abundant particle surface area (details below).

Viewed against prior TD-AMS results, the volatility of Seoul’s winter MO-OOA presents a unique case, particularly in the nature of its O:C–volatility relationship. Prior urban studies have commonly reported substantial SVOC-OA, consistent with high photochemical activity or elevated loadings; for example, prior TD-AMS studies in Mexico City, Los Angeles, Beijing, and Shenzhen have all reported substantial SVOC–IVOC contributions during polluted periods, indicating that high OA volatility is a common feature of urban environments across seasons (Cappa and Jimenez, 2010; Xu et al., 2019; Cao et al., 2018) ~~Mexico City/Los Angeles showed pronounced SVOC–IVOC contributions during warm seasons (Cappa and Jimenez, 2010), and summertime Beijing and wintertime Shenzhen likewise exhibited strong overall OA volatility (Xu et al., 2019; Cao et al., 2018).~~ While these comparisons establish that volatile OA is common, they generally did not report the factor-level inversion observed here, where the highly-oxidized OOA component (MO-OOA) was more volatile than a less-oxidized OOA (LO-OOA). This behavior is distinct from findings in colder, lower-loading regimes; wintertime Paris, for instance, maintained the conventional hierarchy where the more-oxidized OOA was comparatively less volatile (Paciga et al., 2016). Furthermore, seasonal context within Seoul showed springtime OA with lower oxidation levels than our winter MO-OOA despite similar SVOC contributions (Kang et al., 2022). This comprehensive comparison

underscores the unusual nature of the O:C-volatility relationship observed under the specific winter haze conditions in Seoul.

3.3.1 High-volatility nature of MO-OOA in Seoul wintertime

MO-OOA exhibited high O:C ratios and high apparent volatility, characteristics that were further amplified during haze episodes—periods marked by reduced ozone levels, low solar radiation, and elevated aerosol mass concentrations (Fig. 67 and Fig. S6, yellow shading). Spectrally, MO-OOA was defined by a consistently high f_{44} (CO_2^+) signal and a comparatively stable f_{43} ($\text{C}_2\text{H}_3\text{O}^+$) signal relative to LO-OOA (Fig. S8b6). Notably, when MO-OOA concentrations intensified during haze, only f_{44} was significantly enhanced, while f_{43} remained nearly unchanged (Fig. 6). This trend is corroborated by the haze–non-haze comparison (Fig. S12), where haze periods (including high MO-OOA intervals) showed elevated contributions from oxygenated fragments (m/z 28, 29, 44) and higher O:C ratios. In contrast, non-haze periods were characterized by larger fractional contributions from hydrocarbon-like fragments (m/z 41, 43, 55, 57). The observed temporal pattern—elevated f_{44} without corresponding changes in f_{43} —is a typical signature of highly oxidized and fragmented organic aerosol (Figs. 6 and 7), suggesting that aging was dominated by fragmentation rather than functionalization (Kroll et al., 2009). These spectral patterns collectively indicate that MO-OOA is highly oxidized yet remains relatively volatile compared to LO-OOA.

The elevated volatility of MO-OOA despite its high O:C (~ 1.15) indicates that oxidation under these haze conditions did not follow the classical multi-generational OH-driven aging pathway, which typically increases molecular mass and reduces volatility. Instead, the data align with fragmentation-dominated aging, where highly oxygenated but lower-molecular-weight compounds (e.g., small acids or diacids) are formed. Prior field and laboratory studies using online AMS/FIGAERO-CIMS and EESI-TOF have similarly reported high-O:C yet volatile product distributions characterized by high f_{44} and stable f_{43} (Kroll et al., 2009; Ng et al., 2010; Chhabra et al., 2011; Lambe et al., 2012; Lopez-Hilfiker et al., 2016; D’Ambro et al., 2017).

While direct mechanistic measurements were not available in this study, we hypothesize that the formation of this volatile, high-O:C component may be driven by specific low-light oxidation pathways consistent with the observed environmental conditions. The suppressed ozone levels during haze likely indicate a low-OH oxidation regime (Fig. 67). Under such conditions, radical chemistry involving NO_3 (which is longer-lived in low light) or particle-phase autoxidation could preferentially produce highly oxygenated but relatively small organic fragments (Ehn et al.,

595 2014; Zhao et al., 2023). Although haze suppresses photolysis, HONO concentrations—maintained via
596 heterogeneous conversion or surface emissions—could still provide a non-negligible source of OH (Gil et al., 2021;
597 Kim et al., 2024; Slater et al., 2020). Furthermore, the high aerosol mass loadings during haze (C_{OA}) provide
598 abundant surface area for absorptive partitioning (Pankow, 1994; Donahue et al., 2006). This increased partitioning
599 mass allows even relatively volatile, oxidized compounds to condense into the particle phase, contributing to the
600 high apparent volatility and oxidation state observed (Jimenez et al., 2009; Ng et al., 2016). Consequently, these
601 results underscore the need for SOA models to incorporate fragmentation-dominated pathways to accurately
602 represent wintertime haze evolution.

603 Proposed Mechanism: Fragmentation and Condensation under Low OH Haze

604 ~~MO-OOA concentrations increased during haze episodes—characterized by reduced ozone levels, low solar~~
605 ~~radiation and elevated aerosol mass concentrations (Fig. 6 and Fig. S6, yellow shading). high f_{44} (CO_2^+) MO-~~
606 ~~OOA in this study was characterized by a consistently high f_{44} (CO_2^+) signal and a relatively stable f_{43} ($C_2H_3O^+$)~~
607 ~~signal compared to LO-OOA (Fig. S8b). During specific periods when MO-OOA concentrations increased,~~
608 ~~only f_{44} was noticeably enhanced, while f_{43} remained flat (Fig. 6). This temporal pattern—elevated f_{44} without~~
609 ~~corresponding changes in f_{43} —is a typical signature of highly oxidized and fragmented organic aerosol and~~
610 ~~suggests advanced aging dominated by fragmentation rather than functionalization (Kroll et al., 2009).~~
611 ~~Consistent with this, the haze–non-haze comparison, including the high MO-OOA interval (Fig. S12), shows~~
612 ~~larger oxygenated fragments (m/z 28, 29, 44) and higher f_{44} and O:C during haze, whereas non-haze periods~~
613 ~~exhibit relatively larger fractional hydrocarbon fragments (m/z 41, 43, 55, 57). Unlike classical OH initiated,~~
614 ~~multi generational aging—which increases molecular mass and reduces volatility—fragmentation dominated~~
615 ~~oxidation can cleave larger precursors into smaller oxygenated compounds, resulting in higher volatility~~
616 ~~despite elevated O:C. Consistent with this interpretation, online AMS/FIGAERO-CIMS and EESI-TOF, as~~
617 ~~well as offline HRMS/GC-MS, have reported high O:C yet more volatile product distributions accompanied~~
618 ~~by elevated f_{44} with comparatively stable f_{43} under fragmentation dominated aging (Kroll et al., 2009; Ng et al.,~~
619 ~~2010; Chhabra et al., 2011; Lambe et al., 2012; Lopez-Hilfiker et al., 2016; D’Ambro et al., 2017). Furthermore,~~
620 ~~high aerosol mass loadings during haze events provide abundant surface area for the uptake of semi-~~
621 ~~volatile/intermediate volatility organics via absorptive partitioning, so that higher C_{OA} enhances condensation~~
622 ~~(Pankow, 1994; Donahue et al., 2006; Hallquist et al., 2009; Robinson et al., 2007). We also note that~~
623 ~~**functionalized, low molecular weight compounds** can reside in the SVOC–IVOC range and thus contribute~~
624 ~~to the high apparent volatility of MO-OOA (Ng et al., 2010; Chhabra et al., 2011; Lopez-Hilfiker et al., 2016;~~

D'Ambro et al., 2017). . This facilitates the condensation of even relatively volatile, oxidized compounds onto particles (Fig. 6). The net result is an apparent increase in both oxidation state and volatility of OA, as reported in aging studies under stagnant and polluted conditions (Jimenez et al., 2009; Ng et al., 2016).

The suppressed ozone likely indicates lower OH radical production via O₃ photolysis, leading to a low OH oxidation regime. We note that haze also suppresses HONO photolysis; however, HONO concentrations can be elevated at night and early morning via heterogeneous NO₂ conversion and surface emissions, so net OH from HONO may remain non-negligible even as photolysis rates are depressed (e.g., Gil et al., 2021; Kim et al., 2024; Slater et al., 2020). Under such conditions, particle-phase autoxidation involving RO₂ radicals can become the dominant oxidation pathway. Even under low OH conditions, NO₃ formed via NO₂ + O₃ can initiate RO₂ production through addition to alkenes, while reduced photolysis at night/low light extends NO₃ lifetimes; these RO₂ then participate in particle-phase autoxidation, yielding highly oxygenated yet relatively volatile products. These processes tend to produce highly oxidized but relatively low molecular weight products (Ehn et al., 2014; Zhao et al., 2023). Unlike classical OH-initiated, multi-generational aging which increases molecular mass and reduces volatility, fragmentation-dominated oxidation can cleave larger precursors into smaller oxygenated compounds, resulting in higher volatility despite elevated O:C. Consistent with this interpretation, online AMS/FIGAERO-CIMS and EESI-TOF, as well as offline HRMS/GC-MS, have reported high O:C yet more-volatile product distributions accompanied by elevated *f_n* with comparatively stable *f₃* under fragmentation-dominated aging (Kroll et al., 2009; Ng et al., 2010; Chhabra et al., 2011; Lambe et al., 2012; Lopez-Hilfiker et al., 2016; D'Ambro et al., 2017). Furthermore, high aerosol mass loadings during haze events provide abundant surface area for the uptake of semi-volatile/intermediate volatility organics via absorptive partitioning, so that higher C_{OA} enhances condensation (Pankow, 1994; Donahue et al., 2006; Hallquist et al., 2009; Robinson et al., 2007). We also note that functionalized, low molecular weight compounds can reside in the SVOC-IVOC range and thus contribute to the high apparent volatility of MO-OOA (Ng et al., 2010; Chhabra et al., 2011; Lopez-Hilfiker et al., 2016; D'Ambro et al., 2017). . This facilitates the condensation of even relatively volatile, oxidized compounds onto particles (Fig. 6). The net result is an apparent increase in both oxidation state and volatility of OA, as reported in aging studies under stagnant and polluted conditions (Jimenez et al., 2009; Ng et al., 2016).

In line with these reports, our results also revealed a decoupling between O:C and volatility, with MO-OOA showing high volatility despite its elevated O:C ratio (~1.15). While this behavior has been observed in other urban environments, this study provides one of the first detailed thermodynamic assessments of this decoupling under winter haze conditions in Seoul using real-time TD-AMS measurements. Supporting this interpretation,

MO-OOA in this study was characterized by a consistently high f_{44} (CO_2^+) signal and a relatively stable f_{43} ($\text{C}_2\text{H}_5\text{O}^+$) signal compared to LO-OOA (Fig. S8b). During specific periods when MO-OOA concentrations increased, only f_{44} was noticeably enhanced, while f_{43} remained flat (Fig. 6). This temporal pattern—elevated f_{44} without corresponding changes in f_{43} —is a typical signature of highly oxidized and fragmented organic aerosol and suggests advanced aging dominated by fragmentation rather than functionalization (Kroll et al., 2009). Consistent with this, the haze non haze comparison, including the high MO-OOA interval (Fig. S12), shows larger oxygenated fragments (m/z 28, 29, 44) and higher f_{44} and O:C during haze, whereas non haze periods exhibit relatively larger fractional hydrocarbon fragments (m/z 41, 43, 55, 57). These spectral contrasts indicate that the elevated volatility of MO-OOA reflects advanced oxidation via autooxidation and the condensation of small oxygenated fragments rather than enrichment of high molecular weight ions, particularly under conditions of limited OH and high particulate surface area.

This unexpected volatility behavior of highly oxidized MO-OOA highlights the need for secondary organic aerosol (SOA) models to incorporate fragmentation dominated oxidation pathways—especially under haze conditions where conventional assumptions linking O:C to volatility may break down. Including such mechanisms could improve model accuracy in representing OA aging and volatility in urban air quality simulations.

4 Conclusions

This study provides a comprehensive characterization of wintertime submicron aerosols (PM_{10}) in Seoul, integrating chemical composition, volatility measurements, and source apportionment to reveal critical insights into urban OA evolution. The two most significant findings are the robust real-time identification of a nitrogen-containing organic aerosol (NOA) factor and the observation of unexpected volatility behavior in highly oxidized OA. The NOA factor, spectrally dominated by low-molecular-weight alkylamine fragments, was successfully resolved primarily due to the accumulation of pollutants during wintertime stagnation, which sufficiently enhanced the spectral signals of these semi-volatile species for identification. Its temporal and chemical characteristics point to a mixed primary/secondary origin: driven by direct combustion emissions (e.g., residential heating) but significantly enhanced by the rapid gas-to-particle partitioning of semi-volatile amines under cold, humid conditions. Concurrently, the volatility analysis revealed a striking decoupling between oxidation state and volatility for the More-Oxidized Oxygenated OA (MO-OOA). Despite its high O:C ratio (~ 1.15), MO-OOA exhibited elevated volatility, a deviation from classical aging models that typically associate high oxidation with low volatility. This behavior is attributed to the specific conditions of winter haze—reduced photolysis and high aerosol mass

loadings—which favor fragmentation-dominated aging pathways and the absorptive partitioning of volatile oxygenated products.

These results revise our understanding of wintertime aerosol dynamics and underscore the limitations of current models in representing reduced-nitrogen species and non-canonical oxidation pathways. To address the remaining uncertainties, future research should prioritize evaluating the seasonal variability of NOA to better disentangle the influence of meteorological drivers from specific emission sources. Concurrently, there is a critical need to directly probe radical oxidation mechanisms, such as RO₂ autoxidation and NO₃ chemistry, particularly under haze conditions. Integrating these field inquiries with laboratory studies and advanced molecular-level measurements (e.g., FIGAERO-CIMS, EESI-TOF) will be essential for constraining the formation, lifetime, and climate impacts of these complex organic aerosol components in polluted megacities.

~~These results revise our understanding of wintertime aerosol dynamics and underscore the limitations of current models in representing reduced-nitrogen species and non-canonical oxidation pathways. To address the remaining uncertainties, future research should focus on three key areas: (1) evaluating the seasonal variability of NOA to disentangle the influence of meteorological drivers from specific emission sources; (2) quantifying the specific roles of heterogeneous reactions and aminium salt formation in NOA persistence; and (3) directly probing radical oxidation mechanisms (e.g., RO₂ autoxidation, NO₃ chemistry) under haze conditions. Combining these field inquiries with laboratory studies and advanced molecular-level measurements (e.g., FIGAERO-CIMS, EESI-TOF) will be essential for constraining the formation, lifetime, and climate impacts of these complex organic aerosol components in polluted megacities.~~

~~This study offers a detailed characterization of wintertime submicron aerosols (PM₁) in Seoul by integrating chemical composition, volatility behavior, and source apportionment to better understand their formation and atmospheric evolution. Organic aerosols (OA), particularly secondary organic aerosols (SOA), were the dominant PM₁ component, highlighting the significance of oxidative processes even during cold seasons. A notable result is the successful real-time resolution of a nitrogen-containing organic aerosol (NOA) factor, enabled by cold, humid meteorological conditions that enhanced the partitioning and stabilization of amine-derived compounds. The NOA factor was characterized by tracer ions associated with low-molecular-weight alkylamines such as TMA, DMA, and MA, which likely originated from biomass combustion.~~

712 Volatility analysis revealed distinct thermodynamic behavior across OA sources. Primary OA factors such as
713 HOA, BBOA, and COA exhibited relatively high volatility, while LO-OOA showed low volatility and a higher
714 oxidation state, consistent with aged, low volatility material. The similarity in volatility distributions and
715 diurnal patterns between BBOA and NOA suggests that biomass combustion under wintertime conditions is a
716 likely contributor to both primary organic and nitrogenous aerosol formation.

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718 Interestingly, MO-OOA—despite its high oxygen to carbon (O:C) ratio—exhibited elevated volatility, diverging
719 from the expected inverse relationship between oxidation state and volatility. This suggests that under stagnant,
720 polluted conditions with suppressed ozone and OH radical levels, particle phase autooxidation and fragmentation
721 pathways may dominate over traditional OH-initiated aging, yielding highly oxidized yet semi-volatile products.
722 These findings highlight the importance of coupling high-resolution chemical and physical aerosol measurements
723 to better understand OA formation processes and properties within urban air quality frameworks. Consequently,
724 air quality models should incorporate diverse oxidation mechanisms and avoid assuming a direct link between
725 oxidation state and volatility.

726 **Data availability.**

727 Data presented in this article are available upon request to the corresponding author.

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732 **Author Contributions**

733 Hwajin Kim designed and prepared the manuscript. Jiwoo Jeong operated the TD-AMS and analyse the data. Jihye Moon
734 analyse the data. Hyungu Kang analyse the volatility of OA.

736 **Competing interests.**

737 The authors declare that they have no conflict of interest.

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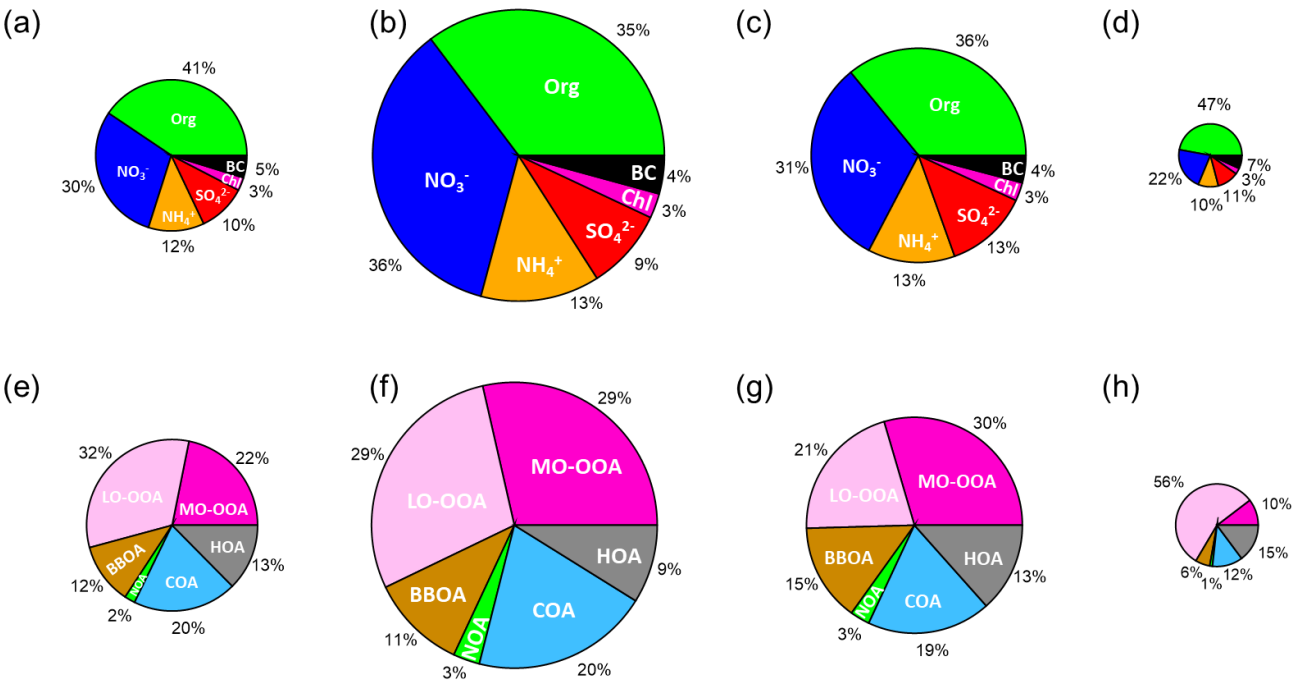
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Tables and Figures



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	Period	Standard	Avg. Mass conc.($\mu\text{g m}^{-3}$)
Total	2019.11.28 ~ 2019.12.28		Avg PM ₁ = 26.37
Clean	2019.12.04 ~ 2019.12.06	Daily PM ₁ < 10.00 $\mu\text{g m}^{-3}$	Avg PM ₁ = 9.98
Haze 1	2019.12.07 ~ 2019.12.11	Daily PM ₁ > 30.00 $\mu\text{g m}^{-3}$	Avg PM ₁ = 51.88
Haze 2	2019.12.21 ~ 2019.12.25	Daily PM ₁ > 30.00 $\mu\text{g m}^{-3}$	Avg PM ₁ = 37.71

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Figure 1. Compositional pie charts of PM₁ species for (a) the entire study period, (b) haze period 1, (c) haze period 2, and (d) a clean period; and of each OA source for (e) the entire study period, (f) haze period 1, (g) haze period 2, and (h) the clean period. Table. Standard and average PM₁ mass concentrations during the entire study period, haze period 1, haze period 2, and the clean period.

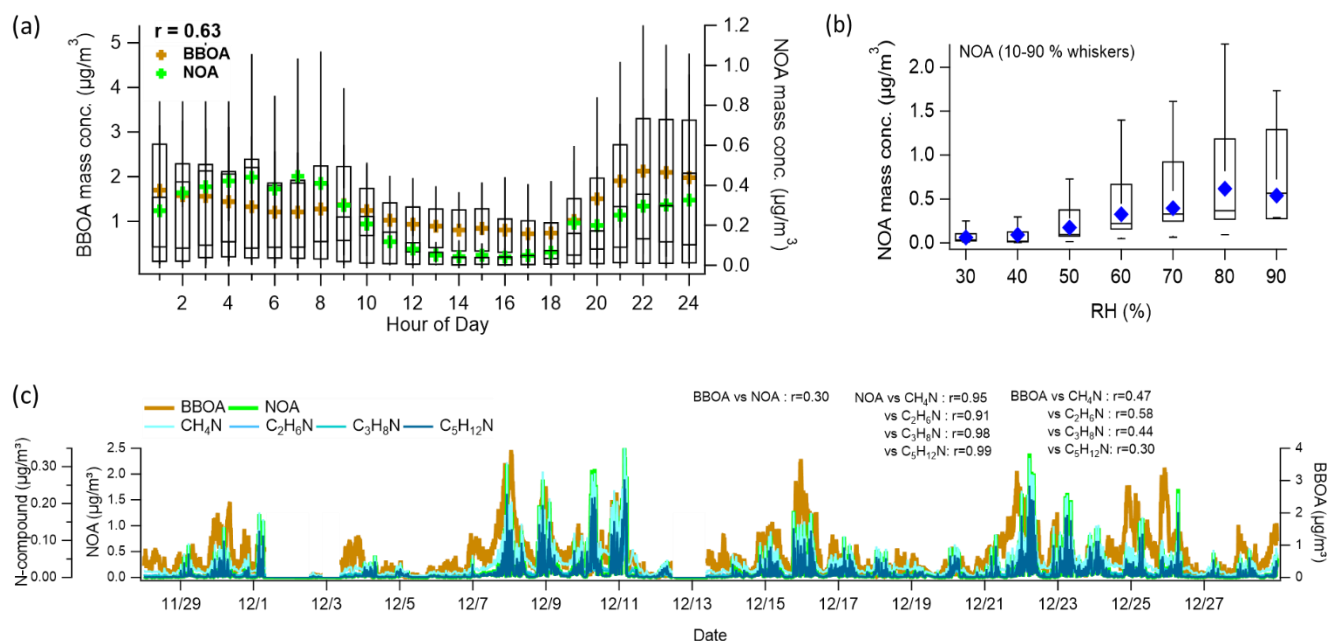


Figure 2. (a) Diurnal mean profiles of NOA and BBOA. Whiskers denote the 90th and 10th percentiles; box edges represent the 75th and 25th percentiles; the horizontal line indicates the median, and the colored marker shows the mean. The diurnal correlation between NOA and BBOA mean values is 0.63. (b) Relative humidity (RH)-binned nighttime (19:00–05:00) profile of NOA. Box and whisker definitions are the same as in panel (a). (c) Time series of NOA, BBOA, and amine-related ions (CH_4N^+ , $\text{C}_2\text{H}_6\text{N}^+$, $\text{C}_3\text{H}_8\text{N}^+$, $\text{C}_5\text{H}_{12}\text{N}^+$), along with their correlations with NOA and BBOA.

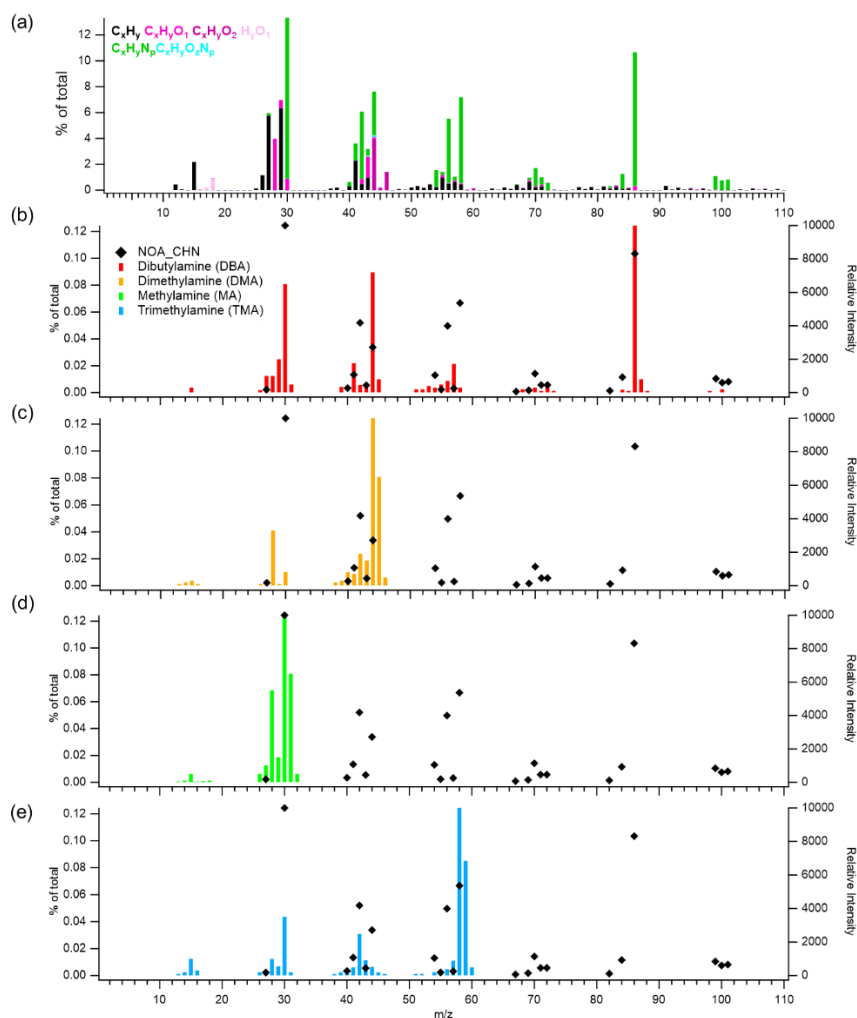
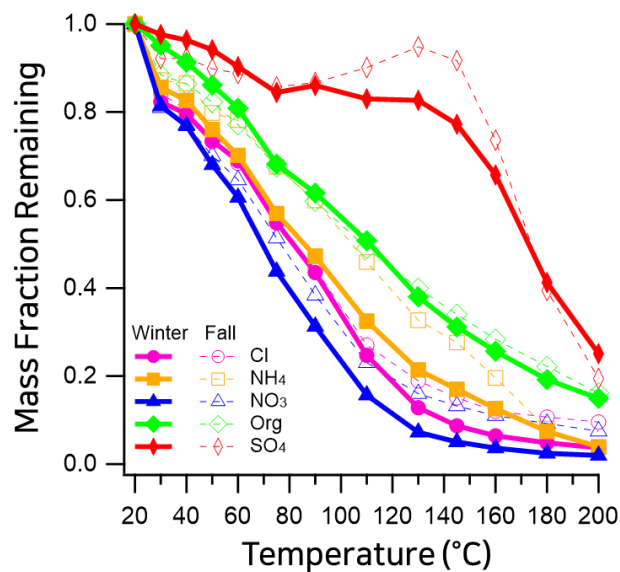


Figure 3. Mass spectra of (a) the NOA factor resolved by PMF analysis in this study, and reference spectra of amines from the NIST library: (b) dibutylamine (DBA), (c) dimethylamine (DMA), (d) methylamine (MA), and (e) trimethylamine (TMA). In panels (b)–(e), the left y-axis indicates the contribution of CHN-containing ions in the NOA factor (% of total), while the right y-axis shows the relative intensity of each compound’s mass spectrum from the NIST library.



796 **Figure 4.** Mass fraction remaining (MFR) of non-refractory (NR) aerosol species measured in Seoul using a thermodenuder
 797 coupled to a high-resolution time-of-flight aerosol mass spectrometer (HR-ToF-AMS). Winter 2019 (this study; dashed) is
 798 compared with fall 2019 (previously reported; solid) (Jeon et al., 2023). Species include organics (magenta), nitrate (blue),
 799 sulfate (orange), ammonium (green), and chloride (red).

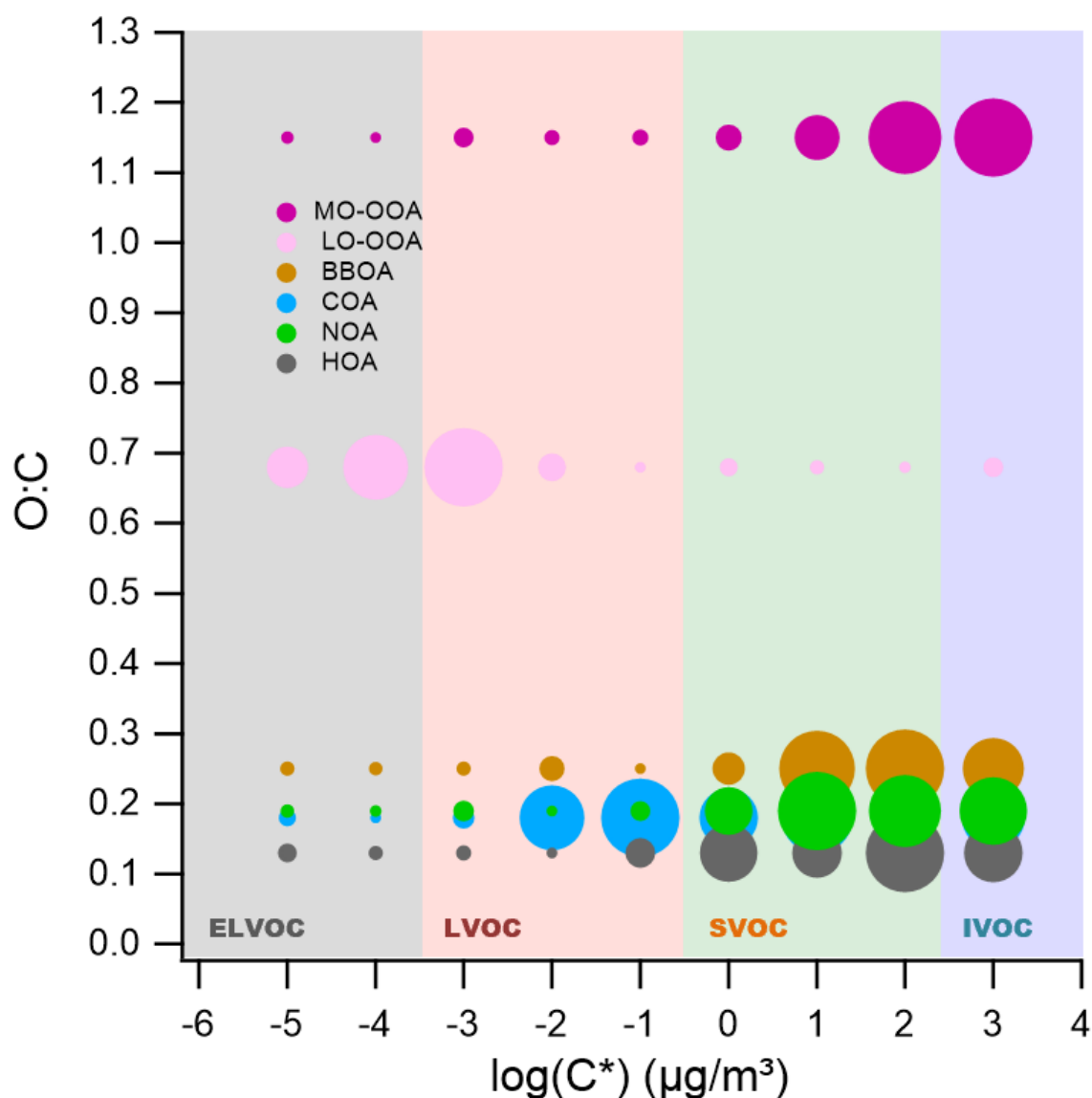
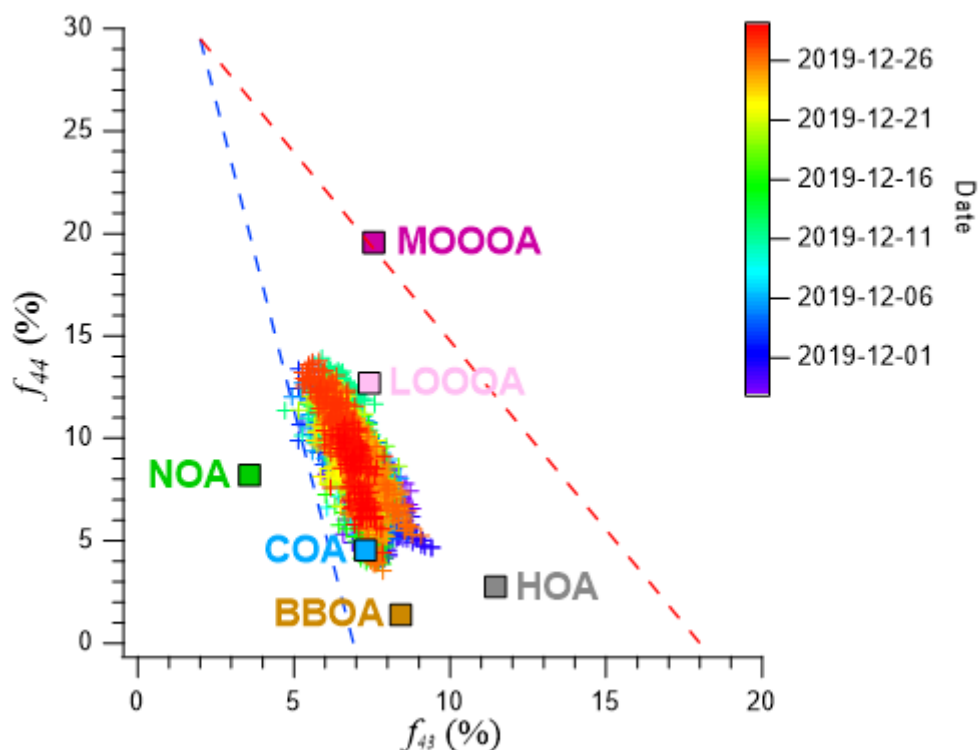


Figure 5. Two-dimensional volatility basis set (2D-VBS) representation of organic aerosol (OA) sources identified in winter 2019 in Seoul. The plot illustrates the relationship between the oxygen-to-carbon (O:C) ratio and the effective saturation concentration (C^*) for each OA source resolved via positive matrix factorization (PMF). Solid circles represent the volatility distribution across C^* bins, with marker size proportional to the mass fraction within each bin for the given source. Shaded regions correspond to different volatility classes: extremely low-volatility organic compounds (ELVOCs), low-volatility organic compounds (LVOCs), semi-volatile organic compounds (SVOCs), and intermediate-volatility organic compounds (IVOCs), delineated by their C^* values.

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Figure 6. scatterplot of f_{44} (CO_2^+) versus f_{43} ($\text{C}_2\text{H}_3\text{O}^+$). for the measured organic aerosol. The data points are color-coded by date to illustrate the temporal variation in OA composition throughout the observation period. The separated OA factors (HOA, COA, BBOA, NOA, LO-OOA, and MO-OOA) are also shown to enable comparison of source contributions and oxidation characteristics. The dashed line represents the typical f_{60} threshold associated with biomass-burning influence, while the triangular boundary indicates the conventional oxidative aging trend in the f_{44} – f_{60} space.

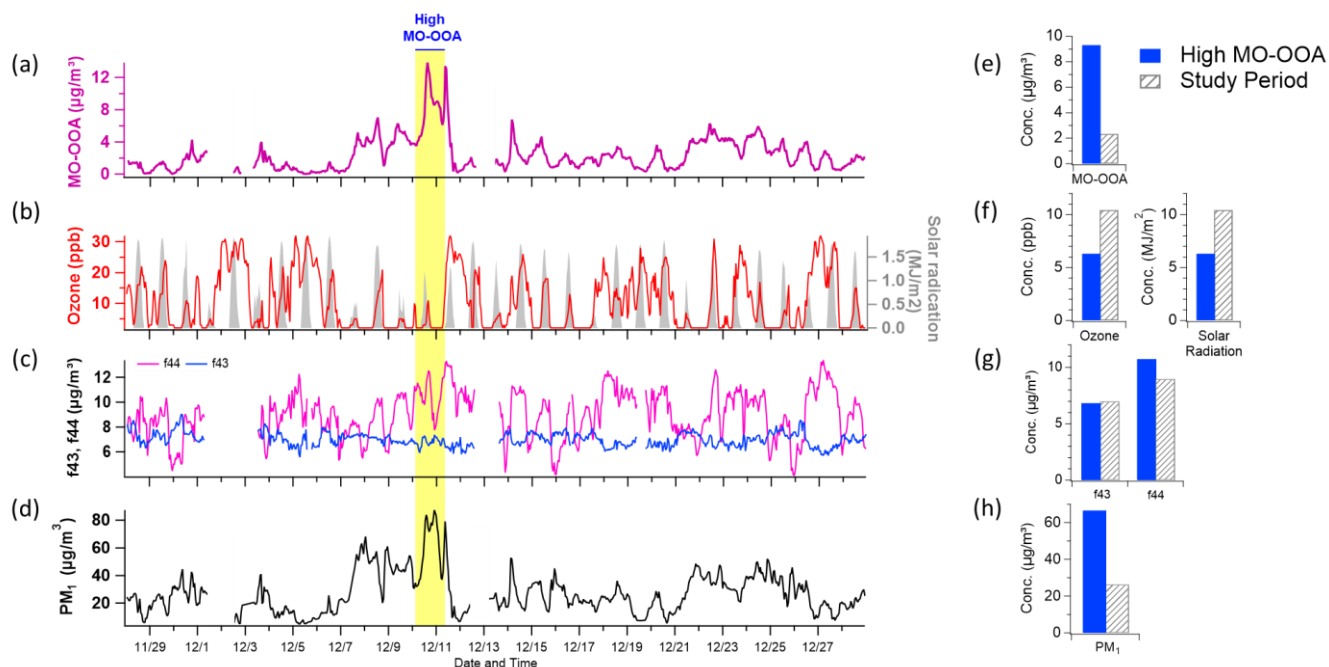


Figure 67. Time series plots of (a) MO-OOA concentration, (b) ozone (O_3) and solar radiation, (c) f_{44} and f_{43} (indicative of oxidation state), and (d) total PM_{10} concentration. The period characterized by elevated MO-OOA levels is highlighted in bright yellow. Panels (e)–(h) present comparative distributions of these variables—MO-OOA, O_3 and solar radiation, f_{44} and f_{43} , and PM_{10} —between the high MO-OOA period (shaded in blue) and the entire measurement period (indicated by gray hatching).

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