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Photodegradation of naphthalene-derived particle oxidation products†

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While photochemical aging is known to alter secondary organic aerosol (SOA) properties, this process remains poorly constrained for anthropogenic SOA. This study investigates the photodegradation of SOA produced from the hydroxyl radical-initiated oxidation of naphthalene under low- and high-NO_x conditions. We used state-of-the-art mass spectrometry (MS) techniques, including extractive electrospray ionization and chemical ionization MS, for the in-depth molecular characterization of gas and particulate phases. SOA were exposed to simulated irradiation at different stages, *i.e.*, during formation and growth. We found a rapid (*i.e.* >30 min) photodegradation of high-molecular-weight compounds in the particle-phase. Notably, species with 20 carbon atoms (C₂₀) decreased by 2/3 in the low-NO_x experiment which was associated with particle mass loss (~12%). Concurrently, the formation of oligomers with shorter carbon skeletons in the particle-phase was identified along with the release of volatile products such as formic acid and formaldehyde in the gas-phase. These reactions are linked to photolabile functional groups within the naphthalene-derived SOA products, which increases their likelihood of being degraded under UV light. Overall, photodegradation caused a notable change in the molecular composition altering the physical properties (*e.g.*, volatility) of naphthalene-derived SOA.

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Environmental significance

Naphthalene is a major anthropogenic volatile organic compound (VOC) and an important contributor to secondary organic aerosol (SOA) formation in urban environments. However, the evolution of the chemical composition of naphthalene-derived SOA remains uncertain. In this study, SOA were generated from the OH-initiated oxidation of naphthalene and exposed to UV radiation to investigate the influence of photochemical aging on the chemical composition. We observed that photodegradation induced a loss of SOA mass and high-molecular-weight dimers that resulted in the production of smaller unsaturated dimers and the emission of volatile species into the atmosphere. These observations underline the importance of photochemistry on urban SOA physicochemical properties and lifetime.

Introduction

Organic aerosols (OA) are nefarious to human health,^{1–3} influence the global climate by scattering and absorbing solar radiation,^{4,5} and modify the lifetime and albedo of clouds by acting as cloud condensation nuclei.^{6–10} OA are a key component of atmospheric aerosols and account for 20 to 90% of the

tropospheric submicron aerosol mass.^{11,12} The OA sources include anthropogenic sources; *i.e.*, traffic emissions, industrial activities, energy production,^{13,14} and biogenic sources such as biomass burning and emissions from vegetation.^{15–18}

From these important emission sources, the oxidation of volatile organic compounds (VOCs) leads to oxidation products of lower volatility than the precursor VOC and greatly participates in SOA formation and growth. This growth occurs either by the reaction of an oxidized gas molecule with a particle after collision (*i.e.*, reactive uptake),^{19,20} by absorptive partitioning^{21,22} or by the condensation of low-vapor pressure compounds.^{12,18,23,24} Due to the importance of SOA on our environment and health, investigating the chemical processes involved in SOA growth and aging is necessary, especially given that organic species experience substantial chemical changes over their lifetime (usually a couple of days).²⁵

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SOA aging is influenced by a variety of chemical processes, such as oxidative processes, (*e.g.* heterogeneous oxidation from ozone (O₃), N₂O₅, or hydroxyl radicals (OH), which can be formed in the aerosol itself),²⁶ or non-oxidative processes, *e.g.*, hydrolysis, aldolisation, esterification, peroxyhemiacetal/hemiacetal formation.^{27,28} These chemical reactions lead to different outcomes (*e.g.*, fragmentation, functionalization, or oligomerization) in the condensed phase.^{12,29} Such changes not only alter the SOA chemical composition but also ultimately influence their optical and physical properties (*e.g.*, volatility, hygroscopicity, viscosity).^{30–32} Functionalization and oligomerization tend to decrease the volatility of the molecules by increasing their polarity or carbon number, thus changing their lifetime in the aerosol phase. Fragmentation and carbon bond cleavage result in higher vapor pressure products, which can cause a reduction of the SOA mass if fragmentation continues unabated.^{31,33} Photochemical processes have also been shown to greatly impact the SOA chemical composition during aging,^{34–38} and are hypothesized to be a key driver of SOA physicochemical properties. More specifically, previous work showed that photochemical aging of biogenic-derived SOA alters their morphology from well-mixed fresh to phase-separated aerosols.³⁹ This process may be caused by an increase in oligomer content increasing particle viscosity. In addition to impacting SOA composition and morphology, photodegradation is a major SOA removal mechanism that needs to be considered.^{40–44} This mass loss is associated with the presence of photolabile functional groups that are susceptible to be photodegraded and form volatile species that can outgas from the condensed phase. Among them, organic peroxides and carbonyls represent an important fraction of the SOA mass^{45–50} and have been identified as photochemically active species.^{40,42–44} Krapf *et al.*⁴⁰ suggested that carbonyls determine the photolability of α -pinene SOA rather than peroxides while Pospisilova *et al.*³⁰ showed that moderately oxygenated molecules are the most photolabile species. These uncertainties underline the need to perform additional investigations on the SOA evolution under various environmental conditions.

In urban atmospheres, polycyclic aromatic hydrocarbons (PAHs) represent an important group of anthropogenic precursors capable of forming SOA.^{51–54} Composed of fused aromatic rings, PAHs can be rapidly oxidized by the different atmospheric oxidants, predominantly OH radicals.^{55–59} PAH-derived SOA are also known to constitute a considerable health hazard because of their deleterious effect on human health (carcinogenic, embryotoxic, oxidative potential).^{2–4,60–64} Studies have shown that PAH-derived SOA are sensitive to photochemistry.⁶⁵ As a result, evaluating the impact of photodegradation on PAH-derived SOA is important to understand the evolution of aerosol particles notably in urban environments.

In this study, we investigated the formation and evolution of secondary organic aerosol (SOA) produced from the oxidation of naphthalene, one of the most abundant gaseous PAH in the atmosphere,^{63,66} by OH radicals under various UV and environmental conditions. Experiments were conducted in a Teflon atmospheric simulation chamber. Gas and particle-phase chemical characterization was achieved using state-of-the-art online mass spectrometry techniques.

Experimental section

Experiments were conducted at the Paul Scherrer Institute in Switzerland in a 9 m³ atmospheric chamber made of Teflon® fluorocarbon film described by Platt *et al.*⁶⁷ They were performed in batch mode (*i.e.*, static), with no dilution once the particle formation started, except during the first part of the low-NO_x experiment, in which an airflow was unintentionally kept open for 2.5 hours before being closed. The container housing the chamber was temperature-regulated and maintained at a temperature of 24 °C (±2 °C). The chamber was flushed overnight at a flow rate of 35 standard liter per minute (SLPM) with dry purified air obtained from an AADCO (AADCO 250 series, AADCO Instruments, Inc., USA) system. At the start of the experiments, humid air was produced by bubbling dry clean air through ultrapure water. Several instruments were used to monitor the experimental conditions. O₃ was obtained by passing pure air over a UV light source inside a stainless-steel cylinder and was measured by a UV photometric O₃ analyzer (model 49C, Thermo). A dew point hygrometer (SC-05, Rotronic) was used to monitor the temperature and relative humidity (RH). NO_x concentrations were not available, but significant differences in NO_x levels were expected depending on the OH production process. OH radicals were produced within the chamber by either the ozonolysis of tetramethylethylene (TME) (IUPAC name: 2,3-dimethyl-2-butene), a reaction yielding OH, HO₂ radicals and acetone with low NO_x levels,⁶⁸ or by the photolysis of nitrous acid (HONO), resulting in OH radicals and nitric oxide (NO).⁶⁹ In the dark experiment, NO_x concentrations would remain below limit of detection since there was no NO_x injections into the chamber. Historically, generating OH radicals with HONO results in NO_x concentrations below 2 ppb without the injection of either NO or NO₂.^{69,70} Photodegradation was initiated using a bank of black light (set of 40 Cleo Performance solarium lamps from Philips with a power of 90–100 W each, emission spectrum available in Fig. S1†). Another study using this chamber estimated the UV-A lights J_{NO_2} to be $3 \times 10^{-3} \text{ s}^{-1}$.³⁰ OH radical concentrations were estimated by monitoring the decay of butanol-d9 (1 μL –30 ppb) as previously described by Barmet *et al.*⁷¹ Naphthalene was injected into the chamber by passing an airflow through the headspace of solid naphthalene and was monitored with a total hydrocarbon counter (THC monitor APHA-370, Horiba).

The impact of aging and photochemistry-driven aging on naphthalene-derived SOA was investigated by performing two types of experiments: (i) particles grown in the dark before being exposed to the lights, (ii) particles grown with the lights on and aged in the dark. SOA formation in the dark was performed by mixing naphthalene with OH radicals generated from TME ozonolysis, resulting in oxidation under low-NO_x levels. Therefore, this experiment is referred to as the low-NO_x experiment. This experiment was conducted at high relative humidity (RH = 71%) with a concentration of 1.25 ppm and 106 ppb of O₃ and naphthalene, respectively. 10 μL of TME (227 ppb) was injected twice through a septum into a heated line which was flushed with dry air to introduce the VOC into the



chamber. The first injection was diluted by the airflow that remained open. After closing the airflow, the second injection was performed to form more SOA. During the high-NO_x experiment, OH radicals were formed from the photolysis of HONO. This experiment was conducted at high relative humidity (RH ~ 82%) in the presence of 131 ppb of naphthalene with a background level of 0.09–0.1 ppm of O₃. HONO was obtained by passing an airflow through a reactor where both sulfuric acid (H₂SO₄) and sodium nitrite (NaNO₂) were added dropwise.⁷²

The chemical composition of gaseous species was retrieved with an atmospheric pressure chemical ionization (CI) inlet coupled to a Q-Exactive Orbitrap mass spectrometer (Thermo Scientific, USA), hereafter referred to as CI-Orbitrap, and a Vocus proton transfer reaction time-of-flight mass spectrometer (Vocus 2R PTR-TOF, Tofwerk), later referred to as Vocus. An in-depth description and characterization of the inlet was performed by Riva *et al.*,⁷³ so the CI inlet was only described succinctly here. A coaxial sample flow of 10 SLPM was supplied to the instrument and was kept in the center of the drift tube by adding a 20–40 SLPM sheath flow controlled by a mass flow controller (MFC) to minimize wall losses. By flushing 2 standard cubic centimeters per minute (scm) of dry pure air over a 1% liquid ammonia solution, ammonia vapors (NH₃) were generated and mixed with the sheath flow. NH₃ vapors passed through a soft X-ray photoionizer (Hamamatsu, L9491) to generate ammonium ions (NH₄⁺). NH₄⁺ was chosen as the reagent ion due to its low selectivity (*i.e.*, suitable for a large variety of oxidation products⁷⁴).

The Vocus reagent ion source consisted of a hollow cathode producing a plasma between two conical surfaces.⁷⁵ Like the CI-Orbitrap, the Vocus was operated in ammonia mode. The generated NH₄⁺ ions entered the focusing ion-molecule reactor (FIMR), which replaced the drift tube of traditional PTR-TOF.^{75–79} NH₄⁺ ions and the analytes were mixed within the FIMR where adduct ions were formed. Using both instruments allowed us to study compounds with an extended volatility range^{74,80} and to be more robust toward possible RH variations. Indeed, the NH₄⁺-CI-Orbitrap has exhibited a RH dependence, which was not the case for the Vocus-PTR.^{74,75}

Particles in the experiment were monitored using a scanning mobility particle sizer (SMPS, TSI Incorporated, USA) over a 15–638 nm diameter range. Particles were chemically characterized using an extractive electrospray ionization inlet coupled to an atmospheric pressure interface time-of-flight mass spectrometer (API-TOF, Tofwerk AG, Thun, Switzerland). This instrument was previously described by Lopez-Hilfiker and referred to as EESI-TOF from henceforth.⁸¹ The aerosol flow (0.8–1.0 SLPM) before entering the EESI-TOF went through a multi-channel carbon denuder (Ionicon) to remove >99.9% of gas-phase compounds.^{81,82} Periodical blanks (~every 5–15 min) were performed by passing the aerosols through a HEPA filter to obtain a background-corrected signal. The EESI electrospray solution consisted of ultrapure water doped with 100 ppm of sodium iodide (NaI), pushed through an untreated fused silica capillary (360 μm outer diameter, 75 μm internal diameter, BGB Analytik AG, Boeckten, Switzerland) at 100–200 mbar. The fluid flow was controlled by a high-precision pressure regulator (MFCS-EZ

1000 mbar, Fluigent, Inc., Lowell, MA, USA) coupled to the electrospray liquid bottle. 2700–2900 V was applied to ensure spray stability and particle extraction.^{81,83} The instrument was operated in positive mode, resulting in the formation of [analyte + Na]⁺ adduct ions. The EESI-TOF and Vocus signal were normalized using the method detailed in the ESI.[†]

Results and discussion

Particle formation and growth

Fig. 1 shows the SOA formation and the evolution of the mass concentration and chamber conditions throughout the experiments, with particle densities assumed to be 1.5 g cm⁻³ (Kautzman *et al.* 2010).⁸⁴ Both the total SOA mass concentration (orange circles) and the mass loadings for individual particle sizes (color scale) are indicated. An overview of the experimental conditions was compiled in Table S1.[†] While SOA can form from naphthalene ozonolysis,⁸⁵ particles were not observed before the production of OH radicals in both experiments. Presumably, this was due to the weak naphthalene reactivity with ozone. The first injection of TME resulted in particle formation up to ~102 μg m⁻³. Due to the unintentional dilution flow in the chamber, OH levels were not estimated and SOA mass rapidly decreased. Therefore, a second injection of 10 μL of TME was performed after ~2.75 hours (*t* = -1 hour in Fig. 1A) to increase particle mass, with RH = 33% and 0.6 ppm of O₃ present in the chamber. This second injection resulted in an OH radical concentration of 2.7 × 10⁷ molecules per cm³, which was significantly higher than tropospheric OH levels (1.1 × 10⁶ molecules per cm³).⁸⁶ This high OH level rapidly increased particle mass concentration to 122 μg m⁻³. While the first injection raised the number of particles from 646 particles per cm³ (*i.e.*, background level) to 1.2 × 10⁶ particles per cm³, the 2nd injection only showed an increase from 5.4 × 10⁴ to 2 × 10⁵ particles per cm³. This difference indicates that existing particles acted as a condensation sink for gaseous oxygenated species produced from naphthalene oxidation, and inhibited part of the new particle formation. The fast SOA formation from the 1st injection was followed by a significant mass concentration decrease, from ~102 to 42 μg m⁻³ (59% decrease).

The 2nd injection raised the mean mass geometric diameter to ~140 nm, indicating that gaseous species condensed onto preexisting particles instead of participating in new particle formation and allowed to produce sufficient SOA mass to study aging processes. The mean diameter stayed within 140–155 nm for the rest of the experiment. Lights were turned on ~50 min after the second injection for 65 min to study the impact of the photodegradation processes on dark-grown SOA. When lights were turned on, a net decrease of both the mean mass geometric diameter (from 150 to 140 nm) and particle mass concentration (12% loss after wall-loss corrections, details about the correction in Pospisilova *et al.*)⁸⁷ was observed. This is indicative that photodegradation led to particle mass reduction due to a release of volatile compounds, as observed in previous studies.^{30,41–44}

During the high-NO_x experiment, UV lights were turned on for 1.5 hours. OH radicals concentration reached 4.1 × 10⁸



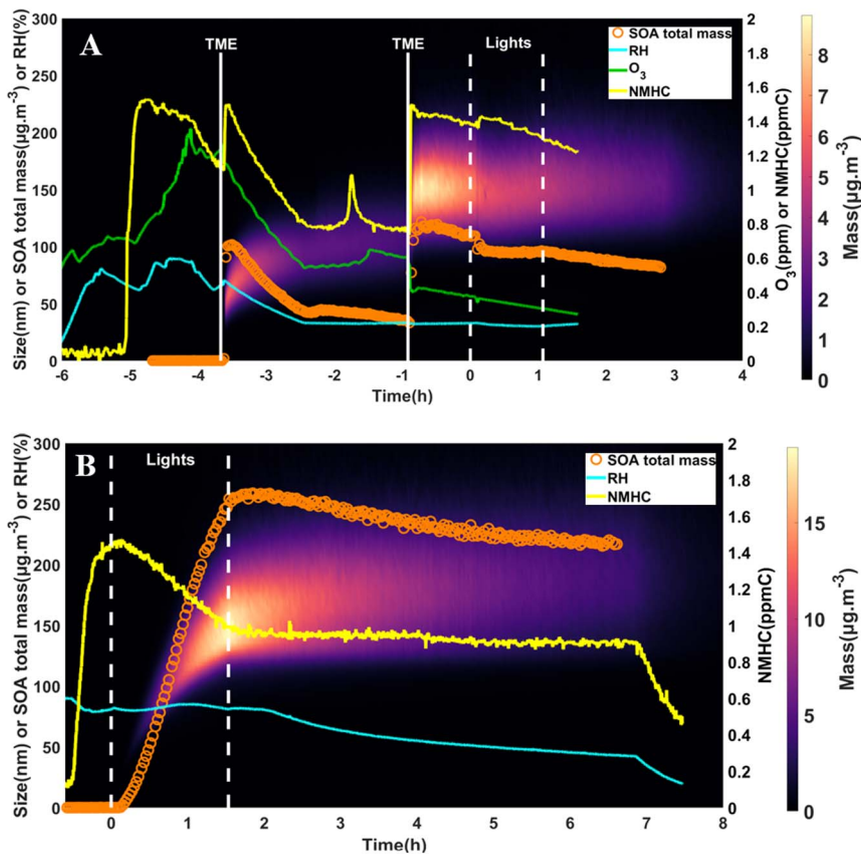


Fig. 1 Evolution of SOA mass concentration measured by the SMPS and chamber conditions for (A) low- NO_x and (B) high- NO_x experiment. The mass loadings of the various sizes of particles are indicated by the color scale. $t = 0$ hours marked the start of the UV lights, and dotted white bars indicate the period when UV lights were turned on. The chamber conditions monitored include the total SOA mass (wall-loss corrected, orange circles), RH (blue), ozone (green), and non-methane hydrocarbon (NMHC, yellow). NMHC was used to monitor the naphthalene concentrations as well as the TME injected. The measurement unit was ppmC, meaning that the real concentrations of naphthalene and TME were respectively 10 times and 6 times lower than what the NMHC suggests.

molecules per cm^3 and up to 1.8×10^5 particles per cm^3 were formed. SOA formation peaked at $254 \mu\text{g m}^{-3}$, with a geometric mean diameter of 160 nm. The mass loss associated with photodegradation cannot be observed in this case, as it was overshadowed by the SOA formation. Over the next 5 hours, particles continued to grow due to either the slow condensation of organic vapors or coagulation, and the geometric mean diameter increased by $\sim 20\%$ up to 190 nm.

Gas- and particle-phase species identification

Naphthalene oxidation from OH radicals occurred mostly ($\sim 95\%$) from the OH addition pathway,⁸⁸ with the remaining fraction of the reaction taking place through the hydrogen abstraction pathway. The resulting RO_2 radicals were $\text{C}_{10}\text{H}_7\text{O}_y$, $\text{C}_{10}\text{H}_9\text{O}_y$, and $\text{C}_{10}\text{H}_{11}\text{O}_y$, and the main dimers formed from the bimolecular reaction of these RO_2 radicals were $\text{C}_{20}\text{H}_{16}\text{O}_y$, $\text{C}_{20}\text{H}_{18}\text{O}_y$ and $\text{C}_{20}\text{H}_{20}\text{O}_y$.^{88–90} The suggested reaction pathways were given in Fig. S2,† and the resulting closed-shell oxidation products observed were compiled in Fig. S3 and Table S2.† The contribution of the oxidation pathways to SOA formation was estimated by comparing the ratios of C_{10}H_6 oxidation products

(formed through the H abstraction pathway) to $\text{C}_{10}\text{H}_{10}$ compounds (formed through the OH addition pathway). At the peak of SOA formation of the low- NO_x experiment, particle-phase $\frac{\text{C}_{10}\text{H}_6}{\text{C}_{10}\text{H}_{10}}$ ratio amounted to 0.07 while gas-phase $\frac{\text{C}_{10}\text{H}_6}{\text{C}_{10}\text{H}_{10}}$ ratio was 0.25, showing that $\text{C}_{10}\text{H}_{10}$ species were more abundant (relatively to C_{10}H_6) in the particle-phase than in the gas-phase. Similar observations from the beginning of SOA formation in the high- NO_x experiment were noted, with particle-phase and gas-phase $\frac{\text{C}_{10}\text{H}_6}{\text{C}_{10}\text{H}_{10}}$ amounting to 0.11 and 0.84, respectively.

This suggests that species produced from the OH addition pathway partitioned more efficiently towards particle-phase, although it is important to stress that these observations were qualitative.

In the low- NO_x experiment, the SOA chemical composition remained similar across the two injections (Fig. S3-A†). Moreover, most of the gaseous and particulate compounds detected were consistent with what was observed in previous studies on naphthalene oxidation.^{52,65,84,91} Table S2† depicts both N-containing and non-N-containing species identified in previous studies. However, to ease the comparison between the



different experiments, this section focuses on the non-N-containing compounds. Important non-N-containing oxidation products were identified such as phthalaldehyde ($C_8H_6O_2$), 1,4-naphthoquinone ($C_{10}H_6O_2$), and 2-formylcinnamaldehyde ($C_{10}H_8O_2$). Generally, condensed-phase non-N-containing compounds were highly oxidized (*i.e.*, $O \geq 4$ for monomers and ≥ 6 for dimers), with no visible difference between the low- and high- NO_x conditions. As shown in Fig. S3-A and Table S2,† some of the most abundant particle-phase compounds identified in the low- NO_x experiment included $C_8H_6O_4$ ($m/z = 166.026$, number 7 in Fig. S3 and Table S2†), $C_{10}H_8O_3$ ($m/z = 176.047$, number 10), $C_{10}H_8O_4$ ($m/z = 192.042$, number 13), $C_{10}H_{10}O_3$ ($m/z = 178.062$, number 11), $C_{10}H_{10}O_4$ ($m/z = 194.057$, number 14), and $C_{10}H_{10}O_5$ ($m/z = 210.052$, number 17). The three former compounds ($C_8H_6O_4$, $C_{10}H_8O_3$, and $C_{10}H_8O_4$) were ring-opening products. Contrariwise, $C_{10}H_{10}O_3$ was a ring-retaining product and $C_{10}H_{10}O_4$ and $C_{10}H_{10}O_5$ were also most likely ring-retaining compounds. High- NO_x particulate oxidation products (Fig. S3-C and Table S2†) exhibited a lesser fraction of non-N-containing ring-retaining species, suggested by the larger concentration of $C_{10}H_8O_4$ compared to $C_{10}H_{10}O_4$. The same trend was observed for the gas-phase oxidation products, where $C_{10}H_{10}O_2$ ($m/z = 160.052$, number 6) dominated in the low- NO_x environment (Fig. S3-B and Table S2†) while $C_{10}H_8O_2$ ($m/z = 162.068$, number 5) dominated in the high- NO_x environment (Fig. S3-D and Table S2†). These results agree with previous studies,^{52,84} which noted a higher relative intensity in gas-phase ring-retaining products for low- NO_x conditions compared to high- NO_x conditions.

EESI signal variations

Fig. S4-A† compares the variation of the wall-loss corrected SOA mass recorded by the SMPS with the evolution of the normalized total signal recorded by the EESI for $C_{8-10}H_xO_y$ and $C_{18-20}H_xO_y$ species during the low- NO_x experiment. The light period caused a 12% loss of the particle mass loadings, while the EESI normalized signal originally did not show a considerable drop. After that, the particle mass remained constant, while the EESI signal continued to decay quickly. A possible explanation can be the dependency of the EESI sensitivity as a function of the particle size. Indeed, as shown by Lee *et al.*⁹² the EESI signal depends not only on the mass loading but also on the geometric mean aerosol diameter. This is caused by smaller particles diffusing and coalescing more efficiently with the electrospray droplets, therefore resulting in greater extraction efficiency. However, this effect is mainly important for particles with a diameter lower than 100 nm. In our experiment, naphthalene-derived SOA exhibited a mean diameter of ~ 150 nm (Fig. 1A). The variation in the mean diameter during the low- NO_x irradiation period was roughly 3%, *i.e.*, 145 nm. This variation was negligible and should not have influenced the extraction efficiency. Furthermore, Pospisilova *et al.*³⁰ observed a similar trend, *i.e.*, a larger decrease in the EESI signal compared to the SOA mass loading. In their experiment, the mean diameter of the SOA did not vary significantly under irradiation, which was consistent with our

observations. Hence, the EESI signal decrease was most likely not caused by the EESI sensitivity dependence on the particles' diameter. Instead, Pospisilova *et al.*³⁰ proposed that the EESI-TOF was less sensitive to the composition of the aged α -pinene SOA, which were composed of a higher fraction of highly oxygenated compounds after irradiation. Lopez-Hilfiker *et al.*,⁸¹ who characterized the EESI-TOF, showed that the instrument was the most sensitive for species with a relatively low oxygen number ($O = 3-4$ for carboxylic acids, $5-6$ for saccharides and $4-6$ for α -pinene SOA), while more oxygenated species were detected less efficiently. Bell *et al.*⁹³ also found that the EESI-TOF had a positive bias towards the less oxygenated monomers, and a negative bias towards highly oxygenated monomers and dimers. A possible explanation is that more oxygenated species are less volatile, which makes them less likely to evaporate and be measured by the instrument. On the other hand, less oxygenated species could more readily evaporate from the EESI droplets. Therefore, the formation of more oxygenated species at the expense of lower oxygen-containing monomers could explain the loss of EESI-TOF sensitivity. In contrast to the photodegradation period, the decay rates of the mass and EESI signal during the dark aging period were consistent, both decreasing by $\sim 10\%$ by the end of the low- NO_x experiment and by $\sim 18\%$ by the end of the high- NO_x experiment (Fig. S4†). Dark aging did not change the chemistry of the SOA enough to cause a change in the EESI sensitivity, underlying the importance of photochemical aging.

Photolysis rates of bulk SOA

During the low- NO_x experiment, photolysis rates of the total wall loss-corrected particle mass and the total EESI signal were estimated to be $3.5 \times 10^{-3} s^{-1}$ and $2.7 \times 10^{-3} s^{-1}$, respectively. These photolysis rates were obtained using the method presented by Pospisilova *et al.*³⁰ In that previous study, photolysis rates of $6.5-7.6 \times 10^{-4} s^{-1}$ for the particle mass and $4.1-5.6 \times 10^{-4} s^{-1}$ for the EESI total signal were estimated. The differences in photolysis rates are likely caused by the fact that α -pinene-derived SOA are more photorecalcitrant than naphthalene-derived SOA. This is further supported by Romonosky *et al.*⁹⁴ who showed that naphthalene-derived SOA were more likely to be lost through photodegradation than α -pinene-derived SOA. Additionally, naphthalene-derived SOA in our low- NO_x experiment were formed under higher RH than Pospisilova *et al.*³⁰ Hence, naphthalene-derived SOA photolytic losses were most likely enhanced by the higher RH, as shown by Wong *et al.*⁴⁴ Indeed, photodissociation products in less viscous SOA (high RH) could more efficiently diffuse out of the particle, while recombination could have more time to happen in more viscous particles (low RH).

Global impact of photodegradation on SOA chemical composition

The time evolution of SOA species is depicted in Fig. 2A and B while the evolution of gaseous species is presented in Fig. 2C and D ($t = 0$ hours indicates when UV lights were turned on).



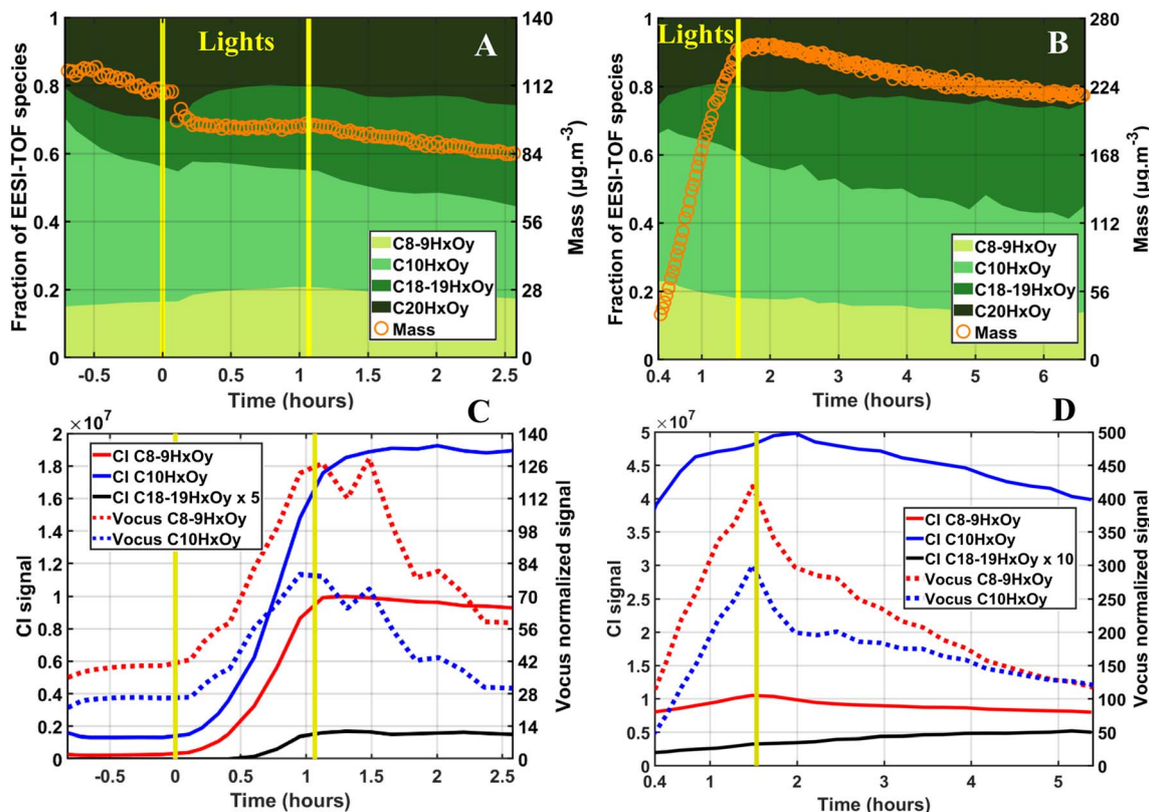


Fig. 2 Time evolution of the relative intensity of species (the sum of all compounds amounting to 1) and wall-corrected organic mass measured by the EESI-TOF and SMPS, respectively, for (A) the low- NO_x experiment ($t = -0.7$ hours to 2.6 hours) and (B) the high- NO_x experiment ($t = 0.4$ hours to 6.6 hours). The evolution of the signal for the gaseous species, measured by the CI (full lines) and the Vocus (dotted lines), was shown in (C) and (D) for the low- NO_x and high- NO_x experiments, respectively. The irradiation period started at $t = 0$ hours and is delimited by the yellow lines, ending just over $t = 1$ hour for (A) and (C), and ending at $t \sim 1.5$ hours for (C) and (D).

The time scale during the low- NO_x experiment showed species formed in the dark (Fig. 2A and C starting at $t = -0.7$ hours) before they were exposed to UV lights ($t = 0$ hours). During the high- NO_x experiment, the time scale of the experiment started after irradiation began (Fig. 2B and D starting at $t = 0.4$ hours), as species could not be detected by the EESI-TOF before that point. EESI-TOF figures (Fig. 2A and B) show the evolution of the relative intensity of the species, with all compounds combined amounting to 1. Gas-phase figures (Fig. 2C and D) show the evolution of the signals of compounds measured by the CI-Orbitrap and the Vocus. The compounds were grouped as $\text{C}_{8-9}\text{H}_x\text{O}_y$, $\text{C}_{10}\text{H}_x\text{O}_y$, $\text{C}_{18-19}\text{H}_x\text{O}_y$, and $\text{C}_{20}\text{H}_x\text{O}_y$, the first two groups being monomers and the last two groups being dimers. These compounds are expected to contain functionalities such as a peroxide group or an ester group as the linking functionality between the two monomeric units.^{18,95} Dimeric compounds such as ester accretion products can also be produced from particulate-phase reactions.^{95,96} Unfortunately, the instruments used in this study do not provide information about the chemical structures.

Fig. 2A shows that photodegradation-induced SOA mass loss correlated with a change in particle composition during the low- NO_x experiment. This change was seen for dimeric species, which evolved differently under irradiation depending on their

carbon numbers. The main dimers produced from gas-phase chemistry, *i.e.*, $\text{C}_{20}\text{H}_x\text{O}_y$ species, experienced a net loss during photodegradation. On the other hand, the fraction of dimeric compounds with lower carbon numbers, *i.e.*, $\text{C}_{18-19}\text{H}_x\text{O}_y$ species, increased when the lights were on. The difference in dimer composition ($\text{C}_{20} \rightarrow \text{C}_{18,19}$) suggest that dimer photodegradation altered specifically $\text{C}_{20}\text{H}_x\text{O}_y$ species. Similar results were observed in Fig. 2B for the high- NO_x experiment. It should be noted that Lee *et al.*⁹¹ showed a decrease in the double bond equivalent (DBE) which they associated with the fragmentation of larger naphthalene-derived SOA compounds.

Regarding the monomers during the low- NO_x experiment, the main species ($\text{C}_{10}\text{H}_x\text{O}_y$) initially increased when the lights were turned on and then slowly decreased under irradiation, while the $\text{C}_{8-9}\text{H}_x\text{O}_y$ molecules steadily increased with irradiation. It is worth mentioning that the fraction of monomers and dimers remained at similar levels (55% and 45% respectively) before and after the lights were on. This indicates that accretion reactions were not the main reaction pathway. The gaseous monomers and dimers species also increased, as seen in Fig. 2C. Comparing the evolution of gaseous C_{20} dimers was not possible since the instruments were not able to detect $\text{C}_{20}\text{H}_x\text{O}_y$. A possible reason is the prompt formation of new particles which reached high mass loadings and provided an additional



condensation sink that could have favored the partitioning of species with low volatility, such as $C_{20}H_xO_y$ oxidation products. However, the evolution of the gaseous monomers did not show significant differences between $C_{8-9}H_xO_y$ and $C_{10}H_xO_y$ where the gas-phase species were delayed for those formed in the particulate phase.

Finally, dimeric species also behaved differently than monomers in the dark. In the dark, both experiments showed that the fraction of particulate $C_{10}H_xO_y$ and $C_{8-9}H_xO_y$ started decreasing once irradiation stopped, while the fraction of dimers increased. This process was probably not linked to partitioning, as the gas-phase monomers remained stable or decreased in the dark. Species detected by the Vocus seemed to decrease faster than those measured by the CI-Orbitrap, most likely due to the instrument's selectivity toward different species.^{74,80}

Photodegradation of C_xH_y compounds

To elucidate which compounds were optically active or formed from the photodegradation, oxidation products were sorted out according to their carbon and hydrogen numbers as well as their evolution. Before photodegradation in the low- NO_x experiment, $C_{20}H_{16-22}O_y$ had high relative signals (see Fig. 3 or Table S3[†] for numeric values). As shown in Fig. S2,[†] such compounds can be expected from bimolecular RO_2 reactions and can rapidly condense and partition to the particle-phase due to their low vapor pressure. However, these species could also have been formed from particle-phase reactions. The current data does not allow to confirm the dominance of one pathway to another. Nevertheless, these species increased in the

dark before being depleted by the photodegradation process which led to the production of species with a different number of carbon, hydrogen, and oxygen atoms.

During the low- NO_x experiment, photodegradation was not uniform across all dimeric compounds and appeared to be dependent on the hydrogen number. The most saturated C_{20} species (*i.e.*, $C_{20}H_{18-22}O_y$), which were expected from radical reactions as described before, experienced a greater loss under irradiation than the compounds having larger DBE (*e.g.*, $C_{20}H_{14,16}O_y$). Similarly, the increase of relative signal for C_{18-19} species was more important for the most unsaturated species (*e.g.*, $C_{18-19}H_{14-16}O_y$) than for the ones with a higher hydrogen-content. This indicated that the unsaturated species were mainly produced when the UV lights were on. It is worth noting that they were moderately formed from gas-phase chemistry and then remained constant in the condensed phase when the UV lights were off.

This difference in reactivity could also be explained by the nature of the functional groups present in the molecule. It is known that carbonyl and peroxy moieties are photolabile and likely to be depleted under irradiation.⁹⁷ It has also been shown that photodegradation results in an increase of species carrying a carboxylic acid group.^{30,65,97} Replacing an organic peroxy functional group with a carboxylic acid would be associated with the loss of at least 2 hydrogen atoms, which aligns with the increase of unsaturated dimers. Additionally, dimers with a high H number probably did not bear functional groups associated with a reduced H content, such as carbonyls and carboxylic acids. Instead, they were more likely to bear organic peroxide moieties or hydroxyl groups. Finally, similar observations can be made for monomeric species such as $C_{10}H_xO_y$, decreasing under the UV

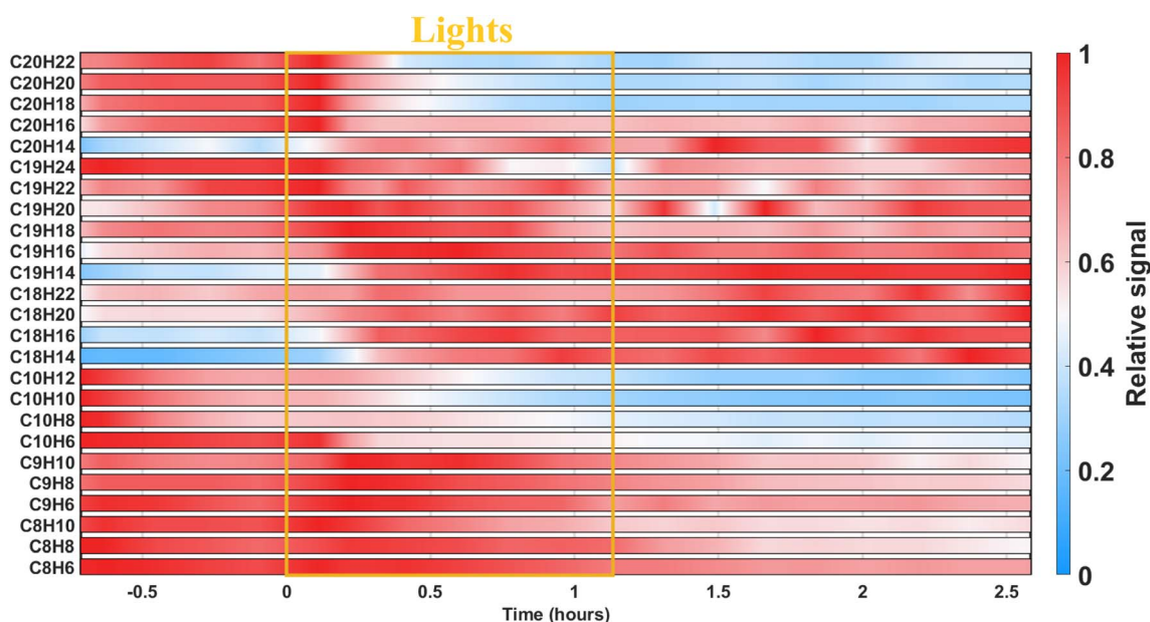


Fig. 3 Evolution of the relative signal of particle-phase species during the low- NO_x experiment, with compounds grouped by carbon and hydrogen numbers. Red indicates that the species are close to their maximum relative intensity, while blue signals a low relative intensity. UV lights are indicated by the yellow box.



lights due to photodegradation, with the most saturated species (*i.e.*, $C_{10}H_{10-12}O_y$) decreasing more than the ones with lower H content ($C_{10}H_{6-8}O_y$). $C_8H_xO_y$ behaved similarly, although the variations were not as important as for $C_{10}H_xO_y$. Interestingly, these changes appeared to be exclusive to the particulate phase, as all gas-phase monomers increased homogeneously under irradiation as depicted in Fig. S6.† Similar results were found in the high- NO_x experiment, as shown in Fig. S7,† for both non-N-containing and N-containing dimers. This suggests that

autoxidation reactions resulted in oxidation products bearing photolabile functional groups (such as carbonyls)^{40,42-44} under both low- and high- NO_x levels.

Evolution of individual species under irradiation

After having determined how the families of species evolved under UV lights, variations of the individual particulate monomers and dimers during the low- NO_x experiment were

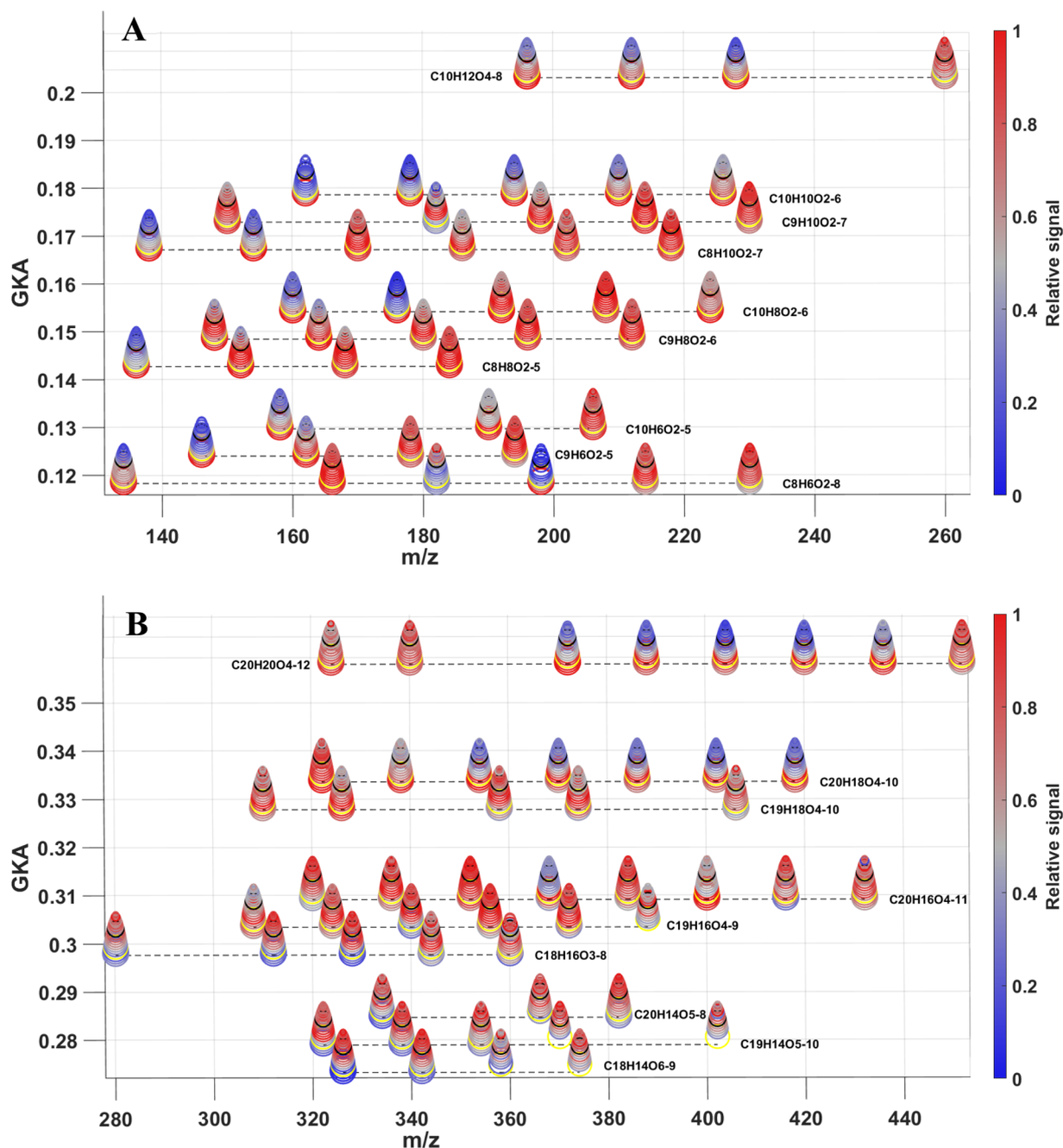


Fig. 4 Stacked generalized Kendrick analysis plots for (A) monomers and (B) dimers during the low- NO_x experiment. The multiple rings show the elapsed time from the start of the experiment (largest and lowest ring) to the end of the experiment (smallest and highest ring). The relative intensity of each compound is indicated by the color (blue = low intensity, red = high intensity, grey = medium intensity). The yellow and black rings indicate the beginning and the end of the irradiation period respectively.



further explored as shown in Fig. 4A and B respectively. This was done using stacked general Kendrick analysis (GKA) plots. GKA plots are related to traditional Kendrick mass defect analysis but use a tunable parameter to contract or expand the mass scale and a base unit that determines the horizontal alignment of the species. In this study, the base unit chosen was oxygen, so species were plotted horizontally according to their oxygen content. To generate the stacked figures, GKA values were calculated for different experimental times using the method presented by Alton *et al.*,⁹⁸ and were then stacked on top of each other to represent the time evolution of the relative intensity of the compounds. Many low-oxygenated species ($O = 2-4$), both for $C_{8-9}H_xO_y$ and $C_{10}H_xO_y$, decreased under irradiation. This suggests that photolabile functional groups were already present even for low-oxygenated species. Therefore, the OH-initiated oxidation of naphthalene must have resulted in at least one photolabile moiety early on. This was supported by previous studies showing monomers with a low oxygen content bearing carbonyl functional groups.^{84,85,99} However, it cannot be ruled out that the loss of $C_{8-10}H_xO_{2-4}$ could also be due to repartitioning from the particle-phase to the gas-phase, as proposed by Pospisilova *et al.*³⁰ As a result, the less oxygenated particulate monomers were then possibly depleted both by photodegradation and/or evaporation. More oxidized species should also have been sensitive to photodegradation but seemed to increase (*e.g.*, $C_8H_8O_{3-5}$, $C_9H_6O_{4-5}$, $C_9H_8O_{4-6}$, and $C_9H_{10}O_{5-7}$) or remain stable during irradiation, as also observed in previous studies.^{30,97} The lack of loss could be explained by the presence of photorecalcitrant functional groups such as carboxylic acids or through the production of oxidants in the particle-phase during irradiation producing highly oxygenated monomers.²⁶ Another source of those species would be gas-phase compounds condensing after being formed under UV lights (Fig. S8-B†). The decay in the dark of the compounds within one family seemed to also depend slightly on the O content of the species, as the most oxidized particulate compounds decayed more slowly (*e.g.*, $C_9H_{10}O_7$ compared to $C_9H_{10}O_2$). Higher oxygenated compounds have lower volatility, meaning that even with photodegradation they could continue to have sufficiently low volatility to remain in the particle-phase.

The evolution of the dimers is presented in Fig. 4B. Similarly to the monomers, the photodegradation efficiency of the $C_{20}H_{16-20}O_y$ compounds was dependent on the oxygen number. However, whereas many low-oxygenated monomers were lost upon photodegradation, this was not the case for the dimers, the low-oxidized dimers seeming less inclined to be photodegraded. Contrariwise to the monomers, dimers with a low oxygen content ($O = 3-5$) were less likely to evaporate from the particle phase. Additionally, $C_{20}H_xO_y$ species with a low number of oxygen atoms were less photodegraded. A possible explanation is that less oxygenated $C_{20}H_xO_y$ species possessed fewer functional groups, which decreased the chance of carrying a photolabile functional group. Furthermore, low-oxygenated dimers could also have been formed from the photodegradation of the more oxygenated dimers. Almost all the $C_{18-19}H_{14-18}O_y$ showed an increase in production during irradiation. The clearest increase concerned $C_{18-19}H_{14-16}O_{\approx 7}$ species, *i.e.*,

unsaturated low-oxidized dimers. This was consistent with the results of the $C_{20}H_{16-20}O_y$, whose photodegradation would have resulted in $C_{18-19}H_xO_y$. Many of the $C_{18-19}H_{14-16}O_{>7}$ compounds were not present at first, but were formed during irradiation. Overall, this showed that the photodegradation of the particles was not only dependent on the hydrogen and carbon numbers but also on the oxygen number of a molecule. The global O : C ratio slightly increased after the photodegradation (from 0.408 to 0.417), which was similar to the observations in Bateman *et al.*,⁹⁷ who showed an increased O : C ratio after the irradiation of D-limonene-derived SOA. Results from high- and low- NO_x experiments were consistent as presented in Fig. S10.†

Emission of volatile species from particle- to gas-phase following photodegradation

The photodegradation of $C_{20}H_{16-20}O_y$ to $C_{18-19}H_{14-16}O_y$ would most likely result in the emission of formic acid (HC(O)OH), formaldehyde (HCHO), acetic acid ($CH_3C(O)OH$), or acetaldehyde (CH_3CHO) to the gas-phase, as proposed in Fig. 5A. The emission of these species is a distinctive sign of photodegradation that has been shown in multiple SOA photodegradation studies.^{35-37,44,97} We investigated the production of these species using the Vocus and found that formic acid, acetic acid, and acetaldehyde increased during photodegradation (Fig. 5B and C). This confirms the photodegradation pathway and has important atmospheric implications. Indeed, these species are also highly important in the atmosphere due to their abundance, the acids' ability to contribute to the free atmospheric acidity, and the aldehydes' role as transition species leading to the formation of HO_2 or CO in the atmosphere.^{100,101}

To estimate the concentrations of acetic acid, formic acid, and acetaldehyde emitted, previous experiments from the same campaign using known amounts of monoterpene injected as well as the Vocus sensitivity towards these monoterpenes were used. Furthermore, we estimated the sensitivity of the Vocus towards the photodegradation volatile species by comparing the differences in instrumental sensitivity between monoterpene and organic acids/carbonyls from another study.¹⁰² The resulting concentrations are shown in Table 1 and reached in the low- NO_x experiment up to 3 ppb, 0.5 ppb, and 0.07 ppb of acetic acid, formic acid, and acetaldehyde, respectively. During the high- NO_x experiment, emissions reached up to 10 ppb, 1 ppb, and 0.08 ppb for acetic acid, formic acid, and acetaldehyde, respectively. The amounts of volatile products emitted were consistent with the loss of SOA mass in the low- NO_x experiment, but it should be noted that these results remain qualitative.

Evolution of the volatility pre-to post-photodegradation

To characterize the change in the particles' physical properties that is expected from the changed chemical composition, the effective saturation mass concentration (C^*) of each particulate oxidation product has been calculated. The estimation was made using the equation from Li *et al.*¹⁰³ and followed their structural and formulaic estimations. Particle-phase oxidation products were categorized as intermediate volatile organic compounds (IVOCs), semi-volatile organic compounds (SVOCs),



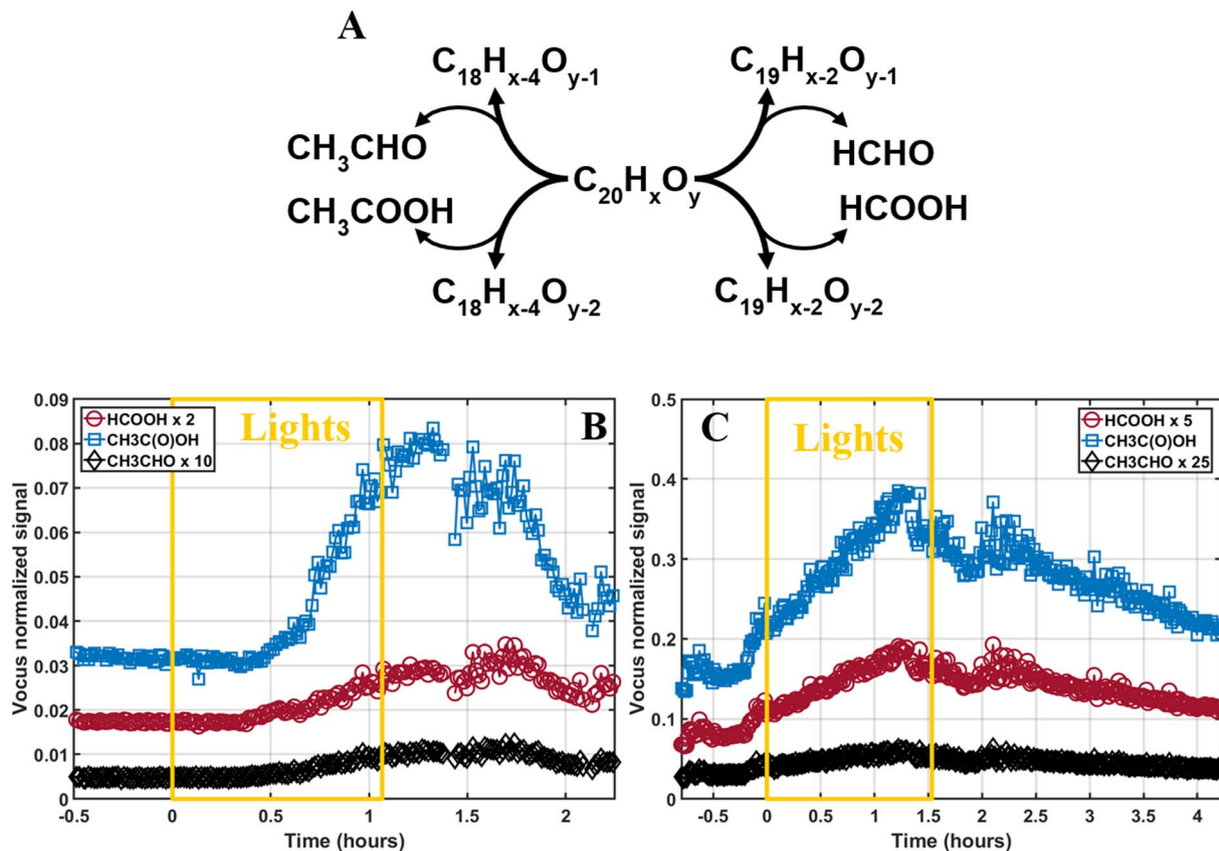


Fig. 5 (A) Photodegradation mechanism of C₂₀H_xO_y dimers. The time evolution of formic acid (HC(O)OH), acetic acid (CH₃C(O)OH), and acetaldehyde (CH₃CHO) are shown during (B) low- and (C) high-NO_x experiments. The intensity of the HC(O)OH and CH₃CHO signals was adjusted for easier visualization. $t = 0$ hours indicate the start of lights, and the yellow box indicates the irradiation period.

Table 1 Maximum estimated amounts of volatile species emitted during photodegradation during the low- and high-NO_x experiments. To estimate these amounts, the normalized signals to injected masses ratios of monoterpene were used (corrected for the instrument sensitivity)

	ppb	$\mu\text{g m}^{-3}$	SOA mass loss	% Of SOA loss from the volatile species
Low-NO _x	Acetic acid: 3	Acetic acid: 7.5	14 $\mu\text{g m}^{-3}$	Acetic acid: 54%
	Formic acid: 0.5	Formic acid: 0.93		Formic acid: 7%
	Acetaldehyde: 0.07	Acetaldehyde: 0.12		Acetaldehyde: 0.9%
High-NO _x	Acetic acid: 10	Acetic acid: 25	X	X
	Formic acid: 1	Formic acid: 2.3		
	Acetaldehyde: 0.08	Acetaldehyde: 0.15		

low volatile organic compounds (LVOCs), extremely low volatile organic compounds (ELVOCs), and ultra-low volatile organic compounds (ULVOCs) according to their estimated C^* . Fig. 6 shows how the fraction of these species evolved during the low-NO_x experiment pre- (Fig. 6A) to post-photodegradation (Fig. 6B). The fraction of the species during the high-NO_x experiment at the beginning of the photodegradation and after photodegradation ended are displayed in Fig. 6C and D, respectively. Both experiments showed that species with low volatilities (*i.e.* LVOCs, ELVOCs and ULVOCs) slightly increased in both experiments after the particles were photodegraded. Additionally, the mean C^* was calculated and showed that irradiated particles were less volatile after photodegradation

(from 0.74 to 0.67 $\mu\text{g m}^{-3}$ for low-NO_x and from 1.53 to 1.04 $\mu\text{g m}^{-3}$ for high-NO_x). Low-NO_x particles globally showed a lower C^* than high-NO_x particles, but low-NO_x C^* also did not decrease as much as high-NO_x C^* after irradiation. This was consistent with the larger decrease of IVOC post-photodegradation for the high-NO_x experiment. Although C^* variations were not very significant, they still agreed with the results from Pospisilova *et al.*³⁰ who showed that α -pinene SOA presented lower volatility after its exposure to UV lights by performing isothermal evaporation experiments. However, our results may have been influenced by changes in the parametrizations used to calculate C^* . Indeed, these parametrizations may have differed from pre-photodegradation to post-



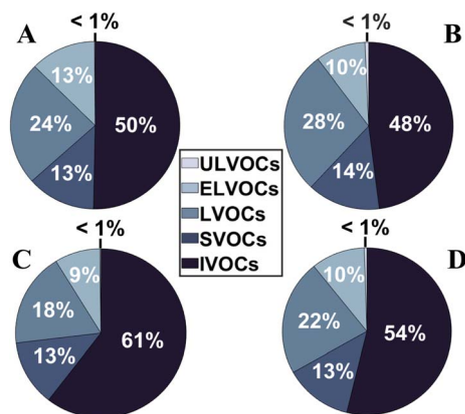


Fig. 6 (A) Volatility distribution of the particulate oxidation products for the low-NO_x experiment before photodegradation ($t = -0.1$ h) and (B) after photodegradation ($t = 1.13$ h). (C) Volatility distribution for the high-NO_x experiment at the beginning of photodegradation ($t = 0.37$ h) and (D) after photodegradation ($t = 1.73$ h). Colors indicate the fraction of ultra-low volatile (ULVOCs), extremely low volatile (ELVOCs), low volatile (LVOCs), semi-volatile (SVOCs) and intermediate volatile (IVOCs) organic compounds respectively.

photodegradation as the functional groups in the particle-phase compounds could have changed. The fraction of non-N-containing and N-containing species in the high-NO_x experiment remained unaffected by photodegradation. It went from a 0.76/0.24 ratio for non-N-containing and N-containing species respectively at the beginning of photodegradation ($t = 0.4$ hours) to a 0.75/0.25 ratio after photodegradation ($t = 1.7$ hours).

Conclusion

This study investigated the composition of naphthalene SOA and the influence of photodegradation as a marker for anthropogenic VOC/PAH aerosol photodegradation due to its abundance and role in urban environments. Our results showed that the changes observed in particles aging under irradiation were significantly greater than the variations caused by other chemical processes. These other processes were controlled by letting particles age under a similar timescale in the dark. Prominent degradation (2/3 loss) of the C₂₀ dimers during photodegradation was revealed through an in-depth molecular characterization of naphthalene-derived SOA. The degradation of the dimers yielded particulate-phase fragmentation products with C₁₈₋₁₉ while releasing C₁₋₂ gaseous compounds. Photodegradation was found to be dependent on the chemical composition and functional groups with hydroperoxide and carbonyl species being the most photolabile groups. SOA physical properties were also affected by photodegradation, as particles showed a decrease in volatility. Further work coupling the EESI-TOF to a thermal denuder as done by Bell *et al.*⁹³ would provide more accurate information about the particles changing volatilities.

The photochemical process was accompanied by a loss of particle mass under irradiation (−12%) which corresponded to

the release of volatile products. Among them, three products playing a major role in global atmospheric chemistry were identified: formic acid, acetic acid, and acetaldehyde. These results highlight the need to consider photochemical activity as one of the main drivers of SOA aging as naphthalene-derived SOA and other anthropogenic-sourced aerosols are an important factor affecting health, visibility, and the environment. Predicting their impact on any of these subjects requires information about their chemical composition and physical properties, both of which are modified due to UV irradiation, showcasing the complexity of considering SOA photodegradation in broader atmospheric issues.

Data availability

The data that support the findings of this study are available on request from the corresponding authors, D. B. at: david.bell@psi.ch and/or M. R. at: matthieu.riva@ircelyon.univ-lyon1.fr.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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