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The effect of aging conditions at equal OH exposure in an oxidation flow reactor on the composition of toluene-derived secondary organic aerosols†

Hendryk Czech, **D**abc** Pasi Yli-Pirilä, **Petri Tiitta, **; ** Mika Ihalainen, **D** Anni Hartikainen, **D** Eric Schneider, **ad Patrick Martens, **§ Andreas Paul, **D** ae Thorsten Hohaus, **D** e Christopher P. Rüger, **ad Jorma Jokiniemi, **D** Ralf Zimmermann** and Olli Sippula**

Oxidation flow reactors (OFRs) have been increasingly used to conduct research on secondary aerosol formation potential and composition in laboratory and field studies by exposing aerosols to high levels of oxidants in short time periods. In order to assess the atmospheric relevance of the triggered chemical reactions, kinetic models have been developed to reveal the production of atmospheric oxidants and the fate of volatile organic compounds (VOCs). However, it is unknown how different OFR conditions generating the same OH exposure affect the chemical and physical properties of the secondary aerosol particle phase because the model is based on gas phase chemistry. Toluene as a well-investigated precursor of secondary organic aerosols (SOAs) was aged in the high-volume OFR "PEAR" at three different external OH reactivities (OHRext) and in an environmental chamber at the same OH exposure of (1.09 \pm 0.09) \times 10¹¹ s cm⁻³. These specific OFR conditions altered the majority of the investigated chemical and physical properties of the toluene-derived SOA (tol-SOA). However, OFR-aging at low OHR_{ext} associated with atmospherically relevant conditions did not lead to physical and chemical SOA properties most similar to chamber-generated SOAs at the same low OHR_{ext}. Particularly, for the most detailed chemical analysis by electrospray (ESI) high-resolution Orbitrap mass spectrometry (HRMS), tol-SOA from the PEAR at high OHR_{ext}, deviating from atmospherically relevant conditions, and the chamber were most different to tol-SOA from the PEAR with low and medium OHR_{ext}. Our study challenges the concept of atmospherically relevant OFR aerosol aging and motivates further exploration of OFR conditions on secondary aerosol composition.

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Environmental significance

Oxidation flow reactors (OFRs) became a widely used device for laboratory atmospheric aging studies, especially for bridging the gap between the maximum age achievable in an environmental chamber and aerosol particle lifetime in the atmosphere. In OFRs, aerosols are exposed to high concentrations of oxidants within a short time in order to cover exposures to OH radicals equivalent to up to two weeks. This approach has been criticized to induce irrelevant chemical reactions, so kinetic models have been developed to define constraints for atmospherically relevant OFR operation. Experimental results from toluene photooxidation challenge the concept of atmospherically relevant OFR conditions and explore which physical-chemical properties of secondary organic aerosols from toluene photooxidation may be reproduced from complementary experiments in an environmental chamber.

^aChair of Analytical Chemistry, University of Rostock, 18059, Rostock, Germany. E-mail: hendryk.czech@uni-rostock.de

^bFine Particle and Aerosol Technology Laboratory, Department of Environmental and Biological Science, University of Eastern Finland, 70211, Kuopio, Finland

[&]quot;Cooperation Group "Comprehensive Molecular Analytics" (CMA), Helmholtz Zentrum München, 81379, München, Germany

^dDepartment "Life, Light and Matter", University of Rostock, 18059, Rostock, Germany

^eInstitute of Energy and Climate Research: Troposphere (IEK-8), Forschungszentrum Jülich, 52425, Jülich, Germany

Department of Chemistry, University of Eastern Finland, 80101, Joensuu, Finland † Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d4ea00027g

[‡] Now at: Envineer Oy, Kuopio, Finland.

[§] Now at: Desert Research Institute (DRI), Reno, NV, USA.

1. Introduction

Secondary organic aerosols (SOAs) from the oxidation of volatile organic compounds (VOCs) largely contribute to the ambient organic aerosol burden from both anthropogenic and biogenic sources. ^{1,2} Depending on the detailed chemical composition and optical properties, SOAs may affect the climate³⁻⁵ and human health. ⁶⁻⁹ The oxidative gas-to-particle conversion of VOCs to SOAs is initiated by reactions with hydroxyl (OH) radicals, ozone (O₃), and nitrate radicals (NO₃) or *via* photolysis, with OH being the dominant atmospheric oxidant during daytime. ¹⁰

Laboratory atmospheric aging of VOCs, VOC mixtures including plant emissions, and combustion aerosols is conducted to understand atmospheric processes and composition as a bottom-up approach.11 Environmental chambers are wellestablished and regarded as the most realistic model for laboratory atmospheric aging. Both aerosol and atmospheric oxidants are injected or generated at near-ambient levels, so the aerosol residence in the environmental chamber appears close to the atmospheric residence time of an aerosol under equivalent conditions.12 Losses to the inner chamber walls continuously decrease aerosol concentrations, which can be corrected by inert tracers or experiments with model aerosols without aging. Thus, the uncertainty of aerosol measurement increases with longer residence in environmental chambers. Consequently, atmospheric aging in environmental chambers of common size has an inherent upper limit of about two equivalent days in the atmosphere. Therefore, some environmental chambers with large volumes were built to minimize wall effects for studies on atmospheric aerosol concentrations at remote sites (SAPHIR with 270 m³)¹³ to the upper level like in dense wildfire plumes (PHOTO-LAC with 1800 m³).14 Nevertheless, residence times of aerosol particles in the troposphere of up to two weeks cannot be reached.

Oxidation flow reactors (OFRs) expose aerosols to high levels of OH radicals on the time scale of a few minutes.15 In OFR254 mode, OH radicals are generated from the photolysis of O₃ at 254 nm to singlet oxygen O(1D), which subsequently reacts with water vapor to give two OH radical equivalents. Alternatively, ozone may be generated in situ by first photolysis of molecular oxygen (O₂) to triplet oxygen atoms O(³P) (OFR185 mode) and subsequent reaction with O2. Moreover, OH radicals may be formed by the photolysis of water vapor. 15 In these ways, high OH exposures (time \times OH concentration) of 1×10^{12} s cm⁻³ equivalent to up to 14 days and beyond may be reached, and were used to study OH-initiated SOA formation from VOCs, 16,17 combustion aerosols,18,19 ambient air20,21 and also heterogeneous oxidation of carbonaceous particulate matter.22,23 However, concerns have been raised that OFR chemistry may be generally irrelevant to tropospheric chemistry. 15

Peng *et al.* (2016, 2017, and 2019)^{24–27} developed a model for the potential aerosol mass (PAM) OFR^{28,29} to define the space of atmospherically relevant reactions and beyond, for example by the classes "safer", "transition" and "riskier" concerning the contribution of non-OH oxidation reactions under low-NO_x conditions, or the fate of peroxyalkyl radicals (RO₂). This model

is based on gas phase chemistry and assesses the atmospherically relevant space using the ratios of atmospheric oxidants and established reaction pathways. The crucial parameters of OFR aging are the external OH reactivity (OHRext), which is the sum of the product of reactive species concentrations with their rate constants with OH radicals, the photon flux, ozone addition or in situ generation, and the level of humidity as well as the level of NO_x. Combined with the OFR residence time, the OFR fate of individual reactive species, such as by reactions with OH radicals, ozone or photolysis, may be calculated and compared to the fate under real atmospheric conditions. However, it is unknown how OFR conditions of different atmospheric relevance affect the physical and chemical properties of the formed secondary aerosol particles, which has consequences for toxicological studies with SOAs and aged aerosols, 30-33 because the atmospheric representativity of the formed aerosol particle characteristics presumably plays a critical role.

In this study, the SOA precursor toluene was photochemically aged in the high-volume OFR "PEAR", ³⁴ operated in OFR254-7 mode (external feed of 7 ppm ozone into the OFR and photolysis at 254 nm) without NO_x addition. ¹⁵ Toluene concentrations and photon flux were adjusted in order to obtain "safer", "transition" and "riskier" OFR conditions at an equal OH exposure, *i.e.* an equal photochemical equivalent age. ²⁴ Furthermore, photochemical aging of toluene was conducted in an environmental chamber as a reference for the most realistic laboratory model for atmospheric processing. The physical and chemical properties of a toluene-derived SOA (tol-SOA) under all conditions were analyzed by several online and offline analytical techniques and discussed concerning the effect of OFR conditions and the similarity of OFR-generated tol-SOA to chamber-generated tol-SOA.

Materials and methods

2.1. Generation of tol-SOA

Different concentrations of toluene vapor ("low", "medium", and "high") were generated by gentle heating of a reservoir filled with toluene. The reservoir was connected by stainless steel tubing, so toluene vapor entered the main flow of $100 \, \mathrm{L} \, \mathrm{min}^{-1}$ of purified air (737-15 zero air generator; Aadco Inc., VT, USA) to the "Photochemical Emission Aging flowtube Reactor" (PEAR), so the mode of the residence time distribution appeared at \sim 70 s. The PEAR is a high-volume oxidation flow reactor made of stainless steel with a volume of 139 L and an inner surface to volume ratio of 16.4 $\, \mathrm{m}^{-1}$. It consists of a conical inlet diffuser, a cylindrical center part and a conical outlet. Inside the cylinder center part, four UV lamps are mounted, which may be turned on and off individually. More information on the PEAR reactor may be found elsewhere.³⁴

Prior to entering the PEAR, water vapor, ozone, and toluene vapor were mixed (OFR254-7 mode) to obtain 50% relative humidity, 7 ppm of ozone and three different toluene concentrations. The three different settings for the PEAR were chosen to achieve "safer", "transition" and "riskier" according to the "OFR Exposures Estimator v3.1"^{24,27} by varying toluene concentrations (and consequently OHR_{ext}) and UV lamp intensity but keeping the same OH exposure of $(1.09 \pm 0.09) \times 10^{11}$ s cm⁻³ (Table 1). OH exposure was determined online using the ratio of

Table 1 Settings of individual toluene aging experiments with the initial toluene concentration [tol]_{in}, external OH reactivity (OHR_{ext}), photon flux at 254 nm and OFR conditions based on the "OFR Exposures Estimator v3.1".24.27 All OFR experiments (low, medium, and high) were conducted under 7 ppm ozone and 50% relative humidity at approximately 20 °C. Chamber experiments (cham) were conducted at 50% relative humidity with H_2O_2 as the source of OH radicals without the addition of ozone. No seed aerosol was used in any experiment and the blank particle mass (derived from SMPS assuming a density of 1.5 g cm⁻³) was below 0.3 μ g m⁻³

Experiment	[tol] _{in} [ppb]	$OHR_{ext}\left[s^{-1}\right]$	OH _{exp} [s cm ⁻³]	Photo flux \textcircled{a} 254 nm [cm ⁻² s ⁻¹]	OFR conditions	Y _{SOA} [%]
low1	316	37	9.94×10^{10}	8.71×10^{14}	Safer	4.4
low2	335	39	1.05×10^{11}	8.71×10^{14}	Safer	4.3
low3	298	35	1.16×10^{11}	8.71×10^{14}	Safer	3.1
med1	680	79	1.07×10^{11}	1.76×10^{15}	Transition	5.0
med2	673	78	1.10×10^{11}	1.76×10^{15}	Transition	4.5
med3	663	77	1.17×10^{11}	1.76×10^{15}	Transition	3.8
high1	3840	445	1.08×10^{11}	7.06×10^{16}	Riskier	6.2
high2	3850	446	1.09×10^{11}	7.06×10^{16}	Riskier	5.8
high3	4010	465	1.14×10^{11}	7.06×10^{16}	Riskier	4.8
cham1	309	36	9.99×10^{10}	0	_	39.7
cham2	303	35	1.07×10^{11}	0	_	40.5
cham3	313	36	1.07×10^{11}	0	_	39.9

introduced toluene into the PEAR and toluene at the PEAR outlet, according to the concept of "photochemical clock",35 which was measured by single-photon ionization time-of-flight mass spectrometry (SPI-TOFMS, see Section 2.2.1.). Particle measurements and sampling after the PEAR were corrected for wall losses using the loss function for silver particles at 100 L min⁻¹ as provided in Ihalainen et al. (2019).34 Due to the assumed relatively high volatility of tol-SOA, true losses are likely higher, so the wall losscorrected results denote a lower limit of particle concentrations.

Complementary to OFR-aging, the experiment was repeated under similar conditions in the ILMARI smog chamber of 27 m³ volume36 with a target concentration as for "low" OFR conditions (Table 1). As an OH source without NO_x generation, H₂O₂ was added to the chamber with an air flow over one hour by vaporizing 1 mL of an aqueous H2O2 solution in a flask. The resulting concentration of H₂O₂ was estimated to be <2.7 ppm, which is similar to other chamber experiments.37 Subsequently, toluene was injected into the chamber to reach the same concentration as used in "safer" OFR aging, which required about 5 h of residence. Finally, 340 nm blacklights were turned on with full power for approximately five hours until the same OH exposure as in OFR experiments was reached. The background concentrations of PM, NO2 and O3 in the chamber were below 0.3 μ g m⁻³, 4 ppb and 2.4 ppb, respectively. All chamber experiments were conducted at a relative humidity of 50% and a temperature of approximately 20 °C. Particle measurements and sampling from the chamber were corrected for wall losses using the loss function for polydisperse ammonium sulfate particles as provided in Leskinen et al. (2015).36 Due to the assumed relatively high volatility of tol-SOA, true losses are also likely higher in the chamber, so the wall loss-corrected results denote a lower limit of particle concentrations. Online data on chamber aging were obtained by data acquisition for 30 min after UV lamps had been switched off.

A summary of the experimental conditions is listed in Table 1.

The assessment of non-OH chemistry^{24,27} revealed that the reactions of methane, ethane, and benzene with O(1D) and O(3P) would be comparable to ambient air conditions, while the reactions of terpenes α -pinene and γ -terpenene as well as isoprene would be underestimated in all three OFR conditions. The photolysis of toluene at 254 nm becomes slightly overestimated in "high" with 10% atmospheric fate, while acetylacetone photolysis accounts for 55%, which appears clearly outside the considered atmospherically relevant range. However, the model proposed by Peng et al. (2016)24 assumes photodissociation quantum yields of unity, so the model prediction denotes an upper limit for the fate of photolysis. Therefore, the fates of toluene in the PEAR may be closer to the fates in the atmosphere and other processes determine potential deviations from atmospheric SOA formation. Experiments with toluene exposed to different intensities of 254 nm UV light without the addition of ozone may reduce this bias in the model result.

Organic peroxy radicals (RO2) may be formed from the reaction of O₂ with carbon-centered radicals. The subsequent atmospheric fate of RO2 involves possible reactions with hydroperoxyl radicals (HO₂), NO or previous RO₂ isomerization. In the case of primary RO₂ and acyl RO₂, reactions between different RO₂ (RO₂ + RO₂) may be significant in the atmosphere but are often underrepresented in OFR aging. In OFR aging with high levels of OH and high OHR_{ext}, the RO₂ fate is dominated by OH (RO₂ + OH), deviating from atmospheric processing.26 Furthermore, sufficiently long RO2 lifetimes for a relevant contribution of isomerization to the RO2 fate may be achieved by conditions with high UV intensities, but then photolysis at 254 nm becomes a relevant fate of toluene, which is not anymore in the atmospherically relevant range either. Thus OFR254-7 conditions cannot lead to overall good, i.e., atmospherically relevant aging.²⁶ Even in "low" belonging to "safer" OFR conditions (Table 1), RO₂ + OH accounted for 56.3%, which increased in "high" to 87.9%, with accordingly minor contributions from RO2 + HO2 and negligible contributions from RO₂ + RO₂ and RO₂ isomerization (Table S1†). However, the fates of individual RO₂ radicals depend

on the inherent RO_2 structure and may significantly deviate from the model result.

2.2. Online measurements

2.2.1. Gas phase. Toluene concentrations were determined using a single-photon ionization (SPI) time-of-flight mass spectrometer (TOFMS; compact time-of-flight II, mass resolution $m/\Delta m$ of 1000 at m/z 78, Firma Stefan Kaesdorf, Germany). For SPI, a Nd:YAG laser (Spitlight400, 20 Hz repetition rate, radiant power at 1064 nm of 6 W, and pulse duration of 7 ns; Innolas GmbH, Germany) was used as a light source for 1064 nm radiation, 38 which was frequency multiplied to 118 nm VUV photons by isotropic monopotassium phosphate crystals and a xenon cell at 5 mbar.39 The energy (10.49 eV) of 118 nm VUV photons slightly exceeds the ionization energy of most organic compounds including toluene, thus yielding predominantly molecular ions with low fragmentation. A 2-way-8-port switching valve (Vici Valco, Switzerland) was installed in order to conduct automated sampling before and after the PEAR, while the switching time was matched with the residence time of the PEAR. Furthermore, an isotope-labelled standard of D3toluene (m/z 95; toluene methyl-D3, 98% isotopic purity, Cambridge Isotope Laboratories Inc., MA, USA) was continuously added from a pressure vessel to the sampling line with a targeted concentration of approximately 500 ppb. The exact D3toluene concentration was determined by comparing the peak intensity at m/z 95 with a calibration standard of 1 ppm toluene in N2 (Linde AG, Germany). With D3-toluene, both wall losses of toluene and changes in laser performance were corrected.

The background concentrations of reactive gases O_3 , NO and NO_2 were analyzed using a trace-level chemiluminescence NO– NO_2 – NO_x analyzer (Thermo 42i-TL; Thermo Fisher Scientific Inc., USA) and a UV photometric ozone analyzer (Thermo 49i; Thermo Fisher Scientific Inc., USA).

2.2.2. Particle phase. The particle size distribution (14.6–685.4 nm) was monitored with a scanning mobility particle sizer (SMPS 3082; TSI GmbH, Germany) with a time resolution of 5 min. From the size-resolved particle number concentration, the total particle mass was obtained by integrating the size distribution over the particle size, assuming spherical particles with a density of 1.5 g cm $^{-3}$. Combined with the concentration of reacted toluene (Δ tol), the SMPS-derived SOA mass concentration was used to calculate the SOA yield ($Y_{\rm SOA}$) according to Odum *et al.* (1996):⁴⁰

$$Y_{SOA} = \frac{SOA}{\Delta tol} \tag{1}$$

The optical properties of OFR-derived tol-SOA were analyzed using a seven-wavelength aethalometer (AE33; Magee Scientific, USA). Wavelength pairs 370 and 520 nm were used to derive the Ångström Absorption Exponent (AAE),

$$AAE = -\frac{\ln\left(\frac{b_{\text{abs,370 nm}}}{b_{\text{abs,520 nm}}}\right)}{\ln\left(\frac{370 \text{ nm}}{520 \text{ nm}}\right)}$$
(2)

which describes the wavelength dependence of the absorption coefficient $b_{\rm abs}$. ⁴¹ This pair of wavelengths was used because of too low $b_{\rm abs}$ at longer wavelengths.

The chemical composition was analyzed with a soot-particle time-of-flight aerosol mass spectrometer (SP-AMS, Aerodyne Research Inc., USA), operated in tungsten vaporizer mode only because of the absence of soot particles in the experiments. The vaporizer temperature was set to 600 °C and volatiliz non-refractory particle components such as organic aerosols (OAs), nitrate, ammonium, sulfate and chloride. Intensities at m/z 44 were corrected for interfering carbon dioxide with a trace gas analyzer (GMP 343; Vaisala Oy, Finland). Elemental ratios O:C and H:C of OAs were determined by the "improved ambient method", 42 which allows the calculation of the average carbon oxidation state (OS_C) according to

$$OS_C \approx 2 \cdot O : C - H : C$$
 (3)

Further details of the SP-AMS operation can be found in Hartikainen *et al.* (2020).¹⁸

2.3. PM filter sampling

Samples of PM were collected after the PEAR on quartz fiber filters of 47 mm in diameter (Munktell, Filtrak; Sweden) through 6 mm stainless steel tubing at a flow rate of 20 L min. In order to reduce differences in filter loads, PM was collected for 3.5 h, 2 h and 0.5 h under "low", "medium" and "high" OFR conditions with total carbon filter loads from 20 to 90 μ g cm⁻². At the end of each chamber experiment, PM samples were collected for 1 h at a flow rate of 60 L min⁻¹ to obtain the same filter loading as for "low". All filter samples were immediately stored after collection at $-20~^{\circ}\text{C}$.

2.4. Offline PM analysis

Filter samples were analyzed by multi-wavelength thermal optical carbon analysis (MW-TOCA) hyphenated to SPI-TOFMS. The MW-TOCA consisted of a DRI Model 2001 A carbon analyzer, which was retrofitted with seven laser diodes of wavelengths (λ) in visible UV and near-infrared ranges, replacing the He–Ne laser for optical correction of charring.⁴³ This modification enables the measurement of laser transmittance over a spectral range and connects the optical properties of PM to their carbon content. Due to the low filter load and inherent absence of elemental carbon in tol-SOA, no correction for the shadow effects was applied⁴⁴ and ATN was related to the carbon content of tol-SOA to obtain the mass absorption efficiency MAE for each λ :

$$MAE_{\lambda} = \frac{[ATN]_{\lambda}}{C \cdot TC}.$$
 (4)

where TC denotes the total carbon load on the filter in mass per area and C refers to the correction factor for multiple scattering, which was set to 3.5.⁴⁵ To account for light scattering by the filter, the initial laser transmittances of the filter samples were related to the laser transmittance at the end of the carbon analysis. Because of the overall low absorptivity of the tol-SOA, the MAE $_{\lambda}$ could only be determined for λ of 405 and 450 nm.

The thermal refractiveness of the tol-SOA and its carbon content were determined according to the *Improve_A* temperature protocol, formally defining four fractions of organic carbon (OC1–OC4) and three fractions of elemental carbon (EC1–EC3).⁴⁶ In all samples of this study, the laser transmittance returned to its baseline value not before EC3, and thus no true EC was measured but only pyrolytic OC. Nevertheless, the fractionation of EC was kept in the discussion because it might reveal tol-SOA properties.

Evolved gas analysis was conducted at a modified quartz inlet of the TOCA, so an aliquot of the total flow was taken from the oven and analyzed by SPI-TOFMS⁴⁷ as described in Section 2.2.1. SPI mass spectra were acquired at 1 s time resolution and summed to one spectrum per carbon fraction. In principle, SPI with 118 nm photons may ionize all compounds with an ionization energy below 10.49 eV, mainly generating molecular ions. However, ion yields may vary between different compound classes up to one order of magnitude.⁴⁸

For tol-SOA analysis by high-resolution Orbitrap mass spectrometry, half of each PM filter sample was extracted by adding 5 mL of methanol (LC-MS grade) with gentle shaking for 60 min, and then filtered with a 0.2 µm pore size PTFE membrane (Sartorius, Goettingen, Germany) in a stainless-steel filter using a glass syringe. Mass spectrometric analysis was performed on a Velos Orbitrap (Thermo Scientific, Bremen, Germany) with electrospray ionization in both positive and negative ionization modes (ESI+/ESI-). ESI was operated with a direct-infusion ion source electrospray setup using a spray voltage of 4.2/3.4 kV and a syringe flow rate of 5 µL min. High-resolution data in the frequency domain were collected in the equivalent range of 50 to 1000 Da, resulting in a resolving power of >100 000 at m/z 400 and a mass accuracy below 3 ppm. The exported mass spectra were processed using in-house MATLAB (R2020; The Math-Works, USA) functions driven by the graphical user interface "CERES Processing". For elemental composition and sum formula assignments, the following restrictions were deployed in the range of 70 to 650 Da for the sum formula $C_cH_hO_o$ with 3 $\leq c \leq 35$, $3 \leq h \leq 60$, $o \leq 24$, and a maximum error of 3 ppm. Additional restrictions of 0.4-2.4 for H: C, 0-1.4 for O: C and 0-20 double bond equivalents (DBEs) were applied.

2.5. Statistical data analysis

Statistical testing was conducted with the Statistic Toolbox in MATLAB (R2020; The MathWorks, USA). Analysis of variance (ANVOA) was performed with Bonferroni correction in multiple testing at a significance level of 0.05 using the commands "anova1" and "multcompare". Principal component analysis was performed on the covariance matrix of L_1 -normalized SPI mass spectra using the command "pca". Correlations were either calculated as unadjusted correlation (or congruence, $r_{\rm c}$) for pattern comparison or as the Pearson correlation coefficient (r).

3. Results

3.1. Particle concentration, size distribution and SOA yields

The particle sizes generated under three different OFR conditions showed a shift toward larger particle size modes from

"low" to "high" (Fig. S1†). While "low" and "medium" generated particles with geometric mean diameters (GMD) from 32 to 38 nm and 35 to 42 nm, respectively, "high" OFR conditions increased the GMD to 60 to 63 nm. However, all particle sizes of OFR-generated tol-SOA were significantly smaller than those of chamber-generated tol-SOA, which peaked between 150 and 200 nm. In a mass-based size distribution, the particle size distributions of all OFR conditions interfere more with GMDs of 90 to 97 nm, 99 to 114 nm, and 120 to 126 nm, but still it is evident that the concentration of tol-SOA originates from larger particles generated in chamber experiments with GMDs from 180 to 128 nm (Fig. 1). Furthermore, with a Y_{SOA} of only 3.1 to 6.2%, OFR-aging generated significantly less SOA than chamber-aging with a Y_{SOA} of \sim 40%, which agrees well with the conclusion of Lambe et al. $(2015)^{49}$ on generally higher Y_{SOA} in chamber- than in PAM OFR-aging. YSOA typically increases exponentially with the aerosol concentration in chamber experiments;40 however, when comparing OFR-aging in "low" and "high", the more than one order of magnitude higher concentration of toluene and the resulting SOA caused only an increase in $Y_{\rm SOA}$ from (3.9 \pm 0.7) % to (5.6 \pm 0.7) %.

In SOA formation, organic vapors are oxidized in the gas phase and form low-volatile organic compounds (LVOCs), condensing on existing particles or forming new particles by nucleation. The lifetime of LVOCs is determined by their conversion into the particle phase (τ_{aer}) , further oxidation to products of higher volatility (τ_{OH}) and losses to the walls (τ_{wall}). 18 The vast majority of LVOCs condense on particles under all OFR conditions when using the particle size distribution after OFR processing (Table S2†). However, it is unknown how fast surface area particles are formed, and hence the results give an upper limit. When using only half of the particle number concentration measured after the PEAR, 10 to 19% of the LVOCs exit the OFR in the gas phase under "low" conditions (Table S3†). This tendency is in line with the observation of decreasing Y_{SOA} from "high" to "low", but it would require an even lower condensation sink than half of the measured particle size distribution to

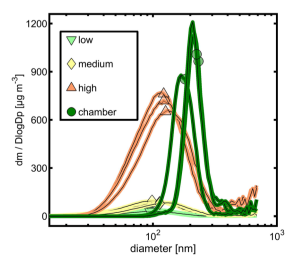


Fig. 1 Size distributions derived from the SMPS based on particle mass concentration assuming spherical particles with a density of $1.5~{\rm g~cm^{-3}}$. Symbols represent the geometric mean particle diameter.

explain quantitative differences in $Y_{\rm SOA}$. In the chamber, the same OH exposure is received by toluene, but with a longer residence time and lower ratio of surface to volume compared to the PEAR. A quantity to compare the fate of LVOCs in the OFR and chamber provides the ratio condensation sink concerning condensation on particles CS_{par} related to losses on the inner wall CS_{wall}. With 409 ± 50 , "high" OFR conditions are closer to 350 ± 65 for chamber-aging, while "low" and "medium" OFR conditions have a CS_{par}/CS_{par} of 30 ± 6 and 65 ± 14 , respectively. Hence, "high" conditions suppress LVOC wall losses on a relative scale, similar to the conditions in the chamber.

Lambe *et al.* (2015)⁴⁹ modeled the time required for a gas phase species to condense on pre-existing particles, and thus when seed particles were used, they provided an initial condensation sink in contrast to our study. With the assumption of a 10–100 μ m² cm⁻³ particle surface, 150 g mol⁻¹ average SOA molar mass and an accommodation coefficient of 1, a substantial fraction of oxidized vapors exit the reactor in the gas phase, this not increasing Y_{SOA} . In our study, the longer residence time of 5 h compared to the PEAR residence time of 70 s cancels out the effect from the 10 times higher OHR_{ext} (and 3 to 4 times higher particle surface area) under "high" OFR conditions compared to "cham" and provides an explanation for the significant differences in Y_{SOA} between the OFR and chamber.

3.2. Optical properties

In the absence of NO_x , toluene is less susceptible to the formation of light-absorbing organic carbon ("brown carbon", BrC) due to the decomposition of aromatic rings as chromophores and the absence of the formation of strongly absorbing nitroaromatics by addition of NO_2 .⁵⁰ The MAE_λ is a measure of light absorption related to the amount of particulate carbon. Because of small attenuation in laser transmittance after filter sample penetration, MAE_λ could be determined only for 405 and 450 nm in the MW-TOCA. The mean MAE_λ values for all OFR and chamber experiments appeared closely together, and the associated experimental and measurement uncertainties do not allow us to conclude an effect from the aging conditions (Fig. 2a).

A widely used approach to calculate the AAE is an exponential fit of the aerosol absorption coefficient νs . all seven wavelengths of the aethalometer, and hence the absolute AAE from other studies may differ compared to the AAE obtained from the wavelength pair 370 nm/532 nm. In contrast to the quantities discussed in Section 3.1., the AAE did not cluster according to OFR conditions, spanning the AAE range from 3 to 5.5. Among all experiments with the same targeted OH exposure, small differences in the exact OH exposure in the order of 5×10^9 cm³ s were found to be determining for the AAE, having a Pearson correlation coefficient of 0.91 (Fig. 2b). Therefore, the OFR conditions of this study do not seem to affect the BrC formed from toluene photooxidation.

3.3. Thermal refractiveness

3.3.1. Thermal-optical carbon analysis. Thermal-optical carbon analysis quantifies the amount of carbon in individual thermal fractions according to the *ImproveA* protocol, providing

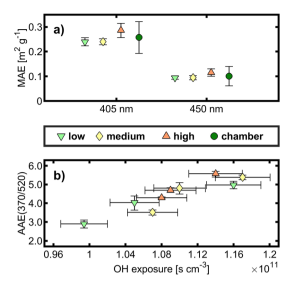


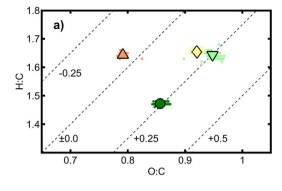
Fig. 2 (a) Mean Mass Absorption Efficiency (MAE) related to total carbon (TC) at 405 and 450 nm derived from MW-TOCA with standard deviation and (b) correlation of the Angström Absorption Exponent (AAE) between 370 and 520 nm, derived from an aethalometer, to OH exposure with a 95% confidence interval for the means of the individual experiments.

a measure of thermal refractiveness. Because of different filter sampling durations and the associated possible "blow-off" of semi-volatile materials, OC1 was excluded from the analysis. Generally, the distributions of the carbon fractions were similar, especially among "low" and "medium" OFR conditions and chamber photooxidation; however, "high" OFR conditions caused a significantly larger contribution of carbon to OC2 and EC1 but less to OC3 and EC2 (Fig. S2†) based on one-way ANOVA with Bonferroni correction. Apparent EC virtually accounted for the organic material because only tol-SOA was on the filter, forming a pyrolytic OC. Altogether, OFR conditions resemble the carbon volatility distribution of chamber-generated tol-SOA, but "high" reveals more charring and a shift towards carbon of higher volatility or less thermal refractiveness. Thus, "high" OFR conditions may be used to generate tol-SOA in sufficient amounts, but with reduced "blow-off" during long sampling, for the application of a volatility basis set.51

3.4. Chemical composition

3.4.1. Bulk composition

3.4.1.1. Aerosol mass spectrometry (AMS). The bulk elemental composition was determined by Aerosol Mass Spectrometry (AMS). The elemental ratios O:C and H:C (Fig. 3a) as well as the molar ratio of organic matter to organic carbon (OM:OC) were significantly different between all individual tol-SOA experiments (Table S4†). The largest O:C of 0.94 was obtained for "low" OFR conditions and the lowest for "high" (O:C of 0.8), with chamber-generated tol-SOA appearing in between (O:C of 0.85), which is in line with a previous study on *m*-xylene and α -pinene photooxidation attributing the higher O:C in the OFR to a possibly faster gas phase oxidation than nucleation. ⁵² OM:OC ranged from 2.4 ("low") to 2.2 ("high") also with



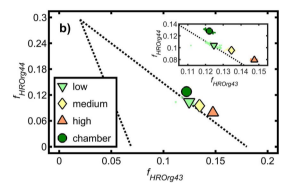


Fig. 3 (a) Van Krevelen diagram with the average carbon oxidation state indicated by dashed lines and (b) triangular space according to Ng et al. (2010)⁵³ with magnification of the narrow space of tol-SOA. Symbols without edges refer to individual data points, and symbols with black edges denote the mean values of one aging condition.

chamber-generated tol-SOA in between (OM:OC of 2.3). However, despite significance, the differences have low physical meaning because they originate from the high repeatability of the experiments and low uncertainties. In contrast to O: C, H: C only exhibited significant differences between all OFR- (H:C of 1.64 to 1.65) and chamber-generated tol-SOA (H:C of 1.47). Consequently, the average carbon oxidation state (OS_C) is affected, showing similarly high OS_C for "low" ($OS_C = 0.25$) and chamber-generated tol-SOA ($OS_C = 0.24$) but lower OS_C of 0.18 and -0.06 for "medium" and "high" OFR-generated tol-SOA.

In AMS data analysis, the relative intensities of organic ions at m/z 43 and m/z 44 have been used to delineate the space of ambient organic aerosols and classify them into hydrocarbonlike, biomass burning-line and oxidized organic aerosols (HOA, BBOA and OOA) in the triangular space. 53 Ions at m/z 43 may belong to C₃H₇⁺ or C₂H₃O⁺, associated with primary aerosol sources spanning the range of the x-axis. Upon atmospheric aging, oxygenated organic species such as carboxylic acids are formed, which increase the intensity of CO_2^+ at m/z 44, thus moving organic aerosols towards the vertex of the triangular space. As expected from previous experiments,34 the data points of tol-SOA were distributed along the hypotenuse of the triangle (Fig. 3b). However, despite lower O:C than "low" and "medium" OFR-generated tol-SOA, chamber-generated tol-SOA had the largest contribution of f_{44} and hence was classified as the most intensely aged organic aerosol. Furthermore, OFR-

generated tol-SOA lined up on the hypotenuse from "high" to "low" OFR conditions with increasing f_{44} , although all tol-SOA received the same OH exposure.

3.4.2. Molecular level composition

3.4.2.1. Single-photon ionization time-of-flight mass spectrometry. Mass spectrometry with soft ionization for evolved gas analysis in analytical pyrolysis provides insights into the molecular composition beyond bulk properties. During MW-TOCA, a small subsample of the evolving tol-SOA is analyzed by single-photon ionization time-of-flight mass spectrometry (SPI-TOFMS), thus giving molecular ions of thermally desorbed tol-SOA constituents or thermal fragments for individual fractions of OCs. Highly oxygenated organic aerosols already start to decompose at temperatures >80 °C in thermal desorption,54 so detecting compounds in evolved gas analysis likely underwent loss of H₂O, CO₂ or other small molecules in addition to larger thermal fragments detected by SPI-TOFMS. Therefore, the fractions OC2 and OC3 were combined to form OC23 for further analysis. OC1 and OC4 were discarded because of sampling artifacts and unspecific pyrolysis products, respectively.

The SPI mass spectra of OC23 revealed multiplets with dispersions of ± 2 u and a distance of 14 u, indicating homologous series with structural increments of -CH2-, -OCH3 or -OH (Fig. 4a-d). The mostly absence of odd m/z emphasizes the low yield of fragment ions, but the large yield of molecular ions in SPI. However, the mass spectra were very similar having an uncentered correlation coefficient larger than 0.935. Therefore, principal component analysis (PCA) was conducted on the centered SPI mass spectra after L1-normalization. Eigenvalue analysis by the broken stick method55 (Fig. S3†) supports the use of only two principal components (PCs), which account for 77% of the total variance. It is evident that the experimental intraclass to total variance was sufficiently low to separate all four types of tol-SOA with some interference between the 50% confidence ellipsoids of "low" and "medium" (Fig. 4e). Based on the PCA scores, "high" and chamber-generated tol-SOA were most different in chemical composition, while "low" and "medium" appeared in between. Nevertheless, with an explained total variance of 31%, the second PC still has a large contribution, so tol-SOA generated in "high" and in the chamber seems to contain mutual molecular features.

Overall, the first two PC loadings did not give a clear picture of the chemical composition, so most correlating m/z values were determined by extracting m/z within $\pm 18.2^{\circ}$ of the line between the origin and the center of the scores, equal to $r_c >$ 0.95. It becomes apparent that those highly correlated thermal fragments increase in mass from "low" to "high" with intensityweighted m/z means of 88 ("low"), 104 ("medium"), 142 ("high") and 88 ("cham") rounded to the nearest integer. Hence, tol-SOA from "low" and chamber-aging were likely more oxidized than tol-SOA from "medium" and "high" with more intense decomposition. However, larger m/z values mainly appear in the negative values of the first two PC loadings, so chambergenerated tol-SOA seems to cover the entire mass spectral range spanned by the three OFR conditions.

3.4.2.2. High-resolution mass spectrometry ESI(\pm). Chemical analysis of methanolic tol-SOA extracts by ESI(\pm)-Orbitrap mass

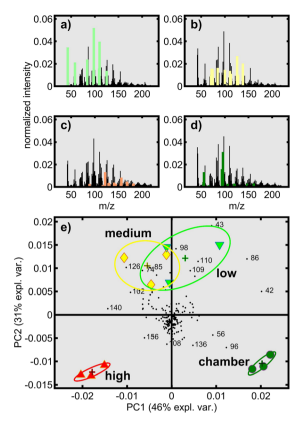


Fig. 4 Average and normalized OC23 SPI mass spectra of tol-SOA generated under (a) "low", (b) medium, and (c) "high" OFR conditions and in (d) the chamber. Colored stems in (a–d) indicate m/z highly correlated ($r_c > 0.95$, angle of ± 18.2 °) with the center ("+" in biplot) of the four tol-SOA classes in (e) principal component (PC) biplot space. For simplicity, only variables with the highest loadings in either PC1 and/or PC2 are labelled.

spectrometry revealed only minor differences in the Van Krevelen space of O:C $\nu s.$ H:C but noticeable differences in the intensity-weighted means of elemental ratios (Table 2). O:C, H:C and OS_C were decreasing from "low" to "high" for both ionization modes, while chamber-generated tol-SOA had elemental ratios and OS_C slightly below those of "high".

Interestingly, among all multiple comparisons only the pairs "low" and "medium" as well as "high" and "cham" gave an insignificant difference at the 5% significance level. Based on the sum formula, the modified aromaticity index (AI_{mod}) may be derived, providing classifications into saturated aliphatic (AI_{mod} = 0), unsaturated aliphatic (0 < AI_{mod} < 0.5), monoaromatic (0.5 < AI_{mod} < 0.67) or polyaromatic (AI_{mod} > 0.5) compounds for individual sum formulae.⁵⁶ Generally, the AI_{mod} values for all types of tol-SOA were low and between 0.04 and 0.11 for ESI+ and between 0.03 and 0.07 for ESI-. However, chambergenerated tol-SOA had significantly higher AImod compared to "low" in ESI+ and generally significantly higher AI_{mod} compared to all other tol-SOA in ESI-. The mean values of the relative number of sodium adducts [M + Na⁺] to total ions indicate that [M + Na⁺] are formed in the order low > medium > high > chamber but without significant differences among the tol-SOA types.

To explore further the differences in intensity-weighted mean elemental ratios and derived quantities, all twelve individual mass spectra were L₁-normalized, set to the same number of variables, i.e. m/z, and merged to one mean mass spectrum for each type of tol-SOA. In both ionization modes, the same main peaks could be observed in all tol-SOA; however, differences appeared in the mass spectrometric pattern (Fig. S4†). In ESI(+), three regions of interest (ROIs) may be defined between m/z 100 and 200 (A), (b) m/z 200 and 300 (B), and m/z 300 and 400 (C). ROIs A and C are present in all tol-SOA, but there is a decrease in the ratio A/C from "low" over "medium" and "high" to "cham". Simultaneously, peaks in ROI B start to vanish in the same order, but with a single distinct peak at m/z 213.038 with the sum formula $C_7H_{10}O_6$, detected as [M + Na]⁺. With lower intensities, the same ion was detected in all OFR-generated tol-SOA, and hence an impurity or artifact of the chamber can be ruled out. Elemental analysis of the main peaks of all ROIs revealed that ROIs A and C contained monomers and dimers of toluene with 7 and 14 carbon atoms and O: C between 0.55 and 0.75. In contrast, peaks in ROI B belong to sum formulae with carbon numbers slightly larger than seven, but distinctly lower than 14, such as $[C_8H_{14}O_7+Na]^+$ (m/z)245.064) or $[C_9H_{16}O_8+Na]^+$ (m/z 275.074). Hence, ROIs A and C

Table 2 Elemental composition (O:C, H:C, and OS_C), modified aromaticity index (AI_{mod}) and rel. intensity of sodium adducts of tol-SOA analysis ESI(+)- and ESI(-)-Orbitrap mass spectrometry

Experiment	(+)O : C	(+)H : C	(+)OS _c	$(+)AI_{mod}$	(+)rel Na	(-)O:C	(-)H:C	$(-)OS_c$	$(-)AI_{mod}$
low1	0.75	1.43	0.06	0.06	0.63	0.86	1.13	0.59	0.04
low2	0.76	1.43	0.00	0.04	0.71	0.85	1.14	0.56	0.04
low3	0.78	1.44	0.11	0.04	0.76	0.88	1.16	0.60	0.03
med1	0.71	1.37	0.05	0.07	0.61	0.79	1.13	0.46	0.05
med2	0.74	1.41	0.07	0.05	0.72	0.81	1.14	0.49	0.04
med3	0.73	1.39	0.07	0.05	0.69	0.83	1.14	0.51	0.04
high1	0.60	1.27	-0.06	0.11	0.51	0.72	1.14	0.31	0.04
high2	0.64	1.31	-0.03	0.07	0.69	0.72	1.13	0.30	0.05
high3	0.63	1.30	-0.05	0.09	0.63	0.73	1.14	0.31	0.04
cham1	0.57	1.28	-0.14	0.11	0.47	0.71	1.09	0.33	0.07
cham2	0.62	1.26	-0.02	0.10	0.55	0.70	1.09	0.31	0.07
cham3	0.65	1.28	0.01	0.07	0.69	0.70	1.09	0.32	0.07

contain monomers and dimers of toluene-SOA, whereas ROI B includes products from the reaction of C₇ units with units of a lower carbon number or possibly further processing of dimers by fragmentation. A more detailed view of dimer formation is provided by the analysis of the carbon number (#C) vs. the average carbon oxidation state (OS_C) together with the relative peak intensities for each #C (Fig. 5). Among the tol-SOA types, "low" contains the largest proportion of species with a #C of 8 and 9, while in all other tol-SOA a local minimum occurs in the #C vs. peak intensity space. Furthermore, two local maxima appear at a #C of 12 and 14, which can be assigned to dimers. However, the ratio of C₁₂ to C₁₄ species decreases in the order chamber > low > medium > high, thus having the highest similarity between "cham" and "low". Nevertheless, the strongest overall correlation between the #C vs. peak intensity patterns of the three OFR-generated SOAs to "cham" was obtained for "medium" with 0.97 (0.95 for "low" and 0.89 for "high").

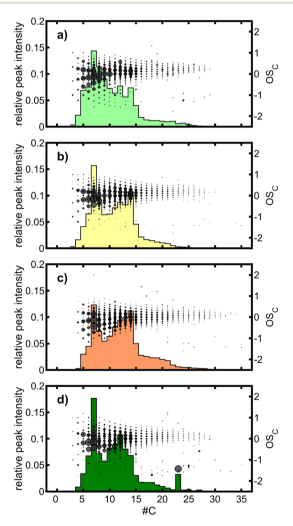


Fig. 5 #C vs. OS_C with summed relative intensities for each #C for ESI(+) of (a) "low", (b) "medium", (c) "high" and (d) "cham". The contribution to #C = 23 in "cham" mainly belongs to $[C_{23}H_{40}O_4 + H]^+$, which could be a common plasticizer from the family of cyclohexanedicarboxylic acid esters.

Due to its high abundance in the atmosphere, several secondary products of toluene from atmospheric aging were identified and partly used as markers, such as 2,3-dihydroxy-4oxopentanoic acid (DHOPA) with the sum formula C5H8O5.57 Such a compound was detected as a sodium adduct [C₅H₈O₅ + Na] in all tol-SOA with relative intensities in the order: "low" $(0.61\%) > \text{"medium"} (0.58\%) > \text{"cham"} (0.11\%) \approx \text{"high"}$ (0.10%). Hence, the yield of DHOPA was the largest under "low" OFR conditions, but those from "high" were most comparable to the chamber-generated tol-SOA. However, the peak of $[C_5H_8O_5 + Na]^+$ may also belong to isomers of DHOPA and does not provide an appropriate marker isolation from the complex mixture of toluene oxidation products.

In ESI(-) mass spectra, larger m/z values increase in relative intensity from "low" to "high" conditions and show qualitatively the same main peaks as in ESI(+). Some of the most intense peaks detected in all tol-SOA appear at m/z 191.0198 ([C₆H₈O₇-H]⁻), m/z 173.0092 ([C₆H₆O₆-H]⁻) and m/z 161.009 ([C₅H₆O₆-H]⁻), indicating highly oxidized or sugar-like components with $O:C \ge 1$. ROIs derived from the pattern in ESI- mass spectra were not as apparent as in ESI+, but matched the possible structures of monomers and dimers derived from elemental analysis. From "low" to "high" OFR conditions, the intensities of m/z < 250 decreased while m/z > 250 increased. Moreover, most intense peaks of m/z > 250 in "high" shifted to larger m/z, agreeing with a shift of the #C vs. peak intensity pattern of "high" to larger #C. Generally, monomers and dimers appeared at #C of 6 and 13 in OFR-generated tol-SOA with favoring monomers in "low", dimers in "high" and none in "medium" OFR experiments (Fig. S5†). In contrast, chamber-generated tol-SOA exhibits a maximum in the #C vs. peak intensity pattern at #C = 11 and a local minimum at #C = 7, which equals the carbon number of toluene. Therefore, acidic tol-SOA components of monomers and dimers underwent fragmentation involving carbon loss. The #C vs. OS_C and peak intensity derived from ESI(-) mass spectra were less similar between chamberand OFR-generated tol-SOA than derived from ESI(+). The largest similarity to "cham" was found for "medium" ($r_c = 0.94$), the lowest for "low" ($r_c = 0.87$) and "high" in between ($r_c = 0.90$).

Peak intensities in direct-infusion ESI mass spectra may suffer from matrix effects, so an alternative approach for data analysis is based on binary treatment of m/z being present or not present in samples, regardless of their intensity. In a Venn diagram, peaks are grouped according to their occurrence in the different types of tol-SOA. To limit the influence of different concentrations associated with a larger number of peaks above a signal-to-noise of six, particularly, in the higher m/z range, the input data for the Venn diagram were restricted to a maximum #C of 15. Of the 798 detected peaks in ESI+, 356 appeared in all tol-SOA, accounting for 57%, 61%, 62%, and 58% of the peaks detected in "low", "medium", "high" and "cham", respectively. Moreover, 75%, 83%, and 77% of the peaks in "low", "medium", and "high" also appeared in "cham". In ESI-, 265 out of 479 peaks were detected in all tol-SOA, accounting for 78%, 76%, 67%, and 70% of the peaks detected in "low", "medium", "riskier" and "cham", respectively. Moreover, 79%, 82%, and 80% of the peaks in "low", "medium", and "high" also appeared

in "cham". Overall, the alternative binary peak analysis demonstrates fairly low differences among the tol-SOA types and the high similarity of OFR-generated to chamber-generated tol-SOA.

4. Discussion

OFRs are widely used to study the fate and impact of primary aerosols released into the atmosphere. Their main advantage lies in the potential coverage of the entire range of atmospheric particle lifetime, but at the cost of different mechanisms of secondary aerosol formation and subsequent multiphase chemistry. Recent work has modelled the range of OFR conditions for the OFR "potential aerosol mass" (PAM), triggering similar gas phase chemistry to that in the troposphere. 15 However, the effect of OFR conditions, regarded as atmospherically relevant or irrelevant, on the physical and chemical properties of SOAs has not been systematically investigated yet, which motivated our case study on toluene photooxidation products. OFR experiments with the PEAR were conducted with settings of OH reactivity and photon fluxes ("low", "medium", and "high"), appearing in the range of "safer", "transition" and "riskier" OFR conditions according to the PAM model. Chamber experiments were conducted as a laboratory reference for atmospherically relevant tol-SOA.

Our results reveal that "low" conditions, representing "safer" OFR conditions, do not stringently prove a higher similarity to atmospherically relevant SOAs when compared to complementary environmental chamber experiments conducted at comparable concentrations to the literature. For the aerosol properties O:C from AMS, elemental quantities and the molecular composition derived from ESI±, as well as the SOA yield, "low" even showed the least comparability to "cham". Moreover, insignificant and fairly small differences were found between both OFR- (all conditions) and chamber-derived tol-SOA for the optical properties AAE and MAE $_{\lambda}$ as well as the distribution of carbonaceous species among the thermal fractions of carbon.

On a quantitative scale, tol-SOA from OFRs was generally produced in lower yields than from chamber aging in agreement with Lambe $et\ al.\ (2015).^{49}$ Here, the approach of constant OH exposure with high OH concentrations and short time shows its disadvantage as oxidized vapor may require more time to condense and form SOA than the residence time typically permitted in most OFRs, so the majority of low-volatile compounds exit the OFR in the gas phase. This may partially be counteracted by using a seed aerosol, which provides an additional condensation sink and consequently enhances $Y_{\rm SOA}$. In the same manner, the shift of the particle size distribution for "low" over "medium" to "high" would be avoided and could appear closer to the chamber-generated tol-SOA.

In order to cover the complex composition of SOAs, bulk properties and parametrizations are used. OFR conditions did not show any effect on H:C, but led to decreasing O:C from "low" to "high" OFR conditions, so OS_C decreased as well. Although "low" and "cham" agree well, similar OS_C originates from different H:C and O:C, pointing towards a different

chemical composition. This is further supported by the parametrization in the triangular space f_{43} vs. f_{44} , ⁵³ where chamber-generated tol-SOA revealed the largest contribution of f_{44} , thus the most advanced aging.

The most substantial differences between "low" and "cham" were observed in the more detailed chemical composition, particularly in ESI high-resolution Orbitrap mass spectrometry. Surprisingly, in pattern analysis "high" and in peak number-based comparison "medium" showed the largest correlation with "cham", which could be attributed to differences in dimer formation. However, thermodesorption SPI mass spectra suggested that chamber-generated tol-SOA contains molecular features of all three OFR conditions.

Overall, our study on tol-SOA challenges the concept of "safer", *i.e.*, atmospherically relevant OFR conditions. However, due to the wide space of parameters, an experimental approach is required to elucidate the effect of OFR conditions on SOA composition and gas phase species, which have not been addressed in this study. This would include aging under high-NO_x conditions,²⁵ SOA precursors with a stronger tendency towards oligomer formation,^{31,61} the application of seed aerosols, different initial O₃ concentrations in OFR254 mode, and the comparison to the OFR185 mode with *in situ* ozone generation. Furthermore, recent process-level modelling enabled the connection between environmental chamber and OFR results,⁶² which might be incorporated in OFR condition assessment.

As an extension of physical and chemical aerosol properties, the toxicity of aged aerosols has been studied using an OFR. Fine particulate matter, including specifically SOAs, is harmful to human health, 6,63 so OFRs enable toxicological studies with in vitro exposure to investigate specific biological endpoints and SOAs from individual precursors. Cytotoxicity and inflammatory responses start with oxidative stress, which has been linked to the concentrations of airborne reactive oxygen species (ROS).8,30,64 In the OFR, high levels of ROS may be substantially formed over the full length of the reactor. Although the majority of ROS is short-lived,65 a significant fraction of the ROS is leaving the OFR because of its short residence time. In environmental chambers, the residence time is orders of magnitude higher, so those short-living ROS would already react during the longer residence in environmental chambers and lower concentrations of short-living ROS are present at the end of the aging experiment. Ozone as part of the ROS class is removed using denuders,8 which likely reduce other ROS as well, but systematic research on different ROS would be beneficial to rule out if cell cultures exposed in the air-liquid interface possibly receive a significantly higher dose of ROS from OFRs than chamber-generated to aged aerosols.

Besides ROS, polycyclic aromatic hydrocarbons (PAHs) are health-relevant ambient aerosol constituents, which become potent after metabolization in the human body. Recently, Hrdina *et al.* (2022)⁹ pointed out the similarity of PAH metabolism in the human body and (short) atmospheric aging, leading both to oxygenated PAHs. Here, a deeper analysis and understanding of OFR chemistry products is required since the detailed chemical structure of aerosol constituents, *i.e.*, the

position of a functional group in the carbon backbone, may be crucial to their toxicity.

The most carcinogenic PAHs, including benzo(a)pyrene, are predominantly located in the particle phase, so direct oxidation by OH is slower compared to homogeneous gas phase oxidation of more volatile PAHs like naphthalene. Additionally, PAHs may be shielded from atmospheric oxidants by condensing SOAs as a coating, further decelerating their degradation. Hence, the underestimation of heterogeneous oxidation would be another task for process-level modelling in order to harmonize PAH degradation in OFRs and environmental chambers and determine the highest carcinogenic/mutagenic potential of PAHs during their atmospheric lifetime, which would be beneficial for OFR studies on aged combustion aerosols.

Conclusion

Paper

OFR-aging conditions by means of different ratios of atmospheric oxidants and photon fluxes but with a similar level of OH exposure affect the physical-chemical properties of tol-SOA. While the relative effect towards less atmospherically relevant conditions was substantial for Y_{SOA} and particle size distribution, only small differences were observed for chemical bulk properties, such as elemental ratios, but there were larger discrepancies in the more detailed chemical composition. Most surprisingly, for many physical-chemical properties of OFRgenerated tol-SOA less atmospherically relevant conditions exhibit the largest agreement with chamber-generated tol-SOA, which resembles photochemical aging more closely than an OFR at the same OH exposure. OFRs may bridge the gap between the upper limit of the equivalent photochemical aerosol age achievable in chambers and the maximum atmospheric particle residence time, but the value from their application depends on the research objective and considered aerosol properties.

Author contributions

HC: conceptualization, methodology, validation, formal analysis, investigation, data curation, writing – original draft, visualization, supervision, and project administration; PYP: investigation and data curation; AH: validation, formal analysis, and writing – review & editing; MI: investigation and data curation; PT: formal analysis and investigation; PM: investigation, formal analysis and writing – review & editing; ES: investigation, formal analysis, and writing – review & editing; AP: formal analysis; TH: writing – review & editing and supervision; JJ: resources and funding acquisition; CR: methodology, software, writing – review & editing, and supervision; RZ: resources and funding acquisition; OS: resources, writing – review & editing, supervision, and funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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