Quantification of environmentally persistent free radicals and reactive oxygen species in atmospheric aerosol particles

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Abstract
Fine particulate matter plays a central role in adverse health effects of air pollution. Inhalation and deposition of aerosol particles in the respiratory tract can lead to the release of reactive oxygen species (ROS), which may cause oxidative stress. In this study, we have detected and quantified a wide range of particle-associated radicals using electron paramagnetic resonance (EPR) spectroscopy. Ambient particle samples were collected using a cascade impactor at a semi-urban site in central Europe, Mainz, Germany in May – June 2015. Concentrations of environmentally persistent free radicals (EPFR), most likely semiquinone radicals, were found to be in the range of \((1 - 7) \times 10^{11}\) spins \(\mu\text{g}^{-1}\) for particles in the accumulation mode, whereas coarse particles with a diameter larger than 1 \(\mu\text{m}\) did not contain substantial amounts of EPFR. Using a spin trapping technique followed by deconvolution of EPR spectra, we have also characterized and quantified ROS including OH, superoxide (\(O_2^−\)) and carbon- and oxygen-centred organic radicals, which were formed upon extraction of the particle samples in water. Total ROS amounts of \((0.1 - 3) \times 10^{11}\) spins \(\mu\text{g}^{-1}\) were released by submicron particle samples and the relative contributions of OH, \(O_2^−\), C-centred and O-centred organic radicals were \(\sim 11 - 31\%\), \(\sim 2 - 8\%\), \(\sim 41 - 72\%\) and \(\sim 0 - 25\%\), respectively, depending on particle sizes. OH was the dominant species for coarse particles. Based on comparisons of the EPR spectra of ambient particulate matter with those of mixtures of organic hydroperoxides, quinones and iron ions followed by chemical analysis using liquid chromatography mass spectrometry (LC-MS), we suggest that the particle-associated ROS were formed by decomposition of organic hydroperoxides interacting with transition metal ions and quinones contained in atmospheric humic-like substances (HULIS).
1. Introduction

Epidemiological studies have clearly shown positive correlations between respiratory diseases and ambient fine particulate matter (Pope and Dockery, 2006; Strak et al., 2012; West et al., 2016). A recent study has estimated that outdoor air pollution leads to 3.3 million premature deaths per year worldwide, which is mostly due to particular matter with a particle diameter less than 2.5 µm (PM$_{2.5}$) (Lelieveld et al., 2015). Plausible reasons include the cytotoxicity of ambient PM$_{2.5}$ and its ability to induce inflammatory responses by oxidative stress causing functional alterations of pulmonary epithelial cells (Nel, 2005; Gualtieri et al., 2009). Oxidative stress is mediated by reactive oxygen species (ROS) including OH, H$_2$O$_2$, superoxide (O$_2^-$), as well as organic radicals (Pryor et al., 1995; Winterbourn, 2008; Birben et al., 2012; Pöschl and Shiraiwa, 2015). Upon PM deposition into the respiratory tract and interactions with lung antioxidants, H$_2$O$_2$ can be generated by redox-active components contained in PM$_{2.5}$ such as transition metals (Charrier et al., 2014; Fang et al., 2015), semiquinones (Kumagai et al., 1997; Cho et al., 2005; Khachatryan et al., 2011; McWhinney et al., 2013), and humic-like substances (Kumagai et al., 1997; Cho et al., 2005; Lin and Yu, 2011; Charrier et al., 2014; Dou et al., 2015; Fang et al., 2015; Verma et al., 2015a). H$_2$O$_2$ can be converted into highly-reactive OH radicals via Fenton-like reactions with iron and copper ions (Charrier et al., 2014; Enami et al., 2014).

Ambient particles have been found to contain large amounts of ROS (mostly H$_2$O$_2$) in the particle phase (Hung and Wang, 2001; Venkatachari et al., 2005; Venkatachari et al., 2007; Fuller et al., 2014). Substantial amounts of particle-bound ROS are found on biogenic secondary organic aerosols (SOA) produced from the oxidation of α-pinene, linalool, and limonene (Chen and Hopke, 2010; Chen et al., 2011; Pavlovic and Hopke, 2011; Wang et al., 2011; Wang et al., 2012). Recently, Tong et al. (2016) have shown that terpene and isoprene SOA can form OH radicals upon interactions with liquid water and iron ions under dark conditions. This can be explained by the decomposition of organic hydroperoxides, which account for the predominant fraction of SOA mass and are generated via multigenerational oxidation and autoxidation (Docherty et al., 2005; Ziemann and Atkinson, 2012; Crounse et al., 2013; Ehn et al., 2014; Epstein et al., 2014; Badali et al., 2015).

In addition, PM$_{2.5}$ contain environmentally persistent free radicals (EPFR) that can be detected directly by electron paramagnetic resonance (EPR) spectroscopy (Dellinger et al., 2001; Khachatryan et al., 2011; Gehling and Dellinger, 2013). EPFR are stable radicals with an e-folding lifetime exceeding one day (Gehling and Dellinger, 2013; Jia et al., 2016). The chemical nature of EPFR is remarkably similar to semiquinone radicals, which can be
stabilized via electron transfer with transition metals in the particle phase (Truong et al., 2010; Vejerano et al., 2011; Gehling and Dellingier, 2013). EPFR are formed upon combustion and pyrolysis of organic matter (Dellingier et al., 2001; Dellingier et al., 2007). The formation of stable radicals can also be induced by heterogeneous and multiphase chemistry of organic aerosols. Heterogeneous ozonolysis of aerosol particles such as polycyclic aromatic hydrocarbons (PAH) and pollen proteins can lead to the formation of long-lived reactive oxygen intermediates (ROI) (Shiraiwa et al., 2011; Shiraiwa et al., 2012; Reinmuth-Selzle et al., 2014; Borrowman et al., 2015; Kampf et al., 2015; Berkemeier et al., 2016).

In this work, ambient particles with a diameter in the range of 56 nm to 3.2 µm were collected using a cascade impactor during May – July 2015 in Mainz, Germany. Size dependences of EPFR concentrations contained in ambient particles have been measured using an EPR spectrometer. Particles were also extracted in water containing a spin-trapping agent followed by EPR analysis to quantify the formation of various radical forms of ROS including OH, superoxide (O$_2^-$) and carbon- and oxygen-centred organic radicals.

2. Methods

Ambient particles were collected using a micro-orifice uniform deposition impactor (MOUDI, 110-R mode, MSP Corporation) on the roof of the Max Planck Institute for Chemistry, Mainz, Germany (49.99 N, 8.23 E). The sampling was conducted every 24 h starting at 5 PM during 28 May - 9 June 2015. Particles were collected with a sampling time of 48 h during 26-27 June and 18-19 July 2015 in order to collect sufficiently high mass loadings for all stages of different particle size ranges. The sampling was conducted with a flow rate of 30 L min$^{-1}$ with the following nominal lower cut-off particle diameters: 56, 100, 180, 320, 560 nm, 1 µm, and 1.8 µm. Note that transmission and bouncing effects might have caused mixing of particles exhibiting relatively different sizes on one stage, particularly for coarse particles (Gomes et al., 1990; Bateman et al., 2014). Particles were collected on 47 mm diameter Teflon filters (100 nm pore size, Merck Chemicals GmbH). Before sampling, each filter was cleaned and sonicated for 10 min with pure ethanol and ultra-pure water and dried with nitrogen gas before weighing. Teflon filters were weighed four times using a balance (Mettler Toledo XSE105DU) and mounted in the MOUDI. After sampling, each filter has been conditioned for at least one hour (22-23°C and 40-50 RH) and weighted four times before being folded and inserted in a 4 mm EPR tube. Particles were extracted by immersing the filter into a solution containing 350 µL of 20 mM 5-tert-Butoxycarbonyl-5-methyl-1-pyrroline-N-oxide (BMPO, high purity, Enzo Life Sciences GmbH) and stirred with
a vortex shaker (Heidolph Reax 1) for 7-9 min. BMPO is an efficient spin-trapping agent for OH, O$_2^-$ and organic radicals (Zhao et al., 2001; Tong et al., 2016). Note that the trapping efficiency of O$_2^-$ and organic radicals might be lower compared to OH radicals, as the recent study has reported that nitrone-based spin traps have the highest reactivity towards OH and somewhat lower reactivity towards organic radicals and superoxide (Sueishi et al., 2015). Extracts were dried for approximately 14-17 min under 1-3 bar flow to reduce the volume of the solution to 50 µL and then 20 µL were used for EPR measurements.

A continuous-wave electron paramagnetic resonance (CW-EPR) X-band spectrometer (EMXplus-10/12, Bruker, Germany) was used for detection and quantification of stable radicals and ROS. Filters containing particles were folded and introduced into a 4 mm i.d. quartz tube and inserted directly into a high sensitivity cavity. EPR spectra were recorded at a room temperature of 23 °C by setting the following operating parameters: a modulation frequency of 100 kHz; a microwave frequency of 9.84 GHz; a microwave power of 2.149 mW (20 db); a modulation amplitude of 1.0 G; a sweep width of 110.0 G; a sweep time of 175 s; a receiver gain of 40 db; a time constant of 40.96 ms; a conversion time of 160 ms; and a scan number of 6. Paramagnetic species are characterized based on their g-factor values. Free electrons have a g-factor value of 2.0023 and organic radicals have higher g-factor values (2.0030 – 2.0060), depending on the number of oxygen atom in the molecule (Dellinger et al., 2007).

The spin-counting method embedded in the Bruker software Xenon was used to quantify detected radicals. The spin-counting method was calibrated using a standard compound 4-hydroxy-2,2,6,6-tetramethylpiperidin-1-oxyl (TEMPOL). The detection limit of EPR was $\sim$1×10$^{10}$ spins µg$^{-1}$. Concentrations of EPFR and ROS are reported in the unit of spins µg$^{-1}$, which indicates the number of spins (or radicals) per µg of particle mass. For better quantification and determination of the relative contributions of OH, O$_2^-$, carbon-centred and oxygen-centred organic radicals, EPR spectra were fitted and simulated using Xenon and the Matlab-based computational package Easyspin (Stoll and Schweiger, 2006).

3. Results and discussion
3.1. Environmentally persistent free radicals

Figure 1 shows EPR spectra of ambient particles in the lower cut-off diameter range of 56 nm – 1.8 µm. Fine particles, with lower cut-off diameters of 56 - 320 nm, show a single and unstructured peak with a g-factor of $\sim$2.003 and with a peak to peak distance ($\Delta H_{pp}$) ranging from 3 to 8 G. Such spectra are characteristic for EPFR, which have been attributed
to semiquinone radicals (Dellinger et al., 2001; Dellinger et al., 2007; Vejerano et al., 2011; Bahrle et al., 2015). Particles with a diameter smaller than 56 nm and larger than 560 nm did not show significant signals, indicating the reduced amount of EPFR in these size ranges. EPR spectra for particles with the lower cut-off diameters of 56 – 320 nm for each sampling day are presented in Fig. A1.

The black line in Fig. 2 shows the size distribution of EPFR concentrations. Particles with different sizes had different radical contents and particles with the lower cut-off diameter of 100 nm contained the highest EPFR concentrations of $7.0(\pm0.7) \times 10^{11}$ spins $\mu$g$^{-1}$. High abundances of EPFR in particles in the accumulation mode is consistent with mass size distributions of combustion-generated particles such as soot or black carbon, which typically have peak concentrations around 100 – 200 nm (Bond et al., 2013). This observation is in line with that EPFR may be often associated with soot particles (Dellinger et al., 2007).

Figure 3 shows the temporal evolution of EPFR concentrations contained in particles with lower cut-off diameters of 100 and 180 nm. During the sampling period of two weeks, there were two rain events (on 30 May 2015 and 1 June 2015) and three sunny days (4-6 June 2015), and the other days were cloudy. The mass concentrations of particles within the diameters of 56 - 560 nm were in the range of 3.9 – 12.8 $\mu$g m$^{-3}$. Maximum values of $\sim 7 \times 10^{11}$ spins $\mu$g$^{-1}$ were reached during sunny days, indicating that photochemistry may be related to EPFR production. For example, heterogeneous reactions of photo-oxidants including O$_3$ and OH with soot or PAH may contribute to the formation of long-lived radicals (Shiraiwa et al., 2011; Borrowman et al., 2015). Radical concentrations were as low as $6.3 \times 10^{10}$ spins $\mu$g$^{-1}$ during rain events, most likely due to low production of EPFR and scavenging by precipitation.

EPFR concentrations contained in particles within the diameter of 50 nm – 3.2 $\mu$m collected for 48 h during 26-27 June 2015 was $\sim 2.2 \times 10^{11}$ spins $\mu$g$^{-1}$. EPFR concentrations contained in particles within the diameter of 56 – 560 nm averaged over the entire measurement period was $2.0(\pm1.3) \times 10^{11}$ spins $\mu$g$^{-1}$. Squadrito et al. (2001) determined the EPFR concentrations to be in the range of $(1-10) \times 10^{11}$ spins $\mu$g$^{-1}$ in PM$_{2.5}$ sampled for 24 h in five different urban sites in the United States. Gehling et al. (2014) reported that the EPFR concentration was in the range of $(7-55) \times 10^{10}$ spins $\mu$g$^{-1}$ at a site in Louisiana near heavy interstate traffic along a major industrial corridor of the Mississippi River. Shaltout et al. (2015) measured radical concentrations in the range of $(2-6) \times 10^{10}$ spins $\mu$g$^{-1}$ in PM$_{2.5}$ collected in industrial-, residential- and traffic-dominated sites in Taif city, Saudi Arabia. The
EPFR concentrations measured in this work are comparable with those previous measurements.

### 3.2. Reactive oxygen species

Figure 4a shows EPR spectra of ambient particles with lower cut-off diameters of 56 nm - 1.8 μm extracted in water with the spin-trapping agent BMPO. Each EPR spectrum is composed of several overlapped lines, originating from different radical forms of ROS. Dashed lines indicate the positions of each peak for each type of trapped ROS including OH (green), superoxide (red), carbon-centred (orange) and oxygen-centred organic radicals (blue). The relative abundance of these radicals was different for each size range, causing the EPR spectral features to be highly variable. For example, spectra from particles larger than 1.0 μm consist mainly of four peaks that are typical for OH radicals, whereas those for smaller particles contain more peaks indicating the presence of multiple radicals.

To estimate the relative amount of each type of ROS, the observed EPR spectra were fitted and simulated using the software Easyspin 5.0 and Xenon. Four types of radicals have been used to fit the spectra: BMPO-OH (hyperfine coupling constants of $a^N = 14.3$ G, $a^H = 12.7$ G, $a^γ = 0.61$ G), BMPO-OOH ($a^N = 14.3$ G, $a^H = 8.1$ G), BMPO-R ($a^N = 15.2$ G, $a^H = 21.6$ G) and BMPO-OR ($a^N = 14.5$ G, $a^H = 16.6$ G). As shown in Fig. 4b, the simulated EPR spectrum reproduced the observed spectrum very well with a small residual. The deconvolution of spectra allowed us to estimate the relative contribution of four types of ROS within each particle size range.

Figure 5 shows the relative contributions of OH (green), superoxide (red), carbon-centred (orange) and oxygen-centred (blue) organic radicals to the total radicals trapped by BMPO in water extracts of particles collected for 48 h during 26-27 June 2015. Carbon-centred radicals are the most abundant type of radicals, contributing ~50 - 72% of total ROS for PM1. It decreases to 41% and 9% for particles with lower cut-off diameters of 1 μm and 1.8 μm, respectively. The OH radical accounts for ~11 – 31% of total trapped radicals for PM1, whereas OH was the dominant species for coarse particles with diameters of 1.8 – 3.2 μm. The least abundant radical for all size ranges was $O_2^-$, with contributions of ~2 - 8% and without any clear size dependence. The amount of oxygen-centred organic radicals ranges between 12% and 25% in particles with a diameter below 1 μm and its contribution was negligible for coarse particles. Note that the contribution of oxygen-centred organic radicals for particles with a diameter of 1 – 1.8 μm might be attributed to the OH radical: the hyperfine coupling constants for BMPO-OR for better fitting the spectrum for this size range
needed to be changed slightly ($a^N = 13.5$ G, $a^H_p = 15.3$ G, $a^H_y = 0.6$ G). These values are similar to constants of a second conformer of BMPO-OH.

The red line in Fig. 2 shows the size-dependent concentrations of radical forms of ROS (e.g., sum of OH, $O_2^-$, C- and O- centred organic radicals). Particles with the lower cut-off diameter of 100 nm have the highest ROS concentrations of $2.7(\pm 0.2) \times 10^{11}$ spins $\mu g^{-1}$. Concentrations are smaller for particles in the coarse mode with a diameter larger than 1 $\mu m$. This is consistent with previous studies, suggesting that particles in the accumulation mode are the most active in ROS generation (Hung and Wang, 2001; Venkatachari et al., 2007; Saffari et al., 2013; Wang et al., 2013; Saffari et al., 2014). The total concentration of radical forms of ROS was measured to be $1.2 \times 10^{11}$ spins $\mu g^{-1}$. Note that $O_2^-$ concentrations might be underestimated as the lifetime of the BMPO-OOH adduct is relatively short (~20 min) (Ouari et al., 2011; Abbas et al., 2014).

Previous studies have measured redox activity and oxidative potential of PM by the dichlorofluorescein (DCFH) and dithiothreitol (DTT) assays. The DCFH assay is mostly sensitive to $H_2O_2$ and other peroxides. For example, Hung and Wang (2001) reported ROS concentrations as $1 \times 10^{13}$ $\mu g^{-1}$ in Taipei, Taiwan. This value is very similar to $H_2O_2$ concentrations contained in ambient PM$_{2.5}$, which has been quantified to be up to $1 \times 10^{13}$ $\mu g^{-1}$ in an urban environment in southern California using HPLC fluorescence (Wang et al., 2012). The DTT assay is based on the decay of DTT due to redox reactions with PM components, reporting the oxidative potential of PM in moles of DTT consumed per unit of time and mass of PM. Verma et al. (2015b) and Fang et al. (2015) reported that PM$_{2.5}$ sampled in an urban environment in Atlanta, Georgia, USA has a DTT activity in the range of 10-70 pmol min$^{-1}$ $\mu g^{-1}$. Assuming an integration time of 20 min needed for the extraction of PM in this work, this value corresponds to $(1-8) \times 10^{14}$ $\mu g^{-1}$ of DTT molecules consumed. Charrier et al. (2012) also reported that PM$_{2.5}$ sampled in an urban environment in Fresno, California USA has a DTT activity of 27 - 61 pmol min$^{-1}$$\mu g^{-1}$, corresponding to $(2-7) \times 10^{14}$ $\mu g^{-1}$ of DTT molecules consumed in 20 min. Assuming that the consumption of one DTT molecule would correspond to the generation of one ROS molecule (e.g., $H_2O_2$), these values are about a few orders of magnitude higher than concentrations of radical forms of ROS measured in this study. This is reasonable as $H_2O_2$ is closed shell and much more stable than open-shell radical forms of ROS.

3.3. ROS formation mechanism
It has been shown that semiquinones and reduced transition metals including Fe(II) and Cu(I) can react with O$_2$ to form O$_2^-$, which can be further converted to H$_2$O$_2$ (Gehling et al., 2014; Fang et al., 2015). Fenton-like reactions of H$_2$O$_2$ with Fe(II) or Cu(I) can lead to the formation of OH radicals (Winterbourn, 2008; Pöschl and Shiraiwa, 2015). OH radicals can also be generated by the decomposition of organic hydroperoxides (ROOH) contained in SOA, yielding RO radicals (Tong et al., 2016). Several studies have reported a metal-independent decomposition of hydroperoxides and organic hydroperoxides driven by substituted quinones producing RO radicals (Sanchez-Cruz et al., 2014; Huang et al., 2015). The presence of Fe(II) or quinones is suggested to enhance ROOH decomposition and the formation of RO and OH radicals (Zhu et al., 2007a; Zhu et al., 2007b; Zhu et al., 2009; Sanchez-Cruz et al., 2014). Organic peroxides (ROOR) do not yield OH and RO radicals even in the presence of iron ions (Tong et al., 2016).

Based on these previous studies and considering that ambient particles may contain quinones, organic hydroperoxides, and transition metals, the observed ROS formation may be caused by interactions of these chemical components. To further investigate this aspect, mixtures of organic hydroperoxides, quinones, and Fe(II) were analysed by EPR and liquid chromatography mass spectrometry (LC-MS). Two standard organic hydroperoxides, cumene hydroperoxide and tert-butyl hydroperoxide, were used. For quinones, p-benzoquinone and humic-like substances are used, as HULIS are known to contain substantial amounts of quinones (Verma et al., 2015c).

Figure 6 shows the comparison of EPR spectra of ambient particles with a diameter of 180 - 320 nm (black) sampled on 26 June 2015 (same as shown in Fig. 4) and the above mixtures of organic compounds. Panel (a) includes EPR spectra of mixtures of all three different components (ROOH, quinone, metal) and panel (b) presents mixtures of two different components. All three of the organic mixtures in panel (a) resemble the EPR spectrum of ambient particles by reproducing almost all of the peaks. Particularly, the EPR spectrum of the mixture containing cumene hydroperoxide, humic acid and Fe(II) closely overlaps with the ambient particle EPR spectrum. Similarity of spectra between p-benzoquinone and HULIS suggests that the chemical nature of quinones and HULIS is very similar. Note that peaks related to the BMPO-OOH adduct at 3497 G and at 3530 G are more prominent in standard organic mixtures compared to ambient particles. This may be due to the relatively short lifetime of BMPO-OOH of ~23 min (Zhao et al., 2001), which is comparable to the extraction and mixing time of BMPO with the atmospheric particles (21 – 28 min), during which BMPO-OOH may have decayed. The trapped radicals have been...
further characterized by LC-MS, confirming the presence of OH and semiquinone radicals as well as carbon- and oxygen centred organic radicals, as detailed in Appendix A and Figs. A1 and A2.

EPR spectra of mixtures containing two compounds in panel (b) reproduce only a part of the observed peaks. These observations strongly suggest that the combination of these three chemical components play an important role in generating ROS species by atmospheric particles. The role of transition metals is crucial to enhance radical formation, most likely via Fenton-like reactions (Tong et al., 2016) and by participating in redox-cycling of quinones (Khachatryan and Dellinger, 2011), as intensities of EPR spectra without Fe(II) (CHP + HULIS, dark blue; CHP + pBq, orange) are small. Carbon-centred radicals may have multiple sources such as the decomposition of the BMPO-OR adduct by scission of the carbon in β position, yielding for example CH$_3$ radicals (Zhu et al., 2007b; Huang et al., 2015) as detected by LC-MS (Fig. A2). They may also be generated by secondary reactions of non-trapped OH radicals with water-soluble organic compounds.

SOA particles, which may contain large amounts of organic hydroperoxides, account for a major fraction in PM1 (Jimenez et al., 2009). SOA compounds may also coat coarse particles such as biological particles (Pöhlker et al., 2012). As shown in Fig. 2, semiquinones are mostly contained in submicron particles but not in coarse particles. Thus, the release of a variety of ROS species are most likely due to the interactions of organic hydroperoxides, semiquinones, and transition metal ions, whereas the dominance of OH radicals in coarse particles may be due to the decomposition of organic hydroperoxides in the absence of semiquinones.

4. Conclusions and implications

In this study particle-associated environmentally persistent free radicals (EPFR) and radical forms of ROS have been quantified using electron paramaganetic resonance (EPR) spectroscopy. Average EPFR concentrations were measured to be ~2×10$^{11}$ spins µg$^{-1}$ in ambient particles collected in Mainz, Germany in May – June 2015. The chemical identity of EPFR is likely to be semiquinone radicals based on the g-factors observed by EPR spectroscopy. We found that particles with different sizes had different radical contents and particles with a diameter of 100 - 180 nm had the highest abundance of EPFR, whereas coarse particles did not contain EPFR. This is consistent with the size distribution of combustion particles such as soot and humic-like substances (HULIS), which may contain substantial amounts of EPFR.
Reactive oxygen species (ROS) are formed upon extraction of particles into water. Particles with the diameter of 100 – 180 nm have released the highest ROS concentrations of $2.7(\pm 0.2) \times 10^{11}$ spins $\mu$g$^{-1}$. By deconvoluting the obtained EPR spectra, four types of radicals, including OH, $O_2^-$, carbon-centred and oxygen-centred organic radicals were quantified. The relative amounts of OH, $O_2^-$, C-centred and O-centred organic radicals in submicron particles were found to be $\sim$11 - 31%, $\sim$2 – 8%, $\sim$41 – 72% and $\sim$0- 25%, respectively, depending on the particle size. OH was the dominant species for coarse particles with a diameter larger than 1 $\mu$m. We suggest that the formation of these ROS species is due to the decomposition of organic hydroperoxides, which are a major component in SOA, interacting with semiquinones contained in soot or HULIS. ROS formation can be enhanced in the presence of iron ions by Fenton-like reactions.

These findings have significant implications for the chemical processing of organic aerosols in deliquesced particles and cloud water. The released OH radicals within particles or cloud droplets can oxidize other organic compounds, producing low-volatility products including organic acids, peroxides, and oligomers (Lim et al., 2010; McNeill et al., 2012; Ervens, 2015; Herrmann et al., 2015). Autoxidation in the condensed phase might be triggered by OH radicals forming highly oxidized compounds (Shiraiwa et al., 2014; Tong et al., 2016). High aqueous oxidant levels may cause fragmentation of organic compounds, resulting in an increased loss of carbon from the condensed phase (Daumit et al., 2016). The formed carbon- and oxygen-centred organic radicals are also expected to enhance chemical aging by participating in particle-phase chemistry involving aldehydes, carbonyls, and organic peroxides (Ziemann and Atkinson, 2012), although the exact role and impact of formed organic radicals are still unclear and subject to further studies.

Previous studies have shown that redox-active components such as transition metals and quinones can induce ROS formation in surrogate lung lining fluid upon interactions with antioxidants (Charrier and Anastasio, 2011; Charrier et al., 2014). This study also implies that ROS can be formed in lung lining fluid upon inhalation and respiratory deposition of atmospheric aerosol particles. Even though some fractions of ROS may be scavenged by antioxidants contained in lung lining fluid, excess concentrations of ROS including OH radicals, superoxide, and potentially also carbon- and oxygen centred organic radicals may cause oxidative stress to lung cells and tissues (Winterbourn, 2008; Pöschl and Shiraiwa, 2015; Tong et al., 2016). Recently, Lakey et al. (2016) have shown that fine particulate matter containing redox-active transition metals, quinones, and secondary organic aerosols can increase ROS concentrations in the lung lining fluid to levels characteristic for
respiratory diseases. ROS play a central role in chemical transformation of biomolecules such as proteins and lipids in lung fluid to form damage associated molecular patterns (DAMPs), which can trigger immune reactions causing inflammation through the toll-like receptor radical cycle (Lucas and Maes, 2013). Due to the important implications to adverse aerosol health effects, further studies are warranted to characterize and quantify EPFR and ROS contained in atmospheric aerosol particles in various locations including highly polluted regions such as in East Asia and India.

Appendix A. LC-MS analysis of organic mixtures

Two solutions of mixtures of standard organic hydroperoxides and quinones were analysed by liquid chromatography mass spectrometry (LC-MS). Solution (1) was the mixture of 200 µL of p-benzoquinone solution at a concentration of 0.2 g L⁻¹ (Reagent grade, ≥98%, Sigma-Aldrich) in water (trace SELECT® Ultra, ACS reagent, for ultratrace analysis, Sigma-Aldrich), 100 µL of Tert-Butyl hydroperoxide solution at a concentration of 8.9 g L⁻¹ (Luperox® TBH70X, 70 wt. % H₂O, Sigma-Aldrich) in water, 2.5 µL of Iron (II) sulfate heptahydrate solution at 0.3 g L⁻¹ (reagentPlus®, ≥99%, Sigma-Aldrich) in water and 1 mg of 5-tert-Butoxycarbonyl-5-methyl-1-pyrroline-N-oxide (BMPO, high purity, Enzo Life Sciences GmbH). Solution (2) was the same as solution (1) but without Iron (II) sulfate heptahydrate. These solutions were stirred with a vortex shaker (Heidolph Reax 1) for 5 minutes.

These solutions were analysed using a 1260 Infinity Bio-inert Quaternary LC system with a quaternary pump (G5611A), a HiP sampler (G5667A) and an electrospray ionization (ESI) source interfaced to a Q-TOF mass spectrometer (6540 UHD Accurate-Mass Q-TOF, Agilent). All modules were controlled by the MassHunter software (B.06.01, Agilent). The LC column was a Zorbax Extend-C18 Rapid resolution HT (2.1 x 50 mm, 1.8 µm) with a column temperature of 30 °C. The mobile phases were 3% (v/v) acetonitrile (HPLC Gradient Grade, Fisher Chemical) in water with formic acid (0.1 % v/v, LC-MS Chromasolv, Sigma-Aldrich) (Eluent A) and 3 % water in acetonitrile (Eluent B). The injection volume was 10 µL. The flow rate was 0.2 mL min⁻¹ with a gradient program that starting with 3 % B for 3 min followed by a 36 minutes step that raised eluent B to 60 %. Further, Eluent B was increased to 80 % at 40 minutes and returned to initial conditions within 0.1 minutes, followed by column re-equilibration for 9.9 min before the next run.

The ESI-Q-TOF instrument was operated in the positive ionization mode (ESI+) with a gas temperature of 325 °C, 20 psig nebulizer, 4000 V capillary voltage and 90 V fragmentor
voltage. During the full spectrum MS mode, no collision energy was used in order to collect species as their molecular ions. During MS/MS analysis employed for the structure determination, the fragmentation of protonated ions was conducted using the target MS/MS mode with 20 V collision energy. Spectra were recorded over the mass range of m/z 50-1000. Data analysis was performed using qualitative data analysis software (B.06.00, Agilent). Blank solutions without BMPO were also prepared and analysed. Background signals were subtracted from the MS spectrum.

Figure A2 shows LC-MS/MS mass spectra of the products formed from the reaction of tert-butyl hydroperoxide, p-benzoquinone, BMPO in the presence of iron (solution 1). Very similar results were obtained for solutions in the absence of iron (solution 2). BMPO adducts with radicals ·OH, ·CH₃ and ·OCH₃ were identified by LC-MS/MS. As shown in Figure A2(a.1), it was observed that ions at m/z 160.0596, 216.1221 and 238.1020 were major ions formed in the positive mode. These protonated ions represent the [BMPO+OH-C₄H₈+H]⁺, [BMPO+OH+H]⁺ and [BMPO+OH+Na+H]⁺ spin adducts, respectively. Figure A2 (a.2) displays the mass spectrum in the MS/MS mode for the fragmentation of the ion m/z 216.1221. Results confirmed the loss of the t-butoxycarbonyl function (-C₆H₅), which is a characteristic fragment of BMPO, to form the ion m/z 160.0585. The observed ion fragment m/z 114.0544, can be formed by the loss of CH₂O₂, as shown in Fig. A2(a.3). In Fig. A2(b.1), the spectrum showed the mass m/z 158.0804 and 214.1431 that can be attributed to the [BMPO+CH₃-C₄H₈+H]⁺ and [BMPO+CH₃+H]⁺, respectively. The most abundant fragment ion (m/z 158.0803) in the MS/MS mode confirmed the formation of BMPO+CH₃ adduct, as shown in Fig. A2(b.2). The peak m/z 112.0752 can be formed by the loss of CH₂O₂ (Fig. A2(b.3)). The spectrum in Fig. A2(c.1) shows major peaks at m/z 174.0752, 230.1378 and 252.1198, corresponding to [BMPO+OCH₃-C₄H₈+H]⁺, [BMPO+OCH₃+H]⁺ and [BMPO+OCH₃+Na+H]⁺, respectively. The formation of BMPO-OCH₃ was confirmed in MS/MS by the loss of the t-butoxycarbonyl functional group of BMPO to form the ion at m/z 174.0749 (panels (c.2) and (c.3)).

In addition, the radicals C₆H₅O₂· or ·C₆H₅O₂ and C₆H₅O₂· or ·C₆H₅O₂ were detected, although it was not possible to determine whether the chemical structure represented carbon- or oxygen-centred organic radicals using the applied method. Figure A3(a.1) shows the formation of protonated ions [BMPO+C₆H₅O₂+H]⁺ and [BMPO+C₆H₅O₂+Na+H]⁺ with m/z 308.1475 and 330.1298, respectively. The fragmentation in the MS/MS mode confirms the formation of BMPO+C₆H₅O₂ (m/z 252.0855) that correspond to the loss of the characteristic t-butoxycarbonyl function as show in Fig. A3(a.2). The ion fragment observed m/z 128.0702,
can be formed by the loss of C$_5$O$_4$ (Figure A3(a.3)). Figure A3(b.1) shows the ion m/z 312.1789, which can be attributed to the BMPO+C$_6$H$_9$O$_2$ spin adduct. Figure A3(b.2) suggests that the fragmentation of m/z 312.1789 to m/z 256.1166 by the loss of -C$_4$H$_8$ (Figure A3(b.3)).

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References.


**Figure 1:** Electron paramagnetic resonance (EPR) spectra of atmospheric aerosol impactor samples with lower cut-off diameters in the range of 56 nm to 1.8 micrometer collected in Mainz, Germany during 26 - 27 June, 2015.

**Figure 2:** Concentrations (spins per microgram of particles) of environmentally persistent free radicals (EPFR) and radical forms of reactive oxygen species (ROS) in atmospheric aerosol samples plotted against particle diameter. The error bars represent standard errors based on uncertainties of the particle mass and signal integration of EPR spectra.
Figure 3: Temporal evolution of concentrations of environmentally persistent free radicals (EPFR) contained in atmospheric aerosol samples with lower cutoff diameters of 100 nm (red) and 180 nm (blue), measured in Mainz, Germany during May – June 2015. The error bars represent standard errors based on uncertainties of the particle mass and signal integration of EPR spectra.
**Figure 4**: a) Electron paramagnetic resonance (EPR) spectra of ambient aerosol impactor samples (Mainz, Germany, 26-27 June 2015) with lower cut-off diameters in the range of 56 nm to 1.8 µm extracted in water mixed with the spin-trapping agent BMPO. Dashed lines indicate the position of each peak for different types of trapped radicals of $\text{O}_2^-$ (red), OH (green), carbon-centred (orange), and oxygen-centred organic radicals (light blue). b) Simulation of the EPR spectrum of the atmospheric aerosol impactor sample with particle diameters in the range of 180-320 nm (lower to upper cut-off) by deconvoluting into $\text{O}_2^-$, OH, O-centred and C-centred organic radicals (Blue = synthesis, grey = residual).
Figure 5: Relative amount of ROS in atmospheric aerosol impactor samples with lower cut-off diameters in the range of 50 nm -1.8 µm (Mainz, Germany, 26-27 June 2015): O₂⁻ (red), OH (green), carbon-centred (orange), oxygen-centred organic radicals (blue) and residual (unidentified, grey).
**Figure 6:** Electron paramagnetic resonance (EPR) spectra of atmospheric aerosol impactor sample with particle diameters in the range of 180-320 nm (lower to upper cut-off) extracted with water and BMPO (black) and of aqueous substance mixtures with the following ingredients: cumene hydroperoxide (CHP), p-benzoquinone (pBq) and Fe(II) (light blue), t-butyl hydroperoxide (TBHP). The dashed vertical lines indicate the main peaks of BMPO adducts with $\text{O}_2^-$ (red), OH (green), carbon- (light blown), and oxygen-centred organic radicals (light blue).
Figure A1: EPR spectra of atmospheric aerosol impactor samples with lower cut-off diameters of 56 nm (black), 100 nm (red), 180 nm (light blue) and 320 nm (green) for the measurement period during 28 May – 9 June 2015.
Figure A2: Mass spectra obtained with LC-MS/MS in the positive ionization mode from the mixture of tert-butyl hydroperoxide, p-benzoquinone and BMPO in the presence of iron (solution (1)). MS spectra of (a.1) BMPO+OH, (b.1) BMPO+CH₃ and (c.1) BMPO+OCH₃. MS/MS spectra of (a.2) BMPO+OH, (b.2) BMPO+CH₃, and (c.2) BMPO+OCH₃. Proposed fragmentation pathways of (a.3) BMPO+OH, (b.3) BMPO+CH₃ and (c.3) BMPO+OCH₃.
**Figure A3**: Mass spectra obtained with LC-MS/MS in the positive ionization mode for solution (1). MS spectra of (a.1) BMPO+C$_6$H$_5$O$_2$ and (b.1) BMPO+C$_6$H$_9$O$_2$. MS/MS spectra of (a.2) BMPO+C$_6$H$_5$O$_2$ and (b.2) BMPO+C$_6$H$_9$O$_2$. Proposed fragmentation pathway of (a.3) BMPO+C$_6$H$_5$O$_2$ and (b.3) BMPO+C$_6$H$_9$O$_2$. 