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<td>Publication date</td>
<td>2017-11-30</td>
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<tr>
<td>Type of publication</td>
<td>Article (peer-reviewed)</td>
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<tr>
<td>Link to publisher's version</td>
<td><a href="http://dx.doi.org/10.1016/j.atmosenv.2016.11.063">http://dx.doi.org/10.1016/j.atmosenv.2016.11.063</a></td>
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PII: S1352-2310(16)30953-0
DOI: 10.1016/j.atmosenv.2016.11.063
Reference: AEA 15056

To appear in: Atmospheric Environment

Received Date: 23 September 2016
Revised Date: 17 November 2016
Accepted Date: 28 November 2016


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Gas- and particle-phase products from the photooxidation of acenaphthene and acenaphthylene by OH radicals

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Abstract

This work is focused on the gas-phase oxidation of acenaphthylene and acenaphthene by OH radicals and associated secondary organic aerosol (SOA) formation under low and high-NO$_x$ conditions. Experiments were carried out in an atmospheric simulation chamber using a proton transfer reaction time-of-flight-mass spectrometer (PTR-TOF-MS) and an aerosol time-of-flight-mass spectrometer (ATOFMS) to chemically characterize the gas- and particle-phase products, respectively. Due to the structures of these two aromatic compounds, the proposed chemical mechanisms exhibit some differences. In the case of acenaphthene, H-atom abstraction from the saturated cyclopenta-fused ring was found to be competitive with the OH-addition to the aromatic rings. During the photooxidation of acenaphthene using nitrous acid (HONO), aromatic ring-opening products such as indanone and indanone carbaldehyde, generated through OH addition to the aromatic ring, were formed in higher yields compared to low-NO$_x$ conditions. In the case of acenaphthylene, OH addition to the unsaturated cyclopenta-fused ring was strongly favored. Hence, ring-retaining species such as acenaphthenone and acenaphthenequinone, were identified as the main reaction products in both gas- and particle-phases, especially under high-NO$_x$ conditions. Subsequent SOA formation was observed in all experiments and SOA yields were determined under low/high-NO$_x$ conditions to be 0.61/0.46 and 0.68/0.55 from the OH-initiated oxidation of acenaphthylene and acenaphthene, respectively.
1. Introduction

Atmospheric fine particulate matter (PM$_{2.5}$, aerosol with aerodynamic diameters less than 2.5 µm) plays a key role in air quality and climate change, and is associated with damaging effects on human health (Pope III and Docherty, 2006; Hallquist et al., 2009). Globally, the largest mass fraction of PM$_{2.5}$ is organic (up to 90% in some locations) and is dominated by secondary organic aerosol (SOA). Formation of SOA mainly results from the formation of gas-phase products with low vapor pressures from the oxidation of volatile organic compounds (VOCs) (Hallquist et al., 2009). The anthropogenic contribution to global SOA formation is estimated to be small, actually around 10% (Kroll et al., 2008). SOA formation from anthropogenic sources is, however, suggested to be higher than currently predicted (Volkamer et al., 2006; Hallquist et al., 2009). In order to explain the current discrepancy between the mass of aerosols measured in different atmospheres and the mass predicted by atmospheric models, some studies have suggested that other sources of SOA are not yet identified or well characterized (Kroll et al., 2008; Hallquist et al., 2009). Among them, heterogeneous chemistry (formation of oligomers and organosulfates) (Surratt et al., 2008; Pye and Pouliot, 2012) and the contribution of intermediate volatility organic compounds (IVOCs) to SOA formation have been proposed (Robinson et al., 2007; Tkacik et al., 2012). It is worth noting that recent studies have also underlined that reactive uptake and/or multiphase chemistry of water-soluble volatile organic compounds, such as glyoxal, in wet acidic aerosols or cloud droplets could be an important source of SOA (Galloway et al., 2009; Ervens et al., 2011; Pye et al., 2013; Marais et al., 2016). Nevertheless, a large part of the underestimation in urban areas, due to the non-considered IVOC contribution to SOA formation, comes from the involvement of alkanes and polycyclic aromatic hydrocarbons (PAHs) (Tkacik et al., 2012). Much of the current efforts in the research community are focused on trying to identify these missing or misrepresented SOA sources.

PAHs are emitted into the atmosphere from incomplete combustion processes of organic materials and have been identified as major components in traffic and wood burning emissions. The PAHs with less than four aromatic rings exist mainly in the gaseous phase and can undergo photooxidation processes with different atmospheric oxidants (Atkinson and Aschmann, 1988; Sasaki et al., 1998; Keyte et al., 2013; Zhou and Wenger, 2013a,b; Riva et al., 2014). When oxidized, these compounds have been shown to produce large range of oxygenated and nitro compounds with high molecular weights (Sasaki et al., 1998; Reisen and
Arey, 2002; Lee and Lane, 2009; Kautzman et al., 2010; Kleindienst et al., 2012; Zhou et al., 2013a,b; Riva et al., 2015a,b,c, 2016). Although PAHs are potentially carcinogenic and mutagenic (Atkinson and Arey, 1994) some of their oxidation products present a larger toxicity than their parent hydrocarbons (Lin et al., 2005). Gas-phase products can partition to the particle phase and participate in SOA formation (Chan et al., 2009; Kautzman et al., 2010; Shakya et al., 2010; Kleindienst et al., 2012; Riva et al., 2015a; Chen et al., 2016; Riva et al., 2016). A few studies have previously reported the importance of naphthalene gas phase photooxidation in SOA formation (Chan et al., 2009; Kleindienst et al., 2012). Other gaseous PAHs have, however received less attention. Acenaphthylene and acenaphthene are fairly unique among PAHs in that they contain a carbon-carbon double bond and a saturated carbon-carbon bond, respectively, in their structure that enables them to react quickly with all atmospheric oxidants including OH and NO\textsubscript{3} radicals, Cl atoms and O\textsubscript{3} (Atkinson and Aschmann, 1988; Reisen and Arey, 2002; Zhou and Wenger, 2013a,b; Riva et al., 2014). Acenaphthylene and acenaphthene were identified in both indoor and outdoor atmospheres and in large concentrations in certain areas (Chang et al., 2006; Ho et al., 2009). The concentration of both PAHs (greater than 20-30 ng m\textsuperscript{-3}) could be even comparable to that of naphthalene in certain urban atmospheres (Dejean et al., 2009; Hanedar et al., 2014) suggesting that acenaphthylene and acenaphthene might contribute to SOA formation in such areas. Large rate constants for the reaction of acenaphthene ((7.69 ± 1.91) × 10\textsuperscript{-11}) and acenaphthylene ((1.14 ± 0.08) × 10\textsuperscript{-10}) in cm\textsuperscript{3} molecule\textsuperscript{-1} s\textsuperscript{-1}) with OH radicals have been previously reported (Atkinson and Aschmann, 1988; Zhou and Wenger, 2013a,b). These previous efforts underline the potential importance of the OH initiated oxidation of acenaphthylene and acenaphthene in the atmospheric chemistry of PAHs. Indeed, it has been shown that the reactivity of aromatic hydrocarbons is dominated by their reactions with the OH radicals in the atmosphere (Calvert et al., 2002; Estève et al., 2003; Atkinson and Arey, 2007). Although the OH-initiated oxidation of PAHs has been studied previously, the impact of NO\textsubscript{x} concentration on the reactivity of PAHs remains poorly documented. Only two studies on the oxidation of naphthalene performed under high- and low-NO\textsubscript{x} conditions, have been reported hitherto (Kautzman et al., 2009; Kleindienst et al., 2012). Previous works have determined the rate constants and oxidation products for the photooxidation of acenaphthene and acenaphthylene in the presence of NO\textsubscript{x} (Atkinson and Aschmann, 1988; Reisen and Arey, 2002; Zhou and Wenger, 2013a,b). In addition, Shakya and Griffin (2010) have, for the first time, reported the SOA yields arising from these reactions. These results highlight the propensity for both PAHs to participate in SOA formation and only partial mechanisms were...
proposed in these previous studies. However, some discrepancies remain and further work is needed to clarify the reaction products as well as the chemical mechanisms leading to SOA formation.

In this work, we performed a series of simulation chamber experiments to identify products arising from the photooxidation of acenaphthylene and acenaphthene under low- and high-\(\text{NO}_x\) conditions using state-of-the-art mass spectrometry techniques for on-line analysis of both gaseous (proton transfer reaction time-of-flight-mass spectrometer) and particulate (aerosol time-of-flight-mass spectrometer) phases. Based on these approaches, extended and improved reaction mechanisms for the OH-initiated oxidation of acenaphthylene and acenaphthene are now proposed. SOA growth and yields are determined in both cases to evaluate the importance of the photooxidation of acenaphthene and acenaphthylene in SOA formation.

2. Experimental section

Experiments were performed at room temperature (293 ± 2 K) and atmospheric pressure in the 3910 L atmospheric simulation chamber at University College Cork, which is described in detail elsewhere (Thüner et al., 2004). Briefly, it is a cylinder consisting of a Teflon FEP foil tube closed with aluminum plates covered with Teflon FEP foil. The chamber is equipped with fans to ensure rapid mixing of reactants and is surrounded by 16 lamps (Philips TL12, 40W) with an emission maximum at 310 nm and 12 lamps (Philips TL05, 40 W) with an emission maximum at 360 nm (Kourtchev et al., 2009). Before each experiment, the chamber is cleaned by flushing with dried purified air and photolysis of added ozone (ca. 1 ppmv) until the particle number concentration is below 200 cm\(^{-3}\). The flushing also reduces the levels of \(\text{NO}_x\) and non-methane hydrocarbons to < 10 ppbv. The mixing ratios of \(\text{NO}_x\) and \(\text{O}_3\) are measured using standard automated gas analyzers (Thermo Model 42i and 49i respectively). The temperature and water concentration in the chamber were monitored by a dew point meter (Vaisala DM70). The relative humidity in the chamber was typically less than 1% for the experiments with HONO and less than 5% for experiments carried out with \(\text{H}_2\text{O}_2\).

Aacenaphthylene (Sigma-Aldrich, 99%) and acenaphthene (Sigma-Aldrich, 99%) were introduced into the chamber by flowing dry purified air throughout a heated Pyrex glass bulb containing a known amount of the solid compound sufficient to produce PAH mixing ratios around 300 ppb (Table S1). The PAHs and gas-phase oxidation products were monitored
during the experiments using a proton transfer reaction - time of flight - mass spectrometer (PTR-TOF-MS, Kore Technology Ltd.). Details of the instrument and its operating principle are given in Cappelin et al. (2012). Briefly, \( \text{H}_3\text{O}^+ \) is produced in a hollow cathode ion source and reacts with organic compounds (M) that have a higher proton affinity than \( \text{H}_2\text{O} \) to generate positively charged ions \((\text{M} + \text{H})^+\), which are subsequently detected using a time-of-flight mass spectrometer. The PTR-TOF-MS was operated over the \( m/z \) range 0-300 using a sampling time of 1 min. The decay of the PAHs was monitored by following the protonated molecular ions: \( m/z \) 153 (acenaphthylene) and \( m/z \) 155 (acenaphthene). Quantification of identified products was not possible due to difficulty obtaining accurate concentrations in the gas phase during the PTR-TOF-MS calibration.

Experiments were carried out in order to study the oxidation of acenaphthylene and acenaphthene under high- and low-\( \text{NO}_x \) conditions. Nitrous acid (HONO) was used as an OH radical precursor and was generated by adding 0.5 mL of 1 wt % aqueous sodium nitrite (\( \text{NaNO}_2 \), Sigma-Aldrich, 97\%) dropwise into 30 mL of 30 wt % sulfuric acid (Sigma-Aldrich, 99.9\%) in a glass bulb (Cox, 1974). Purified air was flowed through the bulb, delivering HONO to the chamber. The injection of HONO was stopped when the concentration of nitrogen dioxide reached about 200 ppb in the chamber. Hydrogen peroxide (\( \text{H}_2\text{O}_2 \), Sigma-Aldrich, 50 wt \%) was used to generate OH radicals without \( \text{NO}_x \). 60 \( \mu \text{L} \) was injected into the glass bulb followed by slight heating. Purified air was passed through the bulb to introduce \( \text{H}_2\text{O}_2 \) into the chamber.

The formation and growth of SOA was monitored using a scanning mobility particle sizer (SMPS, TSI model 3034): size distribution, number and mass concentrations were determined for all experiments, assuming an SOA density of 1.4 g cm\(^{-3} \) (Chan et al., 2009; Shakya et al., 2010). SOA chemical composition was investigated using an aerosol time-of-flight mass spectrometer (ATOFMS, TSI model 3800), which has been described in detail elsewhere (Gard et al., 1997). Briefly, single particles are sampled through a critical orifice and focused into a tight beam in the aerodynamic lens before transmission to the sizing region. The velocity of the particles is measured using two continuous wave diode-pumped Nd:YAG lasers operating at 532 nm. The time between these two scattering events is used to obtain the aerodynamic size. Aerosols are then transmitted to the ionization region of the instrument where desorption/ionization is performed by a Nd:YAG laser at 266 nm. This ablation process generates high densities of charges, in which intermolecular interactions and matrix effects...
often determine the ion distribution, making quantitative calibration of mass spectra extremely difficult (Zelenyuk and Imre, 2005). Due to instrumental issues only positive ions were detected using the ATOFMS. It is important to note, however, that PAHs are mainly detected as positive ions (Silva and Prather, 2000; Zimmermann et al., 2003). The laser is typically operated with an output energy of around 1 mJ per pulse. However, as described elsewhere lower pulse energies were used in this work to reduce fragmentation of the organic constituents in the aerosol (Zimmermann et al., 2003; Gross et al., 2000; 2005).

Based on the measured decay of PAH and known rate constants for reaction with OH (unit: cm$^3$ molecule$^{-1}$ s$^{-1}$): $(7.69 \pm 1.91) \times 10^{-11}$ and $(1.14 \pm 0.08) \times 10^{-10}$ for acenaphthene and acenaphthylene, respectively (Atkinson and Aschmann, 1988; Brubaker and Hites, 1998; Banceau et al., 2001; Reisen and Arey, 2002; Zhou and Wenger, 2013a,b) the initial OH radical concentrations using H$_2$O$_2$ and HONO as precursor were (unit: molecule cm$^{-3}$): $(3.47 \pm 0.32) \times 10^6$ and $(8.34 \pm 1.36) \times 10^6$ respectively.

3. Results and discussion

3.1 Acenaphthylene photooxidation

The PTR-TOF-MS measurements performed in acenaphthylene photooxidation experiments carried out under low and high-NO$_x$ conditions revealed 6 product peaks with $m/z$ values reported in Table S2. The temporal profiles of the identified ions are proposed in Figure 1. Most of the reaction products are similar to those previously reported or for reactions performed in the same chamber but using Cl atoms or O$_3$ to initiate oxidation (Reisen and Arey, 2002; Zhou and Wenger, 2013b; Riva et al., 2015a; Riva et al., 2016). Based on existing knowledge of the gas-phase chemistry of acenaphthylene, the peak at $m/z$ 185 is assigned to the protonated molecular ion (M + H)$^+$ of naphthalene-1,8-dicarbaldehyde (C$_{12}$H$_8$O$_2$) and a dialdehyde formed from OH addition to the aromatic rings. Previous studies have reported large yields of both compounds from the photooxidation of acenaphthylene in the presence of NO$_x$. The protonated ion at $m/z$ 185 may have, however, more than 2 isomers. Indeed, as discussed below and proposed in Scheme 1, formation of a protonated ion at $m/z$ 183 (C$_{12}$H$_6$O$_2$) and tentatively assigned as acenaphthenequinone cannot be explained by solely the gas-phase oxidation of acenaphthenone (C$_{12}$H$_8$O). Indeed, the abundance of protonated ion at $m/z$ 169 is likely too weak to explain the large signal observed foracenaphthenequinone in the particulate phase (Figure 1). Therefore, formation of C$_{12}$H$_6$O$_2$
might also proceed through the gas-phase oxidation of hydroxy-acenaphthenone as tentatively proposed in Scheme 1. Signals for the protonated ions at $m/z$ 169 and 185 decrease significantly after ca. 4000 s reaction time (Figure 1), suggesting that both primary reaction products undergo further gas-phase oxidation and/or photolysis leading to secondary generation compounds (Scheme 1). For instance, naphthalene-1,8-dicarbaldehyde may react further with OH radicals or be photodissociated as observed for other aromatic aldehydes (Wang et al., 2006).

Although the different compounds identified for each set of experiments were identical under high- or low-NO$_x$ conditions, their abundances were very different. As presented in Figure 1, the abundances of ring-opening products (e.g. oxaaacenaphthylene-2-one, 1,8-naphthalic anhydride and 1,8-naphthalaldehydeic acid) differ depending on the OH radical precursor. Moreover, the high signals of small fragment ions detected by the ATOFMS (Figure 2) at $m/z$ 39 ($C_3H_3$)$^+$, 51 ($C_4H_3$)$^+$, 63 ($C_5H_3$)$^+$, 75 ($C_6H_3$)$^+$, 77 ($C_6H_5$)$^+$ and 115 ($C_9H_7$)$^+$ highlight the presence of ring-opening products in the particulate phase, as demonstrated in previous studies using an ATOFMS at low laser energy (Riva et al., 2015a; Gross et al., 2000; 2005). The higher yield of ring opening products such as oxaacenaphthylene-2-one, 1,8-naphthalic anhydride and 1,8-naphthalaldehydeic acid in the gas phase (Figure 1) is consistent with the higher signals for small fragment ions in the ATOFMS spectra (Figure 2), suggesting that formation of ring-opening products is likely enhanced under low-NO$_x$ conditions. This observation is in contrast with experiments performed in the presence of NO$_x$. In that case, ring-retaining species are enhanced, as revealed by the relatively larger abundances of acenaphthenone (or its enol-form) and acenaphthenequinone (Figures 1 and 2) in both phases, during the oxidation of acenaphthylene in the presence of HONO. It is worth noting that no nitro compounds were identified in either gas or particle phases. Previous studies have, however, reported the formation of nitro-products during the photooxidation of acenaphthylene, but with very low yields, which likely explains the absence of nitro-products in the mass spectra collected in this work (Arey et al., 1989; Reisen and Arey, 2002; Zhou and Wenger, 2013b).

An extended reaction mechanism for the gas-phase oxidation of acenaphthylene is proposed in Scheme 1. It is based on the PTR-TOF-MS data obtained in this work and also builds on the results from previous experimental studies (Reisen and Arey, 2002; Zhou and Wenger, 2013b). Oxidation of acenaphthylene is energetically favored at the unsaturated
cyclopenta-fused ring as highlighted in previous experimental (Atkinson and Aschmann, 1988; Reisen and Arey, 2002; Zhou and Wenger, 2013b; Riva et al., 2015a; Riva et al., 2016) and theoretical studies (Dang et al., 2015). This double bond exhibits high reactivity and explains the higher reactivity of acenaphthylene with atmospheric oxidants relative to other PAHs. Hence acenaphthylene photooxidation is governed by the OH addition at the unsaturated cyclopenta-fused ring. Due to the absence in our experiments of any nitro compounds and the very small yields reported in previous work, the OH-adduct is expected to mainly react with O$_2$. OH-acenaphthylene adduct could then lead to a peryo radical (RO$_2$) ($C_{12}H_9O_3$) and to acenaphthenone (and/or its enol-form). It has been reported that for simple aldehydes and ketones, the keto form is the most stable and thus the equilibrium is likely directed to the acenaphthenone (March, 1985). $C_{12}H_9O_3$ could then react with NO or go through its self- /cross-reactions leading to the alkoxy radical ($C_{12}H_9O_2$). Two different pathways are proposed for the fate of $C_{12}H_9O_2$, which appear to be as expected NO$_x$ dependent. Firstly, the ring-retaining channel could produce hydroxy-acenaphthenone, which could further react and lead to acenaphthenequinone (Scheme 1). As discussed above, this pathway appears to be likely favored in the presence of NO$_x$ due to the relatively lower abundance of small fragment ions in the mass spectrum of the ATOFMS as well as the larger presence of ring-retaining products in both gas and particulate phases (Figures 1 and 2). Secondly, $C_{12}H_9O_2$ can proceed through the ring-opening pathway occurring by C$_1$-C$_2$ bond cleavage. As shown in Scheme 1, this reaction pathway leads exclusively to ring-opening products, which appear to be favored under low-NO$_x$ conditions. Naphthalene-1,8-dicarbaldehyde and 1,8-naphthalaldehydic acid ($C_{12}H_8O_3$) are the two main primary products arising from this ring-opening pathway and further react as shown in Figure 2 and proposed in Scheme 1. As discussed above and reported in previous works (Zhou and Wenger, 2013b; Riva et al., 2015a,b), naphthalene-1,8-dicarbaldehyde can yield 1,8-naphthalic anhydride from H-atom abstraction of the aldehydic function, which would result in the formation of an RO$_2$ radical ($C_{12}H_9O_4$). Similarly to phthalaldehyde, naphthalene-1,8-dicarbaldehyde could undergo photodissociation, leading to 1,8-naphthalic anhydride (Wang et al., 2006; Riva et al., 2015b). $C_{12}H_9O_4$ could then react with NO or RO$_2$ and form an acyl-oxy radical ($C_{12}H_9O_3$). The chemistry of the resulting $C_{12}H_9O_3$ radical could proceed through the bicyclic peroxy radical route in a fashion analogous to that proposed in Scheme 1. $C_{12}H_9O_3$ radical could also decarboxylate quickly (Chacon-Madrid et al., 2013) and the resulting radical would react with O$_2$ leading to the formation of another RO$_2$ radical ($C_{11}H_7O_3$). As
presented in Scheme 1 and previously proposed, \( \text{C}_{11}\text{H}_{7}\text{O}_3 \) could undergo cyclization and lead to the formation of oxaacenaphthylene-2-one (Zhou and Wenger, 2013b).

It is important to point out that \( \text{O}_3 \) formation (~20 ppb final concentration) was observed during the experiments carried out with HONO from the photolysis of \( \text{NO}_2 \). Due to the high reactivity of acenaphthylene with ozone ((acenaphthylene + \( \text{O}_3 \)); \( k = 3.99 \times 10^{-16} \text{ cm}^3\text{ molecule}^{-1}\text{ s}^{-1} \)), \( \text{O}_3 \) might have participated in the formation of oxygenated species and acenaphthylene depletion (Zhou and Wenger, 2013b). Due to the large UV-absorption band overlap of acenaphthylene with ozone at wavelengths between 240 and 260 nm (Zander, 1969) it was not possible to determine an accurate contribution of ozone in the oxidation of acenaphthylene. Riva et al. (2016) recently reported that a secondary ozonide (\( \text{C}_{12}\text{H}_8\text{O}_3 \)) is the main gas phase product arising from ozonolysis of acenaphthylene. In the high-\( \text{NO}_x \) experiments, the product ion corresponding to \( \text{C}_{12}\text{H}_8\text{O}_3 \) was observed in very low abundances (i.e. <70 counts) indicating that the contribution of ozonolysis can be considered to be negligible in the photo-oxidation experiments performed in this work.

### 3.2 Acenaphthene photooxidation.

Ten oxidation products from the reaction of OH radicals with acenaphthene were identified by PTR-TOF-MS and the temporal profiles for the most abundant peaks are presented in Figure 3. These product peaks at \( m/z \) 133, 161, 169 and 177 were tentatively assigned to protonated molecular ions of indanone (\( \text{C}_9\text{H}_8\text{O} \)), indanone carbaldehyde (\( \text{C}_{10}\text{H}_8\text{O}_2 \)), acenaphthenone (\( \text{C}_{12}\text{H}_8\text{O} \)) and oxoindan-carboxylic acid (\( \text{C}_{10}\text{H}_8\text{O}_3 \)), respectively. Among these 4 products, indanone carbaldehyde and acenaphthenone have been previously observed from the OH-initiated oxidation of acenaphthene under high-\( \text{NO}_x \) conditions (Reisen and Arey, 2002; Zhou and Wenger, 2013a). For instance, Reisen and Arey (2002) reported significant formation of indanone carbaldehyde and estimated yields at 14-37%. Four other protonated ions were observed at \( m/z \) 171, 183, 185 and 199 but in weaker abundance and were tentatively assigned, based on existing knowledge of the gas-phase chemistry of acenaphthene, to acenaphthenol (\( \text{C}_{12}\text{H}_{10}\text{O} \)),acenaphthequinone (\( \text{C}_{12}\text{H}_6\text{O}_2 \)), naphthalene-1,8-dicarbaldehyde (\( \text{C}_{12}\text{H}_8\text{O}_2 \) and 1,8-naphthalic anhydride (\( \text{C}_{12}\text{H}_6\text{O}_3 \)), respectively. Finally, weak signals at \( m/z \) 187 and 200 measured only in the experiments performed in the presence of \( \text{NO}_x \) were tentatively assigned to a dialdehyde (\( \text{C}_{12}\text{H}_{10}\text{O}_2 \)) and nitroacenaphthene (\( \text{C}_{12}\text{H}_9\text{NO}_2 \)). Proposed products, and the ions used for their identification, are reported in Table S3.
The different products determined in this work are similar to those identified in previous studies or during other experiments on acenaphthene performed in the same environmental chamber but using Cl atoms as radical precursors (Reisen and Arey, 2002; Zhou and Wenger, 2013a; Riva et al., 2015a). As observed in the case of the photooxidation of acenaphthylene, the abundances of the different products were highly dependent on the NO\textsubscript{x} concentrations. The temporal profiles of the main compounds identified in the gas phase were very different as presented in Figure 3. For example, formation of indanone and indanone carbaldehyde during acenaphthene photooxidation under high-NO\textsubscript{x} conditions was higher, while their abundances remained low in the absence of NO\textsubscript{x}. Moreover, the protonated signal for acenaphthenone, which appears to be the main oxidation product from the photooxidation of acenaphthene under low-NO\textsubscript{x} conditions, was significantly higher compared to the experiments carried out in the presence of NO\textsubscript{x}. These observations highlight that oxidation pathways or branching ratios highly depend on the NO\textsubscript{x} concentration. Contrary to the acenaphthylene system, ATOFMS spectra for acenaphthene SOA (Figure 4) reveal high abundances of small fragment ions under low- and high-NO\textsubscript{x} conditions, suggesting a high contribution of aromatic ring-opening products. Thus, abundances of fragment ions in ATOFMS spectra cannot be used to distinguish reaction pathways. Several species have been identified in both the gas and particulate phases, corroborating a significant formation of aromatic ring-opening products (Reisen and Arey, 2002; Zhou and Wenger, 2013a).

A proposed extended reaction mechanism for the gas-phase oxidation of acenaphthene is presented in Scheme 2. It is based on the PTR-TOF-MS data obtained in this work and also builds on the results from previous experimental studies (Reisen and Arey, 2002; Zhou and Wenger, 2013a). The reaction of OH radicals with acenaphthene can, in principle, proceed via three possible pathways: OH addition to the aromatic rings, H-atom abstraction from the aromatic ring and H-atom abstraction from the saturated cyclopenta-fused ring. However, H-atom abstraction from the aromatic rings has been demonstrated to be negligible compared to OH addition for the oxidation of aromatic compounds (Calvert et al., 2002). Formation of aromatic ring-opened dicarbonyls, such as indanone carbaldehyde or dialdehyde, is proposed as shown in Scheme 2 to occur through OH addition to the aromatic rings followed by the reaction of the OH-acenaphthene adduct with O\textsubscript{2} and further ring cleavages (Reisen and Arey, 2002). In this work, indanone carbaldehyde was observed as the major compound arising from this OH-addition pathway. It is important to mention that secondary chemistry has not been discussed in previous work, however, as presented in Figure 3, primary products...
undergo further oxidation and lead to secondary products, including indanone and oxoindan-
carboxylic acid (Scheme 2).

The initial H-atom abstraction by OH radicals is followed by addition of O₂ leading to a RO₂ radical (C₁₂H₉O₂'). Subsequent reactions with NO or RO₂ radicals proceed via two pathways; (i) the molecular channel, which results in equimolar amounts ofacenaphthenone and acenaphthenol, and (ii) the radical channel which produces an RO radical (C₁₂H₉O'). C₁₂H₉O' can then further react with O₂ and lead to acenaphthenone as the sole product. As proposed in Scheme 2 an alternative pathway, i.e. C-C bond cleavage of the C₁₂H₉O' radical followed by further oxidation could lead to the formation of naphthalene-1,8-dicarbaldehyde. As described previously in the case of acenaphthylene, naphthalene-1,8-dicarbaldehyde could further react or be photolyzed and lead to 1,8-naphthalic anhydride from the H-atom abstraction of the aldehydic function. It is important to note that previous studies did not consider this channel and attributed the formation of acenaphthequinone, naphthalene-1,8-dicarbaldehyde and 1,8-naphthalic anhydride to analytical artifacts or from reactions with acenaphthene impurities (Reisen and Arey, 2002; Sauret-Szczechanski and Lane, 2004). Recently, Zhou and Wenger (2013a) investigated these potential analytical artifacts and demonstrated the formation of naphthalene-1,8-dicarbaldehyde during acenaphthene photooxidation. Acenaphthenone, acenaphthequinone and 1,8-naphthalic anhydride were also previously identified, ruling out the formation of these compounds as analytical artifacts. Moreover, the abundances of these products observed in this work cannot be explained by reactions of OH radicals or ozone with small amounts ofacenaphthene impurities (e.g. acenaphthylene, < 1%). Hence, our results complement those obtained by Zhou and Wenger (2013a) highlighting the competition between OH addition to the aromatic rings and H-atom abstraction from the cyclopenta-fused ring.

The competition between these two pathways appears to be different depending on the NOₓ concentration. As shown above, the relative abundances of the gas phase products were very different in the presence and absence of NOₓ (Figure 3). In the absence of NOₓ, the products from the OH-addition pathways (i.e. indanone, indanone carbaldehyde and oxoindan-carboxylic acid) were observed in much lower abundances, suggesting that NOₓ could contribute to the stability of the OH-acenaphthene adduct (Nishino et al., 2012). Conversely, products (i.e. acenaphthenone) from H-atom abstraction from the cyclopenta-fused ring pathway were observed in relatively higher abundance in both gas- and particulate
phases. Although, the (acenaphthene + OH) reaction proceeds through two different pathways under low-NO\textsubscript{x} conditions, it is difficult to distinguish which one dominates the global mechanism presented in Scheme 2. Previous studies have evaluated the competition between OH addition to the aromatic ring and H-atom abstraction (from aromatic rings) pathways: under atmospheric conditions the mechanism of reaction of OH radicals with aromatic hydrocarbons occurs mainly (i.e. > 90\%) via addition to the aromatic ring (Calvert et al., 2002; Atkinson and Arey, 2007). Therefore, the results presented in this work suggest that the competition between OH-addition and H-atom abstraction from the cyclopenta-fused ring is more important than previously expected. Hence, the OH oxidation of acenaphthene could also be governed by H-atom abstraction from the cyclopenta-fused ring depending on the concentration of NO\textsubscript{x}. More work is, however, needed to quantify the branching ratio between both pathways.

3.3 SOA formation.
Rapid SOA formation was observed in all experiments immediately after the formation of OH radicals. Using a semi-empirical model for SOA formation based on the gas-particle partitioning equilibrium of semi volatile products (Odum et al., 1996), the SOA yields (\(Y\)), were determined from the experiments performed under low- /high-NO\textsubscript{x} conditions to be 0.61/0.46 and 0.68/0.55 for acenaphthylene and acenaphthene, respectively (Table 1). The volume concentration was corrected for particle wall loss by applying size-dependent first-order loss coefficients after SOA growth was finished. The indicated uncertainties (2\(\sigma\)) in Table 1 correspond to scatter in particle volume measurements. The aerosol mass was calculated using volume concentrations measured by SMPS and assuming a particle density of 1.4 g cm\textsuperscript{-3} (Chan et al., 2009; Shakya and Griffin, 2010). The high concentrations used in this work probably enhanced partitioning of semi volatile species to the particle phase resulting in SOA yields that are larger than might be expected under more realistic atmospheric conditions. Nevertheless, the high yields determined in this work further illustrate the important potential of PAHs to form SOA from their oxidation by OH radicals. The first study reporting SOA formation yields from the photooxidation of acenaphthene and acenaphthylene under high- and low-NO\textsubscript{x} conditions was proposed by Shakya and Griffin (2010). The SOA yields proposed in the previous work (0.04-0.13 and 0.03-0.11 for acenaphthylene and acenaphthene photooxidation respectively) are much lower than those reported here. It is worth noting that, even though similar initial PAH concentrations were used, large discrepancies exist between the SOA yields reported by Chan et al. (2009) and
Shakya and Griffin (2010) for naphthalene photooxidation. Indeed, Shakya and Griffin (2010) proposed yields in the range of 0.08-0.16, while Chan et al. (2009) determined yields to be 0.19-0.74. As discussed by Chen et al. (2016), differences of SOA yields could be attributed to different chamber conditions such as light intensity, NOx levels, OH radicals, and organic mass loading. In the case of aromatic chemistry, it has been shown that photolysis processes play a major role in the loss of carbonyl products (Wang et al., 2006; Clifford et al., 2011). For instance, aromatic aldehydes can be photodissociated, leading to the formation of more oxidized compounds, which could further contribute to SOA formation (Wang et al., 2006). Furthermore, Warren et al. (2010) have reported the importance of light intensity in SOA formation from the photooxidation of monoaromatics. Therefore, in addition to the high concentration of PAHs, higher photolysis rates could also explain the subsequent SOA yields determined in this work.

Under high-NOx conditions, the photolysis of HONO generated relatively high concentrations of OH (i.e. \(8.4 \times 10^6\) molecule cm\(^{-3}\)), leading to rapid acenaphthene and acenaphthylene consumption. Under low-NOx conditions, aerosol growth was also observed immediately after the lights were turned on. Nevertheless, the PAH consumption appeared to be slower than under high-NOx conditions, owing to the relatively low concentration of OH radicals produced by H\(_2\)O\(_2\) photolysis. Figure S1 presents the time-dependent growth curves (i.e. the mass of organic aerosol generated, noted as \(\Delta Mo\), as a function of reacted PAH, \(\Delta HC\)) for acenaphthene and acenaphthylene reactions with OH radicals under high- and low-NOx conditions. For all experiments, a constant increase of aerosol mass was observed and reached its maximum when acenaphthylene or acenaphthene was totally consumed. No clear difference was found between the high- and low-NOx conditions, indicating that compounds produced during the photooxidation of PAHs likely participate in SOA growth. Similar trends have been observed previously during the photooxidation of biogenic and anthropogenic precursors (Ng et al., 2006; Chan et al., 2009). The fact that aerosol growth stopped when the precursor was consumed indicates that the first oxidation reaction is the rate-determining step in SOA formation. In this case, the primary products could directly condense and participate in SOA formation, and therefore contribute to aerosol growth.

4. Conclusion

In this work, both gas and particle phase products from the OH oxidation of acenaphthylene and acenaphthene were characterized using two on-line techniques (i.e. PTR-TOF-MS and...
The impact of NO$_x$ on product distributions and SOA formation was explored for these compounds for the first time. Based on these results, extended mechanisms for the gas-phase oxidation of acenaphthylene and acenaphthene are proposed suggesting additional pathways and previously unidentified compounds under the conditions used in this work. The OH oxidation of acenaphthylene under high- and low-NO$_x$ conditions is mainly governed by OH addition. The significant production of products such as acenaphthenequinone or 1,8-naphthalic anhydride suggests preferential OH addition to the unsaturated cyclopenta-fused ring. On the contrary, acenaphthene oxidation begins with OH radical addition to the aromatic ring followed by the reaction of the resultant OH-adduct with O$_2$ and further ring cleavages in the presence of NO$_x$. H-atom abstraction from the saturated cyclopenta-fused ring appears, however, not to be negligible as stipulated in previous work, especially in the absence of NO$_x$. Indeed, products from the H-atom abstraction pathway, such as acenaphthenone, exhibit a higher importance in the experiments performed in absence of NO$_x$. In addition, secondary chemistry was identified in all experiments and is now proposed in the extended mechanisms. The oxidation of acenaphthene and acenaphthylene formed SOA in large yields and suggest that the reaction of OH radicals with PAHs could contribute to anthropogenic SOA formation. It is shown that SOA growth is relatively linear and completes when the precursor hydrocarbon is consumed. Such observations illustrate that SOA formation is mainly due to the first generation products proceeding through gas-to-particle conversion processes. SOA formation yields are higher under low-NO$_x$ conditions than under high-NO$_x$ conditions as previously reported for the photooxidation of other aromatic compounds. Further experimental works on SOA formation from the photooxidation of acenaphthylene and acenaphthene is, however, needed to better evaluate the impact of light intensity and NO$_x$ mixing ratio on the different oxidation pathways.
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ACKNOWLEDGEMENTS

Research at University College of Cork was supported by the EU-FP7 ‘European Simulation Chambers for Investigating Atmospheric Processes’ (EUROCHAMP-2, grant number 228335). The authors wish to thank the French Agency for Environment and Energy Management (ADEME) and the Aquitaine Region for their financial support.
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Figure 1. Temporal profiles of major ions identified by PTR-TOF-MS from the gas-phase oxidation ofacenaphthylene initiated by OH radicals (full and open markers represent experiments performed under high- (a) and low-NO\textsubscript{x} (b) conditions, respectively).
Figure 2. ATOFMS average mass spectra (positive ion mode) using a laser pulse of 0.2 mJ for SOA formed from the photooxidation of acenaphthylene under high- (red) and low-NO_x (green) conditions.
Figure 3. Main gas-phase compounds identified during the photooxidation of acenaphthene using PTR-TOF-MS (full and open markers represent experiments performed under high- (a) and low-NOx (b) conditions, respectively).
Figure 4. ATOFMS average mass spectra (positive ion mode) using a laser pulse of 0.2 mJ for SOA formed from the photooxidation of acenaphthene under high- (red) and low-NOx (green) conditions.
Scheme 1. Proposed mechanism for the photooxidation of acenaphthylene (green and red boxes represent the favored products formed under low- and high-NO$_x$ conditions respectively).
Scheme 2. Proposed mechanism for the photooxidation of acenaphthene (green and red boxes represent the favored products formed under low- and high-NOx conditions respectively).
HIGHLIGHTS

Identification of competitive pathways for the OH-initiated oxidation of Acenaphthene

Formation of ring-opening products favored from the OH oxidation of Acenaphthylene

Impact of NOx on product distributions and SOA formation have been observed

Proposition of identified secondary chemistry in the extended mechanisms

SOA yields in the range 46-68% have been measured