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Ozonolysis of surface-adsorbed methoxyphenols: kinetics of aromatic ring cleavage vs. alkene side-chain oxidation

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Abstract. Lignin pyrolysis products, which include a variety of substituted methoxyphenols, constitute a major component of organics released by biomass combustion, and may play a central role in the formation of atmospheric brown carbon. Understanding the atmospheric fate of these compounds upon exposure to trace gases is therefore critical to predicting the chemical and physical properties of biomass burning aerosol. We used diffuse reflectance infrared spectroscopy to monitor the heterogeneous ozonolysis of 4propylguaiacol, eugenol, and isoeugenol adsorbed on NaCl and α -Al₂O₃ substrates. Adsorption of gaseous methoxyphenols onto these substrates produced near-monolayer surface concentrations of 3×10^{18} molecules m⁻². The subsequent dark heterogeneous ozonolysis of adsorbed 4-propylguaiacol cleaved the aromatic ring between the methoxy and phenol groups with the product conclusively identified by GC-MS and ¹H-NMR. Kinetic analysis of eugenol and isoeugenol dark ozonolysis also suggested the formation of ring-cleaved products, although ozonolysis of the unsaturated substituent groups forming carboxylic acids and aldehydes was an order of magnitude faster. Average uptake coefficients for NaCladsorbed methoxyphenols were $\gamma = 2.3 (\pm 0.8) \times 10^{-7}$ and $2(\pm 1) \times 10^{-6}$ for ozonolysis of the aromatic ring and the unsaturated side chain, respectively, and reactions on α -Al₂O₃ were approximately two times slower. UV-visible radiation ($\lambda > 300$ nm) enhanced eugenol ozonolysis of the aromatic ring by a factor of $4(\pm 1)$ but had no effect on ozonolysis of the alkene side chain.

1 Introduction

Combustion of biomaterials – including natural and anthropogenic forest and grass fires as well as domestic burning for heating and cooking - represents a major tropospheric source of organic and carbonaceous particulates (e.g., Crutzen and Andreae, 1990; Seiler and Crutzen, 1980; Kasischke and Penner, 2004). Microscopic analysis of biomass burning aerosol reveals complex and diverse chemical composition and structure, including soot, tar balls, internally mixed organic-potassium salt particles, and mineral dust containing organic coatings (Posfai et al., 2003; Li et al., 2003, 2010). Volatile and semivolatile organic compounds (VOC and SVOC) are also emitted by biomass burning and may adsorb onto extant airborne particles, further contributing to the organic composition of tropospheric aerosols (Falkovich and Rudich, 2001). The chemical speciation of biomass aerosols and accompanying SVOCs depend on fuel source (e.g., grasses, softwood and hardwood species) as well as combustion conditions (Kjallstrand et al., 1998; Oros and Simoneit, 2001a, b; Reid et al., 2005; Janhall et al., 2010; Zhang et al., 2013).

Lignin is a structural biological polymer synthesized from monolignols (paracoumaryl alcohol, coniferyl alcohol, and sinapyl alcohol) accounting for 25–35 % of the dry mass of softwood species and 18–25 % of hardwoods (Wool and Sun, 2005). Thus, lignin pyrolysis products constitute major emissions from forest fires and residential heating based on wood combustion. Field studies have detected an array of lignin pyrolysis products in ambient air, especially in urban areas with significant residential wood heating (Hawthorne et al., 1989, 1992; Nolte et al., 2001; Bari et al., 2009) and downwind of forest fires (Hoffer et al., 2006). Shakya et al. (2011) also detected lignin polymers in Houston aerosols attributed to directly emitted biological aerosols, which have estimated annual emissions of 50 Tg (Kiehl and Rodhe, 1995). Field and laboratory measurements suggest that lignin pyrolysis



Fig. 1. Prevalent lignin pyrolysis products with vapor pressures listed in parentheses.

products undergo relatively efficient photooxidation (Hoffer et al., 2006), the products of which may contribute to atmospheric humic-like substances (HULIS) (Gelencser et al., 2003; Hoffer et al., 2004); the extent and mechanisms of this chemistry, however, remain uncertain. Important daytime reactions could include OH-initiated chemistry, ozonolysis, and photoenhanced oxidation.

Dozens of compounds classified as lignin pyrolysis products have been identified in biomass burning emissions (Oros and Simoneit, 2001a, b; Schauer et al., 2001). Given the structures of the monolignols that comprise lignin, it is not surprising that phenols, methoxyphenols, and dimethoxyphenols are prevalent motifs in lignin pyrolysis products. Figure 1 displays 12 lignin pyrolysis products along with their vapor pressures, listed in parentheses, which fall in the range of semivolatile compounds. Catechol and guaiacol are of particular interest because the ozonolysis of these compounds results in efficient secondary organic aerosol (SOA) formation (Coeur-Tourneur et al., 2009, 2010; Ofner et al., 2010, 2011). For these systems, ring-cleaved products, which have lower volatility as a result of additional polar functional groups, contributed to SOA growth and likely play a role in SOA nucleation.

Several recent studies have focused on heterogeneous reactions of surface-adsorbed methoxyphenols with O_3 and/or NO_3 (Knopf et al., 2011; Net et al., 2010a, b, 2011; Liu et al., 2012). For instance, Net et al. (2011) investigated the heterogeneous ozonolysis of several methoxyphenols adsorbed on silica. Many methoxyphenols (i.e., vanillic acid, syringic acid, acetovanillone, and acetosyringone) did not react; two aldehydes (vanillin and syringaldehyde) were oxidized to carboxylic acids, while only coniferyl alcohol, the only methoxyphenol studied containing an alkene side group, reacted via a 1,3-cycloaddition mechanism. No ring-cleaved products were observed. In a separate study, ozonolysis of silica-adsorbed coniferyl aldehyde produced glycolic acid, oxalic acid, vanillin, vanillic acid, and 3,4-dihydroxybenzoic acid (Net et al., 2010a). Knopf et al. (2011) measured a reactive uptake coefficient of $\gamma = 8.3 \times 10^{-5}$ for O₃ on thin film nitroguaiacol, which was more reactive than levoglucosan and abietic acid; reaction products were not identified in this study.

Here we evaluate the relative importance of ozone 1,3cycloaddition to alkene side chains vs. across methoxyphenols by identifying products and measuring reaction kinetics for the heterogeneous ozonolysis of eugenol, isoeugenol, and 4-propylguaiacol (Fig. 1 bottom row). Given the semivolatile vapor pressures of these compounds, the current experimental samples were prepared by adsorbing gaseous methoxyphenols onto inorganic tropospheric aerosol substrates that represent biogenic salt and mineral dust aerosol present in biomass burning emissions. This builds on our prior work investigating the heterogeneous ozonolysis of NaCl- and α -Al₂O₃-adsorbed catechol (Woodill et al., 2013). Infrared spectroscopy, gas chromatographymass spectroscopy, and nuclear magnetic resonance spectroscopy are combined to address the following question: how prevalent are ring-cleaved products? To evaluate the kinetics of these systems under atmospheric conditions, the impact of relative humidity and solar radiation on the ozonolysis of eugenol was also investigated.

2 Methods

2.1 Infrared spectroscopy

We used infrared spectroscopy to characterize gaseous and surface-adsorbed methoxyphenols. Eugenol, isoeugenol, and 4-propylguaiacol (Aldrich, 98%), which all have vapor pressures ca. 1 Pa, were used without further purification. A continuous flow of dry, purified air bubbled through liquid methoxyphenols entraining these semivolatile organic compounds in the gas phase. Gas phase spectra were recorded by directing this flow through a 2 L transmission gas cell, which uses White cell optics to achieve a 10 m pathlength (Thermo Antaris), housed in a Thermo Nicolet 6700 FTIR equipped with an MCT-A detector.

Diffuse reflectance (DRIFTS) and attenuated total reflectance infrared spectroscopy (ATR-FTIR) monitored the adsorption of gaseous methoxyphenols on NaCl or α -Al₂O₃ substrates and the subsequent heterogeneous ozonolysis of these samples. Ball-milled NaCl (Aldrich, 99.999%) had a surface area of $0.12 \text{ m}^2 \text{ g}^{-1}$; α -Al₂O₃ (Alfa Aesar, 99.98%) was used without ball milling and had a surface area of $14 \text{ m}^2 \text{g}^{-1}$. For DRIFTS, inorganic powders were packed into a reaction chamber (Harrick Scientific, Praying Mantis) and heated to 300 °C under the continuous flow of dry air for at least 60 min. After cooling to 25 °C, a background spectrum was recorded at 4 cm^{-1} resolution averaging 256 scans. For ATR-FTIR, a methanol/ α -Al₂O₃ slurry was evenly spread over the ZnSe ATR crystal, which was then placed in an oven at 100 °C for 90 min to remove all methanol. We then attached the ATR flow cell (Pike Technologies, HATR) to the continuous flow manifold and recorded a background after purging the system for 60 min.

The relative humidity (RH) was controlled by adjusting the relative flow rates of dry and wet streams of purified air using two mass flow controllers. RH was measured upstream of the DRIFTS reaction chamber using a Vaisala HMT338 gauge. It should be noted that even under "dry" conditions (referred to as 0% hereafter) the DRIFTS substrates contain residual surface-adsorbed water (Goshal et al., 2004). The dry flow could be diverted through the organic bubbler, thereby exposing the NaCl or α -Al₂O₃ substrates to gaseous methoxyphenols. Spectra recorded continuously with an approximate averaging time of three minutes monitored adsorption of gaseous methoxyphenols, and saturated surface coverage was typically achieved in 90 min or less. After isolating the methoxyphenol bubbler, coated samples were purged with air for a minimum of 30 min with negligible loss of adsorbed organics. Heterogeneous ozonolysis was then investigated using a photolytic ozone generator in the dry air stream, which was connected to the reaction chamber by a polypropylene tube to remove any OH radicals (Woodill et al., 2013). Ozone concentrations were constant during the course of each experiment, and separate experiments were conducted with concentrations ranging from 4×10^{12} to 25×10^{12} molecules cm⁻³ (150 to 990 parts per billion - ppb) as measured by a Jelight 308 ozone monitor. The majority of reactions were conducted under dark conditions; however, a series of eugenol ozonolysis experiments used a xenon lamp solar simulator to test for photoenhanced chemistry. Infrared and UV ($\lambda < 300$ nm) radiation was filtered out of the xenon lamp output using water and borosilicate filters, respectively. The solar simulator spectral irradiance (Supplement, Fig. S1) was determined by correlating the integrated power ($\lambda < 800 \text{ nm}$), measured with a Newport Oriel power meter, and the relative intensity as a function of wavelength, which was measured using an Ocean Optics spectrometer (HR2000). Diffuse reflectance UV-visible spectra of adsorbed organics were recorded in a similar manner as described for DRIFTS using a Thermo Evolution 300 spectrometer.

Quantum chemical calculations of eugenol, isoeugenol, and 4-propylguaiacol identified vibrational normal modes associated with infrared peaks. All calculations were performed using Gaussian 09 (Frisch et al., 2009). Ground-state minima were optimized at the B3LYP/6-31G(d,p) level of theory, and harmonic frequency calculations were scaled by 0.977 to account for neglected anharmonic terms, which allowed for comparison to experimental spectra (Irikura et al., 2005).

2.2 Extraction and analysis of adsorbed organics

To determine adsorbed organic surface concentrations, methoxyphenols were extracted from NaCl DRIFTS samples in 2–3 mL of acetonitrile. (α -Al₂O₃ samples were not extracted due to the monodentate binding of methoxyphenols, discussed below.) Extracted slurries were sonicated for 60 min then filtered using a 0.2 µm syringe filter. Many samples were derivatized with N,O-bis(trimethylsilyl)-trifluoroacetamide (BSTFA, Fluka 99 %) to reduce polarity by silating hydroxyl and carboxylic acid groups. Extracted organics were analyzed on an Agilent Technologies GC-MS (6890N-GC/5973-MS) with a Supelco SLB-5ms column and the GC oven temperature ramped from 70 to 200 °C.

To analyze reaction products using GC-MS and ¹H-NMR, \sim 1 g samples of NaCl were packed into a fritted glass stem attached to a gas manifold. Adsorption of methoxyphenols and their subsequent ozonolysis was executed in a manner similar to that discussed above, except coating and reaction times were increased tenfold. For GC-MS, organics were extracted and sonicated in acetonitrile, filtered, and then derivatized using BSTFA. For NMR, reacted samples were extracted using deuterated acetone (Aldrich, 99%, 0.03% TMS) and then analyzed without filtering using a 200 MHz Bruker NMR, averaging 30 000 scans.

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Fig. 2. (a) Experimental and (b) computational spectra of gaseous eugenol. DRIFTS spectra of eugenol adsorbed on (c) NaCl and (d) α -Al₂O₃.

3 Results and discussion

3.1 Adsorption of methoxyphenols on NaCl and α -Al₂O₃

Comparing experimental and computational spectra of gaseous methoxyphenols facilitates the assignment of vibrational normal modes to the observed infrared peaks (Table 1). Optimization and frequency calculations at the B3LYP/6-31G(d,p) predicted harmonic vibrational frequencies, which were scaled by 0.977 to account for neglected anharmonic terms, and infrared intensities. Figure 2 displays the excellent agreement between the (a) experimental and (b) computational spectra of gaseous eugenol. Notable vibrational modes relevant to the ensuing discussion include phenol (OH) stretching and bending vibrations at 3597 and 1373 cm⁻¹, alkene side-chain CH and C=C stretching vibrations at 3089 and 1645 cm⁻¹, and an intense aromatic ring vibration centered at 1517 cm^{-1} . Excellent agreement was also observed for isoeugenol and 4-propylguaiacol.

Figure 2 also shows DRIFTS spectra for eugenol adsorbed on (c) NaCl and (d) α -Al₂O₃. NaCl-adsorbed eugenol retained most of the spectral features seen for gaseous eugenol with only minor frequency shifts. The most notable difference was the broadened phenol OH stretching vibration, which exhibited two peaks centered at 3189 and 3367 cm⁻¹. The phenol bending vibration at 1363 cm⁻¹ also displayed significant broadening. These features indicate that eugenol physisorbed to the NaCl surface, similar to the adsorption of catechol on NaCl (Woodill et al., 2013). Based on this comparison, we infer that a dominant binding interaction on the NaCl (001) surface involves donation of an electron lone pair on the methoxy oxygen atom to a Na⁺ cation and an interaction analogous to a weak hydrogen bond from the phenol to an adjacent Cl⁻ anion (e.g., Reaction Scheme (R1)). Defect and edge sites, which are prevalent in the employed powder samples, also provide favorable binding sites.



 α -Al₂O₃ adsorbed eugenol exhibits features consistent with monodentate binding: absent phenol stretching and bending vibrations, and a negative peak at 3734 cm⁻¹ associated with the loss of alumina surface hydroxyl groups upon eugenol adsorption. Our previous work noted analogous binding for catechol on α -Al₂O₃, which was identified as binuclear bidentate catecholate (Woodill et al., 2013). As characterized in Reaction Scheme (R1), the alkene side chain likely dangles off each organic-coated substrate, and vibrational frequencies associated with this group were within 8 cm⁻¹ of gaseous eugenol modes. Isoeugenol and 4-propylguaiacol spectra also indicated physisorption on NaCl and monodentate complexes on α -Al₂O₃.

A powerful aspect of DRIFTS is that the samples are large enough ($\sim 100 \,\mathrm{mg}$ inorganic substrates) that the adsorbed organics may be solvent extracted for analysis by GC-MS, allowing for identification and quantification by comparison to standards. Several NaCl DRIFTS samples were sonicated in acetonitrile for 60 min, filtered, and then derivatized with BSTFA for GC-MS analysis. Comparison to eugenol standards quantified the extracted concentration, which combined with the BET surface area determined the NaCl-adsorbed eugenol concentration to be $3(\pm 1) \times 10^{18}$ molecules m⁻². This value is similar to the $3.5 \times 10^{18} \, \text{molecules} \, \text{m}^{-2} \, \text{ surface concentration measured}$ for catechol adsorbed on NaCl (Woodill et al., 2013) and is consistent with the similar physisorption binding proposed above. Since it is difficult to estimate the effective area of an individual adsorbed methoxyphenol, we cannot accurately calculate the fraction of a monolayer. Nonetheless, based on our prior work with surface-adsorbed catechol, we believe these surface concentrations correspond to near monolavers (Woodill et al., 2013). For α -Al₂O₃, the magnitude of the negative surface hydroxyl peak at 3734 cm^{-1} can be compared to the maximum loss of this signal at high humidity (Woodill et al., 2013), implying a surface concentration of $3(\pm 2) \times 10^{18}$ molecules m⁻². We expect all methoxyphenols studied here to have similar near-monolayer surface concentrations for both NaCl and α -Al₂O₃ substrates.

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	Eugenol			Ozonolysis products	
Normal mode	Gaseous	NaCl	α -Al ₂ O ₃	NaCl	α -Al ₂ O ₃
Al-O-H, surface hydroxyl			3734*		3734*
Phenol OH stretch	3597	3189, 3367	_		
Alkene CH stretch	3089	3081	3081		
Aromatic CH stretch	3013	3009	3009		
Carbonyl C=O stretch				1736	1726
Carboxylate stretch					1620
Alkene C=C stretch	1645	1641	1641		
Aromatic ring stretch	1517	1518	1518		
Methoxy CH bend	1471	1466	1466		
Methoxy/aromatic CH bend	1439	1431	1435		
Phenol OH bend	1373	1363	_		
Aromatic ring breathing	1273	1269	v. broad		

Table 1. Vibrational assignments (cm^{-1}) for eugenol and its ozonolysis products.

* Negative peaks indicating loss of surface hydroxyl group upon eugenol adsorption.



Fig. 3. DRIFTS spectra of 4-propylguaiacol adsorbed on (**a**) NaCl exposed to 551 ppb O₃ at $t = 0 \min$ (black) and 500 min (blue), and (**b**) α -Al₂O₃ exposed to 420 ppb O₃ at $t = 0 \min$ (black) and 500 min (red).

3.2 Ozonolysis of 4-propylguaiacol

We studied the heterogeneous ozonolysis of NaCl- and α -Al₂O₃-adsorbed 4-propylguaiacol at 0 % RH with O₃ concentrations ranging from 150 to 990 ppb. Figure 3 shows t = 0 (black) and t = 500 (color) minute reaction spectra on (a) NaCl at 551 ppb O₃ and on (b) α -Al₂O₃ at 420 ppb O₃. Dominant spectral features associated with condensed-phase products include peaks at 1745 and 1722 cm⁻¹ assigned to carboxylic acid or aldehyde carbonyl vibrations (Socrates, 2001). Loss of aromatic CH and ring stretching vibrations at 3056 and 1518 cm⁻¹, respectively, imply that ozonolysis cleaved the aromatic ring. α -Al₂O₃-adsorbed products

also exhibited a strong peak at 1610 cm^{-1} , which was not as significant for NaCl-adsorbed products, consistent with carboxylate vibrations (Socrates, 2001). Similar substrate differences were seen for muconic acid formed from the heterogeneous ozonolysis of surface-adsorbed catechol (Wood-ill et al., 2013).

To identify reaction products, we extracted the products from reacted NaCl-adsorbed 4-propylguaiacol samples in acetonitrile for analysis using GC-MS. After derivatization with BSTFA, the total ion current chromatogram displayed two main peaks corresponding to unreacted 4propylguaiacol and a single product at 12.7 and 13.7 min, respectively (Fig. 4a). Figure 4b displays the mass spectrum of the 13.7 min product peak. Although not prominent in the mass spectrum, we identify the parent ion for this product as m/z = 270 (see extracted ion chromatogram in Fig. 4a). Without BSTFA derivatization, the parent ion m/z was 198, indicating the presence of a single alcohol or acid group. Major fragmentation peaks in the mass spectrum of Fig. 4b include 255 (M - CH₃), 211 (M - CO₂CH₃), 153 (M -CO₂SiMe₃), and 73 (SiMe₃), which are consistent with the ring-cleaved product included in Reaction (R2).



¹H-NMR analysis of NaCl-adsorbed 4-propylguaiacol + O₃ reaction samples extracted in d_6 -acetone confirmed this primary product structure (Supplement, Fig. S2; $\delta_{\rm H}$ (200 MHz, d_6 -acetone, Me₄Si) 0.9 (3H, *t*, CH₃), 1.5 (2H, *m*, CH₂CH₃), 2.4 (2H, *t*, CH₂), 3.6 (3H, *s*, OCH₃), 5.7 (1H, *s*, gem-CO₂CH₃), 5.9 (1H, *d*, gem-CO₂H), 7.1 (1H, *d*, gem-C=C)). In total, both the GC-MS and NMR data indicate that the



Fig. 4. (a) Total ion current gas chromatogram (black) and m/z = 270 extracted ion chromatogram (red) of products extracted from 4-propylguaiacol ozonolysis sample. (b) Mass spectrum of 13.7 min peak assigned as ring-cleaved product.

DRIFTS reaction spectra discussed above correlate with the ozonolysis ring cleavage depicted in Reaction (R2).

In contrast to our results, cleavage of the aromatic ring was not observed in previous investigations of the heterogeneous ozonolysis of surface-adsorbed methoxyphenols (e.g., Net et al., 2011). Substrate effects and sample preparation might explain this discrepancy. These prior studies coated silica particles using a liquid-solid adsorption method where the organic and substrate were mixed in a solvent, which was then evaporated off. Here, we used semivolatile methoxyphenols that could be coated on NaCl and α -Al₂O₃ substrates via gas-solid adsorption. We previously noted that under dry conditions, thin film catechol did not react with ozone (Barnum et al., 2011), while NaCl- and α -Al₂O₃-adsorbed catechol readily reacted with ozone (Woodill et al., 2012), indicating that these substrates can greatly enhance this ozonolysis chemistry. Attempts to coat silica with catechol and various methoxyphenols via our DRIFTS technique resulted in reversible adsorption, where purging with dry air dramatically decreased the organic surface concentration. This suggests that substrate–methoxyphenol interactions are considerably weaker for silica compared to NaCl and α -Al₂O₃, which in turn may impact the 1,3-cycloaddition of ozone to these adsorbed aromatic rings.

To evaluate the reaction kinetics for the ozonolysis of surface-adsorbed 4-propylguaiacol, DRIFTS reaction spectra were recorded continuously with averaging times of approximately 3 min. Integration of the key infrared spectral features described above as a function of time facilitated the kinetic analysis of 4-propylguaiacol consumption and product formation. Figure 5a reports one such analysis for NaCl-adsorbed 4-propylguaiacol exposed to 665 ppb O₃, showing the normalized integrated area of the 3058 cm⁻¹ peak (i.e., 4-propylguaiacol) decaying concurrently with the growth of the 1745 cm⁻¹ product peak. A general rate equation for this reaction can be written as

$$Rate = k[O_3]^x [4PG]^y.$$
(1)

Due to the continuous flow of carrier gas through the DRIFTS reaction chamber, the ozone concentration is assumed to be constant during the course of an experiment. Therefore, the experimental rate can be approximated as

$$Rate = k_{exp} [4PG]^{y}.$$
 (2)

An exponential decay function fit the loss of aromatic CH stretching vibrations, indicating the reaction was first order with respect to 4-propylguaiacol with a pseudo-first-order rate constant of $k_{exp} = 1.95 \times 10^{-4} \text{ s}^{-1}$. A first-order product-integrated rate expression fit the carbonyl formation kinetics with a rate constant $k_{exp} = 1.50 \times 10^{-4} \text{ s}^{-1}$. The similarity of these rate constants supports a direct correspondence between 4-propylguaiacol consumption and carbonyl product formation. Figure 5b collates the experimental rate constants for all experiments plotted against ozone concentration. Linear fits to this data are included as solid lines and correspond to an Eley–Rideal-type mechanism, where gaseous ozone would react directly with adsorbed 4-propylguaiacol,

$$O_3(g) + 4-PG(ads) \rightarrow products,$$
 (R3)

$$k_{\exp} = k_{\mathrm{O}_3} \cdot [\mathrm{O}_3]. \tag{3}$$

Linear best-fit parameters find $k_{O_3} = 1.0 \times 10^{-17} \text{ cm}^3 \text{ mol-}$ ecules⁻¹ s⁻¹ and 7.0 × 10⁻¹⁹ cm³ molecules⁻¹ s⁻¹ for 4propylguaiacol adsorbed on NaCl and α -Al₂O₃, respectively.

The heterogeneous ozonolysis of many similar surfaceadsorbed organics follow a Langmuir–Hinshelwood mechanism (Dubowski et al., 2004; Lu et al., 2011; Kwamena et al., 2004, 2007; Zhou et al., 2012; Rosen et al., 2008) in which gaseous ozone first adsorbs to the surface before reacting with adsorbed 4-propylguaiacol:

$$O_3(g) \leftrightarrow O_3(ads),$$
 (R4)



Fig. 5. (a) Kinetics integration of 3058 cm^{-1} peak (4propylguaiacol aromatic CH stretch) and 1745 cm^{-1} peak (product carbonyl vibration). Exponential fits find $k_{exp} = 1.95 \times 10^{-4} \text{ s}^{-1}$ and $1.50 \times 10^{-4} \text{ s}^{-1}$ for 3058 and 1745 cm^{-1} peaks, respectively. (b) Plot of k_{exp} vs. ozone concentration for 4-propylguaiacol adsorbed on NaCl (circles) and α -Al₂O₃ (triangles). Solid line fits correspond to Eley–Rideal-type mechanism; dashed line fits correspond to Langmuir–Hinshelwood mechanism.

$$O_3(ads) + 4-PG(ads) \rightarrow products.$$
 (R5)

For the Langmuir–Hinshelwood mechanism, the experimental rate constant would be

$$k_{\exp} = \frac{k_{\max} \cdot K_{O_3} \cdot [O_3]}{1 + K_{O_3} \cdot [O_3]},$$
(4)

where K_{O_3} is the ozone adsorption equilibrium constant for Reaction (R4) and k_{max} is the maximum rate constant for Reaction (R5). Langmuir–Hinshelwood fits, which are included as dashed lines in Fig. 5b, are indistinguishable from the linear fits at the relatively low ozone concentrations used in this study, preventing us from conclusively identifying the reaction mechanism. (Langmuir–Hinshelwood bestfit parameters in Fig. 5b are $k_{max} = 0.0102 \text{ s}^{-1}$ and $K_{O_3} =$ $1.0 \times 10^{-15} \text{ cm}^3$ for NaCl and $k_{max} = 0.0083 \text{ s}^{-1}$ and $K_{O_3} =$ $8.4 \times 10^{-17} \text{ cm}^3$ for α -Al₂O₃. Since our data do not show signs of approaching saturation, these parameters are poorly constrained, and only the values of k_{max} times K_{O_3} are known with certainty.) Higher ozone concentrations, which would be required to distinguish between these two mechanisms, are unfortunately not feasible with our current experimental setup.

Although future work would be required to conclusively determine which heterogeneous mechanism is operative for the ozonolysis of adsorbed 4-propylguaiacol, this kinetic analysis allows us to estimate the atmospheric lifetime of these systems. Equations (5) and (6) calculate atmospheric lifetimes assuming the Eley–Rideal and Langmuir– Hinshelwood mechanisms, respectively.

$$\tau = \frac{1}{k_{\mathrm{O}_3}[\mathrm{O}_3]} \tag{5}$$

$$\tau = \frac{1 + K_{\rm O_3}[{\rm O}_3]}{k_{\rm max} K_{\rm O_3}[{\rm O}_3]} \tag{6}$$

Assuming a tropospheric ozone concentration of 40 ppb (Vingarzan, 2004), the atmospheric lifetime for 4-propylguaiacol adsorbed on NaCl is estimated to be 27.7 to 28.2 h, depending on the mechanism, vs. 17 days when adsorbed on α -Al₂O₃.

3.3 Eugenol and isoeugenol + O₃

After conclusively identifying ozone-induced ring cleavage for 4-propylguaiacol, we used identical methods and conditions to investigate the heterogeneous ozonolysis of surface-adsorbed eugenol and isoeugenol. Figure 6 shows initial (black) and final (red) DRIFTS reaction spectra for (a) NaCl-eugenol exposed to 424 ppb O₃ and (b) NaClisoeugenol + 180 ppb O₃. Both systems exhibit the loss of alkene side-chain spectral features and the formation of a dominant product carbonyl vibration near 1730 cm^{-1} . Also evident in these product spectra is the diminishing intensity of the aromatic ring vibration at 1518 cm^{-1} , indicating the loss of some aromatic ring structures.

To monitor the ozonolysis kinetics of the alkene side chain vs. across the methoxyphenol, we integrated vibrational features unique to these two sites of attack. Figure 7 shows one set of integrations for the reaction of NaCl-adsorbed isoeugenol exposed to 180 ppb O₃. The alkene side-chain spectral feature at 3064 cm⁻¹ decayed rapidly with a rate constant of $k_1 = 7.50 \times 10^{-4} \text{ s}^{-1}$. Features associated with the methoxyphenol aromatic structure, including vibrations at 1518 and 1380 cm⁻¹, decayed much more slowly with $k_2 = 0.40 \times 10^{-4} \text{ s}^{-1}$. Growth of the product carbonyl feature at 1726 cm^{-1} could not be successfully fit with a single exponential function; a double-exponential product growth function did fit these data with $k_1 = 4.4 \times 10^{-4} \text{ min}^{-1}$ and $k_2 = 0.30 \times 10^{-4} \,\mathrm{s}^{-1}$, in good agreement with the loss of the alkene side-chain and methoxyphenol spectral features, respectively. The amplitude of the second, slower exponential function was twice that of the first, more rapid function. These amplitudes are consistent with the fact that ozonolysis



Fig. 6. DRIFTS spectra of (a) eugenol adsorbed on NaCl exposed to 424 ppb O₃ at $t = 0 \min$ (black) and 340 min (red), and (b) isoeugenol adsorbed on NaCl exposed to 180 ppb O₃ at $t = 0 \min$ (black) and 800 min (red).

of the alkene side chain is expected to form one condensedphase aldehyde or carboxylic acid functional group, while slower ring cleavage should form two condensed-phase carbonyl functional groups. Reaction Scheme (R6) depicts the condensed-phase products consistent with this kinetic analysis, and an analogous mechanism could be constructed for the ozonolysis of eugenol. It should be noted that the rate constant for cleavage of the aromatic ring, k_2 , actually corresponds to ozonolysis of vanillic acid and vanillin and not isoeugenol since the data were fit after t = 100 min, by which time essentially all the alkene side chains were oxidized.



GC-MS analysis of NaCl-adsorbed isoeugenol exposed to ozone revealed unreacted isoeugenol and three major product peaks (see Supplement, Fig. S3). The largest product peak, which accounted for 54 % of the total product peak areas, was identified with confidence by comparison to the NIST library as vanillic acid (BSFTA derivatized m/z: 312 (M⁺), 297 (M–CH₃), 267 (M–3 × CH₃)). Vanillin accounted

for 29 % of the integrated product peak areas (BSFTA derivatized m/z: 224 (M⁺), 209 (M–CH₃), 194 (M–2×CH₃)), while the third peak, which accounted for 17%, could not be conclusively identified but shows fragmentation patterns consistent with a silated methoxyphenol species. Although several minor peaks appeared in the total ion chromatogram, no peak could be conclusively identified as ring-cleaved products. The total ion current chromatogram for extracted eugenol ozonolysis samples showed unreacted eugenol and five significant product peaks (Supplement, Fig. S4). Only two product peaks were conclusively identified by comparison to the NIST library: homovanillic acid (BSFTA derivatized m/z: 326 (M⁺), 209 (M-CO₂SiMe₃), 179 (M- $CO_2SiMe_3-2 \times CH_3$) and homovanillin (BSFTA derivatized m/z: 238 (M⁺), 209 (M–COH–2 × CH₃), 179). Two of the remaining peaks are tentatively assigned to artifacts resulting from the BSTFA derivatization of homovanillin (see Supplement), while the remaining peak could not be conclusively identified but is consistent with a methoxyphenol species. ¹H-NMR spectra of d_6 -acetone-extracted eugenol reaction mixtures were difficult to interpret since most of the product peaks overlapped with unreacted eugenol features. Two discernible product features are noteworthy: aromatic peaks between $\delta_{\rm H} = 6.5$ and 7.0 ppm and an aldehyde peak at $\delta = 9.7$ ppm, consistent with the formation of homovanillin and homovanillic acid. No evidence of higher molecular weight products was detected, although admittedly our GC-MS analysis is not ideally suited to their detection due to reversible dissociation during extraction (Jang et al., 2002). Prior investigations have identified gaseous formaldehyde for the ozonolysis of liquid eugenol (Sachdev et al., 1972), which is likely also produced in these systems.

Although GC-MS and NMR analysis did not conclusively identify ring-cleaved products for the ozonolysis of eugenol and isoeugenol, we believe the similarities with 4-propylguaiacol combined with the kinetic analysis discussed above provide strong evidence that ozone does attack the methoxyphenol of eugenol and isoeugenol, resulting in cleavage of the aromatic ring, albeit at rates an order of magnitude slower than ozonolysis of the alkene side chains. Ozonolysis of both the alkene side-chain and methoxyphenol features appeared to be first order with respect to ozone concentration. Estimates of the relative reaction rates and atmospheric lifetimes, assuming an Eley-Rideal mechanism, for all systems studied here are collated in Table 2. Two general trends emerge from the collective analysis of these results: (1) ozonolysis of surface-adsorbed methoxyphenols occurs at a rate approximately 10 times slower than ozonolysis of alkene side chains, and (2) substrate effects suggest these ozonolysis reactions are faster on NaCl vs. α -Al₂O₃. A comparison to catechol ozonolysis data from Woodill et al. (2013) also suggests that ring cleavage of ortho-diols is 4-10 times faster than ortho-methoxyphenols.

We also conducted a complementary set of experiments using an ATR-FTIR reaction cell to investigate the

Substrate	Organic	$k_{O_3}^a$	Relative Rate ^b	Atmospheric Lifetime ^c	γ
NaCl α -Al ₂ O ₃	catechol ^d catechol ^d	6.3×10^{-17} 1.5×10^{-17}	91 22	4.5 h 18 h	1.5×10^{-6} 3.5×10^{-7}
NaCl α -Al ₂ O ₃	4-propylguaiacol 4-propylguaiacol	$\begin{array}{c} 1.0 \times 10^{-17} \\ 7.0 \times 10^{-19} \end{array}$	14 1	28 h 17 days	2.3×10^{-7} 1.6×10^{-8}
NaCl NaCl α -Al ₂ O ₃ α -Al ₂ O ₃	eugenol, alkene eugenol, ring eugenol, alkene eugenol, ring	$\begin{array}{c} 3.6 \times 10^{-17} \\ 7.9 \times 10^{-18} \\ 5.2 \times 10^{-17} \\ 5.1 \times 10^{-18} \end{array}$	51 11 74 7	7.9 h 1.5 days 5.5 h 2.3 days	$8.3 \times 10^{-7} \\ 1.8 \times 10^{-7} \\ 1.2 \times 10^{-6} \\ 1.2 \times 10^{-7} \\ 1.2 \times 10^{-7$
NaCl NaCl α -Al ₂ O ₃ α -Al ₂ O ₃	isoeugenol, alkene isoeugenol, ring isoeugenol, alkene isoeugenol, ring	$\begin{array}{c} 1.3 \times 10^{-16} \\ 7.9 \times 10^{-18} \\ 2.7 \times 10^{-17} \\ 6.7 \times 10^{-18} \end{array}$	193 11 38 10	2.1 h 1.5 days 11 h 1.7 days	$3.1 \times 10^{-6} 1.8 \times 10^{-7} 6.2 \times 10^{-7} 1.6 \times 10^{-7}$

^a Assumes Eley-Rideal mechanism; cm³ molecules⁻¹ s⁻¹.

^b Estimated using k_{O_3} .

^c Atmospheric lifetimes calculated assuming 40 ppb tropospheric ozone concentration.

^d Catechol rates from data included in Woodill et al. (2013).

heterogeneous ozonolysis of α -Al₂O₃-adsorbed eugenol. A thin film of α -Al₂O₃ was first coated on a ZnSe ATR crystal, then exposed to gaseous eugenol, and finally exposed to 209 ppb O₃ at 0 % relative humidity. Control experiments exposing a clean ZnSe crystal to gaseous eugenol showed no significant condensed-phase organic features, indicating that ZnSe did not act as a reactive substrate in these experiments. Ozonolysis reaction spectra are included in the Supplement, Fig. S5. Inspection of the $t = 0 \min$ ATR spectrum reveals notable differences compared to DRIFTS spectra, with no negative α -Al₂O₃ surface hydroxyl peak at 3734 cm⁻¹, and the presence of OH stretching and bending vibrations at 3420 and $1373 \,\mathrm{cm}^{-1}$, respectively. These features suggest physisorption of eugenol rather than monodentate binding, which is likely due to differences in sample preparation. Subsequent ozonolysis produced spectral features consistent with DRIFTS reaction spectra, including the rapid loss of alkene side-chain peaks at 1639 and 3082 cm^{-1} , slower loss of aromatic features at 1518 and 3015 cm^{-1} , and the formation of carbonyl peak at 1734 cm⁻¹. Reaction rate constants were consistent with DRIFTS experiments.

Reactive uptake coefficients quantify heterogeneous reaction kinetics as the fraction of O₃-surface collisions that result in product formation relative to the total number of O₃surface collisions. Reactive uptake coefficients can be calculated from the ATR kinetic data by calibrating the FTIR spectral features assuming a eugenol surface concentration of 3×10^{18} molecules m⁻² at t = 0 min. The initial loss rate was then determined by linear fits to the first 20 min for the key spectral features discussed above. These loss rates were then compared to the frequency of total O₃-surface collisions, which was calculated using the kinetic theory of gases. Analyzing the rate of decay of the alkene side-chain spectral features yielded an initial reactive uptake coefficient of $\gamma = 1.2 \times 10^{-6}$. The heterogeneous ozonolysis of long-chain aliphatic alkenes, such as oleic acid, has been extensively studied. Uptake coefficients from these studies span a vast range, from 1.6×10^{-3} for oleic acid-generated particles (Morris et al., 2002; Hearn et al., 2005) to 8×10^{-4} for liquid oleic acid coatings (Moise and Rudich, 2002; Thornberry and Abbatt, 2004) down to 2.6×10^{-6} for oleic acid monolayers spread on aqueous droplets (Gonzalez-Labrada et al., 2007). Dubowski et al. (2004) studied the ozonolysis of C3 and C8 vinyl terminated self-assembled monolayers (SAMs) and measured initial reaction probabilities around $\gamma \sim 10^{-5}$ for ozone concentrations below 10^{13} molecules cm⁻³; uptake coefficients decreased with increasing ozone concentrations, consistent with a Langmuir-Hinshelwood mechanism. This wide range of uptake coefficients is likely the result of ozone diffusion into and reaction with the bulk of liquid coatings and particles. SAMs, on the other hand, necessarily only involve surface reactions, although experimental evidence suggests that ozone still has residence times on the order of 7 s (Dubowski et al., 2004). The surface-adsorbed organics employed in the present study, with near-monolayer surface concentrations, have uptake coefficients at the lower end of this range consistent with SAMs and oleic acid monolayers.

Combining the alkene side-chain uptake coefficient ($\gamma = 1.2 \times 10^{-6}$) with the relative rates implies that ozonolysis across the methoxyphenol has a reactive uptake coefficient of approximately 1×10^{-7} , while the ozonolysis of NaCl-adsorbed catechol is predicted to be an order of magnitude



Fig. 7. Reaction kinetics for the ozonolysis of NaCl-adsorbed isoeugenol at 180 ppb O_3 depicting (top) rapid consumption of the alkene side chain, (middle) slower decay of the aromatic ring, and (bottom) product carbonyl formation, which required a double-exponential function with rate constants consistent with competitive alkene side-chain vs. methoxyphenol ozonolysis.

faster with $\gamma = 1.6 \times 10^{-6}$ (Table 2). Knopf et al. (2011) measured $\gamma = 8.3 \times 10^{-5}$ for O₃ on thin film nitroguaiacol, almost three orders of magnitude faster than the current estimate. Barnum et al. (2012) found thin film crystalline catechol to have $\gamma = 5.6 \times 10^{-5}$ at 81 % relative humidity. These differences may be due in part to the diffusion of ozone several layers into the solid thin films allowing for reaction with subsurface compounds. For instance, frozen oleic acid films have uptake coefficients of $\gamma = 5.2 \times 10^{-5}$ to 7.1×10^{-4} , which are less than those of liquid oleic acid films due to limited reaction with the bulk (Moise and Rudich, 2002; Thornberry and Abbatt, 2004), but still more than an order of magnitude greater than for SAMs and monolayers. For the reaction of 4-nitroguaiacol, it is also possible that product channels other than ring cleavage are involved, and additional experiments identifying reaction products are required to evaluate this potential.

3.4 Impact of relative humidity

Given the importance of co-adsorbed water in atmospheric heterogeneous chemistry, we investigated the heterogeneous ozonolysis of eugenol as a function of relative humidity. DRIFTS samples were baked and then equilibrated at the selected relative humidity by adjusting the flow rates of dry and wet carrier gas. Once equilibrated, the dry flow was directed through the eugenol bubbler such that coating of the samples occurred at the selected relative humidity. Spectral features for both NaCl- and α -AL₂O₃-adsorbed eugenol at higher RH were identical to those discussed above for dry conditions, although the overall intensity decreased with increasing RH. This indicated that co-adsorbed water competed with eugenol for surface binding sites, resulting in lower eugenol surface concentrations at higher humidity. This effect was more significant for NaCl, where increasing the humidity to 21 % RH decreased the adsorbed eugenol surface concentration by one-half (i.e., half a monolayer coverage, $\theta = 0.5$). In comparison, α -Al₂O₃-adsorbed eugenol maintained $\theta = 0.85$ surface coverage at 25% and did not decrease to half a monolayer ($\theta = 0.54$) until the humidity was increased to 75 %. Finally, for both NaCl and α -Al₂O₃, increasing the relative humidity had no discernible effect on the pseudo-first-order rate constants, within the experimental uncertainty, determined for the heterogeneous ozonolysis of surface-adsorbed eugenol. Similar humidity-independent results were observed for the heterogeneous ozonolysis of liquid oleic acid, linoleic acid, and linolenic acid films (Thornberry and Abbatt, 2004). Barnum et al. (2012) found that higher humidities increased the reactive uptake coefficients for the heterogeneous ozonolysis of thin film solid crystalline catechol as a result of hydrogen bonding interactions between adsorbed water and the diol. A similar effect is probably not operative in the current systems since the methoxyphenol binding interactions with the substrates outweigh interactions with co-adsorbed water. These results suggest that the heterogeneous reaction rates listed in Table 2 apply to both dry and humid conditions.

3.5 Impact of solar radiation

Surface-adsorbed eugenol exhibits an absorption maximum in the range of tropospheric actinic radiation ($\lambda > 300$ nm). Figure 8 shows the diffuse reflectance UV–visible spectrum of NaCl-adsorbed eugenol (i.e., t = 0 min; black spectrum) with an absorption band centered at 341 nm. After 100 min of exposure to 240 ppb O₃ under dark conditions, this maximum absorbance shifted to 318 nm (blue spectrum). Continued exposure to ozone under dark conditions resulted in dramatically stronger absorbance in the UV region, with approximately 35 % of 314 nm radiation absorbed following 1000 min of ozone exposure, and an absorption profile that tails out beyond 400 nm (red spectrum). The strong UV–visible absorption profiles of eugenol and its dark



Fig. 8. Diffuse reflectance UV–visible absorbance spectra of NaCladsorbed eugenol exposed to 240 ppb O₃ at t = 0, 100, and 1000 min.

ozonolysis products raise the potential for photoenhanced ozonolysis chemistry under atmospheric conditions. Exposing surface-adsorbed eugenol to the solar simulator in the absence of ozone showed no discernible changes after several hours. We therefore investigated the heterogeneous ozonolysis of surface-adsorbed eugenol irradiated with the output of a xenon lamp solar simulator. Control experiments flowing O₃ through an uncoated NaCl sample showed no changes in measured O₃ concentration with the solar simulator on or off, indicating that photolysis of O₃ at $\lambda > 300$ nm was negligible.

Figure 9 reports the reaction kinetics of key vibrational features during an experiment where NaCl-adsorbed eugenol was exposed to $495 \text{ ppb } O_3$ while the solar simulator was repeatedly turned on and off. The rate of decay of features associated with the alkene side chain (i.e., 3083 and $1638 \,\mathrm{cm}^{-1}$) displayed no significant difference between light and dark conditions. Aromatic features, on the other hand, decayed at a faster rate when irradiated than under dark conditions. Experimental rate constants were determined for each segment of this plot by linear fits to a ln (integrated absorbance) vs. time plot, finding the photoenhanced rate to be $4(\pm 1)$ times faster than dark conditions. Growth of the product carbonyl (1745 cm^{-1}) , which contains contributions from both ozonolysis of the alkene side chain and the aromatic ring, also exhibited an increased rate when irradiated. Since neither NaCl nor α -Al₂O₃ absorb radiation in the region of the solar simulator output, the observed photochemistry is likely the result of ozone reacting with photoexcited eugenol. Ozonolysis of coniferyl aldehyde adsorbed on silica particles was independent of irradiation conditions, although a slow aldehyde photolysis channel was also noted (Net et al., 2010a). The dominant ozonolysis reaction for coniferyl aldehyde involved attack of the alkene substituent, and is therefore consistent with our observa-



Fig. 9. Reaction kinetics for NaCl-adsorbed eugenol exposed to 495 ppb O_3 as a function of UV–visible radiation. Xenon lamp "on" and "off" labels indicate light and dark conditions, respectively. Phenol bend (1341 cm⁻¹) and carbonyl (1745 cm⁻¹) were photoenhanced, while the alkene side chain (1638 cm⁻¹) was not.

tions for eugenol's side chain. In contrast, the ozonolysis of surface-adsorbed aromatic aldehydes (verataldehyde and 3,4,5-trimethoxybenzaldehyde) showed significant photoenhanced oxidation by an order of magnitude producing benzoic acid products (Net et al., 2010b, c). Our results demonstrate a similar, albeit smaller, photoenhancement for cleavage of the aromatic ring.

4 Atmospheric implications and conclusions

Biomass burning aerosol is a major source of atmospheric brown carbon, and laboratory studies have demonstrated that oxidation of select lignin pyrolysis products can generate UV–visible absorbing compounds. For instance, the OH oxidation of aqueous 3,5-dihydroxybenzoic acid produces oligomers that closely resemble HULIS and absorb strongly from the UV out to 500 nm (Hoffer et al., 2004). Secondary organic aerosol formed from the reaction of ozone with gaseous catechol and guaiacol also exhibit visible absorption profiles that stretch out beyond 500 nm (Ofner et al., 2011). Understanding the atmospheric fate of aerosol-bound lignin pyrolysis products is thus essential for understanding how oxidative processing affects aerosol chemical and physical properties, including UV–visible absorption. Heterogeneous ozonolysis represents one potential path for the oxidative processing of surface-adsorbed methoxyphenols (Net et al., 2010a, b, 2011; Knopf et al., 2011).

The present study quantifies the importance of aromatic ring cleavage as a result of ozone 1,3-cycloaddition across methoxyphenols adsorbed on NaCl and α -Al₂O₃ substrates. Assuming that the kinetics of this process are similar for 4propylguaiacol, eugenol, and isoeugenol, the average uptake coefficient for ring cleavage of NaCl-adsorbed methoxyphenols under dark conditions is $\gamma = 2.3 (\pm 0.8) \times 10^{-7}$, which corresponds to an atmospheric lifetime of 1.2 days assuming 40 ppb O₃. The average uptake coefficient for α -Al₂O₃adsorbed methoxyphenols was $\gamma = 9.7 (\pm 7.3) \times 10^{-8}$, a factor of two slower, corresponding to an atmospheric lifetime of 2.8 days. UV-visible radiation ($\lambda > 300 \text{ nm}$) increased the rate of these reactions by a factor of $4(\pm 1)$, which will shorten these atmospheric lifetimes to under 24 h. In comparison, ozonolysis of the alkene side chains for eugenol and isoeugenol, which was independent of UVvisible irradiation, reacted faster: NaCl-adsorbed alkene attack, $\gamma = 2(\pm 1) \times 10^{-6}$, lifetime = 3.3 h; and α -Al₂O₃adsorbed alkene attack, $\gamma = 9 (\pm 4) \times 10^{-7}$, lifetime = 7.2 h. Although the ring cleavage reactions are slower, their estimated atmospheric lifetimes are shorter than the average lifetime of fine-mode particulates suspended in the troposphere. Thus, with aging of biomass burning aerosol, especially when transported through polluted urban environments, this heterogeneous ozonolysis chemistry will alter the chemical nature of adsorbed methoxyphenols.

Diffuse reflectance UV-visible spectroscopy demonstrates that these surface-adsorbed methoxyphenols absorb in the UV-A region (e.g., $\lambda_{max} = 341 \text{ nm}$ for eugenol); however, their subsequent ozonolysis did not generate products that absorb strongly in the visible region. While these reactions will not directly produce brown carbon, they may indirectly impact this process. For instance, vanillic acid might react in a manner similar to 3,5-dihydroxybenzoic acid, which generates strong visible absorbers upon aqueous oxidation. The ozonolysis of surface-adsorbed vanillic acid might cleave the aromatic ring, thereby competing with this potential brown carbon formation process. The rate of this ozonolysis is dependent on several factors, including aerosol substrate and the nature of the substrate-organic interaction, as highlighted by the fact that previous laboratory studies did not observe aromatic ring cleavage for silica-adsorbed methoxyphenols, likely due to the different substrate and sample preparation. Thus, the importance of this chemistry in the atmosphere will depend on exact structure of the tropospheric aerosol under consideration, especially the nature of the the nature of the substrate-organic interaction.

Supplementary material related to this article is available online at http://www.atmos-chem-phys.net/14/ 47/2014/acp-14-47-2014-supplement.pdf.

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