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Elizabeth A. Pillar  
*University of Kentucky, epillar86@gmail.com*

Marcelo I. Guzman  
*University of Kentucky, marcelo.guzman@uky.edu*

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Oxidation of Substituted Catechols at the Air–Water Interface: Production of Carboxylic Acids, Quinones, and Polyphenols

Elizabeth A. Pillar and Marcelo I. Guzman*©

Department of Chemistry, University of Kentucky, Lexington, Kentucky 40506, United States

ABSTRACT: Anthropogenic activities contribute benzene, toluene, and anisole to the environment, which in the atmosphere are converted into the respective phenols, cresols, and methoxyphenols by fast gas-phase reaction with hydroxyl radicals (HO	extsuperscript{•}). Further processing of the latter species by HO	extsuperscript{•} decreases their vapor pressure as a second hydroxyl group is incorporated to accelerate their oxidative aging at interfaces and in aqueous particles. This work shows how catechol, pyrogallol, 3-methylcatechol, 4-methylcatechol, and 3-methoxycatechol (all proxies for oxygenated aromatics derived from benzene, toluene, and anisole) react at the air–water interface with increasing O	extsubscript{3}(g) during contact time and contrasts their potential for electron transfer and in situ production of HO	extsuperscript{•} using structure–activity relationships. A unifying mechanism is provided to explain the oxidation of the five proxies, which includes the generation of semiquinone radicals. Functionalization in the presence of HO	extsuperscript{•} results in the formation of polyphenols and hydroxylated quinones. Instead, fragmentation produces polyfunctional low molecular weight carboxylic acids after oxidative cleavage of the aromatic bond with two vicinal hydroxy groups to yield substituted cis,cis-muconic acids. The generation of maleinaldehydic, maleic, pyruvic, glyoxylic, and oxalic acids confirms the potential of oxy aromatics to produce light-absorbing aqueous secondary organic aerosols in the troposphere.

INTRODUCTION

Chemical and physical processes resulting in the generation of secondary organic aerosol (SOA) particles that scatter and absorb sunlight and serve as ice and cloud condensation nuclei play an important role in the earth’s radiation fluxes. Aromatic species emitted to the atmosphere during fossil fuel combustion and biomass burning (e.g., wildfires) processes provide phenol precursors for SOA formation through competing fragmentation and functionalization oxidation reactions. Previous work has explored the catalytic effect played by interfaces and the role of variable relative humidity (RH) during the atmospheric oxidation of phenols, e.g., by O	extsubscript{3}(g) and hydroxyl radicals (HO	extsuperscript{•}). A combination of analytical methods (infrared spectroscopy, mass spectrometry (MS), and chromatography) employed in our laboratory revealed the importance of competitive oxidation mechanisms. The production of several reactive oxygen species (ROS), including the formation of semiquinone radicals, has been demonstrated, suggesting that aromatics provide a significant missing mechanism for the production of low-volatility species observed in aerosols. Furthermore, the common products identified during offline analysis of reactions lasting a few hours and in situ studies under a few microseconds of contact time (τ	extsubscript{c}) validated a flow-through ultrafast interfacial oxidation MS setup for inspecting reactions at the air–water interface.

Combustion and biomass burning emissions provide the two main aromatic species to the atmosphere, benzene and toluene, that total 5.6 and 6.9 Tg of C/year, respectively. We have previously explained that the oxidative processing of benzene by hydroxyl radicals (HO	extsuperscript{•}) consecutively yields phenol (reaction R1, Scheme 1) and dihydroxybenzenes such as catechol (reaction R2) with gas-phase rate constants k\textsubscript{R1; benzene+HO^•} = 1.2 × 10\textsuperscript{−12} cm\textsuperscript{3} molecule\textsuperscript{−1} s\textsuperscript{−1} and k\textsubscript{R2; phenol+HO^•} = 2.7 × 10\textsuperscript{−11} cm\textsuperscript{3} molecule\textsuperscript{−1} s\textsuperscript{−1}, respectively. The aromatic structures and names, the ring functionalization reactions (e.g., numbered as reactions R1, R2, etc.), and the products formed and their m/z values are introduced in Scheme 1 and Tables S1 and S2 (Supporting Information). By analogy to the reactivity observed for benzene, reactions R1 and R2 also show that toluene (k\textsubscript{R1; toluene+HO^•} = 5.6 × 10\textsuperscript{−12} cm\textsuperscript{3} molecule\textsuperscript{−1} s\textsuperscript{−1}) can generate methylenophenol (p-, m-, and o-cresols) and dihydroxytoluenes (e.g., k\textsubscript{R2.2; methylenophenol+HO^•} = 4.3 × 10\textsuperscript{−11} cm\textsuperscript{3} molecule\textsuperscript{−1} s\textsuperscript{−1}) such as 3- or 4-methylocatechol, respectively. In addition, by consecutive hydrogen abstractions from the −CH\textsubscript{3} group of toluene, reactions R3a and R3b show...
Scheme 1. Proposed Sequential Oxidation Mechanisms of (Black Bold Font) Benzene, (Blue Bold Font) Toluene, and (Red Bold Font) Anisole in the (Purple Font and Reactions) Gas Phase by Hydroxyl Radicals, Which Continue at the (Green Font and Reactions) Air−Water Interface and in Aqueous Particles after Partitioning of the Corresponding Substituted Catechols

The names of the products are color coded to the corresponding precursors. Complete lists of the produced polyphenols and their corresponding quinone redox pairs are available in Tables S1 and S2 (Supporting Information) with the observed m/z values.

In general, phenols and dihydroxybenzenes with methoxy (−OCH3), methyl (−CH3), carbonyl (C=O), and carboxylic acid (−COOH) substituent groups (−R) are ubiquitous aromatic species in the atmosphere, as proved from the analysis of cloudwater that has scavenged combustion and biomass burning emissions. The surfactant properties of these polyfunctional phenols make them especially susceptible to undergo oxidations as they accumulate at the air−water interface or partition into aqueous particles (reaction R4).

In this work, the interfacial oxidations of a series of substituted catechols found in the environment are contrasted to understand their reactivity and potential for electron transfer during ultrafast contact (τc ≈ 1 μs) and detection (τd ≈ 1 ms). Electron-density-donating substituents of the aromatic ring alter the reactivity of each molecule relative to catechol. Mass spectral data confirm the interfacial production of short-lived species can proceed through two pathways: (1) functionalization of the aromatic ring by hydroxylation and (2) fragmentation of the aromatic ring and intermediates by ozone (O3)driven oxidative cleavage. The first pathway implies that semiquinone radicals are formed en route to more hydroxylated aromatic and quinone products. In the alternate pathway, the fragmentation of unsaturated C=C bonds provides polyfunctional low molecular weight carboxylic acids widely found in tropospheric aerosols.

■ EXPERIMENTAL SECTION

Sample Preparation. Solutions of catechol (Sigma-Aldrich, 99.9%), pyrogallol (Acros, 99.7%), 3-methylcatechol (TCI,
99.8%), 3-methoxycatechol (Aldrich, 99.7%), and 4-methylcatechol (Acros, 97.6%) were freshly prepared in degassed ultrapure water (18.2 MΩ cm, Elga Purelab flex, Veolia) and infused into a calibrated ESI-MS instrument (Thermo Scientific, MSQ Plus) operating in the negative ionization mode. Stock solutions of ca. 5.00 mM were diluted to 100.0 μM final concentration after adjustment of the pH with 0.01 M NaOH (Fisher, 99.3%) as measured with a calibrated pH meter (Mettler Toledo). The figures report the percent normalized anion count ($I_{m/z}$) at specific mass-to-charge ($m/z$) ratios for experiments at pH 7.8. While the oxidations were explored in the atmospheric relevant pH range 5–10, reporting the results at pH 7.8 provides an example case that demonstrates the complexity of all the chemistry observed. Anthropogenic biomass burning emissions from agricultural practices often coincide with increased ammonia (NH$_3$) emissions from livestock and fertilizers, which explains why fine particulate matter around agricultural regions can be predominately alkaline. For example, in most of the Midwest Region and the Pacific Coast of the United States, where agriculture is a primary economic activity, NH$_3$ emissions are estimated to be sufficient to neutralize all acidic components. Thus, even fine aerosols can be neutral or slightly basic, which indicates the experiments reported are of atmospheric relevance. All experiments are performed in duplicate.

Oxidation of Substituted Catechols. The ESI-MS flow-through reactor to investigate ultrafast oxidations at the air–water interface has been described in depth. The 100 μM aqueous solution of the selected substituted catechol undergoes pneumatically assisted aerosolization to form a fine mist of micrometer size droplets at ambient pressure. A spark discharge generator (Ozone Solutions) acting over a 0.5 L min$^{-1}$ flow of O$_2$ (Scott-Gross, UHP) produces O$_3$(g), which is quantified in a 10 cm path length cuvette (Starna cell) by UV absorption spectrophotometry (Evolution Array UV–visible spectrophotometer, Thermo Scientific) after dilution with 0.0–5.0 L min$^{-1}$ of N$_2$(g) (Scott Gross, UHP) in a flow-through borosilicate chamber (3.785 L capacity). In the final stage, O$_3$(g) is diluted 61-fold with the N$_2$(g) nebulizing gas (12.0 L min$^{-1}$). The interface of the microdroplets containing substituted catechols encounters a 0.2 L min$^{-1}$ flow of 0 ppbv ≤ [O$_3$(g)] ≤ 3 ppmv during a few microseconds. If oxidized species are produced, they are observed as anions by the mass spectrometer at specific $m/z$ values (reported in the text, figures, and schemes) in less than 1 ms after their formation. The experimental conditions employed were the same used during the previous study with catechol and trihydroxybenzenes: nebulizer pressure, 70 psi; nebulizer voltage, −1.9 kV; cone voltage, −50 V; drying gas temperature, 250 °C. Reported $I_{m/z}$ values correspond to solvent background subtracted raw data acquired at fixed time intervals (e.g., time ≥30 s).

The conversion of terephthalic acid (Acros, 99.0%) to 2-hydroxyterephthalic acid (TCI, 98.3%) with a rate constant $k_{\text{terephthalic acid}}$ = 4 × 10$^9$ M$^{-1}$ s$^{-1}$ was used for in situ quantification of HO$^*$ at the air–water interface. The reaction of 100 μM (1) pyrogallol, (2) 4-methylcatechol, (3) 3-methylcatechol, and (4) 3-methoxycatechol with 2.7 ppmv O$_3$(g) was studied at pH 7.8 in the presence of 5 μM terephthalic acid. The characteristic product peak at $m/z$ 181 for 2-hydroxyterephthalic acid was used to quantify the production of HO$^*$ by standard addition after subtraction of the background produced during the controls under 1 atm of N$_2$(g).

## RESULTS AND DISCUSSION

Reactions of Substituted Catechols at the Air–Water Interface. The interfacial oxidation of 100 μM substituted catechols was examined in the absence of O$_3$(g) or in its presence for mixing ratios of 0.223, 0.989, 1.760, and 2.420 ppm. For example, Figure 1 shows ESI mass spectra of aerosolized aqueous solutions of (top) pyrogallol and (bottom) 3-methoxycatechol at pH 7.8 exposed to a 0.200 L min$^{-1}$ flow of (purple trace) 1 atm of N$_2$(g), (blue trace) 223 ppm O$_3$(g), and (yellow trace) 2.42 ppm O$_3$(g). Ion count values are normalized percentages relative to the most intense peak in each mass spectrum, I$_{125}$ for pyrogallol and I$_{139}$ for 3-methoxycatechol.
Following the oxidation channels exhibited by catechol, it is possible to assign the products for pyrogallol in Figure 1 to also be oxalic acid \( (m/z\ 89) \), maleinaldehydic acid \( (m/z\ 99) \), 5-oxo-3-pentenoic acid \( (m/z\ 113) \), 3- and 4-hydroxy-quinones \( (m/z\ 123) \), 3,4-dihydroxy-quinone \( (m/z\ 139) \), 1,2,3,4-tetrahydroxynaphthalene \( (m/z\ 141) \), 3,4,5-trihydroxy-quinone \( (m/z\ 155) \), and pentahydroxybenzene \( (m/z\ 157) \). Similarly, Figure 1 shows that 3-methoxycatechol \( (m/z\ 139) \) is oxidized to give the hydroxylated products 4-methoxypyrogallol or 3-methoxy-4-hydroxycatechol \( (m/z\ 155) \) and their corresponding quinone pairs, e.g., 3-methoxy-6-hydroxy-quinone and 3-methoxy-4-hydroxy-quinone \( (m/z\ 153) \), 4-methoxy-5-hydroxypyrogallol or 3-methoxy-4,5-dihydroxycatechol \( (m/z\ 171) \) and their corresponding quinones pairs, e.g., 3-methoxy-4,6-dihydroxy-quinone and 3-methoxy-4,5-dihydroxy-quinone \( (m/z\ 159) \), and 4-methoxy-5,6-dihydroxy-pyrogallol \( (m/z\ 181) \) and its corresponding 3-methoxy-4,5,6-trihydroxy-quinone \( (m/z\ 185) \). When comparing the pairs at \( m/z\ 155 \) and \( m/z\ 153, m/z\ 171 \) and 169, and \( m/z\ 187 \) and 185, for oxidized 3-methoxycatechol (Figure 1), it is obvious that the peak for the dihydroxylated product of 3-methoxycatechol at \( m/z\ 171 \) largely exceeds the expected value from correlation to the peak at \( m/z\ 169 \). For these quinone/hydroquinone pairs (signals separated by 2 amu), it is obvious that the peak at \( m/z\ 171 \) displays a large contribution of another species, which is 2-methoxymuconic acid, the major product from aromatic ring cleavage.

Figure 2 displays the ESI mass spectra for aerosolized solutions of (top) 3-methylcatechol and (bottom) 4-methylcatechol under the same conditions listed for Figure 1. Both traces in the absence of \( O_3(g) \) display the anion for methylcatechol \( (m/z\ 123) \) and a small peak for its dihydrate \( (m/z\ 159) \), a feature also exhibited by catechol. Some of the products observed during the oxidation of both methylcatechols are oxalic acid \( (m/z\ 89) \), maleinaldehydic acid \( (m/z\ 99) \), 4-methyl-o-quinone \( (m/z\ 121) \) (no 3-methyl-o-quinone is registered probably due to steric impediments), 3-methyl-4-hydroxy-, 3-hydroxy-4-methyl-, and 4-methyl-5-hydroxy-quinones \( (m/z\ 137) \), 3-methyl-4-hydroxycatechol, 4-methyl-pyrogallol, and 4-methyl-5-hydroxycatechol \( (m/z\ 139) \), 3-methyl-4,5-dihydroxy- and 3,5-dihydroxy-4-methyl-o-quinones \( (m/z\ 153) \), 2-methyl-cis,cis-muconic acid \( (m/z\ 155) \), 3-methyl-4,5-dihydroxy- and 3,5-dihydroxy-4-methyl-catechols \( (m/z\ 155) \), 4-methyl-3,5,6-trihydroxy-2-quinone \( (m/z\ 171) \), and 3-methyl-4,5,6-trihydroxy- and 3,5,6-trihydroxy-4-methyl-catechols \( (m/z\ 173) \).

Figure 3 gathers information from experiments with increasing \( [O_3(g)] \) displaying the exponential drop of \( Iوض/3 \) for pyrogallol, 3-methylcatechol, 4-methylcatechol, and 3-methoxycatechol. In agreement with our previous work with catechol, the information in Figure 3 suggests the first-order reaction on \( O_3(g) \) for all the substituted catechols studied takes place within \( \tau \approx 1 \mu s \). Indeed, the reaction with \( O_3(g) \) molecules occurs at the outermost layers of the interface of a few nanometers thickness without diffusion limitations while replenishment of the organic molecule from the core is facilitated.\(^{2,16}\)

**Electron and Proton Transfers.** The speciation of the substituted catechols in Figures 1 and 2 can be calculated using both \( pK_{a1} \) and \( pK_{a2} \) for these weak diprotic species (\( H_2Q \) and their intermediate form (\( HQ \)):

\[
H_2Q \rightleftharpoons HQ^- + H^+ \quad pK_{a1}
\]

\[
HQ^- \rightleftharpoons H_2Q^2^- + H^+ \quad pK_{a2}
\]

While this information is available for catechol and pyrogallol,\(^{22}\) the instability of this class of compound precludes access to experimental values of the other acid dissociation constants. However, theoretical methods capable of accurately predicting \( pK_{a1} \) and \( pK_{a2} \) values have been developed.\(^{26-28}\) Thus, Table 1 combines the experimental and theoretical values for \( pK_{a1} \) and \( pK_{a2} \) which can be used to solve the speciation of the \( H_2Q \) species vs pH. For example, at working pH 7.8, the fractions of fully associated catechol, \( \alpha_{1H_2Q} \geq 0.95 \), and their immediate monoprotic acid, \( \alpha_{1HQ} \leq 0.05 \), are reported in Table 1. Clearly, the diprotic catechols are the dominant species available for the oxidations displayed in Figures 1 and 2.

Under the oxidizing experimental conditions, the substituted catechols and/or their products can act as two-electron reductants, getting converted into the corresponding quinones by two coexisting pathways. The first pathway proceeds through initial electron transfer from the more abundant undissociated catechols to \( O_3 \) (reaction R5a). This electron transfer pathway is supported by the dominant fractions in
equilibrium at pH 7.8, \( \alpha_{\text{H}_2\text{Q}} \geq 0.95 \) (Table 1), which continues by a fast proton transfer (reaction R5b). Reactions R5a and R5b are equivalent to the net loss of a hydrogen atom. However, direct hydrogen atom transfer should be unfavorable in these molecular ensembles as exemplified by the ~10-fold larger hydrogen bond dissociation energies, calculated from data in dimethyl sulfoxide, than the thermal energy available for the system. The second mechanism involves initial dissociation of the diprotic catechols followed by electron transfer (reactions R6a and R6b), which is less likely to be the dominant process given \( \alpha_{\text{H}_2\text{Q}} \leq 0.05 \) (Table 1). The equilibrium constant for reaction R6a corresponds to the first acid dissociation constant, \( K_1 \). The self-reaction of two semiquinone radicals (HQ⁺) via reaction R7 finally produces a quinone (Q) while regenerating a substituted catechol: 2HQ⁺ → Q + H₂Q. Analogously, the cross-reaction between two different semiquinone radicals also generates a quinone: HQ⁺ + HQ⁺ → Q + H₂Q (reaction R8).

The electron-donating capacity of the substituents and their position relative to the \(-\text{OH}\) groups should be considered together with molecular specific properties (e.g., redox potentials and first and second proton dissociation constants) to understand why reactions R5 and R5b are favored herein. Table 1 also lists the one-electron redox potentials of the semiquinone radical cation (HQ⁺⁺) and undissociated substituted catechol pairs, \( E_{\text{HQ}^+ /\text{H}_2\text{Q}} \) which except for catechol reported at pH 7.30 are predicted using structure–activity relationships derived from computational methods.29 The linear dependence of the redox potentials vs the normal hydrogen electrode (NHE) on the sum of the Hammett substituent parameters (\( \sigma \)) available in the literature is given by the following equation:29

\[
E_{\text{HQ}^+ /\text{H}_2\text{Q}} = 1.85 \sum \sigma_i + 0.46
\]

Table 1 reports the positive \( \Delta E = E_{\text{O}_3/\text{O}_3^+} - E_{\text{HQ}^+ /\text{H}_2\text{Q}} \) for electron transfer (reaction R5a) between the substituted catechols and dissolved \( \text{O}_3 \) (\( E_{\text{O}_3/\text{O}_3^+} = 1.01 \) V). The \( \Delta E \) values correspond to thermodynamically favorable processes as indicated in Table 1 by the calculated \( \Delta G^\circ = -nF\Delta E \), where \( n = 1 \) and \( F = 96.485 \text{ kJ mol}^{-1} \text{ V}^{-1} \) is the Faraday constant. From the increasing \( \Delta E \) values in Table 1, it is observed that electron transfer occurs more easily following the trend catechol < pyrogallol < 4-methylcatechol = 3-methylcatechol < 3-methoxycatechol. In other words, the decreasing \( E_{\text{HQ}^+ /\text{H}_2\text{Q}} \) observed with these electron-donating groups favors electron transfer from H₂Q to O₃ (reaction R5a). The directly generated semiquinone radicals H₂Q⁺⁺ (\( pK_{a1} \approx 1 \)) promptly release a proton (reaction R5b), while the ozone radical anion O₃⁻ from reaction R5a coexists in equilibrium R9a with very basic O⁻⁻ (\( pK_{a1} = 11.8 \)).32 Therefore, for environmentally relevant atmospheric waters and other aqueous systems (pH < 10) and in the present experiments, O⁻⁻ is quickly converted into HQ⁺⁺ (equilibrium R9b). As a consequence, this work also provides an opportunity to understand how HQ⁺⁺ continues the oxidation of substituted catechols at the air–water interface, in aqueous aerosols, and in other relevant water media.

Reactions of the Substituted Catechols with HO⁻. In addition to reactions R7 and R8, the semiquinone radicals can also be produced by the attack of HO⁻ on the catechols (reaction R10), which generates resonance-stabilized cyclo-hexadienyl radicals capable of undergoing fast acid- or base-catalyzed elimination of water (reaction R11). For instance, reaction R10 for unsubstituted catechol proceeds at pH 9 with...
a large rate constant $k_{\text{H}_2\text{Q}^+ + \text{catechol}} = 1.1 \times 10^{10} \text{M}^{-1} \text{s}^{-1}$. The rate constants for the aqueous reactions of pyrogallol, 3-methoxycatechol, and 3- and 4-methylcatechols with HO$^\ddagger$ should be larger than that for catechol due to the presence of electron-donating groups. For example, the reaction rate constants for 3-methylcatechol and 4-methylcatechol with HO$^\ddagger$ (available in the gas phase) are 1.9 and 1.5 times larger than for catechol. Comparable oxidation conditions are supported by experiments under a fixed 100 μM concentration for each substituted catechol at pH 7.8 reacting with 2.7 ppmv O$_3$(g), which constrain the produced [HO$^\ddagger$]$_{\text{interface}} = (1.1 \pm 0.3) \times 10^{13}$ radicals cm$^{-3}$. Thus, the attack of HO$^\ddagger$ on the activated rings should also be expected to be $\geq 2 \times 10^4$ times larger than for the corresponding reaction with O$_3$ ($k_{\text{O}_3 + \text{catechol}} = 5.2 \times 10^8$ M$^{-1}$ s$^{-1}$) at pH 7). Once HO$^\ddagger$ is produced, it can also directly abstract a hydrogen atom from H$_2$Q to form HOQ$^\ddagger$. Under typical atmospheric conditions, the competitive fate of the cyclohexadienyl radicals is fast O$_2$ addition, forming a peroxy radical that decomposes into a hydroxylated product releasing hydroperoxy radical (HO$_2^\ddagger$) by reaction R12. Thus, reaction R12 represents the formation of pyrogallol (m/z 125) from catechol, 3-methyl-4-hydroxycatechol, 4-methylypyrogallol, and 3-methyl-5-hydroxycatechol (all m/z 139) from 3- and 4-methylcatechols, and 4-methoxy pyrogallol and 3-methoxy-4-hydroxycatechol (both with m/z 155) from anisole. It is particularly interesting to compare the relative ion counts for the first hydroxylated products (m/z 139) of both methylcatechols studied. The fact that $I_{m/z}$ for 4-methylcatechol is about 2 times larger than for 3-methylcatechol in experiments with 2.42 ppm O$_3$ (Figure 2), and that their $\alpha_{\text{H}_2\text{Q}/\text{O}_3}$, $\alpha_{\text{H}_2\text{Q}/\text{H}_2\text{O}}$ values (Table 1) are practically identical (Table 1) for both species, suggests that the efficiency for electron transfer must not be the cause for this discrepancy. Indeed, the presence of a vicinal −CH$_3$ causes steric hindrance that could decrease the reactivity of 3-methylcatechol relative to 4-methylcatechol for this electron transfer initiated process.

The quick production of HOQ$^\ddagger$ and HO$^\ddagger$ is also reflected in Figure 3 showing the profiles of each species for increasing [O$_3$(g)]. For example, the top panel of Figure 3 shows how pyrogallol (m/z 125) generates a large production of (1) 3- and 4-hydroxy-o-quinones (m/z 123) likely from self-reaction R7 and (2) tetrahydroxybenzenes (m/z 141) via the sequence represented by reaction R13. Reaction R7 can also be invoked for the considerable production of 4-methyl-o-quinone (m/z 121) from 4-methylcatechol in Figures 3. Interestingly, reaction R7 is not observed in the cases of 3-methylcatechol and 3-methoxycatechol, showing the steric hindrance introduced by −CH$_3$ and −OCH$_3$ groups in position 3 (next to the bond with both −OH groups). In addition, reaction R13 also explains the large production of (1) 3-methyl-4,5-dihydroxycatechol and 4-methyl-3,5-dihydroxycatechol (both m/z 155) from 3-methyl-4-hydroxycatechol, 4-methylpyrogallol, and 4-methyl-5-hydroxycatechol and (2) 4-methoxy-5-hydroxypyrrogallol and 3-methoxy-4,5-dihydroxycatechol (both m/z 171) to form 4-methoxy-5,6-dihydroxypyrrogallol (m/z 187). The conversion of penta hydroxybenzene into benzenemelalexol (reaction R15) represents a fourth-generation aqueous product from catechol (Table S1, Supporting Information).

The reactions for the production of all the quinones experimentally observed in Figures 1 and 2 can be explained to proceed analogously to self-reaction R7 and/or cross-reaction R8. An alternate pathway for their production is the elimination of H$_2$O$_2$ after O$_3$ attack to vacant positions 3 and 4 of the substituted catechols. These complex reactions likely proceed through a cyclic ozonide intermediate. The structures for the quinones named above for m/z 121, 123, 137, 139, 153, 155, 169, and 171 are provided in Table S2 (Supporting Information) together with the corresponding precursors. Figure 3 shows the profiles of these quinones are slightly less intense than the corresponding H$_2$Q species 2 amu heavier but generally resemble each other’s behavior with increasing [O$_3$(g)].

Oxidative Cleavage of Substituted Catechols. While the substituted catechols react at the air–water interface, generating a distribution of products dominated by the functionalized aromatic rings described in Scheme 1, fragmentation of the bond between C$_1$ and C$_2$ by electrophilic O$_3$ generates polyfunctional carboxylic acids. The small ion count for the representative fragmentation products with m/z 89 and 99 included in Figure 3 is expected because these secondary oxidation products cannot accumulate during the short $t_e$ in the setup. The detailed analysis explaining the mechanism of reaction for catechol should also be applicable to its substituted counterparts. Catechol, pyrogallol, both methylcatechols, and 3-methoxycatechol undergo a series of oxidative fragmentations summarized in Scheme 2 to yield muconic acid (m/z 141), 2-hydroxymuconic acid (m/z 157), 2- and 3-methy1muconic acids (m/z 155), and 2-methylmuconic acid (m/z 171), respectively.

The produced substituted muconic acids result from an ozonide that is sequentially converted into a Criegee intermediate and a hydroperoxide that also releases H$_2$O$_2$. Scheme 2 also shows that further fragmentation by O$_3$ and in situ generated H$_2$O$_2$ produces glyoxylic acid (m/z 73), crotonic acid (m/z 85), pyruvic acid (m/z 87), 3-oxopropanoic acid (m/z 87), oxalic acid (m/z 89), maleinaldehyde (m/z 99), 4-hydroxy-2-butoenoic acid (m/z 99).
Scheme 2. Products and Observed m/z Values from Substituted Catechol Oxidation by O$_3$ and H$_2$O$_2$ Following the Mechanisms from Ref 24

“Names in bold colored fonts match the precursors and products.

- OH groups and enhances fragmentation by electrophilic O$_3$. The previous fact can be visualized in Figure 2 as well as by the brown lines for m/z 99 in the center panel of Figure 3. Consequently, a slight increase in the number of low molecular weight carboxylic acids is observed for 3-methylcatechol relative to 4-methylcatechol. Finally, it is possible to consider the attack of HO$^*$ on the substituted catechols followed by the addition of O$_2$ to form an endoperoxide that will add a second O$_2$. The peroxide formed can decompose, contributing quinones and mainly low molecular weight carboxyls and carboxylic acids. For example, the production of glyoxal, glyoxylic acid, oxalic acid, and pyruvic acid from both methylcatechols can proceed through this channel.

**Implications.** This laboratory study contributes to characterizing the evolution of combustion and biomass burning emissions, an important subject to understand how the surface of carbonaceous materials is affected during interfacial oxidations that can potentially alter the morphology of particles that play a role in the direct radiative effect. The most important process governing the initial oxidation of aromatic species in the atmosphere is controlled by the presence of HO$^*$. The results from ultrafast experiments analyzed by mass spectrometry reveal the identity of low-volatility reaction products. Quinone (Table S2, Supporting Information) and polyphenol reactive intermediates coexist at the air–water interface, where low molecular weight carboxylic acids found in brown clouds are produced. The quick oxidation of all substituted catechols proceeds by competing functionalization and fragmentation reactions. Although for acidic hygroscopic particles containing sulfate a lower amount of hydroxyl radical should be generated in situ by the electron transfer mechanism presented, the reactions of phenols with this radical will proceed as reported here regardless of the source of HO$^*$. Thus, the studied phenols, available on the surface of tropospheric particles, should react with impinging HO$^*$ produced in the gas phase through the same pathways presented.

The proposed pathways agree with observations previously made for catechol at the air–water and air–solid interfaces under variable relative humidity, which generate more hydrophilic products that partition into the aqueous phase. In addition, photochemical oxidation should also contribute to the degradation of aromatics in the aqueous phase. However, the substituted catechols display an enhancement in their reactivity due to the ring-activating functional groups. Similarly, the functionalized rings with more –OH groups are also highly activated toward fragmentation and should be relevant intermediates in the degradation of aromatics emitted during combustion and biomass burning. In addition, this information can also be applied to support the similar chemical processing of particulate matter with a diameter of <1 μm (PM$_1$) from the greater London area in winter 2012. This PM$_1$ contains nonvolatile organics associated with refractory black carbon that could originate from either combustion or biomass burning emissions. Finally, this work also explains typical degradation pathways for aromatic pollutants derived from benzene, toluene, and anisole in water media and at the air–water interface.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.7b00232.
Additional environmental justification and tables giving the parent species and produced hydroxy aromatics expected from Scheme 1 and names, structures, observed m/z values, and precursors of the produced quinones (PDF)

AUTHOR INFORMATION

Corresponding Author
*Phone: 859-323-2892; fax: 859-323-9985; e-mail: marcelo.guzman@uky.edu.

ORCID
Marcelo I. Guzman: 0000-0002-6730-7766

Notes
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REFERENCES


