

Oxidation of the *para*-Tolyl Radical by Molecular Oxygen under Single-Collison Conditions: Formation of the *para*-Toloxy Radical

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ABSTRACT: Crossed molecular beam experiments were performed to elucidate the chemical dynamics of the *para*-tolyl (CH₃C₆H₄) radical reaction with molecular oxygen (O₂) at an average collision energy of 35.3 ± 1.4 kJ mol⁻¹. Combined with theoretical calculations, the results show that *para*-tolyl is efficiently oxidized by molecular oxygen to *para*-toloxy (CH₃C₆H₄O) plus ground-state atomic oxygen via a complex forming, overall exoergic reaction (experimental, -33 ± 16 kJ mol⁻¹; computational, -42 ± 8 kJ mol⁻¹). The reaction dynamics are analogous to those observed for the phenyl (C₆H₅) plus molecular oxygen system which suggests the methyl group is a spectator during *para*-tolyl oxidation and that application of phenyl thermochemistry and reaction rates to para-substituted aryls is likely a suitable approximation.



P olycyclic aromatic hydrocarbons (PAHs) are rapidly synthesized from reactive remnants produced by the incomplete combustion of fossil fuels,¹⁻³ a process continuously scrutinized due in part to the carcinogenic and mutagenic properties of aromatic hydrocarbons.^{4,5} In recent years, tolyl radials (CH₃C₆H₄) (Scheme 1) have been confirmed as central aromatic growth species in the formation of methyl-substituted PAHs, such as 1- and 2-methylnapthalene,^{6,7} and disubstituted naphthalenes.^{8–12} Unlike on Earth, where they are regarded as





toxic byproducts of combustion, their large molecular weights and proliferation throughout interstellar space have placed PAHs at the forefront of a multidisciplinary effort to resolve the chemical evolution of the Milky Way. The ubiquity of interstellar PAHs has been inferred by the ultraviolet bump on the interstellar extinction curve $(217.5 \text{ nm})^{13-15}$ and various features in the diffuse interstellar absorption bands (0.4 to 1.2 μ m)^{16,17} and unidentified infrared emission bands (3 to 14 μ m).^{18–20} Neutral PAHs and their (de)hydrogenated, ionized, and protonated counterparts represent 15-30% of the galactic carbon budget,^{21,22} are building blocks for carbonaceous dust grains,^{23,24} participate in grain-surface chemistry,²⁵ and may play a role in the evolution of prebiotic molecules.^{26–28} The molecular growth processes of (substituted) PAHs involve aromatic radicals such as phenyl $(C_6H_5)^{29}$ and tolyl (CH₃C₆H₄),^{6,7} resonantly stabilized free hydrocarbon radicals like propargyl (C_3H_3) ,³⁰ and unsaturated hydrocarbons like acetylene (C_2H_2) .^{31,32}

During combustion, PAH formation is inhibited by competing oxidation pathways that effectively destroy aromatic radical precursors necessary for rapid PAH growth. Using crossed molecular beams^{33–35} and pyrolytic reactors³⁶

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Figure 1. Angularly resolved time-of-flight spectra detected at m/z = 107 ($C_7H_7O^+$) resulting from the reactive scattering of the *p*-tolyl radical (C_7H_7) with molecular oxygen (O_2). Laboratory (observation) angles are indicated for each distribution. Open circles indicate experimentally measured data and red lines the scattering simulation.

combined with electronic structure calculations, examinations of the simplest aromatic radical, phenyl, with molecular oxygen revealed that at temperatures above 1000 K, oxidation is highly efficient and is initiated by the formation of a rovibrationally excited phenylperoxy radical intermediate $[C_6H_5OO]^*$ that preferentially decomposes to the phenoxy radical (C_6H_5O) via O atom loss. Whereas an understanding of the chemical dynamics of the reaction of the phenyl radical with molecular oxygen has begun to emerge recently, beyond computational endeavors³⁷⁻³⁹ little is known of the oxidation of the simplest methyl-substituted aromatic radical, tolyl (Scheme 1). Recently, the o-tolyl-O₂ system was examined by trapping a distonic otolyl radical cation in an oxygen bath. Ortho-quinone moiety formation via hydroxyl (OH) loss was the only dissociation channel observed, which was also predicted to be the minimum energy pathway by the accompanying (G3SX(MP3)) computational analysis.⁴

In this Letter, we share the recent findings of a combined experimental and computational study of the gas-phase neutral-neutral bimolecular reaction of the *p*-tolyl $(CH_3C_6H_4)$ radical with molecular oxygen to form the resonantly stabilized *p*-toloxy radical $(CH_3C_6H_4O)$ under single-collision conditions. We provide irrefutable evidence that *p*-tolyl radicals are effectively oxidized and thus removed from the pool of available aromatic radicals that would

otherwise lead to the formation of methyl-substituted PAHs. C_7H_7 isomers are produced by the combustion of toluene. While conversion to the benzyl radical is thermodynamically favored, approximately 14% of primary products are tolyl radicals produced by abstraction of toluene's phenylic hydrogen atoms near 900 K.^{41,42} Also, tolyl forms competitively with benzyl $(C_6H_5CH_2)$ in dicarbon (C_2) reactions with isoprene (C_5H_8) ,⁴³ is a pyrolysis product of xylenes (dimethyl-benzenes, $C_6H_4(CH_3)_2$ via methyl-loss (CH₃),⁴⁴ and has been predicted as an intermediate (o-tolyl) in the thermal decomposition of the benzyl radical.⁴⁵ Considering that complex toluene oxidation models still rely on benzene (C_6H_6) and phenyl data to account for the reactivity of the tolyl radicals, $\frac{46-48}{46}$ our investigation provides valuable, experimentally based insight for future combustion models that involve the degradation (oxidation) of alkyl-substituted aromatic radicals.

The oxidation of the *p*-tolyl radical was studied under singlecollision conditions in a crossed molecular beams machine^{49,50} by perpendicularly crossing supersonic beams of neat molecular oxygen (O_2 ; $X^3\Sigma_g^-$) and *p*-tolyl ($CH_3C_6H_4$; X^2A_1)⁶ at an average collision energy (E_C) of 35.3 ± 1.4 kJ mol⁻¹. Reactively scattered products were found using a triply differentially pumped quadrupole mass spectrometer preceded by an electron impact ionizer (2 mA, 80 eV). Time-of-flight (TOF) spectra at mass-to-charge ratios (m/z) of 123 ($C_7H_7O_2^+$; adduct), 122 ($C_7H_6O_2^+$; atomic hydrogen elimination), 121 ($C_7H_5O_2^+$; molecular hydrogen elimination), 107 ($C_7H_7O^+$; atomic oxygen elimination), and 106 ($C_7H_6O^+$; hydroxyl elimination) were recorded at the center-of-mass (CM) angle of 10.0 ± 0.1°. A reactive scattering signal was observed only at m/z = 107 (Figure 1), suggesting that the formation of C_7H_7O isomer(s) via atomic oxygen loss is the *only* open channel under our experimental conditions. The corresponding laboratory angular distribution was obtained by integrating the angular-dependent TOF spectra of m/z = 107 ($C_7H_7O^+$), recorded in 2.5° intervals from 7.25° to 24.75° (Figure 2), and reveals a



Figure 2. (Top) Laboratory angular distribution of C_7H_7O derived from the angularly resolved time-of-flight spectra recorded at m/z =107 ($C_7H_7O^+$) from 7.25° to 24.75°. The black dots are experimental data ($\pm 1\sigma$), and the red line is the best fit. (Bottom) Newton diagram for the $CH_3C_6H_4 + O_2 \rightarrow C_7H_7O + O$ reaction. The red circle has radius equal to the maximum CM velocity of C_7H_7O (derived from fit) and is restricted to a 42° wedge in the laboratory frame (hashed lines).

maximum at the CM angle. A nonreactive scattering signal from p-tolyl beam constituents could be observed at 7.25° ; following a control experiment which substituted the reactive O_2 beam for a nonreactive N_2 beam, the nonreactive signal was subtracted from the TOF spectrum recorded at 7.25° . To summarize, the laboratory data provide evidence on the formation of C_7H_7O isomer(s) plus atomic oxygen in the reaction of the p-tolyl radical with molecular oxygen under single-collision conditions.

To obtain information on the scattering dynamics and to identify the product isomer(s) formed, the laboratory data are transformed into the CM reference frame using a forward convolution routine.⁵¹ This procedure yields CM descriptions of the translational energy $(P(E_T))$ and angular $(T(\theta))$ flux distributions (Figure 3). The laboratory data could be fit with a



Figure 3. Center-of-mass translational energy (top) and angular (bottom) flux distributions *p*-toloxy radical (C_7H_7O ; 107 amu) produced from the *p*-tolyl radical reaction with molecular oxygen. Hatched areas indicate the acceptable upper and lower error limits of the fits. The solid lines define the best fit function.

single channel leading from the *p*-tolyl (CH₃C₆H₄; 91 amu) and molecular oxygen $(O_2; 32 \text{ amu})$ reactants to the formation of the C₇H₇O isomer(s) (107 amu) plus atomic oxygen (O; 16 amu). First, the CM translational energy distribution, $P(E_{\rm T})$, helps to assign the product isomer(s). For molecules formed without internal excitation, the high-energy cutoff, $P(E_{T_{max}})$, of 68 ± 16 kJ mol⁻¹ represents the sum of the reaction excergicity and the collision energy. A subtraction of the collision energy reveals that the reaction is excergic by 33 ± 16 kJ mol⁻¹. This value correlates well within the error bars of the computed exoergicity of 42 \pm 8 kJ mol⁻¹ to form the *p*-toloxy (*p*- $CH_3C_6H_4O$) radical plus ground-state atomic oxygen (O, ³P_i). Also, $P(E_{\rm T})$ peaks close to zero translational energy with a maximum around 10 kJ mol⁻¹ indicating a rather loose exit transition state to form p-toloxy and atomic oxygen from a decomposing [C₇H₇O₂]* intermediate. Therefore, the reverse reaction is reasonably characterized by only a small entrance barrier or even by a barrierless long-range association. Finally, on average, $32 \pm 9\%$ of the total available energy is channeled into the translational degrees of freedom of the products; this order of magnitude also suggests indirect scattering dynamics. The CM angular distribution, $T(\theta)$, reveals additional



Figure 4. Potential energy diagram for the reaction of *p*-tolyl (CH₃C₆H₄, X²A') with molecular oxygen (O₂, X³ Σ_{g}^{-}) forming the *p*-toloxy (CH₃C₆H₄O, X²A') radical plus ground-state atomic oxygen (O, ³P_j) calculated at the G3(MP2,CC)//B3LYP/6-311++G^{**} level of theory.

information on the reaction dynamics (Figure 3). Here, the best fit $T(\theta)$ has maxima at 0° and 180°. The forward–backward symmetry and flux exhibiting over all angles indicate indirect scattering dynamics via a long-lived $C_7H_7O_2$ complex, whose lifetime is greater than its rotational period. The minimum at 90° further implies geometrical constraints on the exit transition state and suggests that the oxygen atom is emitted within the plane of the decomposing complex, nearly perpendicular to the total angular momentum vector.⁵²

Having identified *p*-toloxy $(CH_3C_6H_4O)$ and atomic oxygen as the products in the gas-phase reaction of ground-state molecular oxygen and p-tolyl radicals under single-collision conditions, we now merge these results with computational data to untangle the underlying reaction mechanism(s) (Figure 4). The computation at the G3(MP2,CC)//B3LYP/6-311+ +G** level of theory determined that molecular oxygen adds barrierlessly to the radical center of p-tolyl to form the ptolylperoxy intermediate [i1] (CH₃C₆H₄OO) that is stabilized by 193 kJ mol⁻¹ with respect to the separated reactants. Considering trajectories holding the highest symmetry, molecular oxygen adds at the p-tolyl radical site along tolyl's effective $C_{2\nu}$ axis (CH₃-C1-C4). Intermediate [i1] then undergoes unimolecular decomposition via cleavage of the oxygen-oxygen bond through a transition state located 146 kJ mol⁻¹ above [i1] to a van der Waals complex [vdW]. The latter is stabilized by 31 kJ mol⁻¹ with respect to the separated products and dissociates to p-toloxy (CH₃C₆H₄O, X^2A') plus atomic oxygen (O, ${}^{3}P_{i}$) in an overall excergic reaction (-42 ± 8 kJ mol⁻¹). These energetics are similar to the *o*-toloxy channel found on the da Silva et al. surface $(-46 \pm 8 \text{ kJ mol}^{-1})$.³⁷ The computational predictions agree very well with our experimental findings of indirect reaction dynamics involving a C₇H₇O₂ intermediate and a loose, product-like exit transition state, where O-O bond scission occurs late in the reaction coordinate. The experimental reaction dynamics have been significantly constrained through interpretation of both the $T(\theta)$ and $P(E_{\rm T})$ distributions for an isomer-specific assignment of C_7H_7O to be p-toloxy (p-CH₃C₆H₄O). In the transition state, the leaving oxygen atom slightly deviates from the molecular plane with the OOCC dihedral angle being close to 31°. However, in the p-CH₃C₆H₄O…O van der Waals complex the departing oxygen atom returns to and remains in the molecular plane until dissociation to p-CH₃C₆H₄ + O, which is in accord with experimental observations.

The bimolecular reaction of *p*-tolyl with molecular oxygen forming the *p*-toloxy radical plus ground-state atomic oxygen is energetically and dynamically similar to that of the analogous reaction of phenyl (C_6H_5) with molecular oxygen forming the phenoxy radical (C_6H_5O) and $O({}^{3}P_{j})$, as examined experimentally³³⁻³⁵ and computationally.^{53,54} Here, molecular oxygen also adds to the radical center of the phenyl radical forming a phenylperoxy intermediate [C₆H₅OO], which is stabilized by 194 kJ mol⁻¹ with respect to the separated reactants. This complex also emitted atomic oxygen via a loose exit transition state located 152 kJ mol⁻¹ above $[C_6H_5OO]$ to form a van der Waals complex bound by 5 kJ mol⁻¹ with respect to the products.⁵³ The similarity of the title reaction with $C_6H_5 + O_2$ strongly suggests the *para*-substituted methyl group is a spectator in the formation of *p*-toloxy. Thus, the adaptation of phenyl characteristics (e.g., reaction rates and thermochemistry) is likely suitable for para- and meta-tolyl radicals in combustion settings. Ortho-tolyl represents an exception, due in part to unique pathways involving benzylichydrogen migration to ortho-bound substituents^{37,40} and its relatively short lifetime at combustion relevant temperatures.⁵⁵ In real combustion settings, the primary products of the phenyl and tolyl oxidation undergo secondary reactions, and of note here is the thermal decomposition by carbon monoxide (CO) elimination from phenoxy to cyclopentadienyl (C_5H_5) ,^{56–} and therefore from toloxy to methylcyclopentadienyl (C_6H_7) .⁵⁹ While phenyl oxidation promotes PAH formation through subsequent $C_5H_5 + C_5H_5$ reactions,⁶⁰ unlike cyclopentadienyl, methylcyclopentadienyl can isomerize and proceed directly to benzene via atomic hydrogen loss.^{61,62⁺} In the phenylmolecular oxygen system, the phenylperoxy intermediate eliminates carbon dioxide (CO_2) to form cyclopentadienyl, which overall (both via the CO_2 and CO + O pathways) accounts for up to 30% of products at 1000 K.³⁶ As previously discussed, the methyl group does not participate in the initial addition of molecular oxygen to *p*-tolyl; thus, carbon monoxide elimination from *p*-toloxy and carbon dioxide elimination from *p*-methylphenylperoxy to methylcyclopentadienyl are also expected with similar energetics and branching ratios in comparable environments. In summary, p-tolyl is oxidized by molecular oxygen to form *p*-toloxy and ground-state atomic oxygen in a complex forming, exoergic reaction (experiment, -33 ± 16 kJ mol⁻¹; computation, -42 ± 8 kJ mol⁻¹). These results contribute to the growing discussion surrounding the reactivity of tolyl and to the oxidation of molecular growth

species of polycyclic aromatic hydrocarbons (PAHs) in combustion systems.

METHODS

Experimental Methods. The oxidation of p-tolyl was performed in a crossed molecular beams machine at the University of Hawaii, which enables the study of bimolecular reactions in a single-collision environment through the combined use of supersonic sources, beam skimmers, and differentially pumped regions to maintain a mean free path larger than 10^2 m in the reaction chamber. Detection is accomplished in the plane of the reactant beams by a triply differentially pumped, liquidnitrogen-cooled detector that employs a variable electron impact ionizer and quadrupole mass filter (1.2 MHz),⁶³ and a single particle counter which consists of an aluminum-coated stainless steel target (-22.5 kV) and a scintillator mounted atop a photomultiplier tube (-1.35 kV).⁶⁴ para-Chlorotoluene (p-CH₃C₆H₄Cl, Aldrich, 97.5%) was purified by multiple freeze (77 K)-pump (<30 mTorr)-thaw (293 K) cycles before seeding at 0.3% in helium (99.9999%; Gaspro). The helium seeded precursor was introduced to a 5 mm² photodissociation region by a piezoelectric pulsed valve,⁶⁵ where the halogenated precursor was photolyzed by 193 nm (Coherent Compex 110, 60 Hz, 12 mJ/pulse) to yield p-tolyl radicals.^{66,67} The p-tolyl radical beam was velocity selected by a 4-slot chopper disk. On axis characterization at m/z 45.5 (C₇H₇²⁺) and a 34 eV ionization energy indicate a peak velocity, $v_p = 1542 \pm 14$ m s⁻¹, and speed ratio $S = 10.6 \pm 1.4$ for the *p*-tolyl radicals. It should be noted that previous crossed beam experiments in our group with m- and p-tolyl radicals verified that the initially formed *m*- and *p*-tolyl radicals do not isomerize.^{6,7,9-11,68,69} In the present study, the p-tolyl reactant intercepted a neat, perpendicularly crossed, molecular oxygen (O₂: $X^{3}\Sigma_{g}^{-}$) beam characterized by $v_p = 776 \pm 20 \text{ m s}^{-1}$ and $S = 17.4 \pm 1.0$. Reactively scattered products were ionized at 80 eV (2 mA) and sampled throughout the scattering plane. The dynamics and energetics of the product channel were analyzed in the center-of-mass (CM) frame, a transformation accomplished through the iterative use of a forward convolution routine that assumes separability of the differential cross-section $I(u, \theta) =$ $P(u) \times T(\theta)$. The differential cross section (DCS) is calculated from operator-defined parameters in the CM frame and transformed to the observation frame for comparison with the experimental TOFs and laboratory angular distribution. This process is repeated until the best fit of the experimental data is attained.

Theoretical Methods. To optimize geometries of all local minima structures and transition states we used the hybrid density functional B3LYP^{70–73} method with the 6-311++G** basis set. The same B3LYP/6-311++G** method was applied to calculate vibrational frequencies and zero-point energy (ZPE) corrections for the reactants, products, and all intermediates and transition states. To refine the final energies we applied a modified G3(MP2,CC)//B3LYP^{74,75} composite scheme with single-point energies computed at the G3-(MP2,CC) level using B3LYP optimized structures, according to the following formula

$$E_0[G3(MP2, CC)] = E[RCCSD(T)/6 - 311G^{**}]$$

+ $\Delta E_{MP2} + E(ZPE)$

where $\Delta E_{MP2} = E[MP2/G3large] - E[MP2/6-311G^{**}]$ is a basis set correction and E(ZPE) is the zero-point energy

computed at the B3LYP/6-311++G^{**} level. The spin-restricted RUHF-RCCSD(T) version of the coupled cluster method was employed. T1 diagnostics were checked during coupled cluster calculations to ensure that wave functions do not possess any multireference character. The calculation scheme represents a modification of the original G3⁷⁶ method. All calculations were performed using the *Gaussian* 09⁷⁷ and *MOLPRO* 2010⁷⁸ program packages.

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Notes

The authors declare no competing financial interest.

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Letter

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