Mechanistic Insight into the Aromatic Hydroxylation by High-Valent Iron(IV)-oxo Porphyrin \(\pi\)-Cation Radical Complexes

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Mechanistic studies of the aromatic hydroxylation by high-valent iron(IV)-oxo porphyrin \(\pi\)-cation radicals revealed that the aromatic oxidation involves an initial electrophilic attack on the \(\pi\)-system of the aromatic ring to produce a tetrahedral radical or cationic \(\sigma\)-complex. The mechanism was proposed on the basis of experimental results such as a large negative Hammett \(\rho\) value and an inverse kinetic isotope effect. By carrying out isolated labeling studies, the oxygen in oxygenated products was found to derive from the iron-oxo porphyrin intermediates.

Aromatic hydroxylation is an important chemical process in enzymatic reactions, as evidenced by the detoxification processes and drug metabolism by Cytochromes P450, nonheme iron mono- and dioxygenases, and copper enzymes.1–3 The aromatic hydroxylation has been extensively investigated over the past three decades in Cytochromes P450 and iron porphyrin models, with the intention of understanding their reaction mechanisms and developing artificial catalysis that can be used in drug synthesis and biodegradation of environmental pollutants.4 The mechanism of the aromatic hydroxylation has been proven to be quite complex, as shown in the proposed mechanisms of the aromatic ring hydroxylation by high-valent iron(IV)-oxo porphyrin \(\pi\)-cation radicals (the so-called Compound I) (Scheme 1). Early studies concluded that the hydroxylation of aromatic compounds occurs via an arene oxide formation (Scheme 1, pathway A).4 More recent experimental evidence, however, has revealed that the aromatic oxidation does not proceed via the epoxide intermediate formation but occurs via the generation of a tetrahedral radical or cationic \(\sigma\)-complex (Scheme 1, pathway B).5 Very recently, density functional theory (DFT) calculations of aromatic hydroxylation by Compound I have shown that the aromatic ring activation involves an initial electrophilic attack on the \(\pi\)-system of the aromatic ring to produce a radical or cationic species (Scheme 1, pathway B).6,7 The evidence for the formation of radical cations by direct electron abstraction from aromatic compounds is still ambiguous (Scheme 1, pathway C),8 but DFT calculations do not support the electron-transfer mechanism.6 The hydrogen abstraction mechanism has been ruled out on the basis of experimental results, including a small kinetic isotope effect (KIE) value despite the high bond dissociation energy of aromatic C–H bonds, and computational calculations (Scheme 1, pathway D).6,9,10

Since the previously proposed mechanisms were based on experimental results obtained under catalytic conditions and with an assumption that Compounds I were generated as reactive species in the catalytic reactions, we have investigated the aromatic hydroxylation by generating high-valent iron(IV)-oxo porphyrin \(\pi\)-cation radicals and using the intermediates directly.

in kinetic studies of aromatic hydroxylation under stoichiometric conditions. In this work, we have observed a large negative inverse KIE. By carrying out 18O-labeling experiments, the oxygen in oxygenated products was found to derive from the iron porphyrin hydroxylation investigated with high-valent iron(IV)-oxo porphyrin cation radicals generated in situ.

A high-valent iron(IV)-oxo porphyrin cation radical complex, [(TPFPP)FeIV(O)]+ (I) (TPFPP = meso-tetrakis(pentafluorophenyl)porphinato dianion) (Supporting Information, Figure S1 for the structure of iron porphyrins),11 was prepared by treating Fe(TPFPP)(CF3SO3) with m-Chloroperbenzoic acid (CPBA) in CH2Cl2 containing a small amount of CH3CN at −80 °C. Upon addition of benzene to the solution, the intermediate reverted back to the starting iron porphyrin complex and showed pseudo-first-order decay as monitored by a UV−vis spectrophotometer (Figure 1a). Pseudo-first-order fitting of the kinetic data allowed us to determine the kobs value to be 3.0(3) × 10−3 s−1 (Figure 1a, inset). The pseudo-first-order rate constants increased proportionally with benzene concentration, thus leading us to determine a second-order rate constant to be 1.0(1) × 10−1 M−1s−1 (Figure 1b). Activation parameters for the benzene hydroxylation were determined by plotting first-order-rate constants measured at different temperatures (183−208 K) against 1/T (Figure 2c); the activation enthalpy and entropy values were calculated to be ΔH‡ = 4.6(3) kcal·mol−1 and ΔS‡ = −29(2) cal·mol−1·K−1, respectively. Product analysis of the benzene hydroxylation with GC and GC−MS revealed that benzoquinone was produced as a major product (vide infra).

We then investigated the electronic effect of benzene substrate on reaction rates, by carrying out the reactions of I with various para-X-substituted benzenes at −80 °C. The second-order rate constants were determined under the conditions described above but with different concentrations depending on their relative reactivities (Supporting Information, Figure S2). The data in Table S1 (Supporting Information) indicate that the electron-donating ability of the para substituents influences the reaction rate significantly. By plotting the second-order rates as a function of σp of the para substituents, a good linear correlation was obtained with a large negative Hammett ρ value of −8.0 (Figure 2), indicating an unusually high charge separation and electrophilic nature in the transition state of the benzene oxidation by iron(IV)-oxo porphyrin cation radical species. Further, such a large negative ρ value of −8.0 suggests the generation of a cationic species rather than a radical species in the transition state (see Scheme 1).6

It is worth noting that Hammett ρ values reported previously in the catalytic oxidation of aromatic compounds by Cytochrome P450 and iron(III) porphyrins are much smaller (ρ value of −1) than the value determined kinetically with iron(IV)-oxo porphyrin cation radicals generated in situ.8b,12 Further, it is of interest to compare Hammett ρ values obtained in the aromatic hydroxylation and olefin epoxidation reactions.11,13,14 In olefin epoxidations, ρ values of −1.8 and −1.9 were reported in the competitive epoxidation of para-substituted styrenes by I and in the kinetic studies of para-substituted styrenes by [(TMP)FeIV(O)]+ (TMP = meso-tetramesitylporphinato dianion), respectively.11,13 Such a large difference between the ρ values in the para-substituted benzenes and styrenes might be due to the steric effects of the substituent.
values of aromatic hydroxylation and olefin epoxidation is attributable to the involvement of a more polar species in the transition state of aromatic hydroxylation, as Shaik and co-workers proposed in theoretical studies that a major difference between benzene hydroxylation and olefin epoxidation is the appearance of a cationic pathway in the case of benzene, whereas radical-only pathways operate in the case of olefins. 6, 15

Kinetic isotope effect (KIE) was also investigated kinetically in the hydroxylation of undeuterated and deuterated aromatic compounds by high-valent iron(IV)-oxo porphyrin σ-cation radicals, such as in the reactions of benzene and deuterated benzene by 1, toluene and deuterated toluene by [(TDFPP) + FeIV(18O)]1 2 (TDFPP = meso-tetakis(2,6-difluorophenyl)porphino diuran), and naphthalene and deuterated naphthalene by [(TDFPP) + FeIV(18O)]1 3 (TDFPP = meso-tetakis(2,6-dichlorophenyl)porphino diuran) (Table 1). The calculated kD/kH value of ~0.8 indicates an inverse KIE in the aromatic ring oxidation reactions; the observation of the inverse KIE is consistent with the sp2-to-sp3 hybridization change during the addition of an electrophilic iron-oxo species to the sp2 center of aromatic ring to form a σ adduct. 5, 6, 8, 16 Further, the inverse KIE rules out the hydrogen abstraction mechanism; 9 large KIE values ranging from 5 to 14 are observed in C1 aromatic ring hydroxylation by 1 at ~80 °C. 11

Finally, the source of oxygen in oxygenated products was investigated by carrying out isotope labeling studies in the hydroxylation of toluene by 18O-labeled [(TDFPP) + FeIV(18O)]1 18O-toluene and PhIO under the identical conditions of the toluene hydroxylation. 22 In the latter experiment, we found that cyclohexene oxide contains 75% 18O. This result indicates that 75% of 1 is 18O-labeled. Further, the formation of quinone products in the oxidation of aromatic compounds. 19a

In summary, we have reported mechanistic details of aromatic hydroxylation investigated kinetically with high-valent iron(IV)-oxo porphyrin σ-cation radicals generated in situ. On the basis of the experimental results of a large Hammett p value and an inverse KIE, we have proposed that the aromatic oxidation involves an initial electrophilic attack on the σ-system of the aromatic ring to produce a tetrahedral radical or cationic σ-complex (Scheme 1, pathway B). We have also demonstrated

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**TABLE 1. Kinetic Isotope Effect in the Hydroxylation of Aromatic Compounds by High-Valent Iron(IV)-oxo Porphyrin σ-Cation Radical Complexes**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Benzene Hydroxylation by 1</th>
<th>Toluene Hydroxylation by 2</th>
<th>Naphthalene Hydroxylation by 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>kD/kH</td>
<td>kD/kH</td>
<td>kD/kH</td>
</tr>
<tr>
<td>C6H6</td>
<td>0.8</td>
<td>0.8</td>
<td>0.8</td>
</tr>
<tr>
<td>C6D6</td>
<td>1.0</td>
<td>2.2</td>
<td>0.8</td>
</tr>
<tr>
<td>kD/kH</td>
<td>1.0</td>
<td>2.2</td>
<td>0.8</td>
</tr>
</tbody>
</table>

Reactions were done with [(TDFPP) + FeIV(18O)]1 18O-toluene and substrates (15 mM for benzene, 150 mM for toluene, and 15 mM for naphthalene) in CH2Cl2 at ~80 °C. Rate constants are averaged by three determinations.

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**SCHEME 2**

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(18) When we checked the percent of 1-18O generated in the labeled H218O experiments by carrying out the catalytic epoxidation of cyclohexene with [FeIII(TPPFP)]2+ and PhIO under the identical conditions of the toluene hydroxylation, we found that cyclohexene oxide contains 75% 18O. This result indicates that 75% of 1 is 18O-labeled.


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unambiguously that the oxygen in oxygenated products derives from the iron-oxo intermediates, not from molecular oxygen.

**Experimental Section**

**Kinetic Studies and Product Analysis.** In general, high-valent iron(IV)-oxo porphyrin π-cation radicals were prepared by adding 1.3 equiv of m-CPBA (2 mM, diluted in 50 µL of CH₂Cl₂) to reaction solutions containing their corresponding iron(III) porphyrin triflates (1.5 mM) in a solvent mixture (0.5 mL) of CH₂Cl₂ and CH₃CN (200:1) in a 0.1-cm UV cell at −80 °C. Then, appropriate amounts of substrates (diluted in 100 µL CH₂Cl₂) were added to the UV cell, and spectral changes of the intermediate were directly monitored by a UV–vis spectrophotometer. Rate constants, \( k_{obs} \), were determined by pseudo-first-order fitting of the decrease of absorption bands at 660 nm for 1, 665 nm for 2, and 680 nm for 3. All the rate constants are averages of at least three determinations.

Product analysis was performed by injecting reaction solutions directly into GC and GC–MS. Product yields were determined by comparison with standard curves of known authentic samples and decane as an internal standard. The source of oxygen in oxygenated products was determined by preparing \([\text{TPFPP}]^\text{+}\cdot\text{Fe}^{IV}(\text{18O})\]^+ (1-18O) and using the intermediate directly in the hydroxylation of toluene. To increase the accuracy of the product analysis, the hydroxylation of toluene was performed under catalytic conditions by adding PhIO, which was prepared by treating PhIO (10 mM, diluted in 50 µL CH₃OH) with H₂18O (10 µL, 0.52 mmol, 95% 18O enriched), into a reaction solution containing Fe(II)porphyrin (2 mM) and toluene (0.2 M) in a solvent mixture (0.5 mL) of CH₂Cl₂ and CH₃CN (200:1) at 25 °C. Then, the reaction mixture was stirred for 0.5 h and directly analyzed by GC–MS. 18O and 16O compositions in 2-methylbenzoquinone were analyzed by the relative abundances of the following mass peaks: \( ml/z = 122 \) (M), 124 (M + 2), and 126 (M + 4) for 2-methylbenzoquinone (see Scheme 2). The reaction procedures for cyclohexene epoxidation were the same as described for the toluene hydroxylation, and the 16O and 18O compositions in cyclohexene oxide were determined by the relative abundances of the mass peaks at \( ml/z = 83 \) and 97 for 16O and \( ml/z = 85 \) and 99 for 18O.

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**Supporting Information Available:** General experimental conditions, Figure S1 for the structure of iron(III) porphyrin complexes, and Figure S2 for the determination of second-order rate constants. This material is available free of charge via the Internet at http://pubs.acs.org.

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