### ELECTROPHILIC AROMATIC SUBSTITUTION · ELECTRON TRANSFER ROUTES IN SIDE CHAIN SUBSTITUTION

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In a continuing series of interesting reports, Baciocchi, Illuminati and coworkers have described the side chain substitution of alkyl-aromatics under a variety of electrophilic conditions<sup>1</sup> For example, hexamethylbenzene is converted to pentamethylbenzyl chloride by chlorine in acetic acid in the dark <sup>2</sup> The scope of these non-conventional substitution reactions comprise a variety of electrophilic reactions including halogenation, (chlorine, bromine and iodine), nitration and thiocyanation which proceed generally on polyalkylbenzenes and other electron-rich aroma -



tics such as thiophenes, etc 3-4

The heterolytic mechanism in Scheme I which they propose for side chain substitution under electrophilic conditions include first, the formation of a o-adduct I between the electrophile and arene in eq 2, followed by either a stepwise or synchronous  $\alpha$ -proton loss and rearrangement (eq 3), e  $g$ , <sup>2</sup>

# Scheme I



It has recently been shown, however, that aromatic substitution, either in the nuclear or side chain positions, may proceed by electron-transfer processes<sup>5</sup> The electron transfer steps in Scheme II have been effected either electrochemically or by metal oxidants such as cobalt $(III)$ , manganese(III), cerium(IV) capable of effecting one-electron changes <sup>5,6</sup> The formation of **Scheme II** 



$$
\underline{\mathbf{H}} \quad \xrightarrow{-\mathbf{H}^+} \bigotimes^{\mathbf{H}_2} \quad \xrightarrow{-\mathbf{E}} \quad \bigotimes^{\mathbf{H}_2^+} \quad \text{etc} \tag{5}
$$

**the aromatic cation-radical 11 1s the key to the electron transfer mechanism, since side chain**  substitution follows directly from the loss of an  $\alpha$ -proton as shown in eq 5 Two factors are important in facilitating the electron transfer process in Scheme II a relatively low oxidation po**tential for the aromatic compound and the avallablllty of a one-electron oxidant We feel that**  both conditions are fulfilled in a number of the side chain substitution reactions reported by **Illuminati et al In particular, many of the electrophlles available for aromatic substitution are also capable of performmg as one-electron oxidants Thus, chlorine, bromine and lodlne readily undergo one-electron reduction and are also highly effective electron acceptors m charge transfer complexes with a variety of arenes ' Slrmlarly, mtroruum ions involved directly or indirectly m nitration have a reasonable high reduction potential relative to nitrogen dioxide 8** 

We wish to show that chlorination of hexamethylbenzene under conditions described by Illuminati et al may proceed by an electron transfer mechanism Thus, acetic acid solutions of **chlorine and hexamethylbenzene were nuxed directly In the cavity of an electron spm resonance spectrometer The intense and well-resolved esr spectrum of hexamethylbenzene cation-radical**  shown in Figure 1 was readily obtained at room temperature Other arenes such as pentamethyl**benzene and durene showed similar behavior The facile formation of the radical-catlon 1s conslstent with an electron-transfer mechamsm described in Scheme III Scheme III** 

$$
\sum_{i=1}^{n} \left(1 + \alpha_i \right) = \sum_{i=1}^{n} \left(1 + \alpha_i \right)
$$
 (6)

$$
\underline{II} \quad \xrightarrow{-H^+} \quad \bigotimes_{CH_{2^*}} \quad \xrightarrow{Cl_2} \quad \xrightarrow{Cl_2} \quad \uparrow \quad \text{Cl}^-\quad \text{etc.}
$$
 (7)



Esr spectrum of the hexamethylbenzene radical-cation (central portion) Figure 1 obtained during the reaction of hexamethylbenzene and chlorine in acetic acid at 25 °C. Proton nmr field markers are in kHz.

The electron transfer mechanism in Scheme III involves readily accessible intermediates and avoids the postulation of cyclohexadienes and their rearrangement A rigorous distinction between heterolytic and electron transfer pathways in Schemes I and III, respectively, is difficult to make since electrophilic addition (eq 2) and electron transfer (eq 7) have much the same electronic requirements <sup>5b</sup> In some electrophilic substitutions, such as proton exchange,<sup>9</sup> acceptors are unavailable for an electron transfer process to occur, and the reaction must perforce take a heterolytic pathway

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