Electron-transfer Chain Catalysis of Substitution Reactions. Experimental Evidence for the S_{on}2 Mechanism

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Summary Two possible cases of an oxidative electrontransfer chain catalysis mechanism, the S_{0N} ² mechanism, are presented : the anodic 'oxidation' of 4-fluoroanisole in the presence of acetate ion to give 4-acetoxyanisole, and the CuIII 'oxidation' of chloro- and fluoro-benzene in the presence of water to give phenol.

In the preceding communication,¹ Alder has introduced the concept of electron-transfer chain (E.T.C.) catalysis in which a non-redox process is catalysed or stimulated by an electron-transfer reagent. Foremost among such mechanisms is the S_{RN} l reaction²⁻⁴ where the key intermediates are radical anions.

Alder extended the E.T.C. concept to mechanisms involving radical cations as well, $e.g.,$ the $S_{0N}2$ mechanism, a bimolecular mechanism for what is formally a nucleophilic substitution *via* one-electron oxidation of the substrate, equations (1) - (4) . For such a reaction to take place a good leaving group is essential, and the obvious choice in the type of bond-breaking step **(3)** involved is fluoride ion. It is extremely difficult to oxidize and is very

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-e^-
$$

Initialization:
$$
RX \longrightarrow RX^+
$$
 (1)

$$
\int \mathbf{R} \mathbf{X}^{+} + \mathbf{N} \mathbf{u}^{-} \rightarrow [\mathbf{R} \mathbf{X} \mathbf{N} \mathbf{u}]^{*}
$$
 (2)

Propagation:
$$
\begin{cases}\n\begin{array}{c}\n\text{INX} + \text{IN} & \rightarrow \text{INX}\n\\ \n\text{[RXNu]} & \rightarrow \text{RNu}^+ + \text{X}^- \\
\text{RNu}^+ + \text{RX} & \rightarrow \text{RNu} + \text{RX}^+ \\
\end{array}\n\end{cases}
$$
\n(3)

$$
RNu^{+} + RX \rightarrow RNu + RX^{+}
$$
 (4)

unlikely to leave as fluorine atom or, worse, fluorine cation from the radical in reaction **(3)** or from species resulting from its further oxidation. Moreover, the abstraction of fluoride ion from a cyclohexadienyl radical has been shown to be highly exothermic by thermochemical calculations.⁵

One likely candidate for the S_{0N} ² mechanism is the substitution of fluorine for acetoxy that takes place upon anodic,⁶ Ag¹¹,7 and 12-wolframocobalt(III)ate⁸ 'oxidation' of 4-fluoroanisole [reaction (5), $Ar = 4-MeOC₆H₄$]] in the presence of acetate ion. In the anodic case, coulometry

$$
ArF + AcO^- \rightarrow ArOAc + F^-
$$
 (5)

indicated the consumption of one electron per mol of substrate.⁶ We now report that reaction (5) $(Ar =$ $4-MeOC₆H₄$ can be run catalytically under conditions of low conversion. Moreover, we show that Cu^{III} can induce S_{ON}2 type behaviour, in that Cu^{III} oxidation of chloro- and fluoro-benzene in trifluoroacetic acid-acetic acid-water gives phenol as the predominant product. Finally, a number of suspected cases of the $S_{\text{on}}2$ mechanism have been spotted in the literature.

Anodic oxidation of 4-fluoroanisole in HOAc-KOAc at low conversions (higher conversions, up to 2 F mol^{-1} lowered the yields appreciably)⁶ gave current yields of 4-acetoxyanisole in excess of loo?/, (Table **l),** the best yield *so* far being obtained at 2% conversion and a relatively low substrate concentration. This experiment corresponds to a consumption of 0.37 e⁻ per molecule of product formed. We are somewhat puzzled over the strong decrease in current yield as conversion is increased, but tentatively ascribe it to further oxidation of 4-acetoxyanisole, a substrate likely to undergo 'cascade' electron transfers under oxidizing conditions (at higher conversions, polymeric material is indeed very clearly visible).

TABLE 1. Current yields of 4-acetoxyanisole from the anodic oxidation of 4-fluoroanisole in HOAc-K0Ac.a

[Substrate]/M	Conversion/ F mol ⁻¹	Current yield ^b /%
$2-0$	0.2	84
$2-0$	0 ¹	99
2.0	0.02	128
$1-0$	0.02	212
0.5	0.02	270

a At a platinum anode at **25** "C; current density *5.0* mA cm-2; [AcO^-] 0.5 M. b Calculated for 1 F mol⁻¹.

A second S_{0N} ² type reaction was found during exploratory studies of the oxidative properties of Cu^{III} complexes *vs*. aromatic compounds. Copper(III) compounds have been used previously for the oxidation of organic compounds,⁹⁻¹² but so far such studies have been confined to aqueous media. Table 2 shows that benzene, alkyl-aromatic compounds, and chloro- and fluoro-benzene are oxidized to the usual range of oxidative-substitution and coupling products,¹³ expected for stoicheiometric electron transfer mechanisms, by the copper(III) complex of biuret¹⁴ in trifluoroacetic acid (TFA) or TFA-AcOH mixtures. However, in the presence of small amounts of water, the formation of phenol from the halogenobenzenes becomes predominant. While we have not yet succeeded in obtaining catalytic yields **of** phenols, compounds that can be very easily oxidized further, the reaction is certainly a non-redox process which, according to control experiments, does not take place in the absence of Cu^{III}. It does not take place either with the much weaker oxidant, Cu^{II}, present.¹⁵

A survey of the recent literature on anodic and metal-ion oxidation of chloro- and fluoro-aromatic compounds[†] showed the following suspected cases. Anodic¹⁶ and cobalt($[III]$) trifluoroacetate¹⁷ oxidation of chlorobenzene gave 2 and 6% of phenyl trifluoroacetate, respectively, lead tetra-acetate

f We have restricted ourselves to chloro- and **fluoro-derivatives'because** of the requirement of having a leaving group which is difficult to oxidize; naturally, several other types of $S_{0N}2$ behaviour can be envisaged, such as the 'oxidative' substitution of the methoxy-group by cyanide ion in certain polymethoxyaromatic compounds either anodica *Am. Chern. Soc.,* **1969,91,4181; L.** Eberson and B. Helgke, *Acta Chem. Scand., Ser. B,* **1975,29,451; 1977,31, 813)** or photochemically **(J.** den Heijer, 0. B. Shadid, J. Cornelisse, and E. Havinga, *Tetrahedron,* **1977, 33, 779).**

TABLE 2. Oxidation of some aromatic compounds by the Cu^{III} complex of biuret.⁸

a Conditions used in all experiments, unless otherwise noted: substrate (5 mmol), KCu(biuret)₂ (10 mmol), glacial acetic acid (25 ml),
FA (2·5 ml), reflux temperature, reaction period 2 h. b Reaction medium in experime d Experiment 3: 3,5,2',4',6'-pentamethyldiphenyl-
Experiment 3: 3,5,2',4',6'-pentamethyldiphenyl-TFA (2.5 ml), reflux temperature, reaction period 2 h. TFA (25 ml) and water (5 mmol), and experiment 3 TFA (25 ml) and mesitylene (12.5 ml). C.l.c. yield based on Cu^{III}.
ment 2: 3,5,2',4',6'-pentamethyldiphenylmethane, 5%, and bimesitylene, 0.1%. Experiment 3: 3,5,2', methane, 24 %, and bimesitylene, **7** %.

oxidation of chlorobenzene in TFA gave traces of phenol,¹⁸ and oxidation of fluorobenzene in aqueous solution by peroxydisulphate in the presence of Cu^{II} and Fe^{II} salts gave phenol in yields of up to **96%** at room temperature.19 We also draw attention to the $ipso$ hydroxydehalogenation process observed with p -halogenotoluenes in aqueous alkali at high temperatures;²⁰ it is catalysed by copper salts and with air present as an oxidant the product pattern changes from a m/p cresol mixture (characteristic of a benzyne mechanism) to predominantly p -cresol, indicating a switch to an $S_{\text{ON}}2$ mechanism. If this is so, it would seem to be possible to find more efficient ways of catalysing this industrially important process. Finally, another entry into the *S0,2* mechanism should also be possible *via ipso*radical attack,²¹ the initiation step then being $RX + Nu$.

 \rightarrow [RXNu][.] [cf. equation (2)]. One possible case is the attack of pentafluorobenzoyloxyl radical on chloro- and bromo-benzene to give relatively high yields $(40-50\%$ based on the amount of $ArCO₂$ ^t) of phenyl pentafluorobenzoate,²² in spite of the fact that the reaction conditions employed were far from ideal for the S_{0}^{3} sequence to occur. In other words, it should be possible to induce or even catalyse nucleophilic aromatic substitution of halogenobenzenes by decomposing diacyl peroxides.

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