HETEROAROMATIC AZO-ACTIVATED NUCLEOPHILIC SUBSTITUTIONS. THE REACTION OF $4-$ (p-METHOXYPHENYLAZO) PYRIDINIUM METHIODIDE WITH PIPERIDINE IN DIMETHYL SULPHOXIDE **Ikenna** myido" and Collins I. Ubochi Department of Chemistry, University of Ibadan. Ibadan, Nigerla

Abstract - The azopyridinium-activated phenyl methyl ether, *I,* undergoes nucleophilic attack by piperidme m dimethyl sulphoxide predominantly at the aryl carbon centre in a reaction that is **base** catalyzed and faster than the corresponding reaction of the dinitro-activated analogue, *L,* by more than two decades.

Alkyl aryl ethers are insensitive to hydrolysis and similar nucleophilic substitution reactions, When the aromatic ring is activated through azophenyl substitution these compounds become susceptible to nucleophilic attack in moderately to strong acidic aqueous media¹. In recent work we have shown that azopyridinyl substitution has a dramatic effect on the reactivity of these compounds particularly when the aza substituent is in a position where it can interact **²**conjugatively with the reactmn **centre** . Ia these examples, **one** or **more** protonation equilibria in the acidic media employed **occur** prior to nucleophilic attack. In connection with these studies and in furtherance of our recent interest in azo compounds as dyes we have synthesized

the azopyridinium-activated phenyl methyl **ether.4-(p-methoxypheny1aro)pyridinium** methiodide, *I.* We wish to report that 1 undergoes facile nucleophilic attack by piperidine in the dipolar aprotic' solvent dimethyl sulphoxide (DMSO) at rates that are significantly faster than the corresponding reactions of 2,4-dinitroanisole, 2, where activation is provided by two nitro groups.

Diazotization of 4-aminopyridine followed by coupling with phenol yielded 4-(p-hydroxyphenylazo)pyridine. Treatment of this azopyridinyl-substituted phenol with 0.1 M methanolic H₂SO₄ under reflux **for** 4 h followed by neutraliaatlon with ice-cold NaOH solution afforded the azopyridinylsubstituted phenyl methyl ether which, on refluxing with CH₃I in methanol for 8 h at 74°C, gave 1^3 .

The kinetics of the reaction of 1 with piperidine in DMSO at 30°C were followed spectrophotometrically with **the mine** in large enough **excess** to **ensure** first order kinetics. The reaction gave experimental abrorbance values at infinite time of **the** product of attack at **the** aryl carbon, 3, which was less than the calculated infinity values at low piperidine concentrations, e.g. 7.0-10x10⁻³M piperidine. Good agreement between the two values was obtained at higher piperidine concentrations. This behaviour is due to competing attack by the nucleophile on the $\frac{4}{\sqrt{3}}$ aryl carbon to give $\underline{3}$ and on the alkyl carbon to give $\underline{4}$ and N-methylpiperidine at low piperidine 4 concentrations. As compound %is known from **the** work of Buncel and **Keum** and was identified s pectrophotometrically by us , a rate constant for the s_N^2 reaction at the alkyl carbon of ca . 1.7×10^{-3} $M^{-1}s^{-1}$ was calculated. The value of this rate constant was independent of piperidine concentration. At higher nucleophile concentrations attack on the aryl carbon centre predminates. **The** relative Values of **the** rate constants for the attack on **the** aryl carbon **(see** Table 11 and the S_N2 reaction explains the increasing preponderance of aromatic nucleophilic substitution (S_MAr) with increasing piperidine concentration.

Table 1. Rate constants $(N^{-1}s^{-1})$ for the reaction of 1 with piperidine in DMSO at 30°C (Initial $[1] = 1.50 \times 10^{-4}$ M).

$\left 10^2\right $ (piperidine)/M: 0.70 1.00 2.00 4.00 8.00 10.0				
$\left 10^3\right $			$: 2.35$ $: 3.16$ $: 4.79$ $: 7.83$ 11.4 12.8	

Concurrent $S_N A r$ and S_N^2 reactions have also been observed in the reaction of $\frac{2}{3}$ with piperidine in DMSO, methanol and benzene³.

Figure 1. Plot of **the** second order rate constant, k,, **versus** piperidine concentration.

The S_NAr reaction is catalysed by piperidine (see Table 1). A plot of the observed second order rate constant, k_A , against piperidine concentration is curvilinear (Figure 1). This kinetic form is explicable in terms of the accepted mechanism for base catalysed S_NAr reactions given in the scheme. In **the** steady state rate expression of Eq li) derrved from the mechanism of **the** scheme, the observed kinetic form corresponds to the condition $k_{-1} k_2 + k_3[B]$,

$$
k_{A} = \frac{k_{1} (k_{2} + k_{3} [B])}{k_{-1} + k_{2} + k_{3} [B]}
$$
 (i)

in which case the decomposition of the intermediate 5 is rate limiting at least at low base (piperidinel concentrations. **There** is controversy regarding the detalled mechanism of **the** ba~e

⁶catalysed **piperidinodemethoxylation** of aronatic substrates and **the** present results do not allow for a choice between the proffered alternatives.

what these results highlight is **the** powerful activating influence of the azopyridmium group. A comparison of reactivity of 1 and 2 towards substitution by piperidine^{5a} in DMSO reveals that $\frac{1}{2}$ reacts faster than 2 by factors of ca. 2 **x** 10^2 and 3 **x** 10^2 in the S_N^{Ar} and S_N^{2} reactions respectively. Hence there is a difference in the activation energies of the order of 3-5 **k** cal mol-' in the reactions of 1 and 2. **These** impressive differences in rates **In** both reactions must have their origin in the presence of the azopyridinium function in 1 in which the pyridiniw molety **serves as** an electron sink by **the** extensive delocalization of **the** incaming electrons £ran the nucleophile. This is evident from **the** resonance contributing structure lp. Thus quaternization of the pyridine nitrogen in 1 obviates the need for the protonation(s) in acidic media necessary for **the** activation of azophenyl and aropyridinyl ethers towards nucleophilic ~ ~ substitutionls2. It **1s** also conceivable that transition state effects could **as** well be important in the determination of the rate ratios noted above. A systematic investigation of **the** origin of **these** rate differences **as** well **as** the **use** of & **as** a model -pound in **seekmg** further information regarding the details of the mechanisms of base catalysed S_NAr reactions is under active consideration,

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