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The Feeble Nucleofugality of a Nitronate Leaving Group and its Enhancement by Ring Strain

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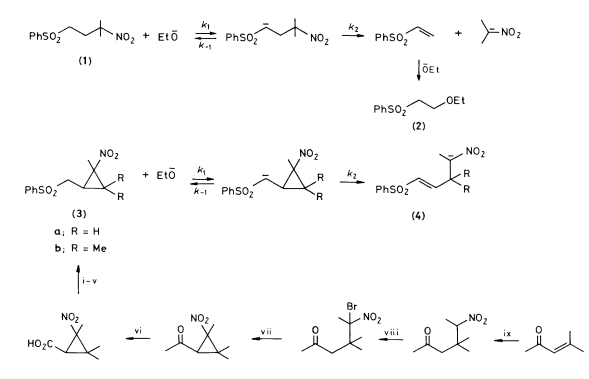
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The rank of a nitronate ion in activated alkene-forming elimination is low (+2.6); incorporation of the leaving group in a cyclopropane accelerates elimination so much that the $(ElcB)_{B-}(ElcB)_{I}$ borderline is traversed but the retro-Thorpe-Ingold effect nevertheless operates.

Carbon leaving groups in alkene-forming eliminations have exceptionally low ranks^{1,2} (= nucleofugalities) which bear no relation to the pK_a of the conjugate acid of the leaving group (Z).² Within a group of three carbon leaving groups of closely

similar pK_a^{ZH} values, the nitronate ion, Me₂CNO₂, came mid-

way in rank between PhCH₂C(CN)₂ and CN for 1,2-elimination activated by a benzoyl group.² We have now measured the rank of this group in sulphonyl-activated 1,2-elimination and studied the enhancement of its nucleofugality on incorporation in a strained ring.



Scheme 1. i, EtOH-H₂SO₄; ii, LiAlH₄-Et₂O; iii, *p*-MeC₆H₄SO₃Cl-pyridine, 0 °C; iv, PhSNa-EtOH; v, H₂O₂-MeOH-(NH₄)₂ MoO₇; vi, Br₂-NaOH; vii, MeCO₂K-EtOH; viii, NaOMe-MeOH-Br₂-CHCl₃; ix, NaOEt-EtNO₂.

Table 1. Eliminations in ethanolic sodium ethoxide.^a

Substrate	$k_{ m obs}{}^{ m b}$	$k_{1^{\mathrm{b,e}}}$	Rank
(1)	1.5×10^{-9d}	0.33	+ 2.6
(3 a)	61.7	3.5	$+12.2^{e}$
(3b)	5.3	1.6	$+11.5^{e}$

^a At 25 °C. ^b Units dm³ mol⁻¹ s⁻¹. ^c Calculated from $\rho^* \sigma^*$ plots (ref. 3). ^d Estimated from reactions at 80 and 90 °C. ^e N.B. a rank >11 shows E2 or, within limits, $(E1cB)_I$ mechanisms.

Substrate (1) (Table 1) obtained by addition of 2-nitropropane to phenyl vinyl sulphone slowly eliminates 2-nitropropane in ethanolic sodium ethoxide at *ca*. 90 °C. Determination by g.l.c. of the ethoxy-sulphone (2) and 2-nitropropane formed allows estimation of the rate constant at 25 °C (Table 1). Calculation of the deprotonation rate constant, k_1^3 allows assignment of rank (+2.6) from the relation⁴ Rank = $k_{obs} - k_1 + 11$.

When $k_1 \ge k_{obs}$ the mechanistic borderline between $(E1cB)_{\rm R}$ processes $(k_{obs} \ll k_1)$ on the one hand, and $(E1cB)_{\rm I}(k_{obs} = k_1)$ and $E2(k_{obs} > k_1)$ on the other, is traversed. Such a situation is revealed when the same nitronate leaving group is incorporated in a strained ring as in (3) (Scheme 1).

The effect of straining the bond to a leaving group on its nucleofugality has recently been quantified⁵ and very large enhancements have been reported. The cyclopropanes (3a) and (3b) have been synthesised (Scheme 1), the routes being modelled on earlier work.⁶ Treatment with ethanolic sodium ethoxide initially gives the alkene (4), the reaction being followed by the u.v. spectral change. The rate constants in Table 1 show once again⁵ the very large enhancement of k_{obs} , in this case by a factor of 4×10^{10} . Calculation of the rank value from k_{obs} and the calculated deprotonation rate shows that for (3a) the value of 11 is greatly exceeded. This points to the E2 mechanism for this substrate, a mechanism

associated with excellent leaving groups such as halide. The effect of strain, therefore, is to convert a very poor into an excellent leaving group. The mechanistic borderline between the $(E1cB)_{\rm R}$ and $(E1cB)_{\rm T}$ mechanisms is traversed as k_2 exceeds k_{-1} , and evidently the very large enhancement of nucleofugality caused by strain calls the E2 mechanism into play.

For substrate (3b), gem-dimethyl substitution produces a small but significant decrease in k_{obs} , of which only a small part can be accounted for by the electronic effect of the methyl groups. Table 1 shows the calculated effect on k_1 were the mechanism a stepwise one. The calculated rank is, however, also above that (11) for the $(E1cB)_1$ process and the lower reactivity of this substrate is consistent with the operation of the *retro*-Thorpe–Ingold effect on the concerted processes of deprotonation and ring cleavage. This is in contrast with the effect of gem-dimethyl substitution on concerted eliminative ring-fission of oxirans.⁷

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