Amidoethylation of Anthracene Hydride by N-Aroylaziridines: Inner-sphere Single Electron Transfer (SET) and Radical Coupling confirmed \dagger P.-Y. Lin and H. Stamm*

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Regioselectivity (near 1:1) of substitutive ring opening of 1-benzoyl-2-methylaziridine by anthracene hydride is incompatible with common nucleophilic attack and thus confirms the radical coupling path.

Reactions of N-aroylaziridines with excess anthracene hydride $(AH⁻)$ may be exemplified by means of 1a (Scheme 1). Aziridino ketyl $4a$ is an essential intermediate³ generated by benzylic fragmentation $(BFR)^4$ of the rapidly formed³ carbonyl adduct 2a. Homolytic ring cleavage of 4a affords the amidatoalkyl radical 5a, a precursor of the main product 6. The second product is 7.

When the aziridine ring of 1a carries substituents, analogues of 7 are obtained³ unless they arise from 2-phenylaziridines and are unstable under usual conditions.^{3,5} The assumption³ that 7 and its analogues are formed by coupling of amidatoalkyl radicals with anthracenide $A^{\bullet -}$ was supported by a regioselectivity of ring opening that seemed to exclude a direct S_N 2-like path to analogues of 7 and hence also to 7. Subsequently it was found⁶ from a study of 1-acyl-2,2-dimethylaziridines that S_N 2-like ring opening may require planarization of the nitrogen pyramid thereby shifting the mechanism to a borderline type whose regioselectivity is compatible with the AH ^{$-$} results. This reopened the mechanistic question since the very fast initial carbonyl attack is reversible.⁷

Ring opening of 1-acyl-2-methylaziridines by strong nucleophiles was recently⁸ shown to strongly prefer cleavage of the N-CH₂ bond. AH^- and 2-methyl-1-pivaloylaziridine provided a mixture of products (total 94%) with an overall regioselectivity isopropylamides:n-propylamides of 35:1. The reaction of xanthenyl anion (oxa analogue of $AH^$ devoid of the BFR path) with 1b yielded 82% of benzoylxanthene and 14.5% of amidoethylated xanthenes with an iso to normal regioselectivity of 28:1. Thus, one may expect a ratio of about 30:1 if i-10 and n-10 (Scheme 2) are formed from $1b$ and AH^- only, or mainly, by nucleophilic ring opening.

Two three-day runs of 10 mmol of 1a with 16 mmol of $AH-Li$ ⁺ in 200 ml of THF provided 58% (47%) of isopropylamide i-9, 14% (18%) of *n*-propylamide **n-9**, 9% (4%) of **i-10** and 9% (5%) of **n-10** (values in parentheses are the yields of the second run). The yields of both 10 are crude yields in the sense that they were estimated by ¹H NMR from fractions containing minor amounts of unknown products, probably isomers of 10, one of them being 11 (see below). But the yield ratios $i:n = 1$ (0.8), determined from the methyl doublets at 1.21 and 0.94 ppm, are sufficiently reliable. These ratios of isomeric 10 are far from the 30:1 ratio expected for an S_N2 mechanism. Both 10 arise consequently only or nearly so from coupling of anthracenide $A^{\bullet -}$ (generated by BFR) with amidatoalkyl radicals i-8 and n-8. Moreover, coupling with position 1 of A^{\bullet} obviously formed traces of 11 (one or two products with isomeric side chains). 11 was identified in the insepar-

Scheme 1

able mixture of isomeric 10 by characteristic $\mathrm{^{1}H}$ NMR signals for the non-aromatic double bond. A doublet $(J 10.4)$ for H-4 at 6.70 ppm shows fine splitting $(ca. 1.1 Hz)$ of the lines from coupling with H-10. A doublet $(J \ 10.4)$ of approximated triplets $(J \text{ ca. 5})$ at 6.08 ppm comes from H-3, the triplets indicating attachment of the amidatopropyl chain to position 1. Olefinic and additional aromatic signals are in accord with those of 2-vinylnaphthalene.¹

There were at least four methyl doublets $(J \text{ ca. } 6.8 \text{ Hz})$, at 1.02, 1.11, 1.31 and 1.42 ppm) in addition to those of both 10. This is compatible with a mixture of i-11 and n-11 when one considers diastereoisomerism. However, part of these signals may come from structural isomers of 11, e.g. Y carried in position 2. Weak signals in the range of $5.9-6.7$ ppm point to isomerism in the non-aromatic ring.

Cleavage $4b \rightarrow i-8$ will be kinetically controlled (*cf.* ref. 9) and reduction of the amidatoalkyl radicals by a second 4b forms probably the primary carbanion faster than the secondary one. It is therefore not surprising to find much more i-9 than n-9.

Experimental

The reactions were performed as described in ref. 4 starting with 17 mmol of dihydroanthracene AH2 and 16 mmol BuLi (hexane). The reactions were quenched with acetic acid. The residue obtained after the usual workup was chromatographed (silica gel Merck, $0.063-0.200$ mm, $40 \text{ cm} \times 4 \text{ cm}$, toluene-ethyl acetate 9:1); compo-

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Scheme 2

site fractions were analyzed by ${}^{1}H$ NMR (CDCl₃, Me₄Si internal). J values are given in Hz.

Run 1 provided hydrocarbons and their oxidation products; 72 mg of unknown products and 120 mg of a 3:1 mixture of i-10 (90 mg) and $n-10$ (30 mg) followed. A crystal of $i-10$ could be manually picked out. Continued elution yielded 482 mg of a 1:1.2 mixture of **i-10** (219 mg, total 309 mg = 9%) and **n-10** (263 mg, total 293 mg = 9%) containing a trace of 11 (¹H NMR data given in the text). Further elution gave 146 mg of i-9 and 1022 mg of a mixture

of 798 mg (total 944 mg = 58%) of **i-9** and 224 mg (14%) of **n-9.**
i-10: mp 192–194 °C; $v_{\text{max}}/\text{cm}^{-1}$ 3303 (NH), 1636 (amide I), 1538 (amide II); δ_H 1.21 (d, J 6.6, Me), 1.87 (m, NCCH₂), 3.88 (d, J 18.4, 10-H pseudo eq), 4.08-4.31 (m, 9-H and NCH), 4.12 (d, J 18.3, 10-H pseudo ax), 5.90 (d br, *J* 8.2, NH), 7.17-7.32 (m, 8 ArH), 7.38 -7.42 (m, m-H and p-H of Ph), 7.63 (m, o-H of Ph).

n-10 (in mixture with i-10): $\delta_{\rm H}$ 0.89 (d, J 6.8, Me) (m, NCCH), 3.35 (dt_{approx}, *J* 13.8 and *ca.* 6.0, 1 H of NCH₂), 3.49 $(dt_{approx}, J_13.8 \text{ and } ca. 6.7, 1 \text{ H of NCH}_2), 3.81 (d, J_74, 9-H),$ 3.85 (d, J 18.3, 10-H pseudo-eq), 4.13 (d, J 18.3, 10-H pseudo-ax), 5.84 (s br, NH), aromatic signals cannot be distinguished from those of i-10.

Mixture of **i-10** and **n-10**: (Found: C, 84.3; H, 6.9; N, 4.0.
C₂₄H₂₃NO requires C, 84.4; H, 6.8; N, 4.1%); $v_{\text{max}}/\text{cm}^{-1}$ 3313 (NH), 1631 (amide I), 1540 (amide II).

Run 2 provided hydrocarbons and their oxidation products; 65 mg of unknown products and 219 mg of a mixture of i-10 (91 mg) and $n-10$ (128 mg) followed. Further elution yielded 182 mg of a mixture containing mainly (more than 90 mg totalling to 279 mg = 9%) 10 in a ratio of 55 mg (total $146 \text{ mg} = 4\%$) of i-10: 35 mg (total $163 \text{ mg} = 5\%$) of n-10. This mixture contained also some 11. Continued elution provided 84 mg of i-9 and 976 mg of a mixture consisting of 687 mg (total 771 mg = 47%) of i-9 and 289 mg (18%) of n-9.

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