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

J. Chem. Research (S), 1998, 646–647<sup>†</sup>

Faculty of Pharmacy, University of Heidelberg, Neuenheimer Feld 346, D-69120 Heidelberg, Germany

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 intermediate<sup>3</sup> generated by benzylic fragmentation  $(BFR)^4$  of the rapidly formed<sup>3</sup> 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 obtained<sup>3</sup> unless they arise from 2-phenylaziridines and are unstable under usual conditions.<sup>3,5</sup> The assumption<sup>3</sup> 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 found<sup>6</sup> 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**<sup>-</sup> results. This reopened the mechanistic question since the very fast initial carbonyl attack is reversible.<sup>7</sup>

Ring opening of 1-acyl-2-methylaziridines by strong nucleophiles was recently<sup>8</sup> shown to strongly prefer cleavage of the N–CH<sub>2</sub> bond. AH<sup>-</sup> 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<sup>-</sup> devoid of the BFR path) with 1b yielded 82% of benzoyl-xanthene 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<sup>-</sup> 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 <sup>1</sup>H 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 I

able mixture of isomeric **10** by characteristic <sup>1</sup>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 *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.<sup>10</sup>

There were at least four methyl doublets (*J ca.* 6.8 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  $AH_2$  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 cm × 4 cm, toluene–ethyl acetate 9:1); compo-

<sup>\*</sup>To receive any correspondence.

<sup>†</sup>This is a **Short Paper** as defined in the Instructions for Authors, Section 5.0 [see *J. Chem. Research* (S), 1998, Issue 1]; there is therefore no corresponding material in *J. Chem. Research* (M).



Scheme 2

site fractions were analyzed by  ${}^{1}$ H NMR (CDCl<sub>3</sub>, Me<sub>4</sub>Si 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** (<sup>1</sup>H 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**.

of 798 mg (total 944 mg = 58%) of **i-9** and 224 mg (14%) of **n-9**. **i-10**: mp 192–194 °C;  $\nu_{max}/cm^{-1}$  3303 (NH), 1636 (amide I), 1538 (amide II);  $\delta_{H}$  1.21 (d, J 6.6, Me), 1.87 (m, NCCH<sub>2</sub>), 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<sub>approx</sub>, J 13.8 and ca. 6.0, 1 H of NCH<sub>2</sub>), 3.49 (dt<sub>approx</sub>, J 13.8 and ca. 6.7, 1 H of NCH<sub>2</sub>), 3.81 (d, J 7.4, 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.  $C_{24}H_{23}NO$  requires C, 84.4; H, 6.8; N, 4.1%);  $\nu_{max}/cm^{-1}$  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 mg = 4%) of **i-10**: 35 mg (total 163 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**.

Received, 29th May 1998; Accepted, 10th June 1998 Paper E/8/04044C

## References

- 1 Arene Hydrides, Part 16. Part 15: T. Mall and H. Stamm, J. Chem. Soc., Perkin Trans. 2, 1997, 2135.
- 2 Aziridines, Part 73. Part 72: Arene Hydrides, Part 15; see ref. 1.
- 3 H. Stamm, A. Sommer, A. Woderer, W. Wiesert, T. Mall and P. Assithianakis, J. Org. Chem., 1985, 50, 4946.
- 4 H. Stamm, T. Mall, R. Falkenstein, J. Werry and D. Speth, J. Org. Chem., 1989, 54, 1603.
- 5 H. Stamm and R. Falkenstein, Chem. Ber., 1990, 123, 2227.
- 6 P.-Y. Lin, K. Bellos, H. Stamm and A. Onistschenko, *Tetrahedron*, 1992, **48**, 2359.
- 7 T. Mall and H. Stamm, Chem. Ber., 1988, 121, 1349.
- 8 P.-Y. Lin, G. Bentz and H. Stamm, J. Prakt. Chem., 1993, 335, 23.
- 9 J. Werry, H. Stamm, P.-Y. Lin, K. R. Falkenstein, S. Gries and H. Irngartinger, *Tetrahedron*, 1989, **45**, 5015.
- 10 C. V. Pouchert and J. R. Campbell, *The Aldrich Library of NMR Spectra*, Vol. IV, Aldrich Chemical Company, 1974.