N-(3,4-Dihydro-4-oxoquinazolin-3-yl)pyridinium Imides: Aziridinating Agents for Alkenes: Measurement of Rotational Barriers around each N-N Bond in an N-N-N System

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Pyridinium imides 3–5 and 8 are obtained as crystalline solids from the reaction of 3-acetoxyaminoquinazolinones 2 and 10 with the corresponding pyridines; barriers to rotation around each of the N–N bonds in 3 have been measured by NMR spectroscopy, and the imides 3–5 react both with styrene and with diethyl fumarate to give the corresponding aziridines.

Oxidation of 3-aminoquinazolinones, e.g. 1, with lead tetraacetate in dichloromethane gives the 3-acetoxyaminoquinazolinone 2 which is stable in solution at -20 °C.¹ Addition of pyridine (10 mol equiv.) gives the pyridinium imide 3 as a pale yellow crystalline solid, mp 134–136 °C (decomp.). Similarly, the pyridinium imides 4 (53%) and 5 (55%) were obtained from reaction of 2 with 2-methylpyridine and 3-methylpyridine, respectively.† These imides are not stable to silica chromatography and were purified by crystallisation after removal of the bulk of the pyridine.

In the NMR spectra of the imides 3 and 5, broadened signals are observed at room temperature for the methyl groups in the quinazolinone 2-isopropyl substituent and from the two *ortho*-protons on the pyridinium ring.

Examination of a number of NMR spectra of 3 at 400 MHz between -95 °C and ambient temperature reveals the involvement of *two* dynamic processes in the broadening of the signals from the two types of proton referred to above. In anticipation of what follows, the preferred stereostructure proposed for this imide ($3' \rightleftharpoons 3''$) is that shown in Fig. 1.

In the NMR spectrum of 3 at -40 °C, the broadened isopropyl methyl signals have separated and sharpened: from their separation and coalescence temperature, a barrier $\Delta G^{\ddagger} = 60 \text{ kJ mol}^{-1}$ was calculated. This barrier is assigned to that for rotation around the N_{α} - N_{β} bond in Fig. 1. (3' \rightleftharpoons 3").

The signals from the *ortho*-protons of the pyridinium ring in 3 are separated at -90 °C (CD₂Cl₂ solution) although not sharpened fully. From their separation and coalescence temperature, a barrier $\Delta G^{\ddagger}=48~kJ~mol^{-1}$ was calculated and this is assigned to that for rotation around the $N_{\beta}\!\!-\!\!N_{\gamma}$ bond in Fig. 1.

Interconversion of the magnetic environments of the two *ortho*-protons of the pyridinium ring in 3 would result from direct *inversion via* 6 at the divalent nitrogen N_{β} in 3' (Fig. 1) in which the three nitrogens are linear. However, the lower energy barrier measured above is *not* that associated with this inversion at N_{β} since such a process (3' \rightleftharpoons 3") would, at the same time, also interconvert the magnetic environments of the isopropyl methyl groups. This inversion at N_{β} via 6 could account for the

R N O R N O N HOAC

1 R = Pri
9 R = Et

10 R = Et

3 R1 = R2 = R3 = H, R = Pri
4 R1 = Me, R2 = R3 = H, R = Pri
5 R1 = R3 = H, R2 = Me, R = Pri
8 R1 = R2 = H, R3 = CN, R = Et

Scheme 1 Reagents and conditions: i, Pb(OAc)₄, CH₂Cl₂, -20 °C

higher energy barrier referred to above; however, this transition state requires a higher energy sp-hybridisation for N_{β} and unfavourable interaction between lone pairs contained in porbitals on N_{α} and $N_{\beta}.\ddagger$ Rotation around this $N_{\alpha}\!-\!N_{\beta}$ bond, on the other hand, can proceed via 7 (Fig. 1) having sp³ hybridisation for N_{β} and less unfavourable interactions between lone pairs on N_{α} and N_{β} .

Support for this interpretation comes from the NMR spectra at various temperatures of the pyridinium imide **8**, prepared from 4-cyanopyridine and the 3-acetoxyaminoquinazolinone **10** derived from **9**. From the separation and coalescence of the *ortho*-protons on the cyanopyridine ring, the barrier for N_{β} - N_{γ} rotation in **8** is higher ($\Delta G^{\ddagger} = 62 \text{ kJ mol}^{-1}$) than that in **3**, corresponding to greater resonance delocalisation with the pyridine ring and hence more double bond character in this N_{β} - N_{γ} bond. The barrier to rotation around the N_{α} - N_{β} is also raised ($\Delta G > 77 \text{ kJ mol}^{-1}$)§ which is consistent with a rotation around this bond which is coupled with the rotation around the N_{β} - N_{γ} bond, *i.e.* the transition state for the N_{α} - N_{β} rotation has an orientation around the N_{β} - N_{γ} bond as illustrated in **7**.

These imides 3–5 function as aziridinating agents: heating 3 with styrene (3 mol equiv.) at 135 °C gave aziridine 11 (44%) identical with a sample prepared by aziridination of styrene using 2. Likewise, aziridination of diethyl fumarate (3 mol equiv.) using 5 at 145 °C gave aziridine 12 (33%); the same aziridine 12 was obtained in 56% yield by heating the 2-methylpyridine-derived imide 4 in diethyl fumarate at 150 °C.¶ Aziridine 12 (81%) was also prepared by aziridination of diethyl fumarate using 2.

Oxidation of 1 in the presence of 3,5-lutidine gave an oily impure product which could not be purified. However, if the oil obtained on work-up was heated directly with diethyl fumarate

Fig. 1

at 145 °C, aziridine 12 was isolated (30%) after chromatography.

A large number of pyridinium imides are known,² the majority having the additional substituent on the divalent nitrogen as CO₂R, COR, SO₂Ar or Ar. Barriers to rotation around the N-N bond in these compounds have not been reported; the symmetry likely to be present in many of them means that the two *ortho*-protons of the pyridinium ring are magnetically equivalent. Some of these pyridinium imides act

Scheme 2

as nitrene precursors and on photolysis bring about the aziridination of alkenes.²

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Footnotes

- † All new compounds have been fully characterised.
- ‡ Some alleviation of the lone-pair interaction in 6 is possible by inversion via a rotamer around the N_{α} - N_{β} bond in which these lone pairs are not eclipsed.
- § The increase in this higher barrier is difficult to account for in terms of inversion at $N_B \nu ia$ a transition state analogous to 6.
- \P The cyano-substituted imide 8 was unreactive towards either styrene or diethyl fumarate even at higher temperatures.

References

- R. S. Atkinson, M. J. Grimshire and B. J. Kelly, *Tetrahedron*, 1989, 45, 2875; R. S. Atkinson, P. J. Edwards and G. A. Thomson, *J. Chem. Soc.*, Chem. Commun., 1992, 1256.
- 2 Reviews: A. Katritzky and J. N. Lam, *Heterocycles*, 1992, **33**, 1011; J. Streith, *Chimia*, 1991, **45**, 65; see also C. R. Arnold, J. Melvin, W. M. Nelson and G. B. Schuster, *J. Org. Chem.*, 1992, **57**, 3316; R. Poe, K. Schnapp, M. J. T. Young, J. Grayzar and M. S. Platz, *J. Am. Chem. Soc.*, 1992, **114**, 5054.