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Cyclisations of Azidoformates. Cyclisation of Aryl Azidoformates

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Summary Phenyl azidoformates give benzoxazolones on 'spray pyrolysis' by direct nitrene attack at the *ortho*-position and α -naphthyl azidoformate gives a naphthoxazolone only by β -attack; biphenyl-2-yl azidoformate gives both 7-phenylbenzoxazolone and an azepine by nitrene attack of the adjacent ring while 2,6-dimethylphenyl azidoformate gives the *endo*-Diels-Alder dimer of 6-isocyanato-2,6-dimethylcyclohexa-2,4-dienone under the same conditions.

WE recently demonstrated that benzyl azidoformates decompose to yield oxazoloazepines and subsequently dimers therefrom by intramolecular nitrene attack.¹ We herein report a preliminary study of the intramolecular nitrene reactions of aryl azidoformates.

R
$$\frac{3}{4}$$
 $\frac{2}{5}$ $\frac{0}{6}$ $\frac{300 - 350 \text{ °C}}{0.1 - 0.2 \text{ mmHg}}$ R $\frac{6}{5}$ $\frac{7}{4}$ $\frac{0}{N}$ CO

(1) a; R = H
b; R = Me
c; R = Cl
d; R = Ph

In an unpublished observation,² German workers noted that phenyl azidoformate (1a) undergoes vapour phase pyrolysis to give benzoxazolone (2a), in high yield. We confirm this result using our 'spray pyrolysis' technique^{1,3} and note that the reaction does not involve a spìro-intermediate (3)⁴ since 4-substituted phenyl azidoformates (1b) and (1c) give the corresponding unrearranged benzoxazo-

lones (2b) and (2c), in good yields. Thus, 4-methylphenyl azidoformate (1b) gave the known⁵ 5-methylbenz-oxazolone (2b) (98%) as confirmed by unambiguous synthesis from the corresponding aminophenol and phosgene. α -Naphthyl azidoformate (4) gave solely the product of β -attack (5; m.p. 237—239 °C, 50%) with no sign of the peri-derived product (6), which we have unambiguously synthesised from 8-amino-1-naphthol and phosgene.

Biphenyl-2-yl azidoformate (1d) has two potential sites for attack: (i) the vacant *ortho*-position and (ii) the 1,2-bond of the phenyl substituent. In fact both pathways are followed since two products (7) and (8) are isolated in 24

and 46% yield, respectively. The former, m.p. 185 °C, shows a typical NH and carbonyl absorption of a benz-oxazolone (3200br, 1765, and 1720 cm $^{-1}$) and an appropriate $^1\mathrm{H}$ n.m.r. spectrum.† The latter azepine (8), an orange crystalline solid (m.p. 86·5—88 °C) shows no NH absorption but a carbonyl signal (1760, 1720 cm $^{-1}$) in its i.r. spectrum and characteristic olefinic absorptions in its $^1\mathrm{H}$ n.m.r. spectrum [δ (CDCl3) 5·4—5·8 (m, 3H), 5·8—6·0 (m, 2H,

azepine ring protons) and 6.9-7.45 (m, 4H, Ar-H)]. The cyclic urethane (8) is rapidly cleaved in cold ethylamine in ether solution to give the azepine (9)‡ as a yellow solid in 90% yield (m.p. 121-122 °C) while acidic hydrolysis converts it into the 3H-azepine (10)‡ (yellow liquid).

† The europium shift reagent Eu(fod), with benzoxazolones appears to complex with the ring oxygen. With the biphenyl derivative (7), an equivalent pair of protons (assigned to 2'-H and 6'-H) with ortho- and meta-coupling are brought to lower field.

‡ For (9); ν_{max} (Nujol) 3400 and 3150br (NH and OH), 1640 cm⁻¹(CO); δ (CDCl₃) 0·92(t, CH₃), 3·13 (quint., CH₂), 4·77br (t, NH), δ ·0—6·6(m, 3H + 2H, azepine protons), 6·6—7·4(m, 4H, Ar-H), and 9·85(br, OH). For (10) ν_{max} (Nujol) 3600—2000br (OH), 2850 and 2925 (CH₂), 1600 (C=N), and 740 cm⁻¹ (o·C₆H₄); δ (CDCl₃) 2·93(d, CH₂, $J_{3.4}$ 7 Hz), 5·36(d of q, 4-H, $J_{4.5}$ 8 Hz), 6·2—6·55(m, 2H, 5-H and another), 6·7—7·1(m, 2H), 7·15—7·5(m, 2H), and 7·68(d of d, 7-H, J 8 and 12 Hz).

Finally, 2,6-dimethylphenyl azidoformate (11) shows another unexpected type of reaction, in which a dimer (13),

derived by endo-Diels-Alder dimerisation of the cyclohexadienone (12), is isolated as shown in the Scheme. Related dimeric cyclohexadienones have been noted particularly from Wessely oxidation of e.g. 2,6-dimethylphenols. Warm ethanol converts the bis-isocyanate (13) into the corresponding bis-urethane. The spectra of the isocyanate (13) are particularly definitive: ν_{max} (Nujol) 2250, 2220 (NCO), 1720 (CO), and 1680 cm⁻¹ (C=C); δ (CDCl₃ at 220 MHz): 1.38, 1.43, and 1.48 (3 \times s, 3-Me, 8-Me, and 10-Me), 1.88 (t, 5-Me, $J_{\text{Me-6}} = J_{\text{Me-7}} = 1.5 \text{ Hz}$), 2.87 (m, 7-H, $J_{\text{6.7}}$ 4.0 Hz), 2.96 (d of d, 2-H, $J_{\text{2.7}}$ 8.5 Hz, $J_{\text{1.2}}$ 1.5 Hz), 3.20 (d of t, 1-H, $J_{\text{1.11}}$ 6.5 Hz, $J_{\text{1.12}}$ 1.5 Hz), 5.64 (d of d, 12-H, $J_{\text{1.112}}$ 8.5 Hz), 6.28 (dd, 11-H), and 6.33 (br d, 6-H).

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⁴ Cf. 5-membered spiro-intermediates: J. I. G. Cadogan, Acc. Chem. Res., 1972, 5, 303. For 6-membered spiro-intermediates see ref. 3. ⁵ W. J. C. Burris, J. Am. Chem. Soc., 1949, 71, 1266.

⁶ This assignment corrects an erroneous statement in a lecture summary: O. Meth-Cohn, Heterocycles, 1980, 14, 1497.

⁷ For a review ('Cyclohexadienones') see A. J. Waring, Adv. Alicyclic Chem., 1966, 1, 129.