Benzoheterocycles via Aryne C-C Cyclisation: 2-Methylisoindole

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Summary 2-Methylisoindole is obtained in high yield by the reaction of N-methyl-2-chlorobenzylaminoacetonitrile with potassamide in liquid ammonia.

Methods used for the synthesis of isoindoles have recently been reviewed. 1 Most of those which involve simple stages rely upon prior formation and isolation of an isoindoline or a related cyclic compound, elimination from an isoindoline with the leaving group on nitrogen being an example. 1,2

Treatment of N-methyl-2-chlorobenzylaminoacetonitrile† in liquid ammonia with potassamide (4 equiv.) over 0.5 h and quenching with ammonium nitrate gave 2-methylisoindole (89%). The m.p., $^{3-5}$ u.v., 6 and n.m.r. spectra, and the 1,3,5-trinitrobenzene adduct were consistent with 2-methylisoindole.

Its identity was further confirmed by its reduction (NaBH₄-EtOH) to 2-methylisoindoline (identified by i.r., 4 n.m.r., and analysis). Reduction was only efficient with an acid (HCl) work-up, suggesting that the reduction is actually of a protonated species.‡

In the formation of 2-methylisoindole it is considered that cyclisation to 1-cyano-2-methylisoindoline occurs first followed by hydrogen cyanide elimination to yield the isoindole. It is not known whether cyanide is eliminated from an anion following proton abstraction from C-1 and proton shift (or direct abstraction from C-3) or is eliminated first to give a cationic isoindolenine followed by proton loss and rearrangement.

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† Prepared by treating 2-chlorobenzyl chloride with methylaminoacetonitrile.

[‡] Although acetic acid work-up of borohydride mixtures has been shown to cause generation of diborane (J. A. Marshall and W. S. Johnson, J. Amer. Chem. Soc., 1963, 28, 421, 595), because of the presence of ethanol and water in the reaction mixture we favour the suggestion made in the text.

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