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Chemistry of Reduced Pyrazines

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Summary The synthesis and reactions, including a novel isomerisation, of functionalised 1,2,3,4-tetrahydropyrazines, and their attempted conversion into 1,4-dihydropyrazines, are described.

VIRTUALLY no work has been reported on 1,2,3,4-tetrahydropyrazines. Likewise the 8π electron 1,4-dihydropyrazines have not been extensively studied and authentic derivatives of this system have become available¹ only since 1971, most of the earlier work having been shown to be erroneous.

$$SO_{2}Ar$$

$$N$$

$$ArSO_{2}NHCH_{2}CH(OR)_{2}$$

$$SO_{2}Ar$$

$$(2)$$

$$(1)$$

$$a; Ar = Ph$$

$$b; Ar = C_{8}H_{4}Me-p$$

$$R = Me \text{ or Et}$$

The alleged synthesis² of the 1,4-dihydropyrazine (1a) by cyclisation of the aminoacetal (2a) was shown by one of us³ to lead instead to the 1,2,3,4-tetrahydropyrazine (3a)[†] or the 2,5-dihydroxypiperazine (4a),[†] according to conditions. It was found that (4a) was readily dehydrated to (3a) and that both compounds reacted with hot acidic methanol to afford (3b). The latter compound had been prepared earlier.⁴

This work has now been extended to the preparation of the tosyl derivatives $(3c)^{\dagger}$ and $(4c)^{\dagger}$ which were found to be interconvertible by the addition and elimination, respectively, of water under acid conditions. The alkoxy derivatives $(3d,e)^4$ and $(4d,e),^4$ prepared from the aminoacetal (2b) and the corresponding alcohol, behaved similarly. The hydroxy group in (3c) could be replaced by an alkoxy group to give (3d) and (3e); the reverse reaction, replacement of alkoxy by hydroxy, was also achieved. Reaction of (3c) with benzenethiol in acidified acetone afforded the sulphide $(3f)^{\dagger}$ which was oxidised to the sulphoxide $(3g)^{\dagger}$ but could not be converted into the sulphone.

Treatment of (3c) or (4c) with thionyl chloride yielded the highly insoluble (4h);[†] on brief heating in toluene this was converted into (3h).[†] The chlorine atom in the latter was very labile as expected. Thus, reaction of (3h) with water or ethanol gave (3c) and (3e), respectively. A number of nucleophilic substitutions were carried out on (3h) in aprotic solvents to afford the new compounds (3i-m).[†] Attempts to prepare the corresponding iodide or cyanide failed.



The action of bromine on (3c) or (3h) led to additionelimination with the formation of (5a); the intermediate piperazine is presumably too hindered to be isolated. Reaction of (5a) with water and with ethanol readily afforded (5b); and (5c), respectively. Treatment of (5a)with magnesium in dry tetrahydrofuran (THF) led to a

† Satisfactory elemental analyses and spectral data were obtained for all new compounds.

very labile isomer to which the cis-configuration (5d) was assigned. Such an isomerisation appears to be without precedent. The dibromide (5d) was converted into (5e)† and (5f).[†] The stereochemistry of the trans-compounds (5a-c) follows from their mode of preparation. Further evidence was provided by the fact that the cis-diol (5e), but not the trans-diol (5b), reacted⁵ with Me₂NCH(OMe)₂, and that the chemical shift of C-2 and C-3 in the ¹³C n.m.r. spectrum of the trans-compound (5c) was at lower field than that of the cis-isomer (5f) (cf., e.g., the isomeric 1,2-dichlorocyclohexanes⁶).



 $Ts = MeC_{6}H_{4}SO_{2}-p$

The tetrahydropyrazines (3) did not undergo [4+2]cycloaddition with dienes; however, (3m) reacted with tetrachloro-o-benzoquinone to yield the novel ring system

(6).^{\dagger} Oxidation of (3c) and (3m) with Jones reagent surprisingly resulted in hydroxylation of the double bond with the formation of 2,3,5-trihydroxy- and 2,3-dihydroxy-1,4-bis(p-toluenesulphonyl)piperazine,† respectively. All the above reactions point to the high electron density of the double bond in 1,2,3,4-tetrahydropyrazines.

Numerous attempts were made to convert (3) or (5) into the 1,4-dihydropyrazine (1b). Dehydration of (3c), dehydrohalogenation of (3h) or (5d), thermolysis of (3g), and debromination of (5a) all failed. However, reaction of (5a) with Zn/Cu couple in dry THF gave a highly unstable product, presumably (1b), which could not be isolated but which on treatment with water afforded (3c). Oxidation of the selenide (31) with m-chloroperoxybenzoic acid under aprotic conditions yielded (3n); with aqueous hydrogen peroxide (31) gave (3c), and with methanolic KIO_4 it afforded (3d). These reactions presumably proceed via the unstable selenoxide which fragments to (1b); this compound is clearly highly reactive, possibly due to its antiaromatic nature, and adds even quite weak electrophiles (m-ClC₆H₄CO₂H, H₂O, or MeOH) as soon as it is formed, giving rise to the observed products. The addition of methanol to a 1,4-dihydropyrazine has been described previously.18

With one exception⁷ all the known¹ 1,4-dihydropyrazines carry aryl substituents in the ring which presumably stabilise the system by alternative modes of conjugation. This may explain the failure to isolate (1b). Experiments to synthesise this compound are continuing.

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