## ON THE STRUCTURE OF CATALINE

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Eu(fod)<sub>3</sub> further supports structure I proposed for this alkaloid. This was<br>confirmed by an X-Ray analysis.

5-Hydroxyberberines and 4-hydroxyaporphines present an interesting problem concerning the biogenetical origen of the benzylic hydroxy group. To find out the configuration of these alkaloids **1** at the hydroxylated center is in this sense of utmost importance<sup>1</sup>.<br>Regarding the 4-hydroxyaporphines, Ribas et al.<sup>2</sup> have proposed the absolute stereochemistry of

cataline as indicated in the structural formula I, in which the hydroxy group is pseudoaxial and the configuration of the hydrogens at C-4 (pseudoequatorial) and C-6a is cis. Cataline has later been taken as a reference compound in proposing the configuration at the benzylic hydroxy center of other 4-hydroxyaporphines<sup>3</sup>.

Nevertheless, Cassels et al.<sup>4</sup> found the assignement of the configuration at C-4 proposed by Ribas et al. for cataline(I)highly speculative, so that the stereostructures of other 4-hydroxyaporphines would also be dubious. The proposal by Ribas' group<sup>2</sup>was based mainly on the appearance of the proton at C-4 in the pmr spectrum of cataline and its O-acetyl derivative as a poorly resolved triplet with half-band width 4Hz. According to Cassels' group, Dreiding models built in the  $4(S) : 6a(S)$  and  $4(R) : 6a(S)$  configurations with pseudoequatorial C-4 protons do not suggest any clear differences as far **as** potential energy is concerned. Therefore, any of these two stereostructures could be suggested for cataline from the above mentioned **pmr** data.

since C-4 and C-6a protons must be cis in the  $4(R):6a(S)$  configuration and trans in the 4(51:6aIS), a **pmr** study has now been carried out on cataline in the presence of an Europium shift reagent. This forms a complex with the C-4 hydroxyl and therefore must affect the chemical shift of the C-6a proton only in the case of the  $4(S):6a(S)$  configuration.

The pmr spectrum of cataline(I) in the presence of Eu(fod) $_3^5$  revealed the most significant displacements to low field, apart from the expected strong shift of the C-4 proton, two broad doublets (slopes: m=9.0 and 5.0). They were proved to be strongly coupled together (J=12Hz) by spin decoupling experiments. Cataline-d<sub>3</sub> (deuterated at C-6a and C-7)<sup>6</sup>, in the presence of Eu(fod)<sub>2</sub> showed the same two doublets, thus proving that those signals correspond to C-5 protons. The absence of any other signal moving downfield which could be due to H-6a rules out in the case of cataline the 4lS):6a(S) configuration which has a half-boat conformation of ring **8.** rurthennore. high temperature **pmr** spectra of cataline **gave** no evidence of a conformational change in ring B. This would convert the C-4 proton of cataline from pseudoequatorial to pseudoaxial and would result in a shift and a change of the multiplicity of its signal which consequently would be identical to that of epicataline(1II where C-4 proton is pseudoaxial. We have found that C-4 proton in epicataline(II) resonates at  $\delta$  4.90 ppm<sup>7</sup> as a doublet of doublets while in cataline



appears at 4.47 ppm as a poorly resolved triplet. Therefore, these results confirm the  $4(R):6a(S)$ configuration for cataline.

This conclusion is also supported by comparison of the nmr data for C-4 and C-3 protons and  $C-3$  carbon ( $^{13}$ Cnmr) of cataline and epicataline with the corresponding values that were reported for the other known 4-hydroxyaporphines<sup>9,10</sup>, 5-hydroxyberberines<sup>1</sup> and benzylisoquinolines<sup>11</sup>. This shows a good correlation. **me** shift values of the C-4 proton signals for cataline and epicaraline on  $\arctan(1.37$  and 1.12 ppm, respectively)<sup>8</sup> agree also with the configuration proposed by Ribas' group. This structure was confirmed by an X-Ray analysis.

Cataline(C<sub>21</sub>H<sub>25</sub>O<sub>5</sub>N) crystallizes in space group P2<sub>1</sub>2<sub>1</sub><sup>2</sup><sub>1</sub><sup>2</sup><sub>1</sub><sup>2</sup>  $z=4$ , with a=7.3558(9), b=14.869(3), c=17.410(4)  $\lambda$ , D<sub>r</sub>=1.2994 g.cm<sup>-3</sup>. Intensities of 1989 independent Friedel pairs were measured up t O-68" on an automatic four-circle diffractometer. Graphite-monochromated CuKu radiation(l.5418 A1 and w/20 scan technique were used. 1822 Friedel pairs were considered as observed according to the criterion 1>20(11. The absorption effect was ignored and the Lorentz and polarization factors were applied as usual. After these corrections were made the above 1822 mentioned Friedel pairs were used in the calculations<sup>12</sup>. The structure was solved by direct methods and the hydrogen atoms were located on a difference map. An **app~opiateweightingscheme'~ was** selected in order not to give any trends of <wA<sup>2</sup>F>vs.<F<sub>0</sub>> and <sin0/ $\lambda$ >. These weights were applied on several cycles of block-diagonal matrix mixed refinement including both hkl and hkl reflexions, which converged for the right enantiomer to the discrepancy indices R=0.036 and  $R_w=0.045$ . The anomalous effect of the oxygen atoms was used to confirm the absolute configuration of the molecule, shown in Fig. 1.

The values of bond distances and bond angles are in good agreement with those expected for this kind of compounds. Ring B presents half-chair conformation at C(3a)-C(6b) and ring C a diplanar one at C(6b)-C(11b) and C(7a)-C(11a) respectively. The O(4) and H(6a) atoms are placed in trans position. The torsion angle through the ideal bond  $C(4) \ldots C(6a)$  being -161(2)°.

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