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ISOLATION AND STRUCTURE OF SUNGUCINE: A NEW TYPE OF BISINDOLINE ALKALOID

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Abstract: From the roots of Strychnos icaja Baill. an unsymmetrical dimeric alkaloid has been isolated and called sungucine. Its original structure has been established by X-Ray diffraction; UV, IR, MS and NMR data are also given.

Strychnos icaja roots are used in Central Africa for the preparation of arrow and ordeal poisons 1. Early investigations have shown that these roots contained the well-known strychnine and its 12 hydroxy-derivative amongst many other bases 2. We now record the isolation, characterization and structure determination of another major alkaloid: sungucine 3.

The dried root barks were extracted with EtOH, the extract concentrated and the resulting solution made acetic. This acidic solution was then extracted with CHCl<sub>3</sub> and has furnished tertiary crude bases (yield: 6%). After purification by column chromatography over  $\text{Al}_2\text{O}_3$  (activity II-III) and elution with  $\text{Et}_2\text{O-C}_6\text{H}_6$ , this main alkaloid was isolated by crystallization in  $\text{Me}_2\text{CO-CHCl}_3$  as prisms (yield: 2%); m.p. > 350°C.

High resolution mass measurement of the new alkaloid suggested its molecular formula as  $^{\text{C}}_{42}$   $^{\text{H}}_{42}$  $^{\text{N}}_{40}$  [found: 634,3249; required: 634,3306].

X-Ray crystallography established the relative configuration of sungucine. Crystals of  $C_{42}$   $H_{42}N_4O_2 - C_3H_6O$  were orthorhombic, space group  $P2_12_12_1$ , with 4 moleculesina unit cell of dimensions a = 13.779(5), b = 24.516(6), c = 10.640(4) Å. The structure was refined to a final R value of 0.05 for 3.394 reflexions.

The absolute stereochemistry is based on the assumption that the 15  $\alpha$  configuration for C<sub>15</sub>H and C<sub>15</sub>H agrees with the biosynthetic hypothesis<sup>5</sup>. From the X-Ray diffraction data<sup>6</sup> (about cis or trans relationship between asymmetrical carbons), the complete absolute stereochemistry of sungucine could be depicted as that shown overleaf.

Sungucine represents a new type of unsymmetrical bis-indoline alkaloid that is very different from other dimeric Strychnos alkaloids, e.g. toxiferines and strychnobilines which are mainly derived from Wieland-Gumlich-like aldehydic alkaloids. Indeed, sungucine has an original  $C_{23}-C_{5}$ , bond between the two parts of the molecule.

The identification of sungucine can now be performed by spectroscopic methods. The MS with peaks at m/e (relative abundance in % of base peak): 634(95),527(17),512(16),437(37),384(10), 360(11),345(47),317(80),290(54),275(61),251(42),220(38),210(22),196(29),180(18),167(18),144(53),

143(13),135(91),122(98),121(base peak),108(75), supports the structure determination. The UV spectrum  $\left[\lambda_{nm}^{EtOH}(\log \xi) 218(4,87),265(3,84),292(4,02)\right]$  and 305(4,01) indicates a new chromophoric system. The IR spectrum  $\left[\nu_{max}^{KKBr} 3040,2950,2850,1710\right]$  (Me<sub>2</sub>CO of crystallization),1670, 1600,1480,1460,1420,755 cm<sup>-1</sup> shows bands which can be assigned to the N<sub>a</sub>-acyldihydroindole (1670) and ortho-disubstituted benzene ring (755). The NMR spectrum (in CDCl<sub>3</sub>) shows characteristic signals at  $\frac{1}{2}$ 8.28 and 8.03 (d;J=8Hz;H<sub>12</sub>-H<sub>12</sub>,),6.02(d;J=10Hz;H<sub>23</sub>,),5.47 and 5.24(q;J<sub>1</sub>=14Hz,J<sub>2</sub>=7Hz;H<sub>19</sub>-H<sub>19</sub>),4.38 and 4.35 (d;J=7.5Hz;H<sub>2</sub>-H<sub>2</sub>,),1.79 and 1.65(d;J=7Hz;Me<sub>18</sub>-Me<sub>18</sub>).

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## References and complementary notes

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