

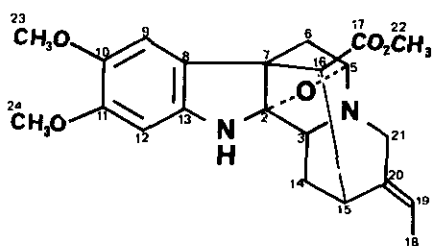
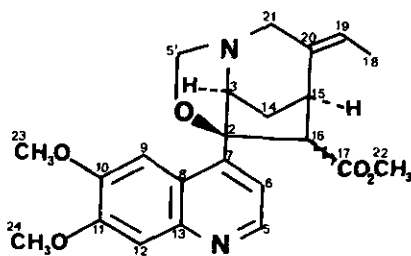
CORIALSTONINE, A NOVEL QUINOLINE ALKALOID FROM *ALSTONIA CORIACEA*

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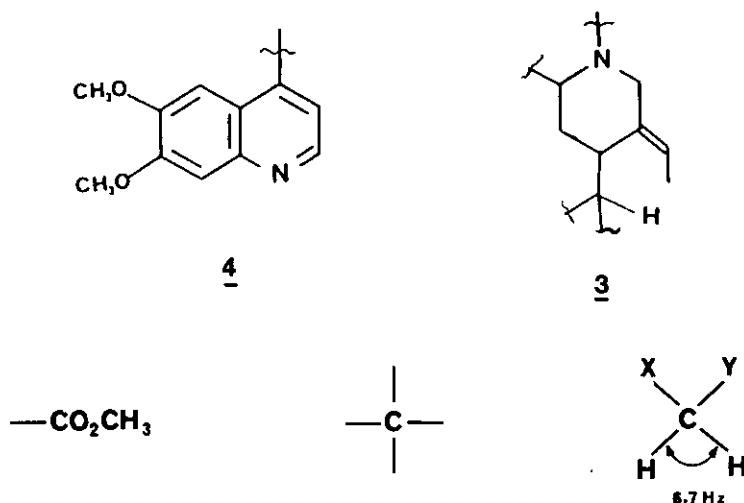
**Abstract** — Corialstonine is a novel quinoline alkaloid isolated from *Alstonia coriacea*. Its structure illustrates a possible new transformation of indoles into quinolines.

The propensity of some indoles to transform themselves into quinolines is exemplified by the *Cinchona*<sup>1</sup> or *Camptotheca*<sup>2</sup> alkaloids and to a lesser extend by lanceomigine from *Alstonia* and *Hunteria* species<sup>3</sup>. We wish to describe a novel quinoline alkaloid, corialstonine **1**, which also illustrates an alternative means of transposition of indoles into quinolines.

**2****1**

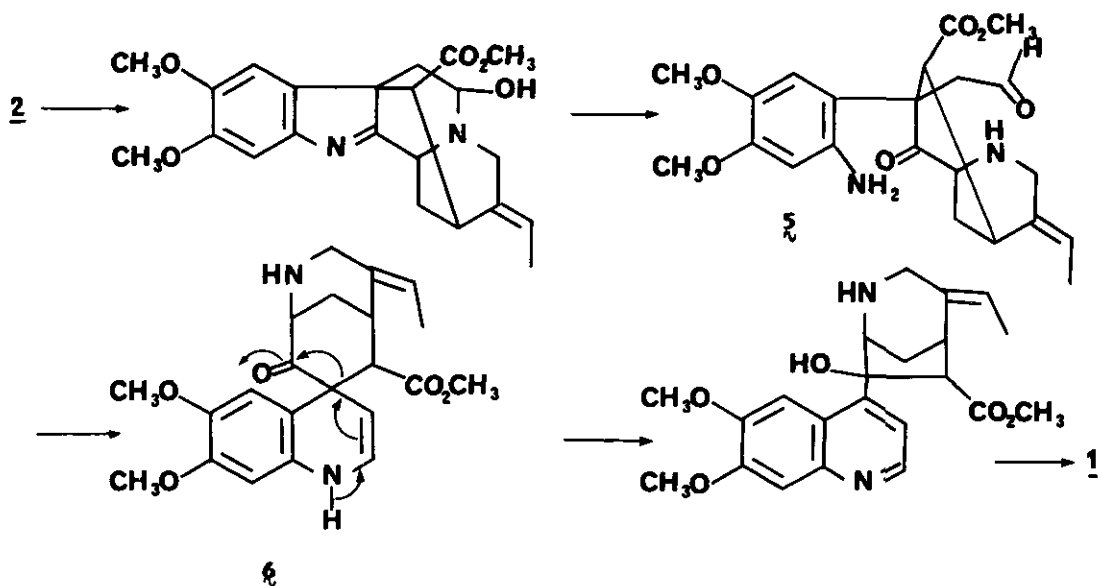
*Alstonia coriacea* is a shrub from New-Caledonia<sup>4</sup> whose stem bark main alkaloid is nor-quaternine **2**<sup>5</sup>. It is accompanied by corialstonine<sup>6</sup>, an amorphous base which is slightly fluorescent on TLC ( $\alpha$ )<sub>D</sub> +102°, CHCl<sub>3</sub>, C=1). Corialstonine does not react with the Ce-IV spray but stains orange after Dragendorff pulverization. Its uv spectrum does not resemble those of the usual indole alkaloids ; two maximum appear at 317 and 330 nm, which are shifted at 354 nm upon acidification : the ms of **1** displays molecular ion at  $m/z$  410, analyzed for C<sub>23</sub>H<sub>26</sub>N<sub>2</sub>O<sub>5</sub> (410.176 ; calc:410.184) ; it is accompanied by an  $m/z$  424 probably due to transmethylation.

$^1\text{H}$  nmr and  $^{13}\text{C}$  nmr spectra for 1 were obtained at 300 and 75MHz respectively. They were uneventfully interpreted by means of homonuclear ( $^1\text{H}$ - $^1\text{H}$ ) and heteronuclear ( $^1\text{H}$ - $^{13}\text{C}$ ) correlated spectroscopy<sup>7</sup>. It is thus found an isolated eight-carbon system 3 corresponding to the core of all type-I indole alkaloids<sup>8</sup> and a quinoline 4. Other systems are a methoxycarbonyl unit, a quaternary carbon and an isolated methylene with an unusually low geminal coupling constant ( $J=6.7\text{Hz}$ ) ; worthy of note is the absence of "tryptamine-like"  $\text{CH}_2\text{-CH}_2$  protons.



Structure 1 was finally deduced from biogenetic considerations and from observation of long range  $^1\text{H}$ - $^1\text{H}$  and  $^1\text{H}$ - $^{13}\text{C}$  couplings. Of particular value are the correlations of and across quaternary carbons such as :  $\text{C-2} \rightarrow \text{H-6}$  ( $3_J$ ) ;  $\text{H-6} \rightarrow \text{H-3}$  ( $5_J$ ) ;  $\text{C-3} \rightarrow \text{H-5'}$  ;  $\text{C-2} \rightarrow \text{H-14}$ . These experiments have also allowed complete assignment of both proton and carbon spectra including methoxyls and quaternary carbons. Configurations of C-15 and of the 19-20 double bond are assumed to be the "biogenetic" configurations ; stereochemistry of the methoxycarbonyl group has not been determined.

From biosynthetic standpoint, 1 may arise from 2 via keto-aldehyde 5 resulting from opening of the carbinolamine ether, ring closure of a six-membered ring 6 and 1,2 carbon shift whose driving force would be aromatization of the quinoline system. Natural or artificial origin of C-5' is still questionable . Other pathways originating from an earlier biosynthetic intermediate such as geissoschizine may also be envisioned ; in this latter case, it is possible to propose mechanisms in which C-5' would be C-17 of the precursors.



|      | <b>1</b> | <b>2</b> | : |                                 | <b>1</b> | <b>2</b> |
|------|----------|----------|---|---------------------------------|----------|----------|
| C-2  | 89.7     | 109.7    | : | C-14                            | 31.4     | 21.6     |
| C-3  | 73.4     | 49.4     | : | C-15                            | 40.2     | 33.2     |
| C-5  | 146.9    | 86.9     | : | C-16                            | 59.4     | 57.9     |
| C-6  | 115.2    | 44.7     | : | C-18                            | 12.9     | 13.1     |
| C-7  | 146.5    | 52.5     | : | C-19                            | 119.5    | 120.0    |
| C-8  | 122.2    | 123.4    | : | C-20                            | 135.8    | 137.8    |
| C-9  | 107.9    | 111.2    | : | C-21                            | 53.1     | 46.6     |
| C-10 | 152.0    | 143.5    | : | C-5'                            | 89.2     | -        |
| C-11 | 148.9    | 145.6    | : | Ar-OCH <sub>3</sub> -23         | 55.9     | 57.0     |
| C-12 | 105.8    | 95.2     | : | Ar-OCH <sub>3</sub> -24         | 55.7     | 57.7     |
| C-13 | 148.0    | 150.0    | : | CO <sub>2</sub> CH <sub>3</sub> | 50.8     | 51.6     |
|      |          |          | : | CO <sub>2</sub> CH <sub>3</sub> | 169.2    | 174.5    |

<sup>13</sup>C nmr data for **1** and **2** (CDCl<sub>3</sub>, 75 MHz)

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3. J.Vercauteren, G.Massiot, L.Le Men-Olivier, J.Lévy, T.Prange, and C.Pascard,  
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H.E.Saad, R.Anton, J.C.Quirion, G.Chauvière and J.Pusset, submitted to Tetrahedron  
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6. Corialstonine (complementary data) : uv :  $\lambda_{\text{max}}$  (MeOH) : 218, 238, 317, 330 nm ;  
(MeOH + HCl) : 220, 246, 354 nm ; ir (CHCl<sub>3</sub>) : 1745, 1620, 1580, 1500, 1480, 1430,  
1345, 1250, 1160 cm<sup>-1</sup> ; ms (electron impact) : m/z (rel.int.) : 424(5), 410(30),  
258(10), 188(12), 135(25), 122(40), 121(100) ; ms (NH<sub>3</sub> chemical ionization) :  
425(30), 411(100) ; <sup>1</sup>H nmr (CDCl<sub>3</sub>, 300MHz) : 8.65(d, J=4.8Hz, H-5), 8.32(s, H-12),  
7.4(s, H-9), 7.08(d, J=4.8Hz, H-6), 5.4(br q, J=7Hz, H-19), 4.7(d, J=6.7Hz, H-5'), 4.35  
(d, J=6.7Hz, H-5'), 4.3(d, J=4.1Hz, H-3), 4.02(s, 3H, OCH<sub>3</sub>-24), 4.00(s, 3H, OCH<sub>3</sub>-23),  
3.95(dq, J=16.5, 2.4Hz, H-21), 3.65(br s, H-15), 3.45(s, 3H, CO<sub>2</sub>CH<sub>3</sub>), 3.2(br d, J=16.5Hz,  
H-21), 2.85(d, J=5.1Hz, H-16), 2.5(d, J=13.2Hz, H-14), 2.1(dt, J=13.2, 4.1Hz, H-14), 1.5  
(dd, 3H, J=7, 2.4Hz, CH<sub>3</sub>-18).
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Received, 6th July, 1987