

STEREOCHEMISTRY OF THE DIBENZOPYRROCOLINE ALKALOIDS
 CRYPTAUSTOLINE AND CRYPTOWOLINE

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Abstract—Stereochemistry of the dibenzopyrrocoline alkaloids, (-)-cryptaustoline (1) and (-)-cryptowoline (2), isolated from the Australian shrub *Cryptocaria bowiei* is deduced both as 7R,13S configuration with *cis* B/C ring structure based on the analysis of proton-nmr spectra of synthetic materials.

In 1952 Ewing, Hughes, and co-workers reported the isolation of two unique quaternary alkaloids, (-)-cryptaustoline (1) and (-)-cryptowoline (2), from the Australian shrub *Cryptocaria bowiei* (Figure 1). These are only known examples of this class of alkaloids² and are believed to be generated in nature from a 1-benzylisoquinoline precursor by intramolecular phenolic oxidative coupling to form the bond between 7-nitrogen and aromatic 7a-carbon.³ Stereochemistry of two alkaloids, however, has been determined only at C-13 center as S,⁴ while both the relative and the absolute configuration of the quaternary 7-nitrogen center remain

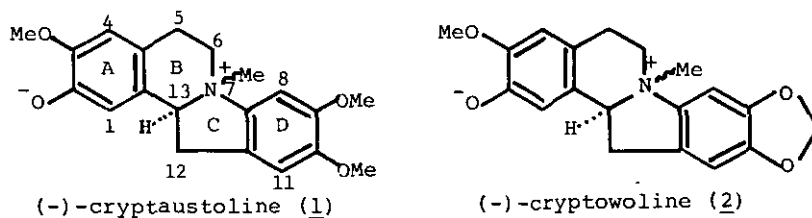


Figure 1

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