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Total Synthesis of 3,14-Dihydroxyisomorphinans and 9α-hydroxy-3methoxyhasubanans

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Summary Total synthesis of 3,14-dihydroxyisomorphinans and 9α -hydroxy-3-methoxyhasubanans via 4a-(2-aminoethyl)-1,2,3,4,4a,9-hexahydro-6-methoxyphenanthrene (Ia) is described.

MUCH attention has been paid recently to the 14-hydroxysubstituted morphine derivatives¹ and their synthetic congeners² as potentially useful pharmacological agents. The corresponding 14-hydroxy-B/C-trans-fused isomers have not been reported, even though potent analgesic³ and narcotic antagonist⁴ activities have been uncovered among some of the isomorphinan structures.

We now report on the total synthesis of 3,14-dihydroxyisomorphinans and 9α -hydroxy-3-methoxyhasubanans via a common intermediate, the unsaturated amine (Ia), which has already been successfully employed in the synthesis of 3,14-dihydroxymorphinans.²

Treatment of (Ia) with ethyl chloroformate-triethylamine afforded the unsaturated urethane (Ib).[†] Oxidation of (Ib) with *m*-chloroperbenzoic acid afforded stereoselectively

[†] Satisfactory elemental analyses and n.m.r. and i.r. spectra were obtained for all new compounds.



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the α -epoxide (II). This was treated with sodium t-pentoxide in boiling benzene to effect intramolecular regioselective opening of the epoxide, and concomitant ring D formation, thus affording the 14-hydroxyisomorphinan skeleton (IVa). Reduction of (IVa) with LiAlH₄ afforded the N-methyl compound (IVb). Hydrolysis of (IVa) with KOH in boiling octan-1-ol afforded 3-methoxy-14-hydroxyisomorphinan (IVc) as an oil, HCl salt, m.p. 269-271°, which was further transformed to the various N-substituted 3,14-dihydroxyisomorphinans by well defined routes.²⁻⁴. The overall yield of (IVc) from (Ia) was 55%.

Alternatively (Ib) was reduced with LiAlH₄ to (Ic) which was then converted into the amide (Id). Oxidation of (Id) with *m*-chloroperbenzoic acid to the α -epoxide (III), and treatment of this with K₂CO₃ in MeOH-H₂O afforded a 1:1 mixture of (IVb) and 9a-hydroxy-3-methoxyhasubanan (V) in an overall yield of 48% from (Ia). The mixture was easily separated by column chromatography.

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