Indole Alkaloids. Enantioselective Synthesis of (-)-Alloyohimbane by a Chemoenzymatic Approach

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The pro-(R) enantiotopic specificity of pig liver esterase-catalysed hydrolysis of the meso-diacetate (2) to give (3) enables this chiron to be used in a highly efficient enantioselective synthesis of the natural (-)-alloyohimbane (1).

There has been considerable interest in yohimbinoid alkaloids (e.g. reserpine, yohimbines) both therapeutically and synthetically, although to date only a few reports deal with the total synthesis of optically active alkaloids embodying this skeletal arrangement. Within this context, Isobe et al. have just disclosed the first enantioselective synthesis of (–)-alloyohimbane (1), based on a chiron approach utilising carbohydrates as immolative chiral auxiliaries. Here we report an alternative and concise route to (–)-(1) involving as the key step a process resulting in dissymmetry (σ - \rightarrow C₁-symmetry) in the mesodiacetate (2).

Our synthesis required the hydroxy ester (1S,2R)-(3) as the pivotal intermediate and this can be produced³ on a preparative scale in good yield (78%) with 96% enantiomeric excess (e.e.) by the pro-(R) pig liver esterase (PLE)-catalysed hydrolysis³ of (2)† to provide the proper chirality at two of the three stereogenic centres of (1). The hydroxy ester (3) was then converted into (-)-(4aR,8aS)-tetrahydroisochroman-3-one (4) (64%) $\{[\alpha]_D^{20} - 5.4^{\circ}$ $(c 2, CHCl_3); \nu_{max}$ 1725 cm⁻¹; δ 4.32 (2H, d, J 3.8 Hz) and 5.65 (2H, m, olefinic protons), \geq 95% e.e. (determined according to the method of Jakovac and Jones⁴)} νia a straightforward sequence [(i) triflic

anhydride, pyridine, CH_2Cl_2 , $-30\,^{\circ}C$; (ii) sodium cyanide, dimethyl sulphoxide (DMSO), $50\,^{\circ}C$; (iii) MeOH-20% aq. NaOH (5:1), 36% hydrogen peroxide, 60°C, then acidic

$$(1) \qquad (2) \quad X = OAc$$

$$(3) \quad X = OH$$

[†] The presence of the double bond in (4) seems to be essential for such a remarkable e.e.

work-up]. Acylation of tryptamine with (4) in refluxing xylene, followed by exposure of the resulting hydroxy amide‡ to N,N-sulphuryl diimidazole in N,N-dimethylformamide (DMF),⁵ then adding sodium hydride at $-40\,^{\circ}\text{C} \rightarrow 0\,^{\circ}\text{C}$, led to the crystalline lactam (5) {81% yield from (4), m.p. 154 °C; $[\alpha]_D^{20}-19.4^{\circ}$ (c 1, CHCl₃); δ 2.33 and 2.43 (2H, AB part of ABX pattern, J 18.0, 6.0, 5.5 Hz, diastereotopic $CH_2\text{CON}$) and 5.57 (2H, m, olefinic protons)}. This was then sequentially converted into (–)-alloyohimbane (1), m.p. 156 °C, $[\alpha]_D^{20}-165.9^{\circ}$ (c 0.5, pyridine) (lit.,⁶ $-166.5^{\circ} \pm 0.8$) as previously reported.²

In summary, the overall efficiency for this synthesis compares favourably with that recorded by Isobe et al. and demonstrates the great potentiality of chemo-enzymatic

methodology for the preparation of optically pure compounds.

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[‡] In our hands elaboration of this intermediate according to Isobe's protocol (ref. 2) gave noticeably poorer yields of (5).