Simple Syntheses of (S)-2- and 4-Amino-5-hydroxypentanoic Acids†

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The title compounds were efficiently prepared by selective reductions of α - and γ -methyl (S)-N-tritylglutamates with LiAlH₄

(S)-2-Amino-5-hydroxypentanoic acid (L- δ -hydroxynorvaline) (4), the next higher homologue of homoserine, and (S)-4-amino-5-hydroxypentanoic acid (8) are competitive inhibitors of γ -cystathionase¹ and γ -aminobutyric acid aminotransferase,² respectively. Furthermore, the amino acid (4) acts as the biological precursor of the polyoxins (nucleoside peptide antibiotics), in which its whole carbon skeleton is found intact,³ and it is also incorporated into the oxazolidine segment of clavulanic acid,⁴ a potent inhibitor of bacterial β -lactamases.

Both compounds (4) and (8) have been prepared previously in low overall yields by rather lengthy and tedious or wasteful synthetic routes.^{2a,5} We now report exceptionally simple syntheses which provide (4) and (8) in good overall yields and excellent purity from commercially available γ -methyl (S)-

glutamate (1) or dimethyl (S)-glutamate hydrochloride (9). In all the synthetic transformations the bulky triphenylmethyl (trityl) group was chosen for α -amino protection for the following reasons: (a) it is easily introduced, and removed by mild acid treatment in excellent yields, 6 (b) it completely suppresses reduction of the α -carboxy function of amino acids by LiAlH₄,7b (c) it offers excellent racemisation resistance even in the case of strongly activated chiral amino acid derivatives, 8 and (d) in contrast to protecting groups of the urethane type, 7a it is compatible with complex metal hydrides. 7b

Our synthetic route to (4) initially involves the tritylation of (1) by the trimethylsilyl ester procedure. 6a The product (2) obtained as the diethylammonium (DEA) salt {m.p. 155—156 °C, $[\alpha]_D^{25}$ +13.7° (c 1, MeOH)} in 95% yield, was further reduced as such with LiAlH₄ in tetrahydrofuran at 0 °C to afford N-trityl- δ -hydroxynorvaline (3), also isolated as the corresponding DEA salt {m.p. 134—135 °C, $[\alpha]_D^{25}$ -14.3° (c 2, MeOH)}, in 87% yield. The reduction proceeds absolutely selectively at the ester function as evidenced by the total

[†] All optically active amino acid derivatives referred to in this communication are of the S-configuration. New compounds gave analytical and spectroscopic data in agreement with the proposed structures.

Scheme 1. Reagents: i, Me₃SiCl/Et₃N; ii, Ph₃CCl/Et₃N; iii, MeOH; iv, LiAlH₄/THF; v, AcOH-H₂O; vi, 1-HOBt, dicyclohexylcarbodi-imide; vii, NaBH₄/(MeO[CH₂|₂)₂O, 0°C; viii, 2M NaOH/MeOH, room temp.

absence of by-products shown by h.p.l.c. Finally, detritylation with aqueous 95% acetic acid at ambient temperature gave, after recrystallisation from aqueous ethanol, the product (4) {m.p. 231.5 °C, $[\alpha]_D^{25}$ +22.6° (c 2, 0.5m HCl) (lit.5m m.p. 220—220.5 °C), $[\alpha]_D^{25}$ +28.2 (c 1.9, 6m HCl)} in 82% yield.

The 4-amino isomer (8) {m.p. 167—168 °C, $[\alpha]_D^{25}$ +24.2° (c 2, H₂O) (lit., 2a m.p. 147-148 °C, no figure for $[\alpha]_{D}^{25}$) was prepared, in 72% overall yield based on (9), in an analogous manner by reduction of α -methyl (S)-N-tritylglutamate (11) followed by detritylation. As depicted in Scheme 1, (11) was prepared by tritylation of (9) followed by selective saponification. Alternatively, the reduction product (7) was obtained in 77% overall yield based on (2) by converting⁹ (2) into the corresponding oily benzotriazolyl derivative (5), followed by selective reduction with NaBH4 and saponification of the resulting methyl 4-tritylamino-5-hydroxypentanoate (6). Neither of the reductions leading to (7) was as selective as that of (3), as evidenced by minor less polar by-products (h.p.l.c.) accompanying its formation. However these by-products are easily separated during crystallisation of the corresponding DEA salt of (7) {m.p. $122 \,^{\circ}$ C, $[\alpha]_{D}^{25} + 42.7 \,^{\circ}$ (c 1, MeOH)}.

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