Enantioselective Synthesis of *(S)-* **and (R)-6-(2,3-Dihydroxypropyl)-l,3-dioxin-4-ones: the Versatile Building Blocks of Four- and Six-carbon Backbones**

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The Sharpless asymmetric epoxidation of 2,2-dimethyl-6-(3-hydroxy-l -propenyl)-I ,3-dioxin-4-one using titanium tetraisopropoxide-diisopropyl tartrate followed by catalytic hydrogenation affords the title compounds as enantiomerically pure compounds, which act as versatile four- and six-carbon building blocks.

1,3-Dioxin-4-ones act as versatile synthons in organic synthesis.1 We have been interested in synthesizing 6-(2,3-di**hydroxypropyl)-l,3-dioxin-4-one 1** by focusing our attention on the utilization of the dioxinone moiety as the corresponding (3-keto acid and acyl ketene equivalents (Scheme 1). Once **1** is synthesized, the following transformations may be expected. Thus, while oxidative cleavage (path *a)* affords **2,** hydrolysis at the acetal function (path \overline{b}) leads to 3. Furthermore, by knowing that the 6-electron cycloreversion (by heating2 or $irradiation³$ at 254 nm) of the dioxinones to acylketenes (path c) takes place readily, their manipulation either to heterocycles **4** by hetero-Diels-Alder reaction4 or to inter-5 and intra-molecular ketene trapping6 compounds by nucleophiles (e.g. formation of *5)* should be expected.

Using readily available 6-methyl derivative **71-** as the starting material, **(S)-1** was synthesized as an enantiomerically pure compound. Though reaction steps are longer (5-step), all reactions except for the first one *(ca. 65%)*‡ proceeded in nearly quantitative yields and are suitable for large-scale preparation. Thus, base-catalysed chloroacetylation to **8** followed by sodium borohydride reduction gave **9** (racemic) . Treatment of **9** with NaOH-ether *(2* mol dm-3) at room temperature gave the ally1 alcohol **11** as the sole product. Presumably, the epoxide **10** was formed first, which was then cleaved to the diol. Epoxidation7.8 of **11** by employing tert-butyl hydroperoxide (TBHP) as an oxygen donor and titanium tetraisopropoxide-diisopropyl D-(-)-tartrate

? This compound is known as diketene-acetone adduct and is commercially available.

(DIPT) as the catalyst, in presence of 4 Å molecular sieves, $9\frac{8}{3}$ gave the epoxide **12.** 1H NMR analysis of the Mosher ester in CDCl_3 indicated $>99\%$ enantiomeric excess (e.e.). Catalytic

Scheme 2 *Reagent and conditions:* i, lithium diisopropylamide (LDA) equiv.), hexamethylphosphoramide $(HMPA)$ Et₂O, then ClCH₂COCl (0.5 equiv.), -78 °C; ii, sodium borohydride (NaBH₄), MeOH; iii, aqueous NaOH (2 mol dm-3); iv, TBHP, diisopropyl $D-(-)$ -tartrate, Ti $(OPrⁱ)₄$, molecular sieves 4 Å, CH₂Cl₂, -20 °C; v, H_2 , Pd/C, AcOEt; vi, O_3 and then Me₂S, -78 °C; vii, CF_3CO_2H , $CH₂Cl₂$; viii, MeOH, conc. $H₂SO₄$; ix, Me₂C(OMe)₂, HClO₄, acetone; x, MeOH, toluene, reflux

^{\$} The yield is based on the consumed **7.**

*^Q*Though the results herein were obtained using the stoichiometric conditions,7 use of the modified conditions⁹ have also given satisfactory results. Details will be reported in a full paper.

hydrogenation of 12, $[\alpha]_{D}^{22} + 36.8^{\circ}$ (c 1.0, CHCl₃), in ethyl acetate afforded the diol (S) -(1a), $[\alpha]_D^{20}$ -22.8[°] (c 2.16, CHC13). The absolute structure of the epoxide was determined by its transformation (ozonolysis followed by treatment with trifluoroacetic acid) to (S)-3-hydroxy-4-butanolide **13.10** Methyl **(S)-3,4-dihydroxybutanoate 14** was also synthesized in the same manner (ozonolysis followed by methylation). Alternative syntheses of **14** and its use in natural products synthesis as well as transformation to other four-carbon building blocks have been carried out by many researchers. 11

The diol **1a** also afforded the protected dihydroxy β -keto ester **15:** the six-carbon building block, which is useful for synthesis of HR 780,12 a synthetic HMG-CoA reductase inhibitor. Though several synthetic methods for **15** are available, none seems to be satisfactory owing to low availability of the starting materials.13 When the route shown in Scheme 2 was carried out by using L-(+)-DIPT in the epoxidation step, the enantiomer $[(R)-6-(2,3-dihydroxy$ propy1)- 1,3-dioxin-4-one] was also synthesized with the same efficiency.

We are currently investigating the use of **1** either according to path c or even as substrates for pericyclic reactions.¹⁴

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