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A General Route to Optically Pure Prostaglandins from a D-Glucose Derivative

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Summary The epoxy-lactone (14), which is a key intermediate in one synthetic route to prostaglandins, has been prepared from a readily available D-glucose derivative. WHEREAS reactions for converting monosaccharide derivatives into functionalised cyclohexanes are well developed,¹ parallel procedures for obtaining cyclopentane analogues are less well known. Several reports of such methods have, however, recently appeared,^{2,3} but few of them are simple

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and efficient. A notable exception is the method reported by Bernet and Vasella³ which results in the bonding of C-1 to C-5 of *aldehydo*-5,6-dideoxyhex-5-enose derivatives and the production of bicyclic compounds containing fused cyclopentane-isoxazolidine ring systems. Our interest in functionalised cyclopentanes obtainable from carbohydrates¹ relates to their potential value as precursors of non-carbohydrate natural products (notably prostaglandins) and we here outline the synthesis of the epoxylactone (14) from a readily accessible glucose derivative. Prostaglandin $F_{2\alpha}$ is obtainable from this compound (the



Scheme $Bz = PhCO-, Ts = p-MeC_{6}H_{4}SO_{2}-.$

synthesis having been effected with the racemic form;⁴ the required enantiomer is known⁵), and the approach represents a carbohydrate-into-prostaglandin conversion of general applicability. Stork's synthesis of prostaglandin $F_{2\alpha}$ from glucose⁶ is a major synthetic achievement and a powerful illustration of the transfer of carbohydrate functionality

and stereochemistry; however, it is specific. Our objective has been to prepare an optically pure compound which can serve as a precursor of a range of members of the series and their analogues.

The cyclopentane ring formation was achieved by use of a modification of the method of Bernet and Vasella.³ Treatment of the methyl 6-deoxy-6-iodo- α -D-glucopyranoside (1)¹ with zinc in refluxing ethanol followed by reaction of the resulting aldehyde (2), without purification, with *N*-methylhydroxylamine hydrochloride in ethanol-pyridine (5:2, 45 °C), gave the isoxazolidine (3) (73%, m.p. 129— 131 °C, $[\alpha]_D - 53^\circ$ in CHCl₃) which was readily isolated. Not only, therefore, can these procedures be applied with 6-bromohexoside derivatives containing base-stable protecting groups,³ but they can be used with iodo-analogues containing benzoyl and toluene-*p*-sulphonyl ester functions.

Reduction of the product (3) with hydrogen over Raney nickel at atmospheric pressure caused not just the expected reductive opening of the heterocyclic ring³ but also intramolecular displacement of the sulphonyloxy-group,7 and the product was the aziridine (4) (74%, m.p. 116-118 °C, $[\alpha]_{\rm p} - 117^{\circ}$ in CHCl₃). Conversion of this into the syrupy alkene (5) (83% after chromatography, $[\alpha]_D - 185^\circ$ in $CHCl_3$) was effected by treatment with *m*-chloroperbenzoic acid in dichloromethane at room temperature,⁸ and the side chain was then elaborated to an acetic acid substituent. Standard esterification gave the toluene-p-sulphonate (6) (94%, m.p. 99—100 °C, $[\alpha]_D$ —115° in CHCl₃); treatment with sodium cyanide in dimethyl sulphoxide produced the nitrile (7) (80% after chromatography, $[\alpha]_{\rm D}$ -110° in CHCl₃). The aldehyde (8) (78%, $[\alpha]_D - 180^\circ$ in CHCl₃) was obtained by way of its 1,2-di-(N-phenylamino)ethane derivative by reduction of the nitrile with Raney nickel and sodium hypophosphite in the presence of the diamine;⁹ mild acid hydrolysis and its oxidation with pyridinium dichromate in NN-dimethylformamide¹⁰ gave the acid (9) (85%).

Iodolactonisation¹¹ of the unpurified acid (9) afforded the bicyclic compound (10) (84%, m.p. 162—164 °C, $[\alpha]_{\rm D}$ + 11° in CHCl₃), the structure and absolute configuration of which were confirmed by single crystal X-ray diffraction analysis. Reduction with tributyltin hydride in benzene¹² afforded the deiodinated diester (11) (88%, m.p. 108—110 °C, $[\alpha]_{\rm D}$ -98° in CHCl₃) from which the diol (12) (84%, m.p. 110—111 °C, $[\alpha]_{\rm D}$ -14° in MeOH) was obtained by use of potassium carbonate in methanol.

Diethyl azodicarboxylate-triphenylphosphine (which together preferentially activate sterically accessible hydroxy-groups)¹³ were applied to the diol (12) in the hope that the required endo-epoxide (14) would be obtained. Instead, they gave the isomeric product (13) (m.p. 68-70 °C, $[\alpha]_{D}$ + 69° in CHCl₃) exclusively, conceivably because the triphenylphosphonium intermediate formed from the endo-hydroxy-group was stabilised by co-ordination with the lactone ring oxygen atoms. Treated with toluene-psulphonylimidazole and sodium hydride in NN-dimethylformamide¹⁴ at 0 $^{\circ}$ C, the diol (12) gave the required epoxide (14), also exclusively. Its ¹H n.m.r. spectrum (80 MHz) was identical to that of an authentic sample, and its melting point was not depressed on admixture (m.p. 76-77 °C, $[\alpha]_{\rm D} = 108^{\circ}$ in CHCl₃; lit.⁵ m.p. 76-77 °C, $[\alpha]_{\rm D} = 115^{\circ}$ in CHCl₃).

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