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ARTICLE TYPE

Diastereo- and Enantioselective Synthesis of 1,3,5,7-Tetraol Structural Units Using a Prins Cyclisation/Reductive Cleavage Sequence†‡

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A stereocontrolled and efficient access to all the diastereomers of 1,3,5,7-tetraol structural units developed using a Prins cyclisation/reductive cleavage tetrahydropyran 10 Furthermore, these tetraols can be selectively functionalized.

Polyols are present in a great diversity of natural products, with different bioactive profiles. 1a Therefore, a number of synthetic methods and strategies have been developed in the past few decades to access these scaffolds.1 The most straightforward 15 strategy involves an iterative sequence, including a step using chiral reagents, to access 1,3,5-triols which relies on onedirectional chain elongation.² Desymmetrisation of meso-polyols and/or derivatives has also been reported.3 For our part, we have reported the synthesis of optically active syn- and anti-1,320 polyols by using highly diastereo- and enantioselective allyltitanations⁴ and we have demonstrated the utility of this method to synthesize a diversity of natural products and/or bioactive compounds.5

Scheme 1 Retrosynthetic analysis

Table 1 Diastereoselective Prins Cyclisation

entry	1	2	major 3	minor 3'	ratio 3/3°	yield (%) ^a
1	OTBDPS TSO H H H	OH OTs (+)-2	TSO H H H OTS	+ TsO H H OTS	$3a/3^{\circ}a = 70:30$	66%
2	OTBDPS TSO H H H 1a	OH OTs	TsO H H H H O H	+ TsO H O H O H O TSO	3b/3'b = 75:25	51%
3	TSO H H	OH OTs	TsO H H H OTS	+ TsO H H H OTS	3c/3°c = 65:35	80%
4	TSO H H H	OH OTs	TSO H H H H O H OTS	+ TsO H H H O H OTS	3d/3'd = 75:25	55%

^a Isolated yield for the mixture of diastereomers 3 + 3.

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Herein, we would like to report the synthesis of 1,3,5,7-tetraol structural units A with excellent diastereoselectivities from bistetrahydropyrans **B** obtained by using a homoallylic alcohol controlled Prins cyclisation applied to tetrahydropyran aldehydes 5 C (Scheme 1).

Enantiopure tetrahydropyrans 1a and 1b⁶ were involved in a Prins cyclisation with enantiopure homoallylic alcohols (+)-2 or (-)-2 in the presence of TFA (26 equiv, CH₂Cl₂, rt, 3 h)⁸. After treatment under aqueous basic conditions (aqueous NaHCO₃, 10 Et₃N), the bis-tetrahydropyrans were obtained as a mixture of diastereomers 3 and 3', in a ratio around 70:30, with a yield in between 50% and 80% (Table 1). We have to point out that compounds 3 and 3' were separated by column chromatography on silica gel. A good control of the stereogenic centers at C8 and 15 C12 was observed and the ratio of the diastereomers 3 and 3' is the result of the moderate control of the stereogenic center at C10.9 In all cases, the substituents at C8, C10 and C12 are all cis in the major isomer. ¹⁰ The results are reported in Table 1.

To transform 3a-3d and 3'a-3'd into the corresponding tetraols 20 A, the bis-tetrahydropyrans 3a-3d and 3'a-3'd were treated with NaI (acetone, μ-wave irradiation, 120 °C) and the obtained iodinated compounds were then treated with zinc (NH₄Cl, EtOH, 90 °C) using the conditions reported by Yadav and co-workers. 8c The corresponding tetraols 4a-4d and 4'a-4'd were isolated in 25 yields around 60% over the two steps, as single diastereomers (Table 2, entries 1-8). The hydroxyl at C10, in the bistetrahydropyrans 3, can be protected with different groups such as a TBDPS group (compound 3e) (Table 2, entry 9) and the resulting ethers were engaged in the iodination/ring-opening 30 sequence (conditions A) to give the corresponding tetraols. It is worth noting that it is necessary to use a different sequence of reactions to introduce other protecting groups such as MOM and Boc groups, due to their instability under the iodination conditions. However, iodination of compounds 3, followed by 35 protection and reductive cleavage of the tetrahydropyrans units (conditions B) gave the doubly protected tetraols 4f-4i in good yields and diastereoselectivities (Table 2, entries 10-12).

To prove the relative stereochemistry of the hydroxyl groups at C5 and C7 in the tetraols, and to confirm that the control is due to 40 the homoallylic alcohol used, compounds 4f and 4g were treated with 2,2-dimethoxypropane (PPTS, acetone, rt) to form the corresponding acetonides 5f and 5g. The relationship between the substituents at C5 and C7 of the acetonides 5 was established by NMR analysis and was confirmed to be trans in compound 5f and 45 cis in compound 5g (Scheme 2).11

Table 2 Access to 1,3,5,7-tetraols

	3) Zn, NH₄CI, EtOH, 90 °C				
entry	3	4 (yield %) ^a			
1 ^b	OTBDPS OH TSO H H H H OTS 3a	TBDPSO OH OH OH 4a (50%)			
2^b	TSO H H H O H OTS	TBDPSO OH OH OH UH			
3^b	TSO H H H H H O H	TBDPSO OH OH OH 4b (57%)			
4^b	TSO H H H H H O H OTS	TBDPSO OH OH OH OH 4'b (62%)			
5 ^b	TsO H O H H OTS	TBDPSO OH OH OH 4c (56%)			
6^b	TSO H H O H H OTS	TBDPSO OH			
7^b	TSO H H H H H H H H H H H H H H H H H H H	TBDPSO OH OH OH			
8^b	TSO H H H H H H O H OTS	TBDPSO OH OH OH			
9 ^b	OTBDPS OTBDPS TSO H H H H OTS 3e	TBDPSO OH OH OTBDPS 4e (67%)			

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^a Overall isolated yields from **3** or **3**°. ^bConditions A. ^cConditions B. ^dReductive cleavage was performed using zinc in a THF/H₂O mixture to prevent Boc migration.

The differentiation of the two terminal olefins can be achieved with compound 4h by applying the Bartlett-Smith iodocarbonate cyclization. ¹² After protection of **4h** as an acetonide, followed by 10 treatment with NIS (CH₃CN, -40 °C - rt, 4.5 h), the cyclic iodocarbonate 6 was isolated in 72% yield as a mixture of two diastereomers in a ratio of 90:10 in favour of the cis isomer. After basic treatment (K₂CO₃, MeOH, rt - 40 °C, 8 h), iodocarbonate 6 was transformed to the hydroxy epoxide 7 in 76% yield in a ratio 15 of 87:13 in favour of the syn isomer (Scheme 3). As all the functionalities are orthogonally protected, this compound can be used to realize the synthesis of complex molecules.

developed have a general summary, we diastereoselective method to prepare functionalized 1,3,5,7-

tetraol structural units, which should provide an access to all the possible diastereomers, iterative using an 25 cyclisation/reductive cleavage sequence. The stereochemistry of the hydroxy groups is controlled during the Prins cyclisation and is dependent on the configuration of the homoallylic alcohol involved. In addition, as the unsaturated Boc-protected 1,3,5,7tetraols can be selectively functionalized, the use of these tetraols 30 in the synthesis of polyketides natural products is ongoing in our laboratory and will be reported in due time.

Notes and references

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This communication is dedicated to Prof. Taylor on the occasion of his 40 65th birthday.

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