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Received 00th January 20xx, Accepted 00th January 20xx

DOI: 10.1039/x0xx00000x

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Carbohydrate biomass represents a potentially valuable sustainable source of raw materials for chemical synthesis, but for many applications, selective deoxygenation/dehydration of the sugars present is necessary to access compounds with useful chemical and physical properties. Selective dehydration of pentose sugars to give tetrahydrofurans can be achieved by treatment of the corresponding N,N-dimethylhydrazones under acidic or basic condiitions, with the two approaches showing complementary stereoselectivity. The dehydration process is readily scalable and the THF hydrazones derived from arabinose, ribose, xylose and rhamnose were converted into a range of useful fragments containing primary alcohol, ketone, carboxylic acid or amine functional groups. These compounds have potentially useful physiochemical properties making them suitable for incorporation into fragment/lead generation libraries for medicinal chemistry. It was also shown that Larabinose hydrazone could be obtained selectively from a crude sample of hydrolysed sugar beet pulp.

Introduction

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The production of chemicals for medicinal chemistry relies heavily on hydrocarbons from petrochemical sources to provide small molecule synthons. Considering the inevitable exhaustion of fossil fuels and the increased restriction of their use due to concerns relating to climate change, the key to future chemical processes will be the exploitation of renewable resources from plant biomass.¹ An important sustainable feedstock is sugar beet, with a production cycle from seeds-to-seeds of only two years. Cultivated predominantly in temperate regions for the production of sucrose, the processing of the beet also generates large quantities of pulp, which is considered as a low-value waste product and is usually used as animal feed.^{2,3} This material is highly enriched in carbohydrates, with the pulp being composed of polymeric glucose (cellulose) and pectin (a copolymer of hexoses and pentoses). The latter fraction contains a high proportion of L-arabinose as well as other sugars, which could potentially be exploited as valuable chemical feedstocks, but processing is required to release the monomeric sugars, and strategies for this have been reported.4,5

A variety of transformations are now available to produce small molecule building blocks from sugar monomers.^{4,5} However,



DOI: 1



Scheme 1: Dehydration of pentose sugars for the synthesis of chiral building-blocks

Recently, some approaches have been developed which take advantage of the intrinsic functionalities and chirality of carbohydrates to develop novel methodology to access small

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Electronic Supplementary Information (ESI) available: Procedures for the preparation of all compounds, together with characterisation data and X-Ray crystallographic data. See DOI: 10.1039/x0xx00000x

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3D-building-blocks.^{11,12} Indeed, we have recently developed a selective method for the dehydration of pentose sugars to afford hydroxy-functionalised THFs without the need for protecting groups.¹² In this paper, we describe the optimisation of this chemistry on a multigram scale, the development of a novel base-mediated cyclic dehydration of sugar hydrazones to access other stereoisomers, and the extension of this methodology to a range of pentoses. We also demonstrate the application of these dehydration reactions to the construction of a small library of chiral THF fragments as useful building blocks for medicinal chemistry,13,14 via routes that offer selectivity, scalability and sustainability. In addition, the direct synthesis of one of our key starting materials directly from sugar beet pulp is described, to demonstrate the feasibility of 'upgrading' waste from biomass into complex chiral building blocks.

Results and discussion

Synthesis of chiral THFs from pentose sugars

The conversion of L-arabinose into 1a-3a was conducted following the procedure described in our preliminary communication for acid catalysed dehydration of pentoses.¹² Hydrazone 1a was obtained quantitatively under acidic conditions and subsequently cyclised using catalytic TFA (Scheme 2). In our previous studies, this reaction had yielded the THF in only moderate yield (67%), but the fate of the remaining material was unclear. However, when the cyclisation was performed on a larger scale (5-20 g), formation of two sideproducts 4 and 5 in 22% and 5% yields respectively was observed. The formation of methyl ether 4 could be rationalised by the reaction of the vinyldiazenium intermediate 2a with methanol. The formation of ketone 5 most likely results from the aerobic oxidation of 1a at the C-2 position.





These issues were readily solved by running the cyclisation in a less nucleophilic solvent (isopropanol) and under an argon atmosphere (see ESI). Under these conditions, 3a was obtained in 81% yield routinely on a 20 g scale. When these reaction conditions were applied to other pentose sugars, the yields were generally superior to the previously reported procedure, with no significant difference in the diastereoselectivity observed (Scheme 3). THFs anti-3a and anti-3b derived from Larabinose and D-ribose could be recrystallised from THF/Petrol to give the pure anti-isomer in each case.







Table 1: Synthesis of chiral THFs syn-3a to syn-3e under basic conditions

Investigations also established that the cyclisation of the sugar hydrazones could be achieved to give THFs under basic conditions, by activating the sugar-hydrazone 1a with

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dimethylcarbonate (DMC).¹⁵ The reaction occured at room temperature and afforded the chiral THF syn-3a from arabinose in excellent yield, with good diastereoselectivity. Importantly, this provides a complementary approach to the acid-catalysed cyclisation in terms of the THF isomer produced. These conditions were also applied successfully to other pentoses, with the major THF isomer in each case having synstereochemistry at the THF carbons C-1 and C-2 (Table 1).

Full conversions were obtained after 4 hours and no change in the diastereomeric ratio was noted during the course of the cyclisation reaction. The presence of dimethylcarbonate and a stoichiometric amount of base were essential for the reaction to proceed in high yield. When a purified sample of the THF anti-3a was submitted to the DMC/K2CO3 reaction conditions, no epimerisation was observed after 24 hours suggesting that there was no interconversion of the two isomers and that the two products anti-3a and syn-3a were likely formed via two different reaction pathways under the basic conditions. This is in stark contrast to the acidic cyclisation conditions under which the two products can rapidly interconvert, leading to production of a thermodynamic mixture of products.

Taking this into account, along with the observed diastereomeric ratios of the products obtained from Larabinose, D-ribose, D-xylose and D-lyxose it is evident that epimerisation of the the C-2 stereocentre must take place prior to irreversible cyclisation, with the observed products resulting from the relative rates of cyclisation of the two diastereomers. Thus, arabinose hydrazone 1a must first be converted into cyclic carbonate 6a (Scheme 4), which can only cyclise relatively slowly to give anti-3a but can undergo epimerisation to 7a which cyclises more rapidly to give syn-3a. Epimerisation may take place via ring opening of the cyclic carbonate to generate zwitterion 8a. In the case of D-ribose hydrazone 1b, the initially formed carbonate 6b (the enantiomer of 7a) cyclises relatively rapidly so less epimerisation to 7b (and subsequent cyclisation to anti-3b) takes place leading to a slightly higher diastereoselectivity in this reaction in comparison to the formation of syn-3a. A similar rationale can be used to explain the observed diastereoselectivities resulting from cyclisation of D-xylose hydrazone 1c and D-lyxose hydrazone 1f. The selective formation of the cyclic carbonate at C-2/C-3 could perhaps be explained by involvement of the hydrazone group as a general base that activates the C-2 alcohol towards reaction with dimethyl carbonate (Scheme 5a). Furthermore, general base assistance could also be invoked in the cyclisation reaction (Scheme 5b), as in each case the cyclisation reaction which proceeds fastest involves reaction of an intermediate in which the hydrazone and nucleophilic primary alcohol are arranged on the same face of the 5-membered carbonate ring. Cyclisation with inversion at C-2 then leads to the THF ring in which the C-3 alcohol and the hydrazone are on the same face of the newly formed ring.







Scheme 5: Base mediated a) cyclic carbonate 6a formation, b) cyclisation of 7a

Chiral functionalised THFs as 3D fragments for medicinal chemistry

The next objective of this study was to synthesise carefully chosen targets for applications in medicinal chemistry, which contained suitable functional groups for derivatisation in order to prepare potential fragments and/or intermediates for incorporation into lead-like molecules. Three families of compounds were selected: (i) THF-triols 9, (ii) THF-carboxylic acids 10 and (iii) THF-amines 11. It was envisioned that these three compound classes would be ideal for generating libraries for fragment-based lead discovery as they are of low molecular

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weight (Mw <150 g/mol), are highly soluble in water, and contain a functional group ready for further elaboration (primary alcohol, carboxylic acid, primary amine). In addition, the targets could be used as strategic reagents¹⁶ for the final derivatisation of core molecules so as to dramatically lower logD by incorporation of the compact polyol moiety. Finally, by using different pentose sugars, we aimed to prepare a range of different stereoisomers, each with a diastereomeric ratio >90:10, suitable for use in these applications.¹⁷



The THF-hydrazone **3** is a versatile platform for accessing several functionalities (e.g. aldehyde, alcohol, nitrile, protectedamine, alkene).¹² As previously reported the hydrazone can be rapidly hydrolysed to the give the aldehyde hydrate **12** using an acidic resin (Amberlyst 15). The reaction was easily scaled-up (>10 g) and extended to the other pentoses (Scheme 6). The reaction could safely be conducted on the mixture of diastereomers obtained after cyclisation, as epimerisation takes place under the acidic hydrolysis conditions. Remarkably, in the case of L-rhamnose the hydrate was obtained with a higher diastereoselectivity than the corresponding hydrazone.



The epimerisation at C-2 during the hydrolysis of hydrazone **3a** to hydrate **12a** offers an opportunity to maximise the value of the material (Scheme 7). Thus, recrystallisation of 6.8 g of **3a** obtained directly from the acidic cyclisation (*dr* 78:22) yielded 2.4 g of *anti*-**3a** as a single isomer. A second crop of high purity *anti*-**3a** was also obtained (1.4 g, *dr* 95:5), leaving the remaining mother liquor as an equimolar mixture of diastereomers. Submitting this material to the hydrolysis conditions led to an increase in the diastereomeric ratio (*dr* 80:20) in the product hydrate **12a**, allowing efficient recycling of the mother liquor from the recrystallisation of *anti*-**3a** (Scheme 7).



The reduction of the hydrate using sodium borohydride provided the alcohol **9a** in excellent yield and without significant change in the diastereomeric ratio (Scheme 8).



Alcohols **9a** and **9b**, respectively derived from L-arabinose and D-ribose could be recrystallised to afford the pure *anti* isomers in high yield. The relative stereochemistry was confirmed through an X-ray structure of ribose derivative **9b** (Figure 2).



Figure 2: X-ray crystal structure of triol **9b**. The thermal ellipsoids are shown at the 50% probability level, while hydrogen atoms are drawn as fixed spheres with a radius of 0.15 Å.[‡]

For the alcohols **9c** and **9d** obtained from the D-xylose and Lrhamnose hydrates **12c** and **12d** respectively, the diastereoselectivity was lower than required, and separation of the two isomers was necessary. In order to discriminate between the two species, formation of a benzylidene acetal on

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the 1,3-diol was performed as selectivity for the syn-diol should be expected.¹⁸ Indeed, the reaction of the D-xylose-triol **9c** with benzaldehyde in the presence of catalytic *p*-toluenesulfonic acid and a dehydrating agent resulted in the selective formation of the acetal 13 along with unreacted anti-9c (Scheme 9a). The stereoselectivity of this acetal protection was confirmed by single crystal X-ray crystallography of the analogous compound 13-Br.§ The triol anti-9c and acetal 13 were easily separated by column chromatography. Finally, hydrolysis of 13 using Amberlyst 15 as a catalyst quantitatively afforded syn-9c. Acetal 13 was also oxidised with a catalytic amount of TEMPO using $PhI(OAc)_2$ as co-oxidant to afford the ketone **14** in high yield. This reaction sequence was also applied to L-rhamnose triol 9d allowing the separation of anti-9d and syn-9d via efficient formation of the acetal 15 (Scheme 9b). Ketone 16 was similarly obtained by oxidation of secondary alcohol 15.



Scheme 9: Separation of the *anti* and *syn* isomers of a) D-xylose triol **9c** and b) Lrhamnose triol **9d**; yields in parentheses are based on the quantity of the relevant diastereoisomer present in the starting mixture; X-ray crystal structure of compound **13-Br** with the thermal ellipsoids shown at the 50% probability level, and hydrogen atoms drawn as fixed spheres with a radius of 0.15 Å.⁺

Oxidation of triol **9a** into the carboxylic acid-THF **10a** required a more conventional synthetic approach, using a protecting group to allow selective oxidation of the primary alcohol moiety. Protection of the *syn* **1**,2-diol was performed with 2,2-dimethoxypropane in the presence of Amberlyst **1**5

(Scheme 10a). The acetal-protected triol 17a was obtained in high yield and was subsequently oxidised following a procedure previously described for the oxidation of nucleosides.¹⁹ The reaction proceeded at room temperature with a catalytic amount of TEMPO as primary oxidant, using PhI(OAc)₂ as the stoichiometric oxidant. The unprotected carboxylic acid 10a was then released under acidic conditions in moderate yields over two steps and with high isomeric purity. The carboxylic acid was also accessible directly from the THF-hydrate 12a thus reducing the number of steps and the stoichiometry of oxidant required (Scheme 10b). Protection of the hydrate-THF 12b resulted in a significant loss of diastereomeric purity. However, after oxidation followed by deprotection, the carboxylic acid **10a** was isolated in moderate yield with a high diastereomeric ratio in favour of the anti isomer. This rise in diastereomeric ratio during the oxidation could be explained by a higher reactivity of the anti-THF hydrate with the hindered oxidant TEMPO in comparison to the syn-THF hydrate.



Finally, we sought an effective method for accessing the THF-amine 11 by direct hydrogenation of the THF-hydrazones **3**. Although the literature is quite rich in precedent for the hydrogenation of hydrazones to hydrazines, the reduction of a hydrazone directly to an amine is rare at low hydrogen pressure.²⁰ In our previous work, we presented a procedure to prepare the THF-amine under 1 bar of hydrogen with Bocanhydride as an additive and Pd(OH)₂/C as catalyst (Scheme 13).12 Reactions were initially conducted in CPME,21 but in subsequent studies better reproducibility was achieved in a mixture of isopropanol/water (2:1). The Boc-protected amine 20 was isolated in moderate yield on a small scale from the hydrogenation of ribose-derived hydrazone **3b**, and standard deprotection using hydrochloric acid in methanol afforded the hydrochloride salt 11a.HCl (Scheme 10). The scalability of the hydrogenation proved to be problematic, however, with a dramatic loss of yield and increased reaction time as the scale was increased to gram quantities of material.

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Me₂NN

pulp



The drop in the conversion can probably be explained by a poor gas-liquid-solid contact in the reaction mixture. Therefore, subsequent reductions were attempted at higher pressures of hydrogen (up to 5 bar) but did not lead to the desired amine, and instead partial reduction to the hydrazine 19 was observed (Scheme 11). Hydrogenation can present serious safety issues in large scale batch reactions with high pressure, flammable solvents and pyrophoric catalysts. We therefore moved to a flow-chemistry set-up which offers several advantages in comparison with more traditional hydrogenation procedures, including the possibility to work at high pressure and temperature with an encapsulated catalyst to reduce the leaching of toxic metal into the reaction mixture.²²⁻²⁴ As a starting point, iPrOH:H₂O (2:1) was used as solvent and Pd(OH)₂/C as catalyst to remain close to the initial set of reaction conditions. However even at high pressure (100 bar), the transformation did not proceed to the amine at room temperature with mostly the hydrazine 22a being formed. Moreover, when the hydrazine 22a obtained was re-submitted to a second hydrogenation-cycle, no amine product was observed. The use of Boc-anhydride as additive did not solve the issue, and as Pd(OH)₂/C did not seem an efficient catalyst under these conditions we moved to Raney Nickel which has been reported previously as a good catalyst for the hydrogenation of hydrazones.²⁵ At room temperature and 100 bar of hydrogen, only reduction of hydrazone 3a to the hydrazine 22a was noticeable. However, at 80 °C, 3a was fully converted to the amine 11a. Importantly no additives were required for this transformation, and the amine was isolated in high yield directly after evaporation of the solvent (Scheme 12). Moreover, no erosion of the diastereoselectivity was observed compared to the initial hydrazone 3a. This method was then applied to the synthesis of amines 11b-11d in 68-94% yield. This direct hydrogenation of the hydrazone provides an alternative route for accessing the amines 11 to our recently reported biocatalytic approach involving reaction of hydrates 12 with transaminase enzymes.²⁶





Scheme 13: Synthesis of L-arabinose hydrazone **1a** from a solubilised sample of sugar beet pulp

Synthesis of sugar hydrazones from sugar beet pulp

Sugar beet pulp contains significant quantities of L-arabinose, so we were interested in its potential application as a feedstock for the synthesis of these chiral THF building blocks. It was therefore important to establish that hydrazone 1a could be prepared from waste pulp material. To determine this, a solubilised sample of sugar beet pulp after steam explosion and acidic hydrolysis treatment was generated (Scheme 13).27-29 Analysis of the solution prepared in this way indicated it contained sodium sulfate (80 g/L) and arabinose (12 g/L), together with galacturonic acid (5 g/L), galactose (3 g/L), rhamnose (1 g/L) and glucose (1 g/L) monomers. Initially, a model synthetic mixture containing these components was treated with N,N-dimethylhydrazine (2 eq. wrt arabinose). Surprisingly, analysis of the crude product by NMR spectroscopy revealed the selective formation of the hydrazone 1a derived from L-arabinose under these conditions in quantitative yield, with only traces of the rhamnose hydrazone present. This selectivity could be explained by (i) the low solubility of galacturonic acid in methanol and (ii) the much slower reaction of hexose sugars with hydrazine compared to the pentoses. Indeed, formation of the hydrazone from galactose was

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previously observed to require harsher conditions for an extended period of time (3 days) to reach only moderate conversion.¹² Reaction of the heterogeneous mixture derived from sugar beet pulp under similar conditions also led to selective formation of hydrazone **1a**. The concentration (>1 M) was found to be a crucial parameter for achieving high conversion in this transformation, so concentration of the crude solution was necessary prior to reaction. After purification, the arabinose hydrazone **1a** was isolated in high purity in 30% yield. The discrepancy in yield between the two experiments suggests that partial decomposition of the sugars is taking place during concentration of the crude starting material, and/or that impurities present in the pulp lead to loss of material during the work-up and purification of hydrazone **1a**.



Figure 3: Chiral THFs prepared in this work. All compounds shown above were obtained in >95:5 dr (except **11c/11d**).

Conclusions

In conclusion, we have oprimised procedures for the selective dehydration of pentose sugars on multigram scales, including: (i) improvement and scale-up (20 g) of the acid-catalysed cyclisation for L-arabinose and other pentoses; (ii) development ARTICLE

of a novel cyclisation procedure under basic conditions which preferentially provides THFs with *syn* stereodfleffister access to a range 3. These two complementary procedures offer access to a range of chiral-THFs with varied stereochemistries. The THFhydrazones were subsequently transformed into THF-triols **9**, bicyclic ketones **14/16**, THF-carboxylic acids **10** and THF-amines **11** to build a library of functionalised THFs (Figure 3) for potential application in medicinal chemistry. Finally, we were able to prepare arabinose hydrazone **1a** selectively from a pretreated sample of sugar beet pulp, thus demonstrating the feasibility of preparing many of these chiral THF building blocks from an abundant renewable feedstock.

Acknowledgements

We would like to thank the Engineering and Physical Sciences Research Council (EPSRC) and the University College London Impact Acceleration Account for providing PDRA funding to L.B. (EP/K503745/1) for a collaboration with AstraZeneca and GlaxoSmithKline. We would like to thank David Andrews (AstraZeneca), Tony Dean and Rob Young (GlaxoSmithKline) for their help and support on this project. We would also like to thank GlaxoSmithKline for providing an EPSRC CASE studentship for RWF. Finally, we thank Maciej Kucharski and Lee Edwards (GlaxoSmithKline) for assistance with the flow hydrogenation experiments.

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^{*}Tables of selected crystallographic data for **9b** and **13-Br** can be found in the supplementary information. CCDC 1877937 and 1877938 contain the supplementary crystallographic data for compounds **9b** and **13-Br** respectively. The data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/structures.

[§]The selectivity of the acetal protection was confirmed by obtaining an X-Ray crystal structure of **13-Br**,[‡] an analogue of **13** prepared from 4-bromobenzaldehyde (see supplementary information for full details).

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DOI: 10.1039/C9GC00448C

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