Easy Access to Modified Cyclodextrins via an Intramolecular Radical Approach


Abstract: A simple methodology to modify the primary face of cyclodextrins (CDs) is described. The 6'-O-yl radical of $\alpha$-, $\beta$-, and $\gamma$-CD abstracts regioselectively the HS$^1$, located in the adjacent D-glucose unit, by an intramolecular 1,8-hydrogen atom transfer reaction through a geometrically restricted nine-membered transition state to give a stable 1,3,5-trioxocane ring. The reaction has been extended to 1,4-diols of $\alpha$- and $\beta$-CD to give the corresponding bis-trioxocanes. The C$_2$-symmetric bis-trioxocane corresponding to $\alpha$-CD is a stable crystalline solid whose structure was confirmed by X-ray diffraction analysis. The calculated geometric parameters confirm that the primary face is severely distorted toward a narrower elliptical shape for this rim.

The design of nanosystems as novel drug carrier systems that allow controlled release is currently a topic of great relevance in medical research to treat a wide variety of diseases, including cancer therapy.[1] In this regard, cyclodextrins (CDs) are of interest as nanocarriers because of their ability to encapsulate biomolecules.[2] These naturally-occurring macrocyclic oligosaccharides composed of 1,4-$\alpha$-linked glucose units are conical in shape and have an internal cavity creating a hydrophobic microenvironment, while the outside is hydrophilic due to the presence of the hydroxyl groups.[3] These properties are responsible for their aqueous solubility and ability to form host-guest or inclusion complexes with a wide range of molecules, making them one of the most important supramolecular host families. These features illustrate the potential utility of CDs as drug- and gene-delivery vehicles,[4] enzyme mimics,[5] or catalysis,[6] among others.

Native CDs are rarely well-suited to pharmaceutical applications, so their modification is necessary to tune their chemo-physical properties such as to enhance their solubility, change the size cavity or improve their complexing ability. Taking advantage of the different accessibility and reactivity of the hydroxyl groups, efficient and selective chemical modifications have been carried out by allowing for instance grafting substituents to different positions to access amphiphilic CD,[7] and hence to fascinating molecular architectures.[8] It is also worthwhile to mention the synthetic reports that generate significant changes in the cavity shape by altering the conformation of glucose residues.[9] Despite the great deal of methodologies employed, radical processes are practically unknown in these macromolecules because in simple carbohydrates they mostly involve the anomeric position, which is committed in the cyclic CD structure.[10] In fact, we have found a single example in the literature concerning intermolecular radical attack on CDs and it is based in EPR experiments with reactive oxygen-centered free radicals generated in aqueous solutions.[11]

As part of our ongoing research program on radical chemistry in carbohydrates, we have recently described a novel 1,8-hydrogen atom transfer reaction (1,8-HAT) between the two pyranose units in Hexp-(1→4)-Hexp disaccharide systems through a nine-membered transition state (TS).[12] The process is initiated by a primary $6^1$-O-yl radical (I) generated under oxidative conditions, which abstracts regioselectively the hydrogen at C5$^0$ to give a C5-radical (II) which, after one-electron oxidation, collapses into the 1,3,5-trioxocane (IV) via an oxacarbenium ion intermediate (III) (Scheme 1). Alternatively, the $6^1$-O-yl radical (I) may abstract the H1$^1$ through a seven-membered TS to generate a C1-radical (V) which finally stabilizes via the oxacarbenium (VI) to give the spiro ortho ester (VII). We have demonstrated that this regioselectivity is not only 

![Scheme 1. HAT reactions in Hexp-(1→4)-Hexp systems.](image-url)
dependent on the configuration of the four chiral centers implicated in the cyclization step (C5\textsuperscript{α}, C1\textsuperscript{α}, C4\textsuperscript{α}, and C5\textsuperscript{β}) but also on the conformations of the glycosidic (ϕ = H1\textsuperscript{α}–C1\textsuperscript{β}–O4\textsuperscript{β}–C4\textsuperscript{β}) and aglyconic (ψ = C1\textsuperscript{α}–O4\textsuperscript{α}–C4\textsuperscript{α}–H4\textsuperscript{α}) bonds.

Scheme 2. 1,8-HAT for 6\textsuperscript{1}−ol−α−CD 1, 6\textsuperscript{1}−ol−β−CD 4, and 6\textsuperscript{1}−ol−γ−CD 7. DIB = (diacetoxyiodo)benzene.

In a disaccharide with an arrangement of α−D-Glc-(1→4)-β-D-Glcp (β-maltose) the 6\textsuperscript{1}−O-yl radical abstracts exclusively the H5\textsuperscript{II} to give a trioxocane ring in a stable boat-chair conformation.\textsuperscript{13a} In the 1,8-HAT TS involved, the glycosidic bond adopts an exo-syn conformation with torsion angles (ϕ = −32.7, ψ = −37.3°) leading to an ideal distance of 3.1 Å between the alkoxyl radical and the abstractable H5\textsuperscript{II}. We wondered whether this radical methodology might be suitably deployed for the remote functionalization in more complex carbohydrate systems such as CDs, where the D-glucose units are linked in a similar fashion (cyclo-[α−D-Glc-(1→4)]\textsubscript{n}). Preliminary observations made on the X-ray structure of permethylated β-CD show that the macrocyclic ring has sufficient interresidue flexibility to permit the glycosidic and aglyconic bonds adopt an exo-syn conformation with values of ϕ and ψ similar to β-maltose.\textsuperscript{13b} Analogous situation is obtained over a minimized structure of permethylated β-CD and therefore, we consider that the 1,8-HAT process between two vicinal units could take place with high regioselectivity.\textsuperscript{13} If this strategy succeeded, it could provide facile access to modified CDs on their primary face, which are difficult to obtain by conventional methods, and which could show significant deformations in their cavities.

In order to test the viability of the radical process, we initiated our investigations by preparing 2\textsuperscript{10},3\textsuperscript{10},6\textsuperscript{10},9\textsuperscript{10}-icosa-O-methyl-β-CD (1).\textsuperscript{13} According to a three-steps literature procedure, where all the hydroxyl groups, with the exception of a primary one, have been transformed into methyl ethers, since these protecting groups usually lead to an increase in solubility both in water and in organic solvents (Scheme 2).\textsuperscript{16} Excitingly, when we treated the alcohol 1 with (diacetoxyiodo)benzene (DIB) (2.2 equiv) and I\textsubscript{2} (1 equiv) under irradiation with visible light at 30 °C, it was found that the reaction proceeded very fast, with complete consumption of starting material after 30 minutes, to give two main products in near quantitative overall yield: the expected trioxocane derivative 2 (36%) together with the acetate 3 (61%) as the major compound. A minor side reaction involving a competitive intermolecular addition of external nucleophilic acetate ion coming from the reagent to the oxacarbenium ions has also occasionally been observed in disaccharide systems.\textsuperscript{13a,14} If we decreased the temperature or the equivalents of DIB or I\textsubscript{2}, the reaction became slower and did not go to completion. However, when we increased the reaction time or the amount of DIB and I\textsubscript{2}, the trioxocane 2 was obtained in 60% yield as the only product.

The NMR data of 2 clearly indicate that oxidation has taken place at C5\textsuperscript{α}. Accordingly, the presence of seven anomeric carbons in the range of δ\textsubscript{C} 96.97–99.39 ppm and the HMBC correlation of H\textsubscript{1}\textsuperscript{α} proton signal at δ\textsubscript{H} 5.06 ppm with the C5\textsuperscript{α} quaternary carbon atom (δ\textsubscript{C} 101.04 ppm) confirm the formation of the trioxocane moiety. Moreover, all D-glucopyranose units are in 4 chair conformations as can be deduced from the values of the coupling constants of the anomeric hydrogens (J\textsubscript{1,2} = 3.6 Hz).

On the other hand, the acetate group at C5\textsuperscript{β} of 3 could be clearly established by 2D HSQC and HMBC experiments, whereas the observed deshielding of the ring hydrogens in this unit VII and the values of their coupling constants (J\textsubscript{1,2} = 1.9, J\textsubscript{3,4} = 2.5, J\textsubscript{5,6} = 1.9 Hz), determined on the basis of 1D TOCSY experiments, deviate considerably from those of the normal 4\textsuperscript{β} C\textsubscript{1} chair and were more consistent with a 1\textsuperscript{C} chair conformation. The stereochemistry of the quaternary C5\textsuperscript{β} was tentatively assigned as R on the basis of NOE interaction between H\textsubscript{1}\textsuperscript{α} and the methyl of the acetate group. These results suggest that the nucleophilic attack of the acetate proceeded with inversion of configuration at C5\textsuperscript{β} and consequently the original α-D-glucopyranose unit has been transformed into a β-L-idopyranose adopting a 1\textsuperscript{C} chair conformation. It is evident that both products are formed by the same C5\textsuperscript{β} radical intermediate promoted by the alkoxyl radical through an intramolecular 1,8-HAT process, no products functionalized at C1\textsuperscript{α} via the alternative 1,6-HAT reaction were detected. As far as we know, the presence of a sugar unit with a structure of β-L-idopyranose as part of a β-CD ring has no literature precedent.
Encouraged by these preliminary results, we decided to extend the reaction to the monoalcohols derived from α- and γ-CD, respectively, which were prepared following the same procedure applied to the 6'-ol-β-CD derivative. Pleasingly, both compounds behaved in an analogous manner to the β-CD derivative albeit generating the corresponding trioxocane derivatives 5 and 8 and acetates 6 and 9, in somewhat lower overall yields (Scheme 2). We soon realized that acetates 6 and 9 were undergoing complete hydrolysis during the standard purification process, giving exclusively the trioxocane derivatives 5 and 8, respectively, and this is most likely responsible for the lower yield observed. Notwithstanding, pure acetates 6 and 9 were obtained avoiding the typical aqueous workup and after careful flash column chromatography using silica gel scraped from commercial coated TLC plates (see Supporting Information). On the contrary, the trioxocane derivatives are stable for extended periods of time at room temperature.

The structures and conformations of the acetyl-derivatives 6 and 9 were analogously confirmed by 1D TOCSY and 2D HSQC and HMBC experiments. Of particular relevance are the differences observed in the conformation of unit VI of compound 6 by changing the solvent from CDCl₃ to C₆D₆. Thus, in CDCl₃ small coupling constants were noted in the 1D TOCSY experiment, between all the vicinal ring protons (J₁₃,₂ = 2.8, J₁,₂ = 3.2, J₂,₃ = 3.2, J₁,₂,₃ = 2.8 Hz) suggesting predominantly a 1C₄ chair conformation while in C₆D₆ these values (J₁₃,₂ = 3.5, J₂,₃ = 8.5, J₁,₂,₃ = 7.9 Hz) preferentially point to a 1C₄ conformation. These observations are in contrast with the acetates derived from β- and γ-CD, 3 and 9 respectively, which show the same 1C₄ conformation in both solvents.

To assess the versatility and scope of this methodology we have investigated the feasibility of applying the same procedure to 1,4-diols of β- and α-CD to determine whether the two hydroxyl groups can participate in two 1,8-HAT reactions simultaneously. The reaction of the 6'-ol-β-CD derivative 10 with DIB (3 equiv) and I₂ (1.7 equiv) afforded four products in good overall yield, bis-trioxocane compound 11 (20%), bis-acetyl-derivative 12 as the major product (49%) and a mixture of monotrioxocane-monooacetol derivatives (14%) (not shown, see the Supporting Information) which could be transformed into 11 after treatment with iodine (Scheme 3). The NMR characteristics of 12 clearly indicate that the introduction of both acetates in the corresponding D-glucose units III and VII has taken place with inversion of configuration at C5. Accordingly, the deshielding of the hydrogens of these units and their small coupling constants (J = 1.9–2.5 Hz) confirmed the presence of two β-L-idopyranose units both in 1C₄ chair conformations. Moreover, the NOE experiments show interaction between H1₁₀ and H1₁₀ and the respective methyl of the acetate group, tentatively establishing the absolute configuration at C5₁₀ and C5₁₀ as R.

Finally, we carried out the reaction with 6'-ol-α-CD derivative 13. The bis- and mono-trioxocanes 14 and 15 were the only compounds obtained in the reaction, no product from the incorporation of acetate in the molecule being detected, revealing the greater steric hindrance in this hexamer (Scheme 3). The structures of both products were determined by 1D and 2D NMR experiments. Furthermore, compound 14 is a stable crystalline solid whose structure was unambiguously confirmed by X-ray crystallographic analysis, showing two 1,3,5-trioxocane rings in boat-chair conformations with a guest n-hexane molecule as an inclusion complex into the CD cavity (Figure...
As expected, the NMR spectra are consistent with the C2-symmetry of the proposed structure.

The comparison of the geometrical parameters of compound 14 with those of the related permethylated α-CD[21] shows that the presence of the two 1,3,5-trioxocane rings does not significantly alter the secondary face of the molecule (see the Supporting Information, Tables S2 and S3). Thus, for instance, the distances and angles between the six interglycosidic oxygen atoms (O4β), as well as the radii of the gravity center of the hexagon formed by these O4β atoms are quite similar in both X-ray structures. Nor do the trioxocane rings seem to affect the C1 chair geometries of the six glucose units, as shown with the Cremer–Pople puckering parameters which describe slightly distorted chairs similar to those found in the permethylated α-CD.[25] The most significant differences are observed in the tilt angles made by the O4β mean plane and the mean planes through the glucose units principally in residues II and V adjacent to the trioxocane rings.

However, the trioxocane rings severely distort the primary face of the molecule. This is clearly observed when comparing the geometrical parameters of the irregular hexagon comprised by the side chain carbon atoms (C6n) which points towards a much narrower elliptical shape for this rim (see the Supporting Information, Tables S2 and S3).

In summary, the remote C-H functionalization initiated by 6-O-yl radicals proceeds with complete regioselectivity in α-, β-, and γ-cyclodextrin models, due to very suitable geometrical requirements which favour the TS of an intramolecular 1,8-HAT process between two contiguous units of the macromolecule. To the best of our knowledge, this is the first time that a radical approach is applied to access modified CDs with significant deformations in the primary face. The mild conditions and high efficiency are the key features of this procedure with potential synthetic and pharmaceutical applications. The possibility to apply this methodology not only to monoalcohols but also to 1,4-diols and carry out two 1,8-HAT processes simultaneously, opens the way to generate more complex CD systems. Work in this sense is now in progress and will be reported in due course.

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**Figure 1.** X-Ray crystal structure of compound 14. C blue, O red, H atoms not shown. Thermal ellipsoids drawn at 50% probability.
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[20] CCDC 1039442 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.


Successful match of cyclodextrins with radical chemistry. The bi-alkoxyxyl radical generated from 6\textsuperscript{1,3,5}-diol-α-CD under oxidative conditions led to a stable crystalline bis-1,3,5-trioxocane with a guest n-hexane molecule where the primary face is severely distorted toward a narrower elliptical shape.

Dimitri Alvarez-Dorta, Elisa I. León, Alan R. Kennedy, Angeles Martín,* Inés Pérez-Martín, Ernesto Suárez*

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