Probing the Lewis Acid Catalyzed Acyclic Pathway of Carbohydrate Conversion in Methanol by In Situ NMR

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Probing the Lewis Acid Catalyzed Acyclic Pathway of Carbohydrate Conversion in Methanol by In Situ NMR

Pernille Rose Jensen,[a] Esben Taarning,[b] and Sebastian Meier*[a]

Abstract: Future bioindustries will rely on the formation of diverse chemicals at high yield through various reaction pathways. These pathways include reactions to a series of alpha-hydroxy esters and acids that can be formed from the conversion of C3-C6 carbohydrates by Lewis acidic catalysts in alcohol and water. Definitive kinetic and mechanistic insights to support the development of carbohydrate conversion processes are arguably not used to their full potential.[4] As the abundant pentose and hexose carbohydrates can occur in various cyclic and acyclic forms, corresponding cyclic or open-chain pathways can play a role in the formation of carbohydrate-derived chemicals.[5] The difficulties in obtaining definitive insight into carbohydrate conversion processes are consequently well witnessed by the long-standing debate on the role of cyclic and acyclic routes of carbohydrate conversion to furanic compounds[1a, 6] which are widely investigated as future precursors for instance of polymers, pharmaceuticals, fuel additives and solvents. Acyclic routes are likely central in the formation of alpha-hydroxy esters and acids by Lewis acidic catalysts in alcohols and water. Alpha-hydroxy esters bear promise as bio-derived solvents and polymer building blocks. While the C3-metabolite lactate can be obtained in attractive yields from chemocatalytic approaches as well,[7] non-metabolite C4- to C6-alpha-hydroxy esters are more difficult to obtain, with decreasing maximum yields from C4 to C6[5a, b, 6] These C4- to C6-alpha-hydroxy esters contain ample functionality to make them prospective substrates for several higher-value chemicals.[9] The C4 variants thus are most promising targets that are not abundantly formed by biocatalysis. Tetrose carbohydrate substrates are rather rare and expensive, but recent approaches have shown the selective accessibility of tetroses from aldol condensation reactions from glycolaldehyde[10] or C3+C1 fragments[11] and from retro-aldol cleavage reactions of C6 sugars.[12] Likewise, C4 alpha-hydroxy esters can be formed directly in situ from larger and smaller than C4 sugars.[7a, 10, 11b, 13] Thus, C4 alpha-hydroxy esters constitute major byproducts in the formation of methyl lactate from hexoses and from disaccharides,[7a, 10] and major products in the conversion of glycolaldehyde by Lewis acidic catalysts enabling C-C bond formation.[10, 8, 10, 13a]

Introduction

A rising demand to accelerate the discovery and development of sustainable processes for biomass conversion to chemicals and fuels (materials and energy) is widely appreciated. Such developments hinge on the rapid and reliable quantification of potentially valuable products by suitable assays. In order to increase the future chances for success of sustainable processes, additional bottom-up approaches should prove valuable for evaluating catalyst function and robustness as well products formation, composition and value. Accordingly, high-resolution spectroscopic methods (especially NMR spectroscopy) have been increasingly applied to follow the progress of biomass-conversion reactions.[1] Arguably, such studies have been more widely applied in the study of lignin conversion[15] than in the carbohydrate field. Mechanistic and kinetic studies of chemocatalytic carbohydrate conversion processes thus also lag behind the widely used analogous metabolic studies of biocatalytic carbohydrate conversion.[3] As a consequence, means to obtain definitive kinetic and mechanistic insights to support the development of carbohydrate conversion are arguably not used to their full potential.[4]

As the abundant pentose and hexose carbohydrates can occur in various cyclic and acyclic forms, corresponding cyclic or open-chain pathways can play a role in the formation of carbohydrate-derived chemicals.[5] The difficulties in obtaining definitive insight into carbohydrate conversion processes are consequently well witnessed by the long-standing debate on the role of cyclic and acyclic routes of carbohydrate conversion to furanic compounds[1a, 6] which are widely investigated as future precursors for instance of polymers, pharmaceuticals, fuel additives and solvents. Acyclic routes are likely central in the formation of alpha-hydroxy esters and acids by Lewis acidic catalysts in alcohols and water. Alpha-hydroxy esters bear promise as bio-derived solvents and polymer building blocks. While the C3-metabolite lactate can be obtained in attractive yields from chemocatalytic approaches as well,[7] non-metabolite C4- to C6-alpha-hydroxy esters are more difficult to obtain, with decreasing maximum yields from C4 to C6[5a, b, 6] These C4- to C6-alpha-hydroxy esters contain ample functionality to make them prospective substrates for several higher-value chemicals.[9] The C4 variants thus are most promising targets that are not abundantly formed by biocatalysis. Tetrose carbohydrate substrates are rather rare and expensive, but recent approaches have shown the selective accessibility of tetroses from aldol condensation reactions from glycolaldehyde[10] or C3+C1 fragments[11] and from retro-aldol cleavage reactions of C6 sugars.[12] Likewise, C4 alpha-hydroxy esters can be formed directly in situ from larger and smaller than C4 sugars.[7a, 10, 11b, 13]

Thus, C4 alpha-hydroxy esters constitute major byproducts in the formation of methyl lactate from hexoses and from disaccharides,[7a, 10] and major products in the conversion of glycolaldehyde by Lewis acidic catalysts enabling C-C bond formation.[10, 8, 10, 13a]

Scheme 1. Details of the acyclic pathway of sugar conversion that were experimentally explored using erythrulose as the probe for conversion by SnCl₂ in short-chain alcohol.

Beyond their prospective roles as substrates for bio-derived polyesters and solvents, tetroses are attractive substrates to study the acyclic pathways of carbohydrate conversion.[16] Tetroses are long enough to enable various intramolecular isomerization, dehydration, cyclization and acetalization reactions, while having high acyclic populations. The ketotetrose erythrulose is an acyclic carbohydrate that cannot form stable cyclic...
conformers. Hence, erythrulose is an obvious probe molecule for studying the Lewis acid catalysed reactions in methanol through reaction progress monitoring. High-field NMR, hyperpolarization and isotope exchange studies were combined to visualize and characterize the acyclic pathway of carbohydrate conversion and to obtain definitive answers on previously hypothesized mechanisms (Scheme 1). Specifically, we sought to obtain (1) insight into the early, rate-limiting and elusive steps of carbohydrate conversion, (2) detection, structural assignment and comprehensive set of $^{13}$C NMR chemical shift assignments for intermediates, byproducts and products of the acyclic pathway, (3) detailed kinetic scrutiny of the rate limiting steps in the pathway from quantitative high-field in situ NMR on reaction progress, and (4) insight into the final, product-forming steps from kinetic isotope effects and distinction of molecules with differing isotope distributions.

Results and Discussion

Hyperpolarized NMR recognizes branching reaction from an elusive 1-oxo-2,3-enol species

A means to gain insight into initial steps of carbohydrate conversion consists in the temporary redistribution of $^{13}$C nuclear spin states to obtain a transient enhancement of the NMR signal by four orders of magnitude.$^{[14]}$ This process is called nuclear spin hyperpolarization and employs the transfer of magnetization from electrons to nuclei at low temperature prior to rapidly bringing the sample to a state that is suitable for liquid state NMR observation. The temporary enhancement of the NMR signal fades with an exponential decay of characteristic decay time on the seconds to minute time scale for non-protonated carbon sites. The vast NMR signal enhancement allows reaction tracking with sub-second to second time resolution. Hence, fast, initial conversions especially at or near the quaternary C2-position in erythrulose could be addressed using this approach (Scheme 2) even at natural isotope abundance, where only 1% of the erythrulose C2-sites carry the NMR detectable $^{13}$C isotope.

Scheme 2. Schematic depiction of the use of erythrulose at natural isotopic abundance to detect early reactions in the the SnCl$_4$-catalyzed conversion upon nuclear spin hyperpolarization. Hyperpolarization temporarily enhances the signal of the carbon backbone (solid spheres). Protonated sites largely lose the hyperpolarization during transfer (light grey spheres), while the chemistry of the enhanced quaternary C2 signal is tracked.

The resultant experimental data are displayed in Figure 1 as a time series of $^{13}$C NMR spectra. Reaction intermediates that were formed from erythrulose by SnCl$_4$ in methanol became observable within few seconds of experiment time. Most notably, elusive 3-deoxythreosone and vinylglyoxal species could be observed as the first intermediates accumulating to significant amounts in the reaction. The 3-deoxythreosone and vinylglyoxal emerged in parallel, which strongly indicated that both molecules are formed from a common precursor in competing (parallel) reactions. The precursor can presumably either be dehydrated to form vinylglyoxal or tautomerize to 3-deoxythreosone. The most plausible precursor form thus validates the previously hypothesized role of a 1-oxo-2,3-enol species as the branch point in the acyclic route of Lewis acid catalyzed carbohydrate conversion and isotope exchange studies were combined to visualize and detectable early reactions in the the SnCl$_4$ catalyzed conversion upon high-field NMR, hyperpolarization and isotope abundance, where only 1% of the erythrulose isotope is detectable.

Figure 1. Times series of molecular species detected from hyperpolarized erythrulose with SnCl$_4$/methanol at 313 K and natural isotopic abundance. Due to the rapid fading of signal from protonated sites, all detected signals arise from the C2 position. The rate constants $k_1$ and $k_2$ are as defined in Scheme 2.

Scheme 3. Reaction scheme highlighting the formation of 3-deoxythreosone and vinylglyoxal from a common elusive precursor (blue, displayed in a bracket). Partial hydrolysis of SnCl$_4$ to active species may occur, as indicated.$^{[15]}$
conversion (Scheme 3). In methanol at mild reaction conditions, both pathways showed comparable influx as witnessed by comparable signals of vinylglyoxal and 3-deoxythreosone species.

Identification and chemical shift assignment of reactants

The three rapidly formed molecules displayed in Figure 1 carry characteristic $^{13}$C chemical shifts of carbonyl groups adjacent to a 3-deoxys group and an aldehyde (~205 ppm) and a carbonyl group adjacent to a vinyl and an aldehyde group (~196 ppm), respectively. The full molecular structures were determined by acquiring two-dimensional assignment spectra on reaction media at 293 K, which slowed reactions sufficiently down to permit the detection of molecules that were formed from erythrulose by SnCl$_4$ catalysis in methanol. Such in situ 2D NMR verified that the carbonyl signal at 194.7 ppm is adjacent to a vinyl group and thus corresponds to vinyl glyoxal. Similarly, the signal at 205.5 ppm was adjacent to a methylene group, followed by a primary alcohol in 3-deoxythreosone. Hence, in situ 2D NMR on stabilized samples validates the assignments shown above (Figures S2-S4). A comprehensive list of C4 species and their corresponding $^{13}$C NMR chemical shift assignments determined in this way is shown in Figure 2.

![Figure 2](image)

Figure 2. Overview of $^{13}$C chemical shift assignments of compounds identified in a stabilized reaction mixture at 293 K with methanol-d$_4$ as the solvent.

Real time NMR of the acyclic pathway at different temperatures

The comprehensive $^{13}$C chemical shift assignments of Figure 2 were subsequently validated by identical amounts and kinetic profiles in in situ NMR experiments. These experiments were used to follow the overall reaction progress beyond the initial branching point that was identified with hyperpolarized NMR. High-field NMR on an 800 MHz instrument equipped with cryogenically cooled detection electronics was used to obtain time series of $^{13}$C NMR spectra. The $^{13}$C NMR signal sensitivity provided by the above system allows the acquisition of $^{13}$C NMR spectra with good time resolution of approximately 5 minutes per spectrum (Figure 3). The $^{13}$C NMR spectra employed inverse-gated decoupling only during data acquisition in order to avoid enhancement of protonated sites. Reactions were conducted using stock solutions of reactants to warrant reproducibility when performing reaction kinetic analyses at 293, 313, and 333 K. Integrals of the C4 positions with the lowest $^{13}$C chemical shifts of carbonyl groups adjacent to a primary alcohol ($~205$ ppm) and an aldehyde ($~205.5$ ppm) were determined by a primary alcohol in 3-deoxythreosone. Hence, in situ 2D NMR on stabilized samples validates the assignments shown above (Figures S2-S4). A comprehensive list of C4 species and their corresponding $^{13}$C NMR chemical shift assignments determined in this way is shown in Figure 2.

![Figure 3](image)

Figure 3. Time series of $^{13}$C NMR spectra at 313 K. Reaction conditions: 30 mg erythrulose, 6 mg SnCl$_4$ 5H$_2$O, 600 µl methanol-d$_4$.

The time series of $^{13}$C NMR spectra of reaction progress as shown in Figure 3 displayed that the overall reaction from erythrulose proceeded as a reaction with HA-MEG as the principal intermediate, which accumulates due to its slower conversion to the main product MMHB. The time series provided excellent sensitivity of the baseline-separated $^{13}$C signals for reliable kinetic analyses of the main reactants. Signal areas in the time series were normalized to the methanol solvent signal to account for theoretically possible NMR instrument instabilities. The reaction of erythrulose was of first order with respect to the substrate concentration, so that substrate decay followed an exponential decay at all three temperatures (Figure 4 and S5). Owing to the low accumulation of other intermediates, the conversion of substrate and of HA-MEG clearly have the lowest rate constants in the acyclic pathway of erythrulose to MMHB. The reaction was therefore treated as a consecutive reaction of two elementary steps. The first of these steps corresponds to a net intramolecular redox reaction with oxidation of C1 and reduction of C3 under concomitant formation of a methyl ether at C4. The second step to MMHB in turn results in the further oxidation of C1 and reduction of C2. Byproducts include the furanose acetals methyl-threoside and methyl-erythrose in addition to the dimethyl acetal of the MEG intermediate. These findings underline that erythrulose can isomerize to some degree to its aldose tautomers$^{[17]}$ under the current reaction conditions and that acetals form poorly-reactive, masked byproducts in the conversion of carboxydrates in alcohols (Figure S1)$^{[15]}$.
The signals of the substrate erythrulose and HA-MEG intermediate both approach zero and the conversions were treated as overall irreversible steps. Resultant fits of reaction progress curves to a consecutive two-step kinetic model are shown in Figure 4 and the obtained rate constants are plotted on a logarithmic scale versus the inverse absolute temperature for Arrhenius analysis in Figure 5. This analysis yields the apparent activation energies of the two main reaction steps converting erythrulose to HA-MEG. The linear trend in the Eyring analysis yielded estimates of activation enthalpies of 66 kJ/mol for the conversion of erythrulose to HA-MEG and of 85 kJ/mol for the subsequent conversion of HA-MEG to MMHB. These values are consistent with previous reports of the apparent activation energy for SnCl₄-catalyzed conversion of erythrulose to MMHB (81 kJ/mol)[8] and are plausible for reactions that progress during few hours in the temperature range that was investigated. The temperature-dependence of rate constants indicates that HA-MEG formation remains the faster step than its conversion over a wide temperature range, which shows that HA-MEG will accumulate as a major intermediate under relevant operation conditions of the homogeneous catalytic system in methanol. Kinetic analysis of hyperpolarized and thermal NMR data in combination showed that the dehydration of erythrulose to vinylglyoxal was followed by approximately 20-fold faster (to HA-MEG) and 20-fold slower steps to the final product MMHB in the homogeneous reaction (Figure 5, bottom).

Figure 4. Integrated signal areas for the three main reactants at 293 K, 313 K, and 333 K. Areas are normalized to the solvent signal. Reaction conditions: 30 mg erythrulose, 6 mg SnCl₄·5H₂O, 600 µl methanol-d₄.

Having identified the main intermediates and visualized their fast and slow conversion in situ, we strived to gain insight into kinetic isotope effects, the reversibility of pathway steps and the stereochemistry of the intramolecular redox reactions through isotope incorporation studies.

Previous low-field NMR analysis of the conversion of erythrulose in deuterated methanol (methanol-d₄) had detected the formation of only one MMHB product species, singly deuterated at C₃[13]. The presence of only one NMR-detectable species would be surprising considering that such a compound should be diastereomeric due to asymmetric mass distribution at C₃ next to a chiral C₂ site. In addition, analogous pathways to methyl lactate had indicated the ability to introduce more than one solvent hydrogen at C₃ due to the reversibility of intermediate enolization/ketonization steps.[18] Also, surprisingly high amounts of products protonated at C₃ in deuterated solvent[19] could raise the question of possible isotope effects in the ketonization of the C₂ position.

Isotope labeling studies and quantitative NMR indicate that the substrate is the major source of labile protons

The amounts of protonated and deuterated product were determined herein using quantitative ¹³C NMR spectroscopy of product mixtures obtained in methanol-d₄ with more than 99.8% nominal degree of deuteration. NMR spectroscopy on the reaction mixture indicated that the majority of labile protons that are incorporated during the acyclic pathway neither arise from spurious water nor from lower-than-nominal deuteration of the solvent, but from the substrate and catalyst hydroxyl groups themselves. Nevertheless, protonation of the reaction products was at least double of what was expected from the fraction of labile protons introduced by the substrate relative to the deuterated solvent. Figure 6 shows the formation of protonated and single deuterated HA-MEG intermediate that is converted to

Figure 5. Eyring plots for the reaction progress kinetic analysis of erythrulose conversion by SnCl₄·5H₂O as displayed in Figure 4. Relative experimental reaction rates for distinguishable steps at 313 K are displayed for comparison.
minor amounts of the dimethylacetal and major amounts of MMHB. The reaction profiles indicate that the surprisingly high amounts of protonated MMHB\textsuperscript{[1]} can be ascribed at least in part to isotope effects that lead to a faster reaction of the protonated than the deuterated intermediate. The intermediates are converted to non-deuterated, single deuterated and double deuterated products as shown in Figure 7. Due to the asymmetric mass distribution at C3, the single deuterated product MMHB yields two different signals for the two pairs of diastereomers.

consistent with reports of multiple deuteration at C3 in the formation of methyl lactate\textsuperscript{[19]} This observation showed the reversible operation of enolization/ketonization steps under the catalytic conditions employed. The kinetic profiles of the different isotopologues differ noticeably, thus supporting the notion that an isotope effect in the ketonization may affect product distribution between the different isotopic forms (Figure 7).

Diastereomeric deuteration indicates low stereoselectivity
The stereoselectivity of the reactions converting erythrulose to MMHB were subsequently addressed. Such investigations are possible with NMR spectroscopy because enantiomeric forms become observable upon diastereomeric deuteration at the C3 position in the main intermediate HA-MEG and the main product MMHB. The relative population of C3 protons in HA-MEG and its dimethylacetal as well as MMHB can be visualized using \textsuperscript{1}H-\textsuperscript{13}C HSQC spectra of the respective spectral regions for these CH groups (Figure 8). The C3 protons in the dimethylacetal are enantiotopic (as C1 carries two identical substituents), yielding only one CH signal. In contrast, the HA-MEG and MMHB methylene protons are diastereotopic due to the presence of a chiral center at C1 and C2, respectively.

Reversibility of enolization and multiple deuteration
Notably, we also identified the double deuterated MMHB product, in contrast to previous reports on the C4 system but

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**Figure 6.** (A) Isotopologues formed in the conversion of protonated erythrulose in deuterated methanol as tracked in real time using a series of \textsuperscript{13}C NMR spectra. (B) Zoom of the highlighted area in (A) showing that deuterated, aliphatic C3 positions are evident from 1:1:1 triplet signals due to coupling to the spin-\textsuperscript{1}H nucleus. (C) Comparison of formation and conversion of C3-deuterated and C3-protonated intermediate showing slower reactions of the deuterated form and fits to equations for intermediates of consecutive reactions. Reaction conditions: 50 mg erythrulose, 25 mg SnCl\textsubscript{4}, 5H\textsubscript{2}O, 600 µl methanol-d\textsubscript{4}, 323 K.

**Figure 7.** (A) Formation of MMHB in the experiment of Figure 6, resolving diastereomeric forms with single-deuteration at C3, in addition to protonated and deuterated forms. (B) Kinetic profiles of the different forms show significant differences, with the curves in the bottom representing normalized fits of the data for the three different isotopic forms to highlight the kinetic disparities.

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Both in HA-MEG and MMHB, the two triplets of the CHD group signals indicated similar propensities of deuterium to enter either of the two methylene group positions. The diastereomeric excess as determined from the HSQC integrals was less than 2.3% both for HA-MEG and MMHB, while values above 45% have been found for the C3 positions in C5 carbohydrates converted by heterogeneous Sn-containing zeolite.\textsuperscript{[19]} The lack of stereoselectivity in the formation of MMHB, indicated that both the ketonization of C2 and the hydride shift converting HA-MEG to MMHB have little stereoselectivity. This finding contrasted the significant stereoselectivity in larger sugars,\textsuperscript{[14, 10, 19]} where higher populations of cyclic intermediates may affect the stereochemistry of the aforementioned reactions due to different reactivities of axial and equatorial hydrogen atoms.\textsuperscript{[20]}

Finally, we observed that the propensity to incorporate deuterium at methylene C3 positions differs measurably in different pathways. Thus, HBL and its open chain equilibrium form MHB (Scheme 3) are not observed to carry significant deuterium at the C3 position upon reactions in deuterated methanol. This finding is consistent with the incorporation of one deuterium atom at C3 in the Michael addition of deuterated methanol to vinylglyoxal intermediate. MMHB was found to incorporate significant amounts of more than one deuterium atom at the C3 position, presumably through reversible enolization of the HA-MEG (and related forms such as pyruvaldehyde)\textsuperscript{[19]} intermediates in catalyst bound forms in parallel to product formation (Figure 7B, inset). As a consequence, a differential propensity for the incorporation of deuterium at the C3 position is found in various steps (Figure S6). These steps include reactions such as the conversion of HA-MEG to MMHB that does not require enolization, but rather is susceptible to competing enolization of the substrate.

Figure 8. Methylene groups at the C3 position when converting erythrulose in deuterated methanol. HA-MEG and MMHB are diastereomers upon single deuteration at C3, yielding non-equivalent sites in the methylene group. Stereoselectivity for deuterium incorporation at the nonequivalent sites is marginal.

In conclusion, we find that the acyclic pathway of carbohydrate can be probed in mechanistic and kinetic detail using hyperpolarized NMR methods, \textit{in situ} NMR at high magnetic field and isotope tracking methods. These approaches give fundamental insight into the initial reaction steps, pathway bottlenecks, byproducts and stereochemistry as well as mechanistic differences in the product forming reactions (Scheme 4). Detailed kinetic and mechanistic studies of C4 products are warranted, as C4 products are accessible not only from C4 carbohydrates, but also from smaller sugars\textsuperscript{[10, 11b, 13a]} and larger\textsuperscript{[12, 13b]} carbohydrates. Competing formation of a 3-deoxyglycosone and vinyl glyoxal species from a transient undetected enol species is the first observable step under the current reaction conditions in methanol as the solvent. Hydrogen incorporation and hydrogen/deuterium isotope effects differ in the pathways following the competing reactions. Two predominant energetic barriers with activation enthalpies of activation of 66 and 85 kJ/mol as well as entropies of activation of -97 and -81 J/(mol·K) are found in the pathway from erythrulose to the main product MMHB. Despite the presence of only one chiral carbon atom in the main intermediate HA-MEG (at C1) and the main product MMHB (at C2), diastereomeric forms due to asymmetric mass distributions at C3 in deuterated solvent reveal that ketonization reactions and final Cannizzaro-type reaction proceed with low stereoselectivity of less than 2.3% diastereomeric excess.
Experimental Section

SnCl\textsubscript{2}·5H\textsubscript{2}O, erythroluse and methanol-d\textsubscript{4} were purchased from Sigma Aldrich (St. Louis, MO, USA). Substrate and catalyst were separately dissolved as stock solutions and were mixed on ice prior to transfer to a 5 mm NMR tube for \textit{in situ} NMR spectroscopy. \textit{In situ} \textsuperscript{13}C NMR spectra were recorded on a Bruker (Fällanden, Switzerland) Avance II 800 MHz spectrometer equipped with an 18.7 T magnet (Oxford Magnet Technology, Oxford, U.K.) and a TCI Z-gradient CryoProbe with sample temperature equilibrated to 293 K, 313 K or 333 K, respectively. Reaction progress under static NMR tube conditions without further mixing was followed by pseudo-2D spectra recording a series of \textsuperscript{13}C NMR spectra employing inverse-gated decoupling, \textit{m} of excitation pulses and inter-scan recycle delays of 4.7 seconds. Kinetic data of reaction progress were integrated in Bruker Topspin 3.0 and fitted to a model of sequential elementary reactions in profit 7 (Quantum Soft, Switzerland). Reactions were slow enough at 298 K to conduct two-dimensional assignment spectra of the reaction mixture on an 800 MHz spectrometer to identify reaction intermediate structures of reactions initiated at 293 K, 313 K or 333 K. Identical reaction profiles and similar amounts of \textsuperscript{13}C signals were used to validate the assignment of the signals to the same molecule. After the reaction, quantitative \textsuperscript{1D} \textsuperscript{13}C NMR spectrum using inverse-gated decoupling was acquired with a recycle delay of 80 seconds and 1024 scans. High-resolution \textsuperscript{1H}-\textsuperscript{13}C HSQC spectra were used to assess the stereoselectivity of reactions \cite{1} in the C1, C2 and C3 positions of the tetrose.

Hyperpolarization of natural abundance erythroluse was performed using trityl radical OX63 (Oxford Instruments, Abingdon, UK) and trimeric gadolinium chelate of \textit{1,3,5-tri-(N-(DO3A-acetamido)-N-methyl-4-aminomethyl-2-methylphenyl)-1,3,5} triazine, 2,4,6-trione (GE Healthcare). Samples contained 100 mg of a self-mixing solution of OX63 (15 mM) and gadolinium chelate (1.5 mM) directly dissolved in erythroluse. The polarization transfer was conducted at 1.2 K by microwave irradiation at 93.89 GHz with 100 mW using a HyperSense (Oxford Instruments, England) polarizer. After 1.5 hours of polarization, the samples were dissolved with 5 mL heated methanol-d\textsubscript{4}, 300 \textmu{}L of the hyperpolarized substrate was injected to a solution of SnCl\textsubscript{2}·5H\textsubscript{2}O (final concentration 18 mg/mL) in methanol-d\textsubscript{4}, placed inside a 400 MHz Varian spectrometer. An array of 128 experiments of 1 second each was started with a nominal flip angle of 10\degree. Data were integrated in Bruker Topspin 3.0 and a non-linear least squares fit was performed in MATLAB to obtain two reaction rates \(k_1\) and \(k_2\) as well as the decay constants of substrate, intermediate and product hyperpolarization (23, 37 and 39 seconds, respectively, consistent with declining proton density near C2). On a separate sample the polarization level was determined to 24% by comparison to a reference NMR spectrum with same pulse and acquisition parameters as for the hyperpolarized sample after addition of 10 \textmu{}L of 0.5 M commercial Gd-complex (Omnispec, GE Healthcare). \(T_1\) of the C2 position in erythroluse was determined to 25 s.

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Keywords: biomass • kinetics • Lewis acid • NMR • reaction pathway

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Elusive intermediates and reactions can be visualized by examining the acyclic pathways of carbohydrate conversion with erythrulose substrate and in situ NMR detection.

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