Showcasing research from Markus Draskovits, Christian Stanetty et al. from the Bioorganic Synthetic Chemistry group, Institute of Applied Synthetic Chemistry, TU Wien, Vienna, Austria.
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Intercepted dehomologation of aldoses by N-heterocyclic carbene catalysis – a novel transformation in carbohydrate chemistry

Exploiting NHCs’ unique selectivity for aldehydes furnished an intercepted dehomologation protocol for reducing aldoses. Principles of substrate governance for intercepted dehomologation or a subsequent redox-lactonisation were identified and with structurally unbiased substrates, catalyst design allowed tuning the selectivity towards either of these two scenarios.

As featured in:

See Christian Stanetty et al., Chem. Commun., 2019, 55, 12144.
Intercepted dehomologation of aldoses by N-heterocyclic carbene catalysis – a novel transformation in carbohydrate chemistry†

Markus Draskovits, Hubert Kalaus, Christian Stanetty* and Marko D. Mihovilovic

The development of an N-heterocyclic carbene (NHC) catalysed intercepted dehomologation of aldoses is reported. The unique selectivity of NHCs for aldehydes is exploited in the complex context of reducing sugars. Examples of strong substrate governance for either intercepted dehomologation or a subsequent redox-lactonisation were identified and mechanistically understood. More importantly, it was shown that catalyst design allowed the tuning of the selectivity of the reaction with structurally unbiased starting materials towards either of the two scenarios.

Carbohydrates are the most abundant class of all biomolecules formed in Nature and are found in all living organisms. They serve as structural components of cells, as energy sources within metabolic cycles and also play a major role in the recognition process of biomolecules.1 Despite their ubiquitous nature, only a limited number of representatives of the family of carbohydrates are readily available, requiring great synthetic efforts to provide the full range of carbohydrates.2 Great effort and progress have been made in the assembly of complex oligosaccharides, often based on elaborative protecting group chemistry to overcome the challenges of both regio- and stereoselectivity with respect to the hydroxyl functionalities.3 Intriguingly, the intrinsically reactive part of an aldose, the aldehyde moiety, is targeted far less in classic carbohydrate chemistry.4 The established methods include addition of organometallics (Mg, In, and Zn)5 or carbanions (Kiliani–Fischer)6 to achieve an elongation. Alternatively, also dehomologation towards shortened carbohydrates (e.g. by the Wohl degradation) initiate by addressing the aldehyde moiety.7 Requirements of stoichiometric amounts of reagents and/or limited compatibility with unprotected aldoses due to solubility and cross-reactivity with the free hydroxyl groups are major challenges in this field.8

In this light, we perceived great potential for transformations triggered by the highly aldehyde-specific interaction of N-heterocyclic carbenes (NHCs) with the (anomeric) carbonyl moiety which we have just started to exploit. After the isolation of the first bench-stable carbene by Arduengo,9 the number of applications of NHCs has vastly increased over the last decades. The main areas of research have been their utilisation as ligands in transition-metal catalysed reactions10 and as organocatalysts.11 The field of organocatalysis is dominated by processes rooted in the umpolung of aldehydes,12 triggering benzoin or Stetter type reactions13 as well as more sophisticated follow-up transformations.14

The applications of NHCs within the realm of reducing sugars, as aldehyde species, are extremely scarce (see Scheme 1).15 Wendelnorn et al. attempted to perform Stetter reactions of fully protected reducing aldoses but instead observed dominant β-elimination towards protected 2-deoxy lactones,16 consistent with earlier studies on α-reducible aldehydes.17 The group of Chi achieved the catalytic activation of fully unprotected aldoses by NHCs generating multiple nucleophilic formaldehyde equivalents.

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via a retro-benzoin reaction in an uncontrolled fashion. The thereby achieved formylation of α,β-unsaturated compounds in a subsequent Stetter reaction was the goal of the study, with the sugars being utilised as a sacrificial feedstock.18

Inspired by these works and building upon the proposed mechanism, we set out to develop an NHC controlled intercepted dehomologation methodology of carbohydrate derivatives. See Scheme 2 for an outline of the mechanistic consideration with 3-O-Bn-glucose 1a as an example. First, the carbene, formed 

**Scheme 2** Proposed mechanism for the NHC intercepted catalyzed dehomologation reaction of aldoses.

of 2-hydroxy-γ-butyrolactone 9a, thus reflecting successful double dehomologation but with concomitant redox-lactonisation via elimination of benzaldehyde. According to our hypothesis the stability of the starting material, intermediates and final products as lactols is responsible for the observed differences in selectivity. Consequently, 3-O-Bn glucose 1a gave mixtures of the dehomologated products 2-O-Bn arabinose 3 and 2-deoxyribonolactone 4a upon NHC-catalysed elimination of BnOH (Scheme 3, bottom) reflecting similar lactol stabilities of 1a/3. To corroborate the above mechanism, we subjected 2-O-Bn arabinose 3 to the standard reaction conditions and confirmed deoxy lactone 4a as the main product. Furthermore, by installing a better leaving group at O3 as in 3-O-(p-nitrophenyl) glucose 10 the subsequent elimination to 4a was dominant under the same reaction conditions (see the ESI†).

Our ultimate goal was to achieve catalyst control over the selectivity between intercepted dehomologation and subsequent redox-lactonisation. Therefore, we selected 3-O-Bn glucose 1a and its 20-epimer 3-O-Bn mannose 1b as model compounds for the re-evaluation of the reaction conditions18 and further catalyst development. In advance, they serve as probes for potential influence of the relative stereochemistry at C2/C3. Both substrates and their common reaction products (3 and 4a) are accessible in a few chemical steps (see the ESI†). A reliable procedure for an efficient and quantitative screening was required, given the complexity of crude reaction mixtures. We developed a method based on a solid phase extraction (SPE) with subsequent derivatisation of all carbohydrate species to allow quantification via a calibrated GC protocol (see details in the ESI†).21

Preliminary studies clearly showed that sub-stoichiometric amounts of base compared to the pre-catalyst (0.20 : 0.25 equiv.) and a sufficient reservoir of chalcone (2.0 equiv.) are mandatory requirements. Under these conditions, the influence of solvent, type of base as well as lower and higher temperatures has been evaluated which is summarized for 1b as a starting material in
Table 1 Screening of reaction conditions of the NHC-catalyzed dehomologation

| Entry | Deviation from standard conditions | 1b\% | 3a\% | 4a\% | Sum\% |
|-------|----------------------------------|------|------|------|-------|
| 1     | None                             | 6    | 48   | 43   | 96    |
| 2     | Solvent:McCN\*                   | 0    | 4    | 63   | 67    |
| 3     | Solvent:DMF\*                    | 0    | 9    | 36   | 45    |
| 4     | Solvent:EtOH\*                   | 0    | 3    | 30   | 33    |
| 5     | Base:Li2CO3                      | 14   | 46   | 18   | 78    |
| 6     | Base:DBU                         | 83   | 14   | 4    | 101   |
| 7\*   | T = 90°C/320 min                 | 7    | 18   | 42   | 67    |
| 8\*   | T = 110°C/80 min                 | 0    | 4    | 44   | 48    |
| 9\*   | T = 150°C/10 min                 | 0    | 2    | 18   | 20    |

\* Reaction conditions: 1b/2/base/5 = 1.00 : 0.25 : 0.20 : 2.00 (molar ratio), 20 min. \* Based on calibrated GC. \* Reactions performed under \(\mu\)W irradiation.

Table 2 Catalyst optimisation towards increased selectivity

| Entry | Starting material | Catalyst | 1b\% | 3a\% | 4a\% | Ratio 3/4a |
|-------|------------------|----------|------|------|------|------------|
| 1     | 1b               | 1b       | 2    | 6    | 48   | 43         |
| 2     | 1b               | 1b       | 12   | 63   | 29   | 9          |
| 3     | 1b               | 1b       | 13   | 43   | 30   | 17         |
| 4     | 1b               | 1b       | 14   | 93   | 2    | 4          |
| 5     | 1b               | 1b       | 15   | 25   | 39   | 10         |
| 6     | 1b               | 1b       | 16   | 7    | 78   | 13         |
| 7\*   | 1b               | 1b       | 16   | —    | 82   | 6          |

\* Reaction conditions: 1b/precatalyst/K2CO3 = 1:0.25:0.20:2.00 (molar ratio), 20 min. Based on calibrated GC. \* Used instead of 1b.

dehomologated to the elimination product, and with diisopropylphenyl substituted thiazolium salt 16, a 6:1 ratio in favour of the dehomologated product 2-O-Bn arabinose 3 was observed. In a separate experiment 3 was reacted with precatalyst 16 (Table 2, entry 7) giving no notable conversion to deoxy lactone 4a, confirming a strong selectivity of 16 for the reaction with 1b over 3. The GC-data obtained in DMSO were validated by a preparative experiment in MeCN and under microwave irradiation, which led to a comparable isolated yield (1 mmol scale, Scheme 4). The fact that the gluco-analogue 1a gave inferior results (ESI\#) for us strongly indicates competing side reactions which dominate in the case of species of high lactol stability (low open chain content). The above data was initially found to show that increasing the steric congestion around the carbene centre increases the selectivity for the intercepted dehomologation. We assumed this to be due to the steric clash between the 2-O-substituent of 3 and the diisopropyl substituent of the catalyst.

Therefore, we next evaluated triphenyl substituted triazolium salts as precatalysts, initially aiming at introducing additional steric bulk around the carbene centre. We discovered that the parent Enders’ triphenyl triazolium salt 17\* led to the exclusive formation of the deoxy lactone 4a from 1a in high GC-yields (Table 3, entry 2) at all obtained time points. We attempted to achieve comparable shifts in the selectivity towards the dehomologated product 3 within this catalyst family. Nevertheless, increasing the steric bulk near the carbene moiety via

Scheme 4 Catalyst controlled divergence in a preparative fashion.
introducing again mesityl or diisopropylphenyl on one adjacent nitrogen atom (precatalysts 18 and 19, respectively) did not lead to a change in chemo-selectivity, as seen in the thiazolium salt series. Instead we observed a decrease in the conversion to 4a, indicating that the electronic properties of the triazolium core dominate the steric effects (Table 3, entries 3 and 4). Again, the reaction with the most promising GC-yield was repeated with MeCN as solvent under microwave irradiation, giving lactone in a good isolated yield (5 mmol, upon acetylation, see Scheme 4). With both the ideal substrate/catalyst combinations it was attempted to decrease catalyst loading which leads to a significant decrease in the conversion (see the ESI†).

In summary, we have delivered a clear proof of concept for the principle feasibility of an NHC-controlled intercepted dehomologization of semi-protected carbohydrate derivatives. Herein, we present the first examples of substrate-dependent and – more importantly – catalyst-controlled divergence between selective intercepted dehomologation based on the retro-benzoin reaction on the one hand and the subsequent β-elimination on the other hand. Screening of our catalysts revealed the influence of both steric and electronic properties of carbenes identifying the first ideal substrate/catalyst combinations. Further studies on the scope and limitations of the current methodology are in progress and will deliver an increased understanding of and give rise to more means of exploitation of the fascinating interaction between NHCs and the aldoses’ aldehyde moieties.

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Conflicts of interest

There are no conflicts to declare.