Aldehyde Coupling Reactions

Enantioselective Organocatalytic Direct Aldol Reactions of α-Oxyaldehydes: Step One in a Two-Step Synthesis of Carbohydrates**

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The growing study of glycobiology^[1] has led to an increased focus upon carbohydrate architecture^[2] as an important platform for reaction design and methodological advancement.[3] Application of the aldol reaction[4] to the synthesis of carbohydrates is well-documented;^[5] however, the attendant need for protection-group manipulations and oxidation-state adjustments has thus far precluded a broadly utilizable protocol. Intriguingly, a highly expedient two-step carbohydrate synthesis can be envisioned based on an iterative aldol sequence using simple α -oxyaldehydes [Eq. (1)]. While attractive in theory, the practical execution of this carbohydrate strategy would require the invention of two new aldol technologies: a) an enantioselec-

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tive aldol union of α -oxyaldehyde substrates (Aldol step 1) and b) a diastereoselective aldol coupling between tri-oxy substituted butanals and an α -oxyaldehyde enolate (Aldol step 2). Herein we report the successful development of the first enantioselective organocatalytic coupling of an α -oxyaldehyde (Aldol step 1). This new aldol reaction provides an operationally simple protocol for the stereocontrolled production of polyol architectures and sets the stage for a two-step enantioselective carbohydrate synthesis. [6]

The development of a direct, enantioselective catalytic aldol reaction between $\alpha\text{-}oxyaldehyde$ substrates (Aldol step 1) is dependent upon three key issues of chemical selectivity. In addition to the traditional requirements of absolute and relative stereocontrol comes the chemoselective constraint that the $\alpha\text{-}oxyaldehyde$ reagent \boldsymbol{A} must readily participate as both a nucleophilic and electrophilic coupling partner while the $\alpha\text{-}oxyaldehyde$ product \boldsymbol{B} must be inert to in situ enolization or carbonyl addition [Eq. (1)]. Recently,

Two-Step Carbohydrate Synthesis: Iterative Aldehyde Aldol

Aldol 1 requires α -oxyaldehyde **A** (reagent) is reactive in aldol union

Aldol 1 requires α -oxyaldehyde **B** (product) is nonreactive in aldol union

Proline-Catalyzed Cross Aldehyde - Aldol Addition

H

Me

$$H$$
 L -Proline

 $DMF, 4 °C$
 C
 H
 C
 Me
 C
 Me
 C
 Me
 C
 Me
 C
 C
 Me

Organocatalytic Aldol 1: Enantioselective α -Oxyaldehyde Coupling

we disclosed an organocatalytic strategy for the highly regioselective, diastereoselective, and enantioselective aldol cross-coupling of α -alkyl-bearing aldehydes [Eq. (2)]. [8] An important feature of this transformation is that the enantioenriched aldehyde products \mathbf{C} do not participate in further aldol reactions (by either enamine formation or carbonyl addition). With this in mind, we hoped that such remarkable catalyst-controlled stereo- and chemoselectivity might be extended to the union of α -oxygenated aldehydes [Eq. (3)], thereby allowing the first step in a two-step carbohydrate synthesis to occur [Eq. 1].

Our enantioselective organocatalytic α -oxyaldehyde coupling was first examined using L-Proline (10 mol%) and a

substrates variety of glycoaldehyde (Table 1). Preliminary studies revealed that the proposed enantioselective aldol union is indeed possible, however, the electronic nature of the oxyaldehyde substituent has a pronounced effect on the overall efficacy of the process. For example, substrates that possess an electron-withdrawing substituent, such as α-acetoxyacetyaldehyde 1a, do not participate in this transformation, while aldehydes bearing relatively electron-rich oxyalkyl groups provide useful levels of enantiocontrol and reaction efficiency (entry 2, R = Bn, 73 % yield, 98 % ee; entry 3, R = PMB, 85% yield, 97% ee). Moreover, aldehydes bearing bulky α-silyloxy substituents can be readily utilized (entry 5, R = TBDPS, 61 % yield, 96 % ee; entry 7, PG = TBS, 50 % yield, 88% ee), with the TIPS-protected glycoaldehyde (entry 6) affording exceptional reaction efficiency (92%), enantioselectivity (95% ee), and a readily separable 4:1 mixture of anti and syn diastereomers. It should be noted that all of the dimeric aldol adducts shown in Table 1 constitute protected forms of the naturally occurring sugar erythrose, a chiral synthon of established utility. [9] More importantly, the α oxyaldehyde products of this new aldol protocol are apparently inert to further proline-catalyzed enolization or enamine addition, a central requirement for the proposed two-step iterative-aldol carbohydrate synthesis [Eq. (1)].[10]

We next examined the ability of proline to catalyze the enantioselective cross-coupling of α -oxy- and α -alkyl-substituted aldehydes (Table 2). The principal issue in this reaction is that the nonequivalent aldehydes must selectively partition into two discrete components, a nucleophilic donor and an electrophilic acceptor. Given that most α -oxy- and α -alkyl aldehydes bear enolizable protons, we anticipated that such catalyst-controlled substrate partitioning would be mechanistically unfavorable. Remarkably, however the glycoaldehyde invariably acts as the electrophile in the presence of alkyl aldehydes that contain αmethylene protons (entries 1-4, 94-99% ee). Surprisingly, even the sterically demanding isovaleraldehyde assumes the role of nucleophile when exposed to proline and α-benzyloxyacetaldehyde or α-silyloxyacetaldehyde (entries 3 and 4). However, both triisopropylsilyl- and benzyl-protected oxyaldehydes can function as aldol donors in the presence of aldehydes that do not

Table 1: Organocatalytic aldol dimerization of α -oxyaldehydes.

Entry	Product	Solvent	Yield [%]	anti:syn	ee [%] ^{[a],[l}
1	O OH OAc OAc Za	DMF	0	-	-
2	O OH OBn	DMF	73	4:1	98
3	O OH OPMB	DMF	64	4:1	97
4	H OMOM OMOM	DMF	42	4:1	96
5	O OH OTBDPS OTBDPS 2e	DMF/dioxane	61	9:1	96 ^[c]
6	O OH OTIPS OTIPS 2f	DMSO	92	4:1	95
7	O OH OTBS OTBS 2g	dioxane	62	3:1	88 ^[c]

[a] Absolute and relative stereochemistry assigned by chemical correlation. [b] Determined by chiral HPLC. [c] Using 20 mol% catalyst. Bn = benzyl, PMB = para-methoxybenzyl, MOM = methoxymethyl, TBDPS = tert-butyldiphenylsilyl, TIPS = triisopropylsilyl, TBS = tert-butyldimethylsilyl.

Table 2: Cross-aldol reactions with protected glycoaldehydes.

O ON
$$R$$
 $OX POINT POIN$

Entry	Aldehyde		Product	Yield [%]	anti:syn	ee [%] ^{[a],[b]}
	α-alkyl	OX				
1	H Me	OTIPS acceptor	H ON OH	75	4:1	99
2	donor	OTBDPS acceptor	Me	84	5:1	99 ^[c]
3	H iPr	OTIPS acceptor OBn	H ON OH OX	54	4:1	99
4	donor	acceptor	Me Me	64	4:1	94
5	H Me Me	OTIPS donor	O OH Me	43	8:1	99
6	acceptor	OBn donor	ŌX Me	33	7:1	96

[a] Absolute and relative stereochemistry assigned by chemical correlation. [b] Determined by chiral HPLC. [c] Determined by Mosher ester analysis.

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readily participate in enamine formation (entries 5 and 6, \geq 33% yield \geq 7:1 *anti:syn*, 96–99% *ee*). It should be noted, however, that significant quantities of the homodimers **2 f** and **2 b** were generated in these respective cases.

These organocatalytic results stand in marked contrast to metal-mediated direct aldol technologies [11] where the increased acidity and nucleophilicity afforded by α -oxygenated aldol donors greatly enhances their effectiveness relative to their all-alkyl counterparts. We are currently investigating the mechanistic origins of such divergent reactivity between metal and organic catalysts in aldol reactions with α -oxygenated substrates.

In summary, we have documented the first direct enantioselective catalytic aldol reaction using α -oxygenated aldehydes as both the aldol donor and the aldol acceptor. Significantly, this method allows direct and enantioselective access to differentially protected polyols and monoprotected *anti*-1,2 diols. A full account of these studies will be presented in due course.

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- a) Glycoscience: Chemistry and Chemical Biology I-III (Eds.: B. Fraser-Reid, K. Tatsuta, J. Thiem), Springer, 2001; b) Glycochemistry: Principles, Synthesis, and Applications (Eds.: P. Wang, C. Bertozzi), Marcel Dekker, 2001.
- [2] While the term carbohydrate can be applied to many hydrated forms of carbon structure, we employ this terminology in the more commonly used and specific sense to describe hexose architecture.
- [3] a) K. M. Koeller, C.-H. Wong, Chem. Rev. 2000, 100, 4465;
 b) K. C. Nicolaou, H. J. Mitchel, Angew. Chem. 2001, 113, 1624;
 Angew. Chem. Int. Ed. 2001, 40, 1576.
- [4] For some reviews of the aldol reaction, see: a) B. Alcaide, P. Almendros, Eur. J. Org. Chem. 2002, 10, 1595; b) T. D. Machajewski, C.-H. Wong, Angew. Chem. 2000, 112, 1406; Angew. Chem. Int. Ed. 2000, 39, 1352; c) R. Mahrwald, Chem. Rev. 1999, 99, 1095; d) D. A. Evans, J. V. Nelson, T. Taber in Topics in Stereochemistry, Vol. 13, Wiley, 1982, p. 1.
- [5] For recent examples of aldol reactions in the syntheses of carbohydrates, see: a) D. A. Evans, E. Hu, J. S. Tedrow, Org. Lett. 2001, 3, 3133; b) S. G. Davies, R. L. Nicholson, A. D. Smith, Synlett 2002, 10, 1637; c) M. P. Sibi, J. Lu, J. Edwards, J. Org. Chem. 1997, 62, 5864; d) for a review of aldolase enzymes in carbohydrate synthesis, see: S. Takayama, G. J. McGarvey, C.-H. Wong, Chem. Soc. Rev. 1997, 26, 407.
- [6] A two-step carbohydrate synthesis has recently been accomplished in our laboratories. Details of this work will be published at a later date.
- [7] For examples of enamine-catalyzed aldol reactions between α-oxyketones and -aldehydes, see: a) W. Noltz, B. List, J. Am. Chem. Soc. 2000, 122, 7386; b) K. Sakthivel, W. Notz, T. Bui, C. F. Barbas III, J. Am. Chem. Soc. 2001, 123, 5260.
- [8] A. B. Northrup, D. W. C. MacMillan, J. Am. Chem. Soc. 2002, 124, 6798.
- [9] For uses of erythrose in synthesis, see: a) W. H. Pearson, E. J. Hembre, J. Org. Chem. 1996, 61, 7217; b) M. Ruiz, V. Ojea, J. M. Quintela, Synlett 1999, 2, 204; c) J. G. Buchanan, A. R. Edgar, B. D. Hewitt, J. Chem. Soc. Perkin Trans. 1 1987, 2371.

- [10] A proline-catalyzed trimerization of propionaldehyde to form nearly racemic tetrahydropyrans in good diastereoselectivities with low yields has been reported: N. S. Chowdari, D. B. Ramachary, A. Cordova, C. F. Barbas III, *Tetrahedron Lett.* 2002, 43, 9591.
- [11] For examples of metal-mediated direct aldol reactions see:
 a) Y. M. A. Yamada, N. Yoshikawa, H. Sasai, M. Shibasaki, Angew. Chem. 1997, 109, 1290; Angew. Chem. Int. Ed. Engl. 1997, 36, 1871; b) N. Yoshikawa, N. Kumagai, S. Matsunaga, G. Moll, T. Oshima, T. Suzuki, M. Shibasaki, J. Am. Chem. Soc. 2001, 123, 2466; c) N. Kumagai, S. Matsunaga, N. Yoshikawa, T. Oshima, M. Shibasaki, Org. Lett. 2001, 3, 1539; d) B. M. Trost, H. Ito, J. Am. Chem. Soc. 2000, 122, 12003; e) B. M. Trost, E. R. Silcoff, H. Ito, Org. Lett. 2001, 3, 2497; f) D. A. Evans, J. S. Tedrow, J. T. Shaw, C. W. Downey, J. Am. Chem. Soc. 2002, 124, 392; g) G. Lalic, A. Aloise, M. Shair, J. Am. Chem. Soc. 2003, 125, 2852.