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De Novo Asymmetric Bio- and Chemocatalytic Synthesis of Saccharides — Stereoselective Formal *O*-Glycoside Bond Formation Using Palladium Catalysis

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The chemical synthesis of carbohydrate domains in saccharides and glycoconjugates such as antibiotics, antitumor agents, glycoproteins, and glycolipids is now recognized as a major frontier for organic chemistry. Fundamental to the synthesis of such carbohydrates and their derivatives is the selectivity of α - or β -O-glycoside bond formation which typically entails the coupling of one nucleophilic (O-donating) glycoside to another electrophilic glycosyl donor; anomeric stereoselectivity is a complex issue usually dependent on the nature of the donor C2 substituent.

Catalytic stereoselective formation of the acetal linkage onto pyranones⁴ of type 1 (Scheme 1) presents a conceptually different solution to this stereochemical problem by providing a stereodefined platform whose chiral information can be relayed around the ring. Such an acetal can be a formal α - or β -glycoside bond depending on the enantiomer of 1, the stereocontrol in the Pd-catalyzed step, and the chemistry used to elaborate the ring.⁵

Scheme 1. Iterative Saccharide Synthesis: Stereoselective Acetal Bond Formation Using Pd Catalysis

Here, we present a novel integrated approach to the de novo catalytic asymmetric synthesis of saccharides uniting two protocols: the enzymatic resolution of racemic acetoxypyranones 16 with a highly stereoselective palladium-catalyzed acetal bond formation onto this embryonic sugar (Scheme 1). Resulting from subsequent steps to elaborate the ring into a diversity of natural and unnatural sugars, a free hydroxyl group can be stereoselectively coupled again to 1, giving rise to an iterative catalytic asymmetric saccharide synthesis. A blank slate for saccharide synthesis, the versatility of this cyclic enone platform has been appreciated for some time.⁷

Despite the widespread use of phenols as nucleophiles in the palladium-catalyzed allylic substitution reaction, ⁸ aliphatic alcohols have received scant attention. ^{9,10} During early investigations, however, we found that the substitution reaction of enantiomerically pure 6-acetoxy-2*H*-pyran-3(6*H*)-one (–)-1⁶ with simple primary and secondary aliphatic alcohols as solvent proceeded with nearly complete retention of stereochemistry. ¹¹

Efforts to improve the viability of this methodology resulted in the coupling depicted in Table 1. The use of 10 mol % $Pd(OAc)_2$ and triphenyl phosphite in DCM at -30 °C¹² was found to convert pyranone (-)-1 into the benzyl alcohol adduct 2A in high yield

Table 1. Stereoselective Acetal Bond Formation Using Pd Catalysis

adduct	donor	% yield	% ee/de	adduct	donor	% yield	% d e ^e
2A	(-)-1	83	94^{b}	2H	(-)-1	65 ^a	91
2B	(-)-1	87	$98^{b,c}$		(-)-3	57^{a}	92
2C	(-)-1	98	99^{b}		(+)-3	61^{a}	96
2D	(-)-1	84	98^b	2I	(-)-1	70^{a}	97
2E	(\pm) -1	69	$\mathrm{nd}^{b,d}$		(-)-3	71^{a}	82
2F	(-)-1	78^{a}	97^{b}		(+)-3	76^{a}	95
2G	(-)-1	77^a	94e	2J	(-)-1	60	f
	(-)-3	88^{a}	94^e				
	(+)-3	96^{a}	98^e				

a Isolated yield of unique stereoisomer.
 b 10% Pd(OAc)₂, P(OPh)₃, DCM,
 −30 °C; stereoselectivities were determined by chiral HPLC analysis.
 c Enantiomeric excess before chromatography.
 d Coupled to racemic 1 only.
 e 5% Pd₂(dba)₃, PPh₃, DCM, −10 °C; diastereoselectivities were determined from ¹H NMR.
 f Mixture of isomers.

and 94% ee. Particularly rewarding were the still higher yields and ee's for anisyl nucleophiles **B** and **C** and the *ortho*-nitrobenzyl alcohol **D**, useful mimics of benzyl linkers^{1b,13} applied to the solid-phase synthesis of saccharides.¹⁴ In preliminary experiments to apply the protocol to the solid-phase, photocleavable **E**, immobilized onto phenolic polystyrene, was also coupled efficiently to racemic **1**. Representative of Mucin-type glycosylation found in the glycopeptides of mammals and other eukaryotes,¹⁵ adduct **2F** was also prepared with excellent stereoselectivity.

Key to the feasibility of the protocol is the success of a first iteration: a stereoselective coupling reaction of enantiopure glycosyl donor with a sugar derivative. The results are illustrated in Table 1. Initial attempts using the $Pd(OAc)_2/P(OPh)_3$ catalyst system failed, but, to our relief, use of $Pd_2(dba)_3/PPh_3$ successfully mediated formation of the desired adducts **2G**–**2J**. Primary alcohol **G**, a 6-deprotected glucopyranose, underwent coupling with (–)-**1** and both (R)-(–)-**3** and (S)-(+)-**3**¹⁶ to afford the stereoisomers of the

products with excellent yield (77–96%) and diastereoselectivity (94–98%). Crucially, similar success was found with the more sterically demanding substrates 4-deprotected glucopyranose **H** and 3-deprotected glucofuranose **I** bearing a secondary alcohol moiety, and good yields (57–76%) and excellent stereoselectivities (82–97%) were obtained during *both R- and S-acetal bond formation*. All adducts were isolated as unique diastereomers by simple column chromatography with the exception of that with **J**, deprotected at the anomeric center.

A preliminary application of our iterative approach is depicted in Scheme 2. Diastereoselective catalytic *cis*-dihydroxylation of enone adduct **2C** was effected by RuCl₃/NaIO₄, ¹⁷ and the resulting diol was protected to the dioxolane **4C** under standard conditions. Subsequent reduction using Zn(BH₄)₂¹⁸ gave **5C**, a β -L-ribose. ¹⁹ Coupling of this sugar under the catalytic conditions previously described successfully afforded the disaccharide precursor **6C** with 96% de. ²⁰

Scheme 2. Preliminary Application of Iterative Saccharide Synthesis^a

^a (i) RuCl₃·3H₂O (20 mol %), NalO₄; (ii) 2,2-DMP, acetone, PTSA; (iii) Zn(BH₄)₂; (iv) (-)-1, Pd₂(dba)₃ (5 mol %), PPh₃, CH₂Cl₂.

Unsuccessful endeavors to alkylate the methylene position of **4C** led to an appraisal of prefunctionalized pyranone substrate **7** in the palladium-catalyzed allylic substitution reaction (Scheme 3). Prepared enantiopure employing a Sharpless dihydroxylation protocol, ^{7j} **7** indeed underwent substitution with complete retention of stereochemistry, giving **8**. 4,4-Dimethyl-substituted pyranone **9**,^{7i,16} applicable to the asymmetric synthesis of L-noviose,²¹ a constituent of the antibiotic novobiocin, also participated with high stereoselectivity to afford **10**.

Scheme 3 . C4-Substituted Glycosyl Donors

Efforts to elaborate on this chemistry by providing a view of an iterative catalytic solid-phase protocol are ongoing.

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Supporting Information Available: Experimental procedures and spectral data for all new compounds (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

References

- (a) Collins, P.; Ferrier, R. Monosaccharides. Their Chemistry and Their Roles in Natural Products, Wiley: U.K., 1995.
 (b) Nicolaou, K. C.; Mitchell, H. J. Angew. Chem., Int. Ed. 2001, 40, 1576.
- (2) (a) Toshima, K.; Tatsuta, K. Chem. Rev. 1993, 93, 1503. (b) Wulff, G.; Röhle, G. Angew. Chem., Int. Ed. Engl. 1974, 13, 157.
- (3) Danishefsky, S. J.; Bilodeau, M. T. Angew. Chem., Int. Ed. Engl. 1996, 35, 1380
- (4) (a) Grynkiewicz, G.; Barszczak, B.; Zamojski, A. Synthesis 1979, 364.
 (b) Mucha, B.; Hoffmann, H. M. R. Tetrahedron Lett. 1989, 30, 4489.
 (c) Grynkiewicz, G. Carbohydr. Res. 1980, 80, 53. (d) Grynkiewicz, G.; Zamojski, A. Z. Naturforsch. 1980, 35b, 1024. (e) Knol, J.; Jansen, J. F. G. A.; Van Bolhuis, F.; Feringa, B. L. Tetrahedron Lett. 1991, 32, 7465.
- (5) The authors recognize that to adopt α- and β-nomenclature for 2 is not strictly appropriate until a second chiral center is introduced to the ring.
- (6) Van den Heuvel, M.; Cuiper, A. D.; Van der Deen, H.; Kellogg, R. M.; Feringa, B. L. Tetrahedron Lett. 1997, 38, 1655.
- (7) For a review, see: Holder, N. L. Chem. Rev. 1982, 82, 287. For selected examples, see: (a) Plaumann, D. E.; Fitzsimmons, B. J.; Ritchie, B. M.; Fraser-Reid, B. J. Org. Chem. 1982, 47, 941. (b) Achmatowicz, O., Jr.; Burzyńska, M. H. Tetrahedron 1982, 38, 3507. (c) Panfil, I.; Chmielewski, M. Tetrahedron 1985, 41, 4713. (d) Georgiadis, M. P.; Couladouros, E. A. J. Heterocycl. Chem. 1991, 28, 1325. (e) López Tudanca, P. L.; Jones, K.; Brownbridge, P. J. Chem. Soc., Perkin Trans. 1 1992, 533. (f) Taniguchi, T.; Nakamura, K.; Ogasawara, K. Synlett 1996, 971. (g) Taniguchi, T.; Ohnishi, H.; Ogasawara, K. Chem. Commun. 1996, 1477. (h) Caddick, S.; Khan, S.; Frost, L. M.; Smith, N. J.; Cheung, S.; Pairaudeau, G. Tetrahedron 2000, 56, 8953. (i) Hoffmann, H. M. R.; Krumwiede, D.; Mucha, B.; Oehlerking, H. H.; Prahst, G. W. Tetrahedron 1993, 49, 8999. (j) Harris, J. M.; Keränen, M. D.; Nguyen, H.; Young, V. G.; O'Doherty, G. A. Carbohydr. Res. 2000, 328, 17.
 (8) For a review on Pd-catalyzed allylic substitution, see: Trost, B. M.; Van
- (8) For a review on Pd-catalyzed allylic substitution, see: Trost, B. M.; Van Vranken, D. L. Chem. Rev. 1996, 96, 395. For examples using phenols, see: Trost, B. M.; Tsui, H.-C.; Toste, F. D. J. Am. Chem. Soc. 2000, 122, 3534 and references therein.
- (9) (a) Cuiper, A. D.; Kellogg, R. M.; Feringa, B. L. Chem. Commun. 1998, 655.
 (b) Trost, B. M.; Schroeder, G. M. J. Am. Chem. Soc. 2000, 122, 3785.
 (c) Kim, H.; Lee, C. Org. Lett. 2002, 4, 4369.
- (10) Glycosides have been allyl protected using a Pd-catalyzed allylic substitution reaction: Lakhmiri, R.; Lhoste, P.; Sinou, D. Tetrahedron Lett. 1989, 30, 4669.
- (11) Van der Deen, H.; Van Oeveren, A.; Kellogg, R. M.; Feringa, B. L. Tetrahedron Lett. 1999, 40, 1755.
- (12) Racemates of compounds 2A-2F were prepared using (±)-1 and 5 mol % Pd(OAc)₂ at ambient temperature. Higher catalyst loading and reduced temperature were found necessary for high stereoselectivities.
- (13) Roussel, F.; Takhi, M.; Schmidt, R. R. J. Org. Chem. 2001, 66, 8540.
- (14) (a) Solid Support Oligosaccharide Synthesis and Combinatorial Carbohydrate Libraries; Seeberger, P. H., Ed.; Wiley: New York, 2001. (b) Osborn, H. M. I.; Khan, T. H. Tetrahedron 1999, 55, 1807.
- (15) For a review on the synthesis of these biomolecules, see: Herzner, H.; Reipen, T.; Schultz, M.; Kunz, H. Chem. Rev. 2000, 100, 4495.
- (16) (\pm)-3 and (\pm)-9 were separated by preparative chiral HPLC.
- (17) Murphy, P. V.; O'Brien, J. L.; Smith, A. B., III. Carbohydr. Res. 2001, 334, 327.
- (18) Gensler, W. J.; Johnson, F. A.; Sloan, A. D. B. J. Am. Chem. Soc. 1960, 82, 6074.
- (19) Achmatowicz, O.; Grynkiewicz, G. *Carbohydr. Res.* **1977**, *54*, 193.
- (20) Preliminary steps toward a disaccharide from glucofuranose adduct 2I proceeded also with complete diastereoselectivity to give 4I, albeit so far with low yield.

(21) (a) Achmatowicz, O., Jr.; Grynkiewicz, G.; Szechner, B. Tetrahedron 1976, 32, 1051. (b) Ferroud, D.; Collard, J.; Klich, M.; Dupuis-Hamelin, C.; Mauvais, P.; Lassaigne, P.; Bonnefoy, A.; Musicki, B. Bioorg. Med. Chem. Lett. 1999, 9, 2881.

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