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Enantiocontrolled Synthesis of Trialkyl-Substituted Stereogenic Carbons. A General Route to *cis-*3,5-Dialkyl γ-Lactones

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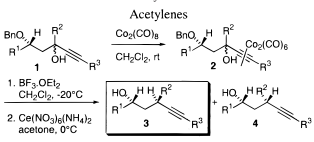
ABSTRACT

Lewis acid treatment of tertiary $\text{Co}_2(\text{CO})_6$ -propargylic alcohols having a stereochemically defined benzyloxy group at the γ -benzyl position yielded after cobalt demetalation sec-dialkyl bishomopropargylic alcohols in good yields. The reaction is highly stereoselective and predictable, providing pure stereoisomers. The use of benzyl- α , α' - d_2 ethers permitted the stereoselective d-labeling of methines and methylenes. Very simple chemical manipulations provided a general methodology to obtain the enantiomers of 3,5-dialkyl- γ -lactones.

Carbon—carbon σ -bond formation is usually the main objective in an organic synthesis. The coupling between an acetylide and an alkylating agent could be considered to be a standard procedure for the synthesis of dialkyl acetylenes. However, due to competitive elimination reactions, this procedure is inefficient for the preparation of sec-dialkyl acetylenes. One alternative procedure that overcomes this difficulty is the reduction of the cobalt complex of α -acetylenic alcohols with sodium borohydride in trifluoroacetic acid. Recently, we described a new procedure for the synthesis of bishomopropargylic alcohols based on the Lewis acid treatment of γ -benzyl-protected $Co_2(CO)_6$ - α , γ -acetylenic diols and further demetalation.

In this paper, we present the possibility of extending such an intramolecular reduction to the stereocontrolled synthesis of *sec*-dialkyl bishomopropargylic alcohols (Scheme 1).⁵ The γ -benzyl-protected α, γ -propargylic alcohols **1** were obtained in accordance with Scheme 2. Chiral 2,3-epoxy alcohols **5**

Scheme 1. Stereoselective Synthesis of *sec*-Dialkyl Acetylenes



⁽¹⁾ Comprehensive Organic Synthesis; Trost, B. M., et al., Eds.; Pergamon Press: Oxford, 1991; Vol. 3.

^{(2) (}a) Brandsma, L. In *Preparative Acetylenic Chemistry*; Elsevier: Amsterdam, 1988. (b) Garratt, P. J. In *Comprehensive Organic Synthesis*; Trost, B. M., et al., Eds.; Pergamon Press: Oxford, 1991; Vol. 3, pp 271–202

Scheme 2. Synthesis of γ -Benzyl Protected α , β -Propargylic Alcohols

Alcohols

Alcohols

Alcohols

OBN O

1. Red-Al®

2. PhCH(OMe)₂, H⁺ R¹

4. PCC

70%

$$\frac{R^{2}MgBr}{81 - 93\%}$$
R¹

OBN OH

R²

1. PCC

2. LiC=C-R³

7

76 - 84%

were regioselectively opened to the corresponding 1,3-diols. Benzylidene protection and regioselective reduction provided the secondary benzyl ether. Oxidation of the primary alcohol furnished the aldehyde 6, which was treated with a suitable Grignard reagent to yield the secondary alcohols 7. Oxidation to the corresponding ketone and lithium acetylide addition provided 1 as a mixture of diastereoisomers.⁶

The $Co_2(CO)_6$ -acetylenic complex **2** was obtained by simple treatment of **1** with $Co_2(CO)_8$ in a CH_2Cl_2 solution. The addition of 1 equiv of BF_3 • OEt_2 to a CH_2Cl_2 solution of the $Co_2(CO)_6$ -acetylenic complex **2**, at -20 °C, provided within minutes the corresponding complexed bishomopropargylic alcohol in a straightforward manner.⁷ In all cases, the $Co_2(CO)_6$ -bishomopropargylic alcohols were satisfactorily demetalized in the standard manner ($Ce(NO_3)_6(NH_4)_2$, acetone, 0 °C) to obtain the free acetylenes. Representative examples with different R^2 groups are outlined in Table 1.⁸

Table 1. Stereoselective Intramolecular Propargylic Reduction in γ -Benzyl-Protected $Co_2(CO)_6$ - α , γ -Acetylenic Diols under Lewis Acid Treatment

entry	2 (R ¹ = C ₁₃ H ₂₇ - n , R ³ = C ₅ H ₁₁ - n)	3:4	(yield, %) ^a
1	$2a, R^2 = CH_3$	100:0	89 (79)
2	2b , $R^2 = C_5 H_{11} - n$	100:0	86 (82)
3	$2c, R^2 = Ph$	100:0	87 (84)
4	2d , $R^2 = Pr-i$	30:1	81 (72)
5	2e , $R^2 = Bu-t$	1:1	79 (70)

 a Yields are not optimized (the overall yield from 1 to 3+4 is given in parentheses).

As can be seen from Table 1, the configuration of the stereogenic center in which the benzyl-protected group was located remains unaffected. However, the most interesting feature of our process was that the reduction of a tertiary propargylic alcohol, when the \mathbb{R}^2 is not very bulky, provided the corresponding sec-dialkyl acetylene as a sole diastere-

oisomer regardless the stereochemistry of the carbinol propargylic center (entries 1 and 2).⁹ Even when the R² substituent was a phenyl group, the reduction provided only one stereoisomer (entry 3). These products represent compounds in which the chirality of the γ-carbon has been completely transferred to the sec-position relative to the acetylene. When the R² increased the steric influence (isopropyl group), a tiny amount of 4d was isolated (entry 4). The reaction lacks stereoselectivity when R² is a tertbutyl group. These facts are consistent with our previously reported mechanism based on a hydride transfer of one benzylic proton to the propargylic carbocation (Scheme 3).⁴

Scheme 3. Proposed Mechanism for the Stereoselective Hydride Transfer to Co₂(CO)₆-Propargylic Cations

The chairlike transition state locates the bulkiest group in a pseudoequatorial position. The cobalt complex substituent is such a group when R² is not highly demanding from a steric viewpoint. However, when the size of R² was very large, as occurred, for example, with a *tert*-butyl group, competition between the group and the complexed acetylene led to poor selectivity.

To establish the stereochemistry of the newly created stereocenter, we envisioned the possibility of transforming 3 into a 3,5-disubstituted γ -lactone. Thus, 3 was selectively hydrogenated to the *cis*-olefin 8 that after acetylation and cleavage of the olefin afforded the corresponding carboxylic acid. Alkaline hydrolysis of the acetate group and further acid treatment afforded in excellent yield the corresponding γ -lactone (Table 2). The methodology was quite general, except for the case for R^2 = phenyl, in which we had to cleave the double bond via the formation of the corresponding *cis*-diol (OsO₄, NMO) and further oxidative fragmentation (KMnO₄, K₂CO₃, NaIO₄). The cis relationship between

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⁽³⁾ Nicholas, K. M.; Siegel, J. J. Am. Chem. Soc. 1985, 107, 4999.

⁽⁴⁾ Díaz, D. D.; Martín, V. S. Tetrahedron Lett. 1999, in press.

⁽⁵⁾ For other examples of "ionic hydrogenations" of alcohols, see: (a) Kursanov, D. N.; Parnes, Z. N.; Loim, N. M. Synthesis 1974, 633 and references therein. (b) Adlington, M. G.; Orfanopoulos, M.; Fry, J. L. Tetrahedron Lett. 1976, 2955. (c) Carey, F. A.; Tremper, H. S. J. Org. Chem. 1971, 36, 758. (d) Olah, G. A.; Arvanaghi, M.; Ohannecian, L. Synthesis 1987, 770. (e) Smonou, I.; Orfanopoulos, M. Tetrahedron Lett. 1988, 29, 5793 and references therein.

⁽⁶⁾ The propargylic alcohol is usually obtained as a mixture in which one of the diastereoisomers slightly predominates (ca. 1.5:1).

⁽⁷⁾ Blank experiments performing the acidic treatment over ${\bf 1}$ gave inseparable mixtures, and in any case, traces of ${\bf 3}$ were detected.

⁽⁸⁾ In our previous work, we have shown that the propargylic reduction is compatible with a broad kind of functional group.

⁽⁹⁾ We have performed the reaction with both diastereoisomers of **2b** leading to **3b** as the sole isolated product.

⁽¹⁰⁾ Prepared by the method shown in: Rodríguez, C. M.; Martín, T.; Ramírez, M. A.; Martín, V. S. J. Org. Chem. 1994, 59, 4461.

⁽¹¹⁾ Evans, D. A. In *Asymmetric Synthesis*; Morrison, J. D., Ed.; Academic Press: New York, 1985; Vol. 4, pp 2–110 and references therein.

⁽¹²⁾ The dibenzyl alcohol **18** was obtained in accordance with the method described in Scheme 2, with the exception of the step relative to the reduction of the benzylidene derivative. In this case, we had to use NaBH₃(CN)—(CH₃)₃SiCl, in CH₃CN, to obtain the secondary benzyl ether since the use of DIBAL provided the benzyl ether in the primary position. Johansson, R.; Samuelsson, B. *J. Chem. Soc.*, *Perkin Trans. I* **1992**, 2371.

Table 2. Stereoselective Synthesis of cis-3,5-Disubstituted γ-Lactones

entry	3 ($R^1 = C_{13}H_{27}-n$, $R^3 = C_5H_{11}-n$)	3 (yield, %)	9 (yield, %)
1	3a , $R^2 = CH_3$	88	72
2	3b , $R^2 = C_5 H_{11} - n$	85	71
3	$3c, R^2 = Ph$	87	64
4	$3d$, $R^2 = Pr-i$	85	63

the 3,5-substituents was clearly determined by NOE experiments over the γ -lactones 9 (Scheme 4). To ensure that such

Stereoselective Synthesis of 3,5-Disubstituted Scheme 4. γ-Lactones

studies were reliable, we prepared the alternative translactone by a different procedure. Thus, the 4-benzoyloxy- α,β -unsaturated ester 10^{10} was submitted to catalytic hydrogenation to yield the saturated diester 11 that was submitted to basic hydrolysis and further acidic treatment to afford the lactone 12. The alkylation under standard basic conditions led mainly to the expected trans-3,5-disubtituted γ -butyrolactone 13.11 The NMR studies were in accord with the stereochemistry.

77%

One exciting option of our method is the stereoselective d-labeling of methines and methylenes. The only preparation

Scheme 5. Stereoselective d-Labeling of Methines and Methylenes

of the suitable benzyl- α , α' - d_2 ethers 14 and further application of our methodology provided cleanly the corresponding deutero compounds 15 (Scheme 5).

From a practical viewpoint, the high regioselectivity of the reaction is another very important feature since only those benzyl ethers located at the γ -position are able to participate in the reduction of the propargylic alcohol. Thus, we prepared the dibenzyl alcohol 18 from S-malic acid in accordance with Scheme 6,12 and it was then submitted to the above-

Scheme 6. Only Benzyl Groups Located in the BisHomopropargylic Position Are Able to Transfer Hydrides

mentioned conditions, yielding only the reduced compound **19**.

In summary, the intramolecular hydride transfer from a secondary γ -benzyloxy group with defined absolute stereochemistry to a Co₂(CO)₆-complexed propargylic cation generated under Nicholas conditions occurs with excellent stereoselectivity, providing a new method to obtain secdialkyl acetylenes with absolute stereochemical control. Since the stereochemistry of the reduction is highly predictable, the judicious choice of the different substituents may provide access to stereochemically defined tricarbon-substituted methines in their enantiomeric forms. Application of this methodology to the synthesis of some natural products is underway and will be published in due course.

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Supporting Information Available: ¹H NMR and ¹³C NMR spectra for 3a-e, 4e, 9a-d, 15, and 19. NOE studies for 9a-d. This material is available free of charge via the Internet at http://pubs.acs.org.

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