

- [2] a) K. Tomooka, H. Yamamoto, T. Nakai, *J. Am. Chem. Soc.* **1996**, *118*, 3317–3318; b) K. Tomooka, M. Kikuchi, K. Igawa, P.-H. Keong, T. Nakai, *Tetrahedron Lett.* **1999**, *40*, 1917–1920.
- [3] Pioneering work on simple systems has been reported: T. B. Grindeley, C. Wickramage, *J. Carbohydr. Chem.* **1988**, *7*, 661–685.
- [4] For a recent review on C-glycosidation, see a) D. Levy, C. Tang, *The Chemistry of C-Glycosides*, Pergamon, Oxford **1995**; b) Y. G. Du, R. J. Linhardt, I. R. Valhov, *Tetrahedron* **1998**, *54*, 9913–9959.
- [5] a) B. M. Trost, E. D. Edstrom, *Angew. Chem.* **1990**, *102*, 541–542; *Angew. Chem. Int. Ed. Engl.* **1990**, *29*, 520–521; b) K. Tomooka, Y. Nakamura, T. Nakai, *Synlett* **1995**, 321–322.
- [6] Crystallographic data (excluding structure factors) for the structures β -(*R*)-**3** and β -(*S*)-**4** reported in this paper have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication nos. CCDC-144556 and -144557, respectively. Copies of the data can be obtained free of charge on application to CCDC, 12 Union Road, Cambridge CB21EZ, UK (fax: (+44)1223-336-033; e-mail: deposit@ccdc.cam.ac.uk).
- [7] Combined yield of C-glycosides **4**, X = SiMe₃ and X = H. These are interconvertible through silylation or desilylation.
- [8] For pioneering studies related to the [1,4] Wittig rearrangement, see a) H. Felkin, C. Frajerman, *Tetrahedron Lett.* **1977**, *39*, 3485–3488; b) M. Schlosser, S. Strunk, *Tetrahedron* **1989**, *45*, 2649–2664.
- [9] The stereochemical assignment was made on the basis of the ¹H NMR analysis and NOE experiment of the product and derivative thereof (see Supporting Information).
- [10] In this case the stereoselectivity at the C1' position is opposite that observed in the pantolactone-derived system. A similar stereochangeover has been observed in the rearrangement of O-glycosides derived from primary alcohols; see ref. [2a].
- [11] The trapping experiment of enolate **A** (SiMe₃Cl, –78 °C) gave an *E*:*Z* mixture (ca. 1:1) of silyl enol ether in 69% yield.

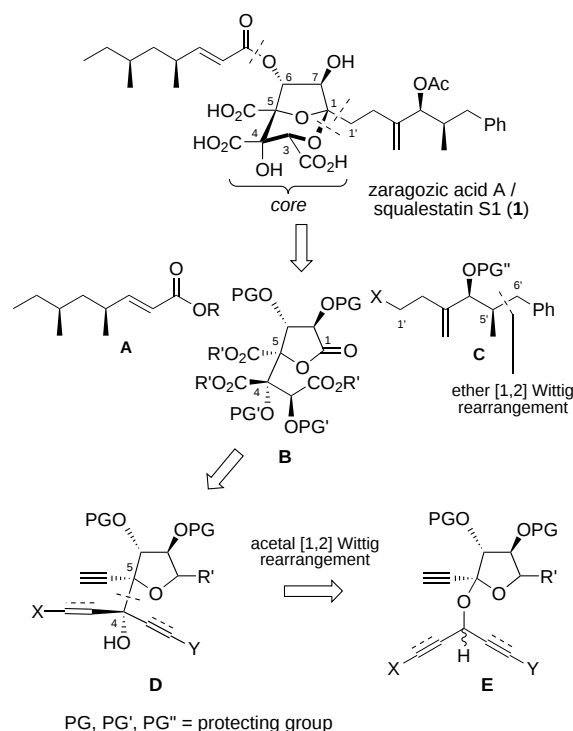
Stereoselective Total Synthesis of Zaragozic Acid A based on an Acetal [1,2] Wittig Rearrangement**

Katsuhiko Tomooka,* Makoto Kikuchi, Kazunobu Igawa, Masaki Suzuki, Ping-Huai Keong, and Takeshi Nakai

The zaragozic acids and squalostatins are a family of naturally occurring fungal metabolites independently isolated and characterized by researchers at Merck^[1] and Glaxo.^[2] These natural products are potent inhibitors of squalene synthase and have potential as therapeutic agents for the treatment of hypercholesterolemia.^[1a, 2a] All the zaragozic acids and squalostatins have a common 2,8-dioxabicy-

clo[3.2.1]octane core with an array of six stereogenic centers including contiguous quarternary carbon atoms, and are different only at the alkyl and acyl side chains at C1 and C6, respectively. Therefore, it is not surprising that these compounds have elicited considerable attention from numerous synthetic chemists.^[3, 4] Zaragozic acid A (squalestatin S1; **1**) is a representative example of this novel class of compounds. Herein, we report the efficient, convergent synthesis of **1** by highlighting the acetal [1,2] Wittig rearrangement^[5] for forming the C4–C5 bond together with the simultaneous creation of the contiguous quaternary carbon atoms as the key step.

Our strategy is shown in Scheme 1. Disconnection of the C6 side chain **A**, unraveling of the C1 acetal, and cleavage of the C1–C1' bond led to lactone **B** and an alkyl anion equivalent



Scheme 1. Retrosynthetic analysis of zaragozic acid A (**1**).

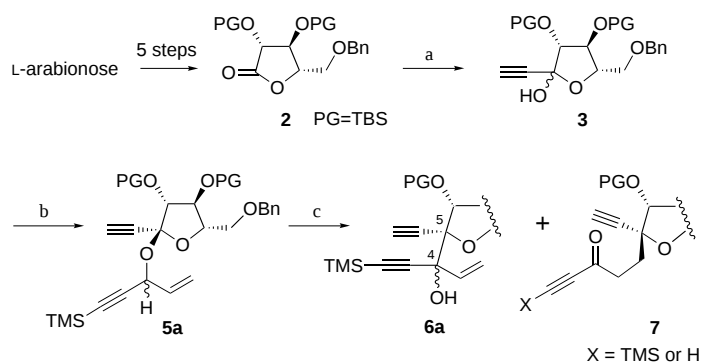
C.^[6] Our interest in the carbanion rearrangement of the sugar derivatives led us to consider a strategy for lactone **B** starting from the C-glycoside **D**, namely an acetal [1,2] Wittig rearrangement product of the O-glycoside **E**. The preceding paper describes the synthesis of this class of highly functionalized C-glycosides based on a rearrangement protocol.^[7] Furthermore, we planned to construct a C5'–C6' bond in segment **C** using the original ether [1,2] Wittig rearrangement.^[8]

Our synthesis begins from the O-glycoside **5a** ($\cong$ **E**), which can be prepared from the hemiketal **3** derived from L-arabino- γ -lactone^[9] and the protected ethynylvinylmethanol **4** (racemate; Scheme 2). Previous work suggested that the carbanion rearrangement of an O-glycoside such as **5a** should produce the [1,2] rearrangement product with high diastereoselectivity and in good yield.^[7, 9] In contrast to this expect-

[*] Prof. Dr. K. Tomooka, Dr. M. Kikuchi, K. Igawa, M. Suzuki, P.-H. Keong, Prof. Dr. T. Nakai
Department of Applied Chemistry
Graduate School of Science and Engineering
Tokyo Institute of Technology
Meguro-ku, Tokyo 152–8552 (Japan)
Fax: (+81)3-5734-3931
E-mail: ktomooka@o.cc.titech.ac.jp

[**] This work was supported by a Grant-in-Aid for Scientific Research from the Ministry of Education, Science, Sports, and Culture, Japan and the Sumitomo Foundation. We thank Merck & Co., Inc. for a generous gift of zaragozic acid A.

Supporting information for this article is available on the WWW under <http://www.wiley-vch.de/home/angewandte/> or from the author.

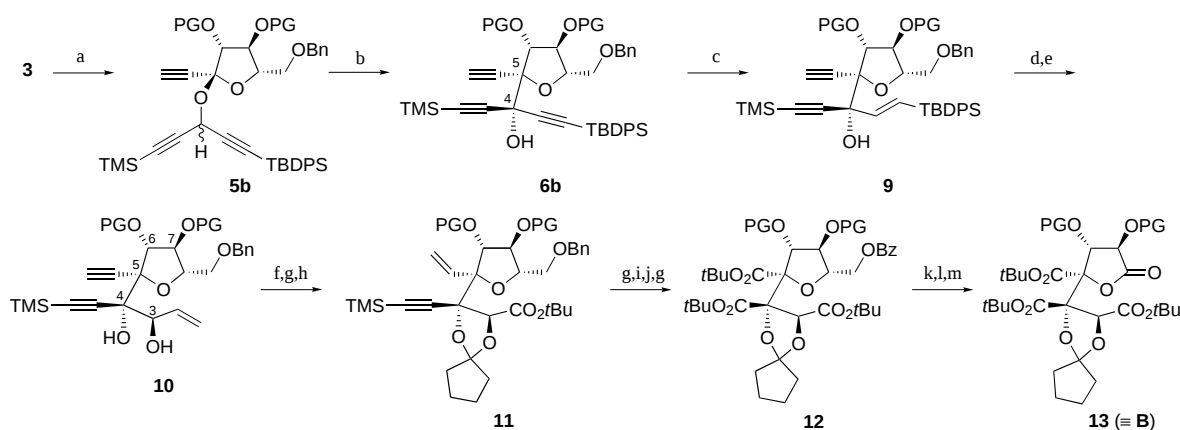


Scheme 2. Synthesis and rearrangement of *O*-glycoside **5a**. a) Lithium acetylide, Et₂O, −78 °C, 75%; b) CH₂=CHCH(OH)C≡C-TMS (**4**), montmorillonite K 10, 4 Å-MS, CH₂Cl₂, RT, 55%; c) *n*BuLi, THF, −78 °C, **6a**: 14%, **7**: 17%, recovered **5a**: 46%. TBS = *tert*-butyldimethylsilyl, Bn = benzyl, TMS = trimethylsilyl.

ation, the reaction of β -**5a** (*n*BuLi, −78 °C, THF) proceeded to form a 1:1 diastereomeric mixture of the desired [1,2] shift product β -**6a** in poor yield, along with a significant amount of the [1,4] rearrangement product **7**. Thus, we tried to redesign the rearrangement system.

Finally, we found that the bis(ethynyl)methanol system satisfied all the requirements as the migrating terminus. The reaction of the *O*-glycoside **5b** ($\cong$ **E**), similarly prepared from ketal **3** and the protected bis(ethynyl)methanol **8** (racemate), exclusively afforded the desired [1,2] Wittig product ($5\beta,4S$)-*C*-glycoside **6b** with high diastereoselectivity (>95% β , 84% d.r. at C4)^[10] in 54% yield. (Scheme 3). Also significant, the anomeric radical, an intermediate in the rearrangement of **5b**, could efficiently discriminate between the enantiotopic faces of the prochiral bis(ethynyl)methanol radical (that is, differentiation of trimethylsilyl and *tert*-butyldimethylsilyl at the γ,γ' positions) during the radical recombination process.

The major diastereomer, separated by column chromatography, was then converted into the key precursor **10**, which has all the stereogenic centers of **B**, by the following sequence.

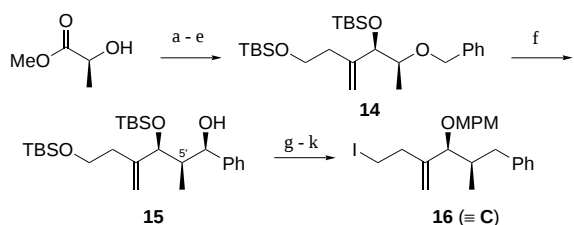


Scheme 3. Synthesis of the lactone **B**. a) TBDPS-C≡CCH(OH)C≡C-TMS (**8**), montmorillonite K 10, 4 Å-MS, CH₂Cl₂, RT, 80%; b) *n*BuLi, THF, −78 °C, 54% 84:16 at C4; c) Red-Al, Et₂O, −15 °C, 83%; d) O₃, MeOH, −78 °C then Me₂S; e) CH₂=CHMgBr, toluene, −78 °C, 49% for two steps; f) cyclopentanone dimethylacetal, *p*TsOH, benzene, RT, 87%; g) O₃, MeOH, −78 °C then Me₂S; NaClO₂, NaH₂PO₄, Me₂C=CHMe, *t*BuOH/H₂O, RT; *N,N'*-diisopropyl-*O*-*tert*-butylisourea, CH₂Cl₂, RT, 73–84% for three steps; h) H₂, cat. Pd/C, MeOH, RT, 88%; i) 0.1N TBAF/THF, THF, 0 °C, 94%; j) H₂, Lindlar cat., MeOH, RT, 93%; k) KOH (aq), MeOH, RT, 78%; l) cat. TPAP, NMO, 4 Å-MS, CH₂Cl₂, RT, 98%; m) O₂, Cu(OAc)₂, 2,2'-bipyridyl, DABCO, DMF, 70 °C, 72%. TBDPS = *tert*-butyldiphenylsilyl, Bz = benzoyl, TBAF = tetrabutylammonium fluoride, TPAP = tetrapropylammonium perruthenate, NMO = *N*-methylmorpholine *N*-oxide, DABCO = 1,4-diazabicyclo[2.2.2]octane.

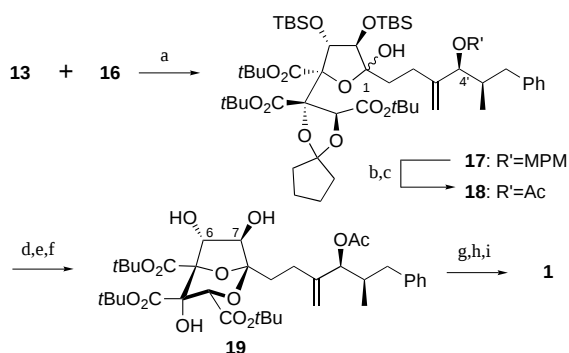
The semireduction of the ethynyl group on **6b** with bis(2-methoxyethoxy)aluminum hydride (Red-Al) in diethyl ether at −15 °C yielded the *tert*-butyldiphenylvinylsilane **9** with high selectivity.^[11] After the ozonolysis of **9**, the addition of vinylmagnesium bromide (toluene, −78 °C) proceeded to give **10** with excellent diastereoselectivity^[12] (>95% d.r.) with the latent C3 carboxy moiety introduced in the form of a vinyl substituent. After protection of the C3 and C4 hydroxyl groups as a cyclopentylidene acetal, the oxidation of the vinyl group at C3 was accomplished through a three-step sequence which involved ozonolysis to the corresponding aldehyde followed by oxidation with buffered NaClO₂ and esterification with *N,N'*-diisopropyl-*O*-*tert*-butylisourea^[13] to give the *tert*-butyl monoester. The tris(*tert*-butyl) ester **12** was obtained in good yield following a similar stepwise oxidation of the ethynyl groups at C4 and C5 to the carboxy moieties. The hydrolysis of the benzoate group at C1^[14] and the two-step oxidation (tetrapropylammonium perruthenate (TPAP)/O₂-Cu(OAc)₂^[15]) of the resulting hydroxymethyl group afforded the lactone **13** ($\cong$ **B**) as a fully elaborated intermediate, to which a nucleophilic side chain equivalent could be added at C1.

Scheme 4 illustrates the synthesis of the C1 side chain equivalent **16** ($\cong$ **C**) using the ether [1,2] Wittig rearrangement as the key step (**14** → **15**). The benzyl ether **15** was synthesized from methyl (*S*)-lactate in five steps. The rearrangement of **14** (*n*BuLi, THF, 0 °C) afforded **15**, which possesses the desired *R* configuration at the C5' position (88% d.r.),^[16] in 32% yield along with 58% of recovered **14**. The eight-step refunctionalization sequence from **15**, which involved removal of the hydroxyl group at C6'^[17] and iodination of the primary alcohol, successfully afforded **16**.

Generation of the alkyl lithium side chain (for C1) derived from iodide **16** followed by the addition of a lactone **13** provided the hemiketal **17** as a mixture of anomers (Scheme 5).^[18] The oxidative cleavage of the MPM ether followed by acetylation of the hydroxyl group at C4' completed the assembly of the hemiketal **18**, the precursor



Scheme 4. Synthesis of the C1 side chain ($\cong$ C). a) $\text{PhCH}_2\text{OC}(\text{=NH})\text{CCl}_3$, cat. $\text{CF}_3\text{SO}_3\text{H}$, cyclohexane, $0^\circ\text{C} \rightarrow \text{RT}$, 90%; b) $i\text{PrMgBr}$, $\text{MeO}(\text{Me})\text{NH} \cdot \text{HCl}$, THF, -10°C , 60%; c) $t\text{BuLi}$, $\text{TBSOCH}_2\text{CH}_2\text{C}(\text{=CH}_2)\text{Br}$, hexane/ Et_2O , -78°C , 92%; d) $\text{Zn}(\text{BH}_4)_2$, Et_2O , $-78 \rightarrow -30^\circ\text{C}$, 93%; e) TBSOTf , 2,6-lutidine, CH_2Cl_2 , $0^\circ\text{C} \rightarrow \text{RT}$, 95%; f) $n\text{BuLi}$, THF, $-20 \rightarrow 0^\circ\text{C}$, **15**: 32% recovered, **14**: 58%; g) TBAF, THF, RT; acetone dimethylacetal, cat. $p\text{TsOH}$, benzene, RT; 1N HCl (aq), THF, 0°C ; TBSCl , imidazole, CH_2Cl_2 , RT, 62% for four steps; h) Li_0 , liq. NH_3 , $-78 \rightarrow -30^\circ\text{C}$, 85%; i) $\text{MPMOC}(\text{=NH})\text{CCl}_3$, cat. $\text{CF}_3\text{SO}_3\text{H}$, cyclohexane, CH_2Cl_2 , 0°C ; j) TBAF, THF, RT, 81% for two steps; k) I_2 , imidazole, PPh_3 , benzene, RT, 88%. MPM = *p*-methoxyphenylmethyl.



Scheme 5. Total synthesis of zaragozic acid A (**1**). a) 3.2 equiv of **16**, 5.0 equiv of $t\text{BuLi}$, 5:1 hexane:diethyl ether, -40°C , then **13**, -40°C , 96%; b) DDQ, CH_2Cl_2 , RT, 96%; c) Ac_2O , DMAP, CH_2Cl_2 , RT, 95%; d) 10.20:1 TFA: CH_2Cl_2 : H_2O , RT; e) N,N' -diisopropyl-*O*-*tert*-butylisourea, CH_2Cl_2 , RT; f) 1N TBAF, THF, RT, 51% for three steps; g) $(t\text{BuOCO})_2\text{O}$, 4-pyrrrolidinopyridine, CH_2Cl_2 , 0°C , 81%; h) **20** ($\cong$ A, R = H), DCC, DMAP, CH_2Cl_2 , RT, 82%; i) TFA, CH_2Cl_2 , RT, 73%. DDQ = 2,3-dichloro-5,6-dicyano-1,4-benzoquinone, DMAP = 4-dimethylaminopyridine, TFA = trifluoroacetic acid, DCC = 1,3-dicyclohexylcarbodiimide.

to the bicyclic core and associated C1 side chain. Hydrolysis/ketalization of the hemiketal **18** with trifluoroacetic acid (TFA), followed by the esterification and TBAF treatment provided the desired bicyclic ketal **19**.^[19] Selective protection of the hydroxyl group at C7 with a *tert*-butoxycarbonyl (Boc) group^[20] followed by acylation with the C6 side chain **20** ($\cong$ A, R = H),^[21] then acid treatment afforded zaragozic acid A (**1**), whose spectroscopic data were in excellent agreement to those of a natural sample (^1H and ^{13}C NMR, IR, HPLC, and $[\alpha]_D$).

In conclusion, we have described the total synthesis of zaragozic acid A, which is highlighted by developing the acetal [1,2] Wittig rearrangement for the construction of the C4–C5 contiguous quaternary chiral centers. This synthetic strategy could be applied to the synthesis of a variety of naturally occurring zaragozic acids as well as designed synthetic analogues.

Received: July 24, 2000 [Z15514]

- [1] a) K. E. Wilson, R. M. Burk, T. Biftu, R. G. Ball, K. Hoogsteen, *J. Org. Chem.* **1992**, 57, 7151–7158; b) C. Dufresne, K. E. Wilson, D. Zink, J. Smith, J. D. Bergstrom, M. Kurtz, D. Rew, M. Nallin, R. Jenkins, K. Bartizal, C. Trainor, G. Bills, M. Meinz, L. Y. Huang, J. Onishi, J. Milligan, M. Mojena, F. Pelaez, *Tetrahedron* **1992**, 48, 10221–10226; c) J. D. Bergstrom, M. M. Kurtz, D. J. Rew, A. M. Amend, J. D. Karkas, R. G. Bostedor, V. S. Bansal, C. Dufresne, F. L. Vanmiddlesworth, O. D. Hensens, J. M. Liesch, D. L. Zink, K. E. Wilson, J. Onishi, J. A. Milligan, G. Bills, L. Kaplan, M. N. Omstead, R. G. Jenkins, L. Huang, M. S. Meinz, L. Quinn, R. W. Burg, Y. L. Kong, S. Mochales, M. Mojena, I. Martin, F. Pelaez, M. T. Diez, A. W. Alberts, *Proc. Natl. Acad. Sci. U.S.A.* **1993**, 90, 80–84.
- [2] a) M. J. Dawson, J. E. Farthing, P. S. Marshall, R. F. Middleton, M. J. O'Neill, A. Shuttleworth, C. Stylli, R. M. Tait, P. M. Taylor, H. G. Wildman, A. D. Buss, D. Langley, M. V. Hayes, *J. Antibiot.* **1992**, 45, 639–647; b) P. J. Sidebottom, R. M. Highcock, S. J. Lane, P. A. Procopiou, N. S. Watson, *J. Antibiot.* **1992**, 45, 648–658; c) A. Baxter, B. J. Fitzgerald, J. L. Hutson, A. D. McCarthy, J. M. Motteram, B. C. Ross, M. Sapra, M. A. Snowden, N. S. Watson, R. J. Williams, C. Wright, *J. Biol. Chem.* **1992**, 267, 11 705–11 708.
- [3] Total synthesis of zaragozic acid A: a) K. C. Nicolaou, E. W. Yue, Y. Naniwa, F. Dericcardis, A. Nadin, J. E. Leresche, S. Lagreca, Z. Yang, *Angew. Chem.* **1994**, 106, 2306–2309; *Angew. Chem. Int. Ed. Engl.* **1994**, 33, 2184–2187; K. C. Nicolaou, A. Nadin, J. E. Leresche, S. Lagreca, T. Tsuru, E. W. Yue, Z. Yang, *Angew. Chem.* **1994**, 106, 2309–2312; *Angew. Chem. Int. Ed. Engl.* **1994**, 33, 2187–2190; K. C. Nicolaou, A. Nadin, J. E. Leresche, E. W. Yue, S. Lagreca, *Angew. Chem.* **1994**, 106, 2312–2313; *Angew. Chem. Int. Ed. Engl.* **1994**, 33, 2190–2191; K. C. Nicolaou, E. D. Yue, S. L. Greca, A. Nadin, Z. Yang, J. E. Leresche, T. Tsuru, Y. Naniwa, F. D. Riccardis, *Chem. Eur. J.* **1995**, 1, 467–494; b) D. Stoermer, S. Caron, C. H. Heathcock, *J. Org. Chem.* **1996**, 61, 9115–9125; S. Caron, D. Stoermer, A. K. Mapp, C. H. Heathcock, *J. Org. Chem.* **1996**, 61, 9126–9134; total synthesis of zaragozic acid C: c) E. M. Carreira, J. Dubois, *J. Am. Chem. Soc.* **1994**, 116, 10825–10826; E. M. Carreira, J. Dubois, *J. Am. Chem. Soc.* **1995**, 117, 8106–8125; d) D. A. Evans, J. C. Barrow, J. L. Leighton, A. J. Robichaud, M. Sefkow, *J. Am. Chem. Soc.* **1994**, 116, 12111–12112; e) H. Sato, S. Nakamura, N. Watanabe, S. Hashimoto, *Synlett* **1997**, 451–454; f) A. Armstrong, L. H. Jones, P. A. Barsanti, *Tetrahedron Lett.* **1998**, 39, 3337–3340.
- [4] Reviews: U. Koert, *Angew. Chem.* **1995**, 107, 849–854; *Angew. Chem. Int. Ed. Engl.* **1995**, 34, 773–778; A. Nadin, K. C. Nicolaou, *Angew. Chem.* **1996**, 108, 1732–1766; *Angew. Chem. Int. Ed. Engl.* **1996**, 35, 1623–1656; K. Tomooka, M. Kikuchi, *Biosci. Ind.* **1999**, 57, 27–30.
- [5] K. Tomooka, H. Yamamoto, T. Nakai, *J. Am. Chem. Soc.* **1996**, 118, 3317–3318.
- [6] During the course of our work, this C1–C1' bond formation approach was successfully utilized in the total synthesis of zaragozic acid C by Evans et al. See ref. [3d].
- [7] K. Tomooka, H. Yamamoto, T. Nakai, *Angew. Chem.* **2000**, 112, 4674–4676; *Angew. Chem. Int. Ed.* **2000**, 39, 4500–4502.
- [8] Reviews: a) U. Schöllkopf, *Angew. Chem.* **1970**, 82, 795–805; *Angew. Chem. Int. Ed. Engl.* **1970**, 9, 763–773; b) R. W. Hoffmann, *Angew. Chem.* **1979**, 91, 625–634; *Angew. Chem. Int. Ed. Engl.* **1979**, 18, 563–572; c) J. A. Marshall, in *Comprehensive Organic Synthesis*, Vol. 3 (Eds.: B. M. Trost, I. Fleming), Pergamon, Oxford, **1991**, pp. 975–1014; d) K. Tomooka, T. Nakai, *J. Synth. Org. Chem. Jpn.* **1996**, 54, 1000–1008; e) K. Tomooka, H. Yamamoto, T. Nakai, *Liebigs Ann.* **1997**, 1275–1281.
- [9] To obtain the retention product (β -C-glycoside) using the acetal [1,2] Wittig rearrangement, a α -configuration at C1 in the *O*-glycoside is essential: K. Tomooka, M. Kikuchi, K. Igawa, P.-H. Keong, T. Nakai, *Tetrahedron Lett.* **1999**, 40, 1917–1920.
- [10] The stereochemistry of the major isomer was assigned by X-ray crystallography of a derivative of **6b**. The diastereomeric ratio at C4 was determined by ^1H NMR analysis.
- [11] It is worth noting that a similar reaction of simple bis(ethynyl)methanol also preferentially proceeds at the TBDPS side over the TMS side. The investigation into the scope and limitation of the present chemoselective hydroalumination as well as its mechanistic study is now in progress.

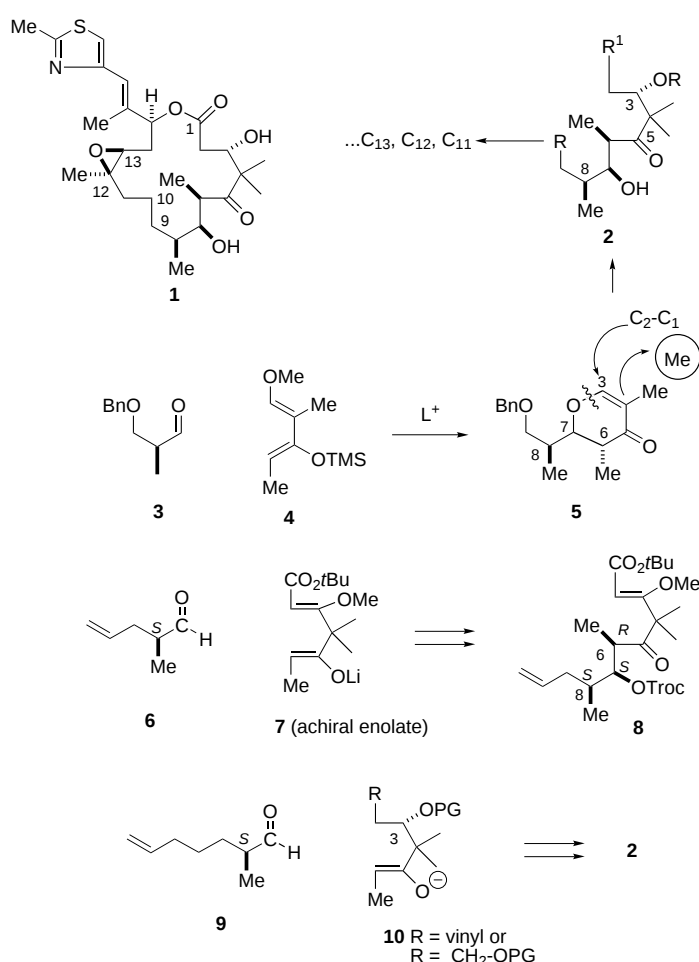
- [12] The stereochemical outcome of this addition reaction is consistent with chelation control through the hydroxyl group at C4.
- [13] L. J. Mathias, *Synthesis* **1979**, 11, 561–576.
- [14] The benzyl ether at C1' was oxidized at the ozonolysis step used for installing the carboxy moiety at C5.
- [15] V. V. Rhee, *Tetrahedron Lett.* **1969**, 985–988.
- [16] The diastereomeric ratio and stereochemistry at C5' were determined by ^1H NMR analysis. See the Supporting Information.
- [17] A. J. Robichaud, G. D. Berger, D. A. Evans, *Tetrahedron Lett.* **1993**, 34, 8403–8406.
- [18] Reaction temperature and the solvent are critical in this step. In fact, the alkyllithium reagent is not reactive to the lactone at -78°C and unstable above -10°C . We have used slightly modified Evans conditions. See ref. [3d].
- [19] The spectral data for **19** was identical in all respects to the authentic sample which was prepared from natural zaragozic acid A.
- [20] This selective protection was reported by Carreira et al.; see ref. [3c].
- [21] Synthesis of the side chain at C6: K. Tomooka, A. Nagasawa, S.-Y. Wei, T. Nakai, *Tetrahedron Lett.* **1996**, 37, 8895–8898.

Subtle Variations in the Long-Range Transmission of Stereochemical Information: Matched and Mismatched Aldol Reactions**

Zhicai Wu, Fei Zhang, and Samuel J. Danishefsky*

In this paper we describe findings with respect to an important question in stereospecific synthesis, that is, how might remote stereogenic loci influence the diastereofacial sense in which covalent bond formation occurs. To address this type of issue in concrete rather than abstract terms, we focused on the very promising antitumor agents, the epothilones (such as epothilone B (**1**), shown in Scheme 1). These compounds have stimulated a great deal of research from the point of view of total synthesis.^[1–6] In this paper we do not directly address the total synthesis problem per se. Rather, we focus on the construction of the C6–C7 bond in the acyl sector en route to the natural products, as a paradigm for accomplishing high orders of selection by long-range induction.

By way of background we note that, originally,^[1a, 1c] the stereochemical information for building the “acyl” sector of the epothilones (see **2** in Scheme 1) was stored in dihydropyrone **5** which itself arose from a cyclocondensation reaction of enantiomerically pure **3** and diene **4**. In our second-generation synthesis,^[7] the enantiomerically homogenous **6** reacted with



Scheme 1. Structure of epothilone B (**1**) and several methods for the construction of the C6–C7 bond in the acyl sector (**2**) of the epothilones. Bn = benzyl, TMS = trimethylsilyl, Troc = trichloroethoxycarbonyl, PG = protecting group.

achiral enolate **7**, giving rise selectively (approximate ratio = 5.5:1) to **8**, with the required *R*, *S*, and *S* configurations at C6, C7, and C8, respectively. The configuration at C3 was subsequently induced by catalytic, reagent-based, asymmetric reduction. System **2** can now be prepared in a highly convergent fashion on a multigram scale by this synthesis.^[6]

The background result which was central to the program described below was reported by Schinzer and co-workers,^[3a] and was subsequently used by others.^[4e, 8] The essence of Schinzer's claim^[9] was that aldol condensation of the enantiomerically defined lithio enolate system (see **10** in Scheme 1) with the enantiomerically defined aldehyde **9** occurs with high stereoselection in the required sense at the emerging C6–C7 bond, leading, eventually, to a product of type **2**. In essence, in the approach of Schinzer et al., the chirality at C3 had induced the required C6(*R*) and C7(*S*) configurations. This is in contrast to our second generation approach where the stereogenicity was induced solely by the *S* aldehyde **6**.

We undertook to revisit this type of aldol condensation for several reasons. The role of the pre-C8 *S* methyl group in **9** in delivering the high diastereofacial selectivity at C6 and C7

[*] Prof. S. J. Danishefsky, Dr. Z. Wu, F. Zhang
Department of Chemistry
Columbia University
Havemeyer Hall, 3000 Broadway, New York, NY 10027 (USA)
E-mail: s-danishefsky@ski.mskcc.org
Prof. S. J. Danishefsky
Laboratory for Bioorganic Chemistry
Sloan–Kettering Institute for Cancer Research
1275 York Avenue, New York, NY 10021 (USA)
Fax: (+1) 212-772-8691

[**] This work was supported by the National Institutes of Health (Grant Nos. CA-28824 and HL-25848).