

**Tetramic Acid Antibiotics: Stereoselective Synthesis of Streptolic Acid and Tirandalydigin\*\***

Yasuhiro Iwata, Naomi Maekawara, Keiji Tanino, and Masaaki Miyashita\*

The tetramic acid family of antibiotics have unique chemical structures composed of the 2,6-dioxabicyclononane skeleton and the characteristic dienoyl tetramic acid moiety. They exhibit potent antimicrobial activities and inhibitory activity against bacterial DNA-directed RNA polymerase.<sup>[1]</sup> The distinctive structural features and potent pharmacological properties render this family of antibiotics worthy targets for synthetic exploration.<sup>[2]</sup> In the tetramic acid antibiotics, two types of 2,6-dioxabicyclononane structures are known. One is the oxabicyclononane structure with an epoxy ketone moiety, as represented by tirandamycin A and B,<sup>[3]</sup> and the other is that involving a vinyl epoxide moiety, such as that in streptolydigin (**1**)<sup>[4]</sup> and tirandalydigin (**2**).<sup>[5]</sup> Tirandamycin A and B, with the chemically stable 2,6-dioxabicyclononane structure, have been extensively studied and their total syntheses have already established by several groups,<sup>[2]</sup> whereas synthetic studies of streptolydigin (**1**) and tirandalydigin (**2**), both of which bear the chemically labile vinyl epoxide moiety, are quite few. Indeed, the only synthesis of streptolic acid (**3**), the degradation product from **1** and **2** and the most potent member of the small family of 3-acyltetramic acid antibiotics, has been reported by Ireland and Smith.<sup>[2f]</sup>

We report herein a new synthetic methodology for streptolydigin (**1**) and tirandalydigin (**2**) that culminates in the first synthesis of the latter antibiotic, as well as a highly stereoselective synthesis of streptolic acid (**3**). Synthetic challenges posed by **1** and **2** include construction of the stereochemically dense 2,6-dioxabicyclononane skeleton, including the extremely acid-labile vinyl epoxide moiety, and synthesis of the distinctive tetramic acid structures. In particular, stereoselective synthesis of the common 2,6-dioxabicyclononane system and construction of the vinyl epoxide moiety are key challenges in the synthesis, since the generally used acid-catalyzed intramolecular acetalization of keto diol precursors has been known not to be effective in the synthesis of the tetramic acid antibiotics.<sup>[2d,h]</sup>

[\*] Y. Iwata, N. Maekawara, Dr. K. Tanino, Prof. Dr. M. Miyashita  
Division of Chemistry, Graduate School of Science  
Hokkaido University, 060-0810 Sapporo (Japan)  
Fax: (+81) 11-706-4920  
E-mail: miyashita@sci.hokudai.ac.jp

[\*\*] Financial support from the Ministry of Education, Culture, Sports, Science, and Technology, Japan, (a Grant-in-Aid for Scientific Research (A) (No. 12304042), a Grant-in-Aid for Scientific Research (B) (No. 16350049), and a Grant-in-Aid for Scientific Research on Priority Areas (A) "Exploitation of Multi-Element Cyclic Molecules" (No. 13029003)) is gratefully acknowledged.



Supporting Information for this article is available on the WWW under <http://www.angewandte.org> or from the author.

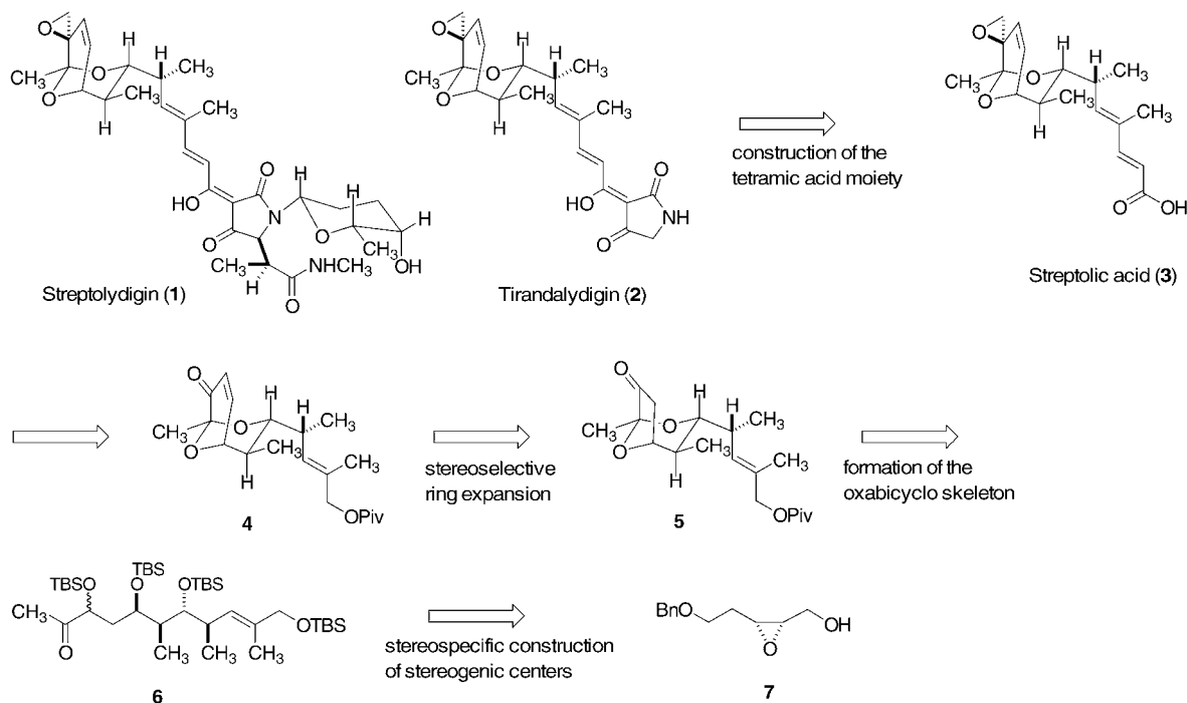
To overcome these difficulties, we designed the synthetic strategy as shown in Scheme 1. This strategy involves a key 2,6-dioxabicyclononen-7-one intermediate **4** that would be derived from a synthetically more accessible dioxabicyclooctanone **5** by a stereoselective ring expansion. We anticipated that the critical intermediate **5** could be efficiently constructed by the acid-catalyzed intramolecular acetalization of the precursor **6**.

Our first objective focused on the stereoselective synthesis of the key precursor **5** for construction of the 2,6-dioxabicyclononane skeleton. At first, the requisite acyclic compound **6** with four contiguous stereogenic centers was synthesized from the known chiral compound **7** in a highly stereoselective manner according to the method shown in Scheme 2. Thus, the epoxy alcohol **7** was converted into epoxy unsaturated ester **8** in 81% yield by a Swern oxidation followed by a Horner–Wadsworth–Emmons reaction. The crucial methylation reaction of **8** occurred stereospecifically with a  $\text{Me}_3\text{Al}$ /water system developed in our laboratory<sup>[6]</sup> to give rise to a single product, **9**, in 96% yield. Protection of the hydroxy group in **9** with TESCl and subsequent reduction of the ester with DIBAL-H in THF furnished allyl alcohol **10** in high yield. When **10** was treated with *m*CPBA in  $\text{CH}_2\text{Cl}_2$ , the single  $\alpha$ -epoxy alcohol **11** was obtained as expected in 88% yield.<sup>[7]</sup> The epoxy alcohol **11** was then transformed into epoxy unsaturated ester **12** by a three-step reaction sequence involving oxidation with PDC in  $\text{ClCH}_2\text{CH}_2\text{Cl}$  to form the corresponding aldehyde, followed by a Wittig reaction in a one-pot operation, and then removal of the TES group with TBAF in THF (79% yield over three steps). The next key methylation reaction of **12** also proceeded stereospecifically upon treatment with a  $\text{Me}_3\text{Al}$ /water system,<sup>[6]</sup> to give rise to a single product, **13**, in 90% yield. Thus, fragment **13** with four contiguous stereogenic centers was synthesized in a straight-

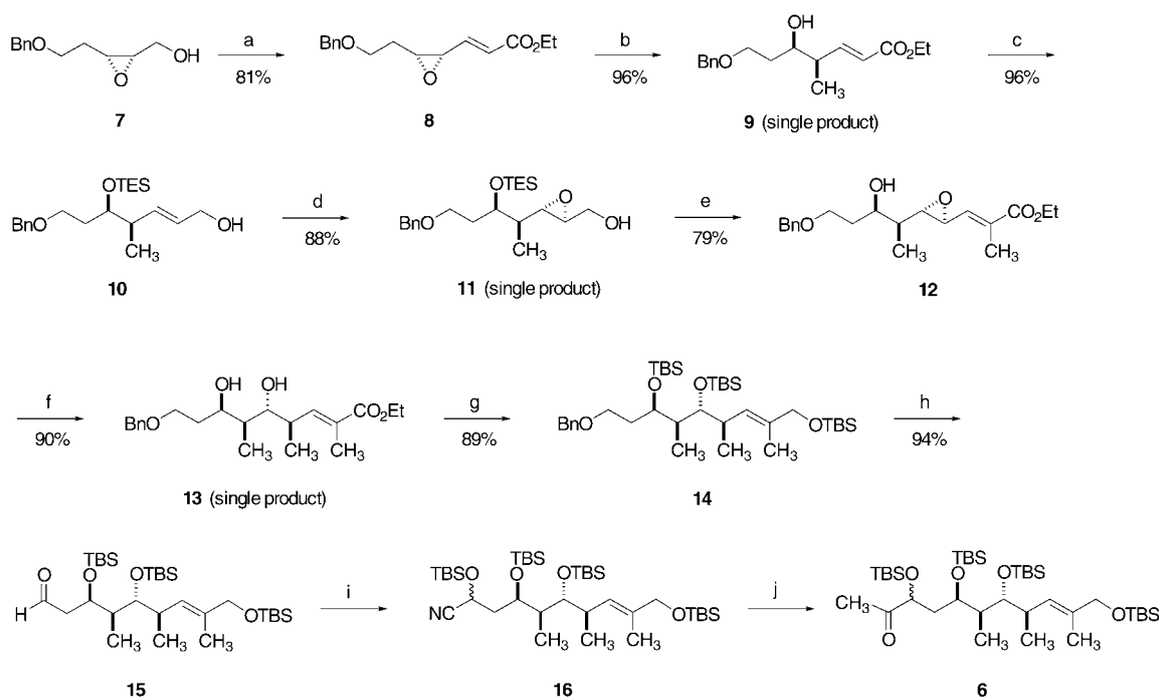
forward and highly stereoselective manner by our original strategy and methodology.

Compound **13** was readily transformed into **14** in three steps: 1) protection of the secondary hydroxy groups with TBSOTf, 2) reduction of the ester with DIBAL-H, and 3) protection of the primary alcohol with TBSCl (89% yield over three steps). When **14** was treated with LDBB in THF and then with Dess–Martin periodinane in the presence of pyridine in  $\text{CH}_2\text{Cl}_2$ , the desired aldehyde **15** was obtained in 94% yield. The crucial acyclic precursor **6** was successfully derived from aldehyde **15** by treatment with TBSCN and  $\text{ZnI}_2$ ,<sup>[8]</sup> which led to cyanohydrin **16**, followed by an addition of MeLi to the nitrile group in THF. The product **6** was a diastereomeric mixture with respect to the configuration of the silyloxy group.

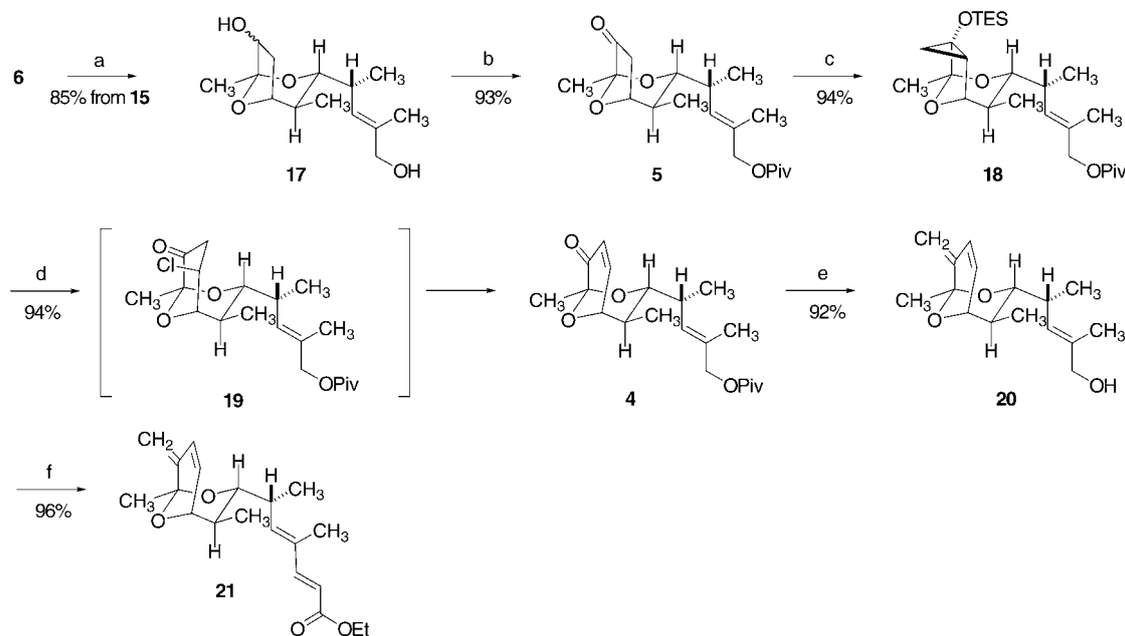
The acid-catalyzed intramolecular acetalization of **6** to form the crucial 2,6-dioxabicyclooctane skeleton proceeded very efficiently as we expected (Scheme 3). Thus, when **6** was treated with aqueous HF in  $\text{CH}_3\text{CN}$  at room temperature, the intramolecular acetalization occurred smoothly and cleanly, to give rise to **17** in 85% overall yield from **15**. After protection of the primary alcohol in **17** as a pivalate moiety, oxidation of the secondary alcohol with Dess–Martin periodinane furnished the desired 2,6-dioxabicyclooctanone **5** in 93% yield. With the critical precursor in hand, we next focused on the ring expansion of **5** to form the dioxabicyclononane skeleton **4**, the key step in the present synthesis. The key transformation was efficiently and highly stereoselectively performed by using the Ito–Saegusa method,<sup>[9]</sup> which involves the following three-step reaction sequence: 1) preparation of the silyl enol ether by treatment of **5** with KHMDs and TESCl, 2) cyclopropanation with  $\text{Et}_2\text{Zn}$  and  $\text{CH}_2\text{I}_2$  to form **18**, and 3) subsequent treatment of **18** with FeCl<sub>3</sub> (88% yield over three steps). Thus, the targeted 2,6-dioxabicyclo-



**Scheme 1.** Retrosynthetic analysis of streptolic acid (**3**) and tirandalydigin (**2**). Piv = pivaloyl, TBS = *tert*-butyldimethylsilyl, Bn = benzyl.



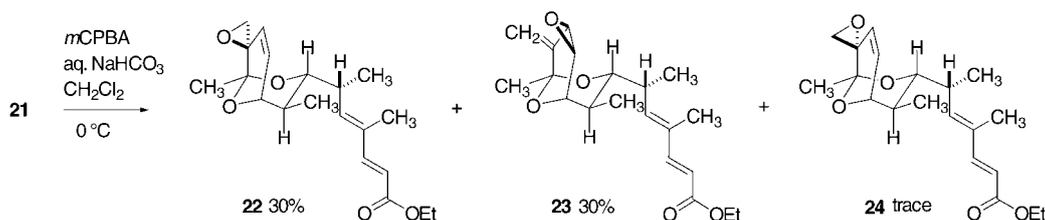
**Scheme 2.** Highly stereoselective synthesis of the acyclic precursor **6**. Reagents and conditions: a) 1. Swern oxidation; 2.  $(\text{EtO})_2\text{P}(\text{O})\text{CH}_2\text{CO}_2\text{Et}$ , NaH, THF,  $0^\circ\text{C}$ , 81% (2 steps); b)  $\text{Me}_3\text{Al}$ ,  $\text{D}_2\text{O}$ ,  $\text{CH}_2\text{Cl}_2$ ,  $-30 \rightarrow -10^\circ\text{C}$ , 96%; c) 1. TESCl, imidazole, DMAP,  $\text{CH}_2\text{Cl}_2$ , room temperature; 2. DIBAL-H, THF,  $0^\circ\text{C}$ , 96% (2 steps); d) *m*CPBA,  $\text{CH}_2\text{Cl}_2$ ,  $0^\circ\text{C}$ , 88%; e) 1. PDC, MS4A,  $\text{ClCH}_2\text{CH}_2\text{Cl}$ ,  $60^\circ\text{C}$ , then  $\text{Ph}_3\text{P}=\text{C}(\text{CH}_3)\text{CO}_2\text{Et}$ ,  $0^\circ\text{C}$ , 79%; 2. TBAF, THF,  $0^\circ\text{C}$ , 100%; f)  $\text{Me}_3\text{Al}$ ,  $\text{D}_2\text{O}$ ,  $\text{CH}_2\text{Cl}_2$ ,  $-30 \rightarrow -10^\circ\text{C}$ , 90%; g) 1. TBSOTf, 2,6-lutidine,  $\text{CH}_2\text{Cl}_2$ ,  $0^\circ\text{C}$ , 93%; 2. DIBAL-H, THF,  $0^\circ\text{C}$ , 98%; 3) TBSCl, imidazole, DMAP,  $\text{CH}_2\text{Cl}_2$ ,  $0^\circ\text{C}$ , 98%; h) 1. LDBB, THF,  $-78 \rightarrow -45^\circ\text{C}$ , 99%; 2. Dess–Martin periodinane, pyridine,  $\text{CH}_2\text{Cl}_2$ , room temperature, 95%; i) TBSCN,  $\text{ZnI}_2$ , room temperature; j) MeLi, THF,  $-78^\circ\text{C}$ , then aq. HCl. THF = tetrahydrofuran, TES = triethylsilyl, DMAP = 4-(dimethylamino)pyridine, DIBAL-H = diisobutylaluminum hydride, *m*CPBA = 3-chloroperoxybenzoic acid, PDC = pyridinium dichromate, MS4A = molecular sieves (4 Å),  $\text{ClCH}_2\text{CH}_2\text{Cl}$  = dichloroethane, TBAF = tetrabutylammonium fluoride, OTf = trifluoromethanesulfonate, LDBB = lithium di-*tert*-butylbiphenylide.



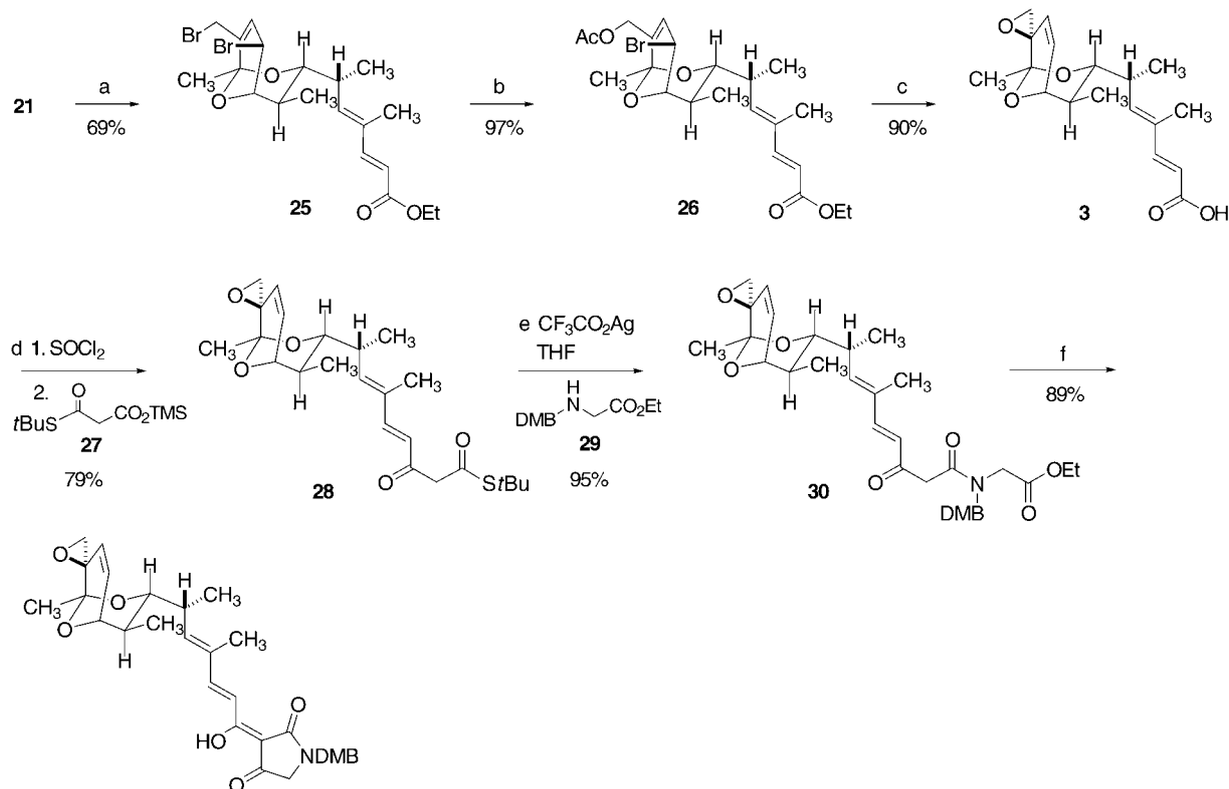
**Scheme 3.** Stereoselective ring expansion leading to the key dioxabicyclo compound, **4**. Reagents and conditions: a) aq. HF,  $\text{CH}_3\text{CN}$ , room temperature, 85% from **15**; b) 1. PivCl, pyridine,  $\text{CH}_2\text{Cl}_2$ ,  $0^\circ\text{C}$ ; 2. Dess–Martin periodinane,  $\text{H}_2\text{O}$ , pyridine,  $\text{CH}_2\text{Cl}_2$ , room temperature, 93% (2 steps); c) 1. KHMDS, TESCl, THF,  $-78^\circ\text{C}$ ; 2.  $\text{Et}_2\text{Zn}$ ,  $\text{CH}_2\text{I}_2$ ,  $\text{Et}_2\text{O}$ , reflux, 94% (2 steps); d) 1.  $\text{FeCl}_3$ , pyridine, DMF,  $100^\circ\text{C}$ , 94%; e) 1.  $\text{Bu}_3\text{SnCH}_2\text{Li}$ , THF,  $-78^\circ\text{C}$ , then MeLi; 2. aq. HF,  $\text{CH}_3\text{CN}$ , room temperature, 92% (2 steps); f) 1.  $\text{MnO}_2$ ,  $\text{CH}_2\text{Cl}_2$ , room temperature; 2.  $(\text{EtO})_2\text{P}(\text{O})\text{CH}_2\text{CO}_2\text{Et}$ , NaH, THF, room temperature, 96% (2 steps). Piv = pivaloyl, KHMDS = potassium hexamethyldisilazide, DMF = *N,N*-dimethylformamide,  $\text{Bu}_3\text{SnCH}_2\text{Li}$  = (tributylstannylium)methylithium.

nonane skeleton **4** could be highly efficiently and stereoselectively secured by the ring-expansion strategy. The next task was introduction of the vinyl epoxide moiety. For this purpose, the enone **4** was initially transformed into diene **20** in three steps: 1) treatment with (tributylstannyl)methyl-lithium<sup>[10]</sup> in THF, 2) then treatment with MeLi to remove the pivaloyl group, and 3) formation of the *exo* methylene group by treatment with HF in CH<sub>3</sub>CN (92% over three steps). Compound **20** was routinely converted into **21** in 96% yield by MnO<sub>2</sub> oxidation followed by a Horner–Wadsworth–Emmons reaction. Notably, every step depicted in Scheme 3 proceeded with excellent (more than 90%) yields.

We had reached a critical stage in the total synthesis of streptolic acid (**3**) and tirandalydigin (**2**), that is, the stereoselective epoxidation of the *exo* double bond and installation of the tetramic acid moiety. To this end, we examined direct epoxidation of **21** to form **22**; however, as shown in Scheme 4,



**Scheme 4.** Unsuccessful epoxidation of **21** with *m*CPBA.



**N**-2,4-Dimethoxybenzyl tirandalydigin (**31**)

**Scheme 5.** Synthesis of streptolic acid (**3**) and *N*-2,4-dimethoxybenzyl tirandalydigin (**31**). Reagents and conditions: a) Bu<sub>4</sub>NBr<sub>3</sub>, 2,6-lutidine, ClCH<sub>2</sub>CH<sub>2</sub>Cl, room temperature, 69%; b) CsOAc, DMF, room temperature, 97%; c) 1. K<sub>2</sub>CO<sub>3</sub>, MeOH, 0°C; 2. NaH, THF, room temperature, 98% (2 steps); 3. aq. NaOH, MeOH, room temperature, 92%; d) 1. SOCl<sub>2</sub>, 2,6-lutidine, CH<sub>2</sub>Cl<sub>2</sub>, room temperature; 2. NaH, **27**, THF, room temperature, 79% (2 steps); e) **29**, CF<sub>3</sub>CO<sub>2</sub>Ag, Et<sub>3</sub>N, MS4A, THF, room temperature, 95%; f) TBAF, THF, room temperature, 89%. DMB = 2,4-dimethoxybenzyl, Bu<sub>4</sub>NBr<sub>3</sub> = tetrabutylammonium tribromide, CsOAc = cesium acetate.

$[\alpha]_{\text{D}}^{22} = +138^{\circ}$  ( $c = 0.55$ , 95% EtOH)<sup>[2f]</sup>, and  $^1\text{H}$  and  $^{13}\text{C}$  NMR, IR, and mass spectra.<sup>[2f]</sup>

The crucial construction of the tetramic acid moiety for the total synthesis of tirandalydigin was performed by employing the protocol of Ley et al.<sup>[11]</sup> Thus, initially, streptolic acid (**3**) was converted into keto thiolester **28** in two steps: 1) treatment with  $\text{SOCl}_2$  leading to the acid chloride and 2) condensation of the acid chloride with **27** by the use of NaH in THF (79% for two steps). Upon treatment of **28** with *N*-2,4-dimethoxybenzyl glycine ethyl ester (**29**),  $\text{Et}_3\text{N}$ , and silver trifluoroacetate in THF, the desired product **30** was obtained in 95% yield; this product was then treated with TBAF in THF to afford the target compound, *N*-2,4-dimethoxybenzyl tirandalydigin (**31**), in 89% yield.<sup>[12]</sup> The synthesis of **31** was unambiguously confirmed by its spectral data, including the  $^1\text{H}$  and  $^{13}\text{C}$  NMR, IR, and mass spectra. The total yield of **31** was 9.8% (an average of 94% yield for each step) in 37 steps from the starting material **7**.

In summary, we have developed a new and promising synthetic methodology for the synthesis of the tetramic acid family of antibiotics and have completed the first synthesis of *N*-2,4-dimethoxybenzyl tirandalydigin (**31**), as well as the synthesis of streptolic acid (**3**), in a highly stereoselective manner.

Received: October 14, 2004

Published online: January 28, 2005

**Keywords:** antibiotics · asymmetric synthesis · methylation · natural products · stereoselectivity

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