## **A Novel Approach toward the Synthesis of Kendomycin: Selective Synthesis of a** *C***-Aryl Glycoside as a Single Atropisomer**

**Stefan Pichlmair, Maria M. B. Marques, Martin P. Green, Harry J. Martin, and Johann Mulzer\***

*Institut fu¨r Organische Chemie, Wa¨hringerstrasse 38, A-1090 Wien, Austria*

*johann.mulzer@uni*V*ie.ac.at*

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**ABSTRACT**



A convergent and concise route to an advanced precursor 2b of kendomycin (1) has been developed by applying a S<sub>N</sub>1 ring cyclization as a **key step. The resulting** *C***-aryl glycoside was initially isolated as a rotameric mixture, but after MOM protection of the** *o***-hydroxyl of the phenol, the conformation was frozen to the desired kendomycin-like atropisomer.**

Kendomycin  $[(-)$ -TAN 2162] (1), a novel ansamycin compound isolated from *Streptomyces* V*iolaceoruber* (strain 3844-33C), was recently described as a potent endothelin receptor antagonist and antiosteoperotic compound with remarkable antibacterial and cytostatic activity.<sup>1</sup> The structure of kendomycin (**1**) features an aliphatic ansa chain with a highly substituted tetrahydropyran ring connected to a unique quinone methide chromophore. Its diverse pharmacological activity and challenging structure have motivated us to embark on a laboratory synthesis of **1**.

A central issue lies in the construction of the pseudo *C*-aryl glycosidic part of the molecule, which is highly sterically congested and shows atropisomeric behavior.2 To reduce steric hindrance, a benzofuran intermediate such as **2** was envisaged that could then be macrocyclized in the C-9/C-11 region and oxidized to the final *p-*quinomethide.

With these considerations in mind, we reasoned that **2** might be obtained from the addition of aldehyde **4** to a carbanion, which could be generated by ortho-directed metalation from **5** (Scheme 1).

The synthesis of aldehyde **4** started with a *syn*-aldol addition of aldehyde **3**, readily available from citronellene, to *â*-keto imide **6** to give ketone **7**. 3

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After stereoselective reduction of  $\overline{7}$  to the  $\beta$ -hydroxy alcohol, $4$  the auxiliary was cleaved by treatment with base, and subsequent acidification with HCl gave the lactone **8** directly. In the presence of 2,2-dimethoxypropane and a catalytic amount of acid, methyl ester **9** was obtained, which

<sup>(1) (</sup>a) Funahashi, Y.; Kawamura, N.; Ishimaru, T. Japanese patent 08231551 [A2960910], 1996; *Chem. Abstr.* **1997**, *126*, 6553. (b) Funahashi, N.; Kawamura, N. Japanese patent 08231552, 1996; *Chem. Abstr.* **1996**, *125*, 326518. (c) Su, M. H.; Hosken, M. I.; Hotovec, B. J.; Johnston, T. L. US patent 5728727, 1998; *Chem. Abstr.* **1998**, *128*, 239489. (d) Bode, H. B.; Zeeck, A. *J. Chem. Soc., Perkin Trans. 1* **2000**, 323. (e) Bode, H. B.; Zeeck, A. *J. Chem. Soc., Perkin Trans. 1* **2000**, 2665.

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<sup>(4)</sup> Evans, D. A.; Chapman, K. T.; Carreira, E. M. *J. Am. Chem. Soc.* **1988**, *110*, 3560.





was first reduced to primary alcohol **10** and then oxidized to aldehyde **4** by Swern oxidation (Scheme 2).

We then tackled the syntheses of two alternative "eastern fragments" from an identical precursor. TBDPS-protected Roche aldehyde **<sup>11</sup>** was subjected to a Horner-Wadsworth-Emmons olefination to afford the corresponding *N*-enoyl sultam ,which was  $\alpha$ -methylated by using Oppolzer's 1,4addition/enolate-trapping protocol.5,6 We obtained **12** in 83% yield with good stereoselectivity (97:3).

The sultam was removed by DIBAL-H reduction and the crude aldehyde was reduced with NaBH4 to give alcohol **13**. Epoxide **15** was obtained from primary alcohol **13** first by conversion to the iodide which was then coupled with isopropenylmagnesium bromide under Schlosser-Fouquet conditions.7 TBDPS deprotection delivered alcohol **14** and Swern oxidation furnished the corresponding aldehyde, which was then transformed directly to **15** by addition of lithiated dibromomethane at low temperature and subsequent warming to room temperature.8 Epoxide **17** was obtained by employing a high-yielding three-step procedure from alcohol **13** (Scheme 3).



 $a$  L-Selectride = lithium tri- $sec$ -butylborohydride, HMPA = hexamethyl phosphoric triamide,  $MCPBA = m$ -chlorperbenzoic  $acid, TBAF = tetrabutylammonium fluoride.$ 

At this stage, epoxide **15** was added to Grignard **19** and epoxide **17** to Grignard **22** to give alcohols **20** and **23**, respectively.<sup>9,10</sup> The alcohols were subjected to a Swern oxidation followed by an acidic catalyzed ring closure. Benzofurans **5a** and **24** were obtained in good yield. Phenol **24** was then reprotected to give the MOM phenol ether **5b** in nearly quantitative yield (Scheme 4).

After careful experimentation, directed ortho-lithiation of the 2-position of benzofurans **5a** and **5b** was performed with

<sup>(5)</sup> For a reference to the preparation of (*S*)-*N*-diethoxyphosphonoacetylcamphorsultam see: Oppolzer, W.; Dupuis, D.; Poli, G.; Raynham, T. M.; Bernardinelli, G. *Tetrahedron Lett.* **1988**, *29* (46), 5885.

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<sup>(7) (</sup>a) Hegedus, L.; Lipshutz, B.; Nozaki, M.; Reets, M.; Rittmeyer, P.; Smith, K.; Trotter, F.; Yamamoto, H. *Organometallics in Synthesis*: *A Manual*; Schlosser, M., Ed.; John Wiley & Sons Ltd: Chichester, UK, 1994; p 344. (b) Tamura, M.; Kochi, J. K. *J. Organomet. Chem.* **1972**, *42*, 205.

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<sup>(9)</sup> Suzuki, S.; Shiono, M.; Fujita, Y. *Synthesis* **1983**, 804.

<sup>(10)</sup> For references to the preparation of **18** see: Corey, E. J.; Gin, D. Y.; Kania, R. S. *J. Am. Chem. Soc.* **1996**, *118*, 9202. For references to the preparation of **21** see: Heckrodt, T. J.; Mulzer, J*. J. Am. Chem. Soc.* **2003**, *125*, 4680.



 $a$  NBS = N-bromosuccinimide, MOMCl = methoxymethyl chloride.

*n*-BuLi in the presence of TMEDA with *n*-BuLi at  $-30$  °C.<sup>11</sup> Addition of aldehyde 4 to these organolithium species at  $-78$ °C gave the desired benzylic alcohols **25** and **26** in 69% and 88% yield as mixtures of diastereomers. Removal of the acetal protection group from **25** and subsequent heating in warm toluene with a catalytic amount of *p*-TsOH resulted in the desired  $S_N1$ -type ring closure as a key step in our synthesis (Scheme 5). In the case of **25**, we were disappointed to find that the resulting tetrahydropyran product **2a** exists as a 3:1 mixture of atropisomers favored to the undesired isomer at room temperature.

Unfortunately, the cyclization of **26** under analogous conditions was thwarted by the formation of a stable acetonide between the oxygens at C1 and C3. Hence, we applied a three-step procedure of acetylation, formation of the triol, and treatment with acid, which furnished **27** in reasonable overall yield. As before, we observed broad peaks in the 1H NMR spectra, implying the formation of a rotameric mixture and that the coalescence temperature is around room temperature. To our delight MOM protection of the phenol **27**



 $^a$  TMEDA =  $N$ , $N$ , $N'$ , $N'$ -tetramethylethylendiamine, MOMCl = methoxymethyl chloride.

resulted in freezing of the conformation to the desired kendomycin-like atropisomer **2b**.

In conclusion, we have developed a convergent and stereoselective route to an advanced intermediate in the synthesis of kendomycin. A novel feature in our sequence is the stereocontrol over the formation of *C*-aryl glycoside atropsiomers, which has been exerted by a proper choice (i.e. MOM and OMe) of the phenolic *o*-hydroxyl groups. The atropisomer thus obtained has the proper geometry for the formation of the macrocyclic ring, which will be the next issue to face.

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**Supporting Information Available:** Experimental procedures and NMR data for compounds **25**, **26**, **2a**, and **2b**. This material is available free of charge via the Internet at http://pubs.acs.org.

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