

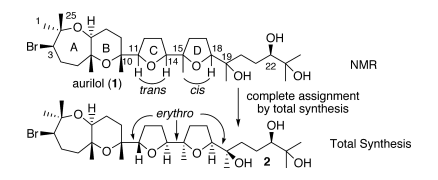
Communication

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J. Am. Chem. Soc., 2005, 127 (16), 5806-5807• DOI: 10.1021/ja050123p • Publication Date (Web): 02 April 2005

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Published on Web 04/02/2005

Total Synthesis and Complete Assignment of the Stereostructure of a Cytotoxic Bromotriterpene Polyether (+)-Aurilol

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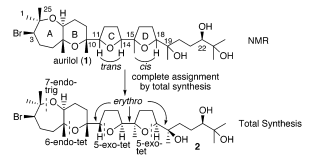
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A bromotriterpene polyether aurilol (1) was isolated from the sea hare, Dolabella auricularia, by Yamada et al. in 1998 and exhibited cytotoxicity against HeLa S_3 cells with IC₅₀ of 4.3 μ g/ mL (Chart 1).1 Although the plane structure and partial stereochemistry of 1 were elucidated by spectroscopic and chemical analyses, determination of the entire stereochemistry has not been reached. There have also been many other types of triterpene polyethers (oxasqualenoids);² however, it is often difficult to determine their stereostructures even by the current highly advanced spectroscopic methods, especially in acyclic systems including quaternary carbon centers such as C10-C11, C14-C15, and C18-C19 in 1. In such cases, it is effective to predict and synthesize the possible stereoisomers.³ 1 also possesses a synthetically challenging 2,8-dioxabicyclo[5.4.0]undecane A,B ring system with a bromine atom at the neopentylic position and 1,3-diaxial dimethyl substituents on the B ring. These contexts have made oxasqualenoids, including bromotriterpene polyethers,⁴ attractive targets for many synthetic organic chemists.³ In this paper, we report that the total assignment of the incomplete stereostructure of (+)-aurilol (1) to 2 has been achieved through its first asymmetric total synthesis featuring biogenetic-like regioselective ether ring formations to secure the stereochemical pathway.

In the course of our structural studies on oxasqualenoids, which could biogenetically be derived by epoxide-opening cyclizations of squalene polyepoxides,^{3b,4b} we predicted the unknown stereochemistry to be *erythro* configuration as shown in **2**. In the retrosynthetic analysis of **2**, we planned to straightforwardly construct all of the A–D ether rings by biogenetic-like cyclizations. The oxepane A ring would be constructed by 7-endo-trig⁵ bromoetherification of the corresponding trishomoallylic alcohol, and the B–D rings would be formed by 6-endo-tet, 5-exo-tet, and 5-exo-tet⁵ epoxide openings of the corresponding bishomoepoxy alcohols, respectively (vide infra).

The synthesis of target molecule **2** began with SEM protection of the hydroxy group in the known optically active epoxy alcohol **3** (98% ee)⁶ (Scheme 1). Selective deprotection of the TBDMS ether in **4** and Sharpless' epoxidation⁷ of the allylic alcohol **5** using L-(+)-DET afforded diepoxide **6**. The C ring formation from **6** under the alkaline condition stereoselectively proceeded via a regioselective 5-exo-tet epoxide opening of the intermediate **A** to provide triol **7** in 88% yield.⁸ The following sequence, (1) mesylation; (2) epoxidation; (3) chain extension with the C₁₀ unit **8**;⁹ and (4) desulfurization,^{3b} uneventfully yielded diol **9** from **7**. Shi's epoxidation of **9** catalyzed by chiral ketone **10**¹⁰ diastereoselectively gave labile bishomoepoxy alcohol **11**. Treatment of **11** with a protic acid underwent a regioselective 5-exo-tet cyclization to produce diol **12** in 98% yield.¹¹

With the highly stereocontrolled **12** in hand, we embarked on the elaboration of the formidable A,B-ring system. Deprotection Chart 1. Structures of Aurilol

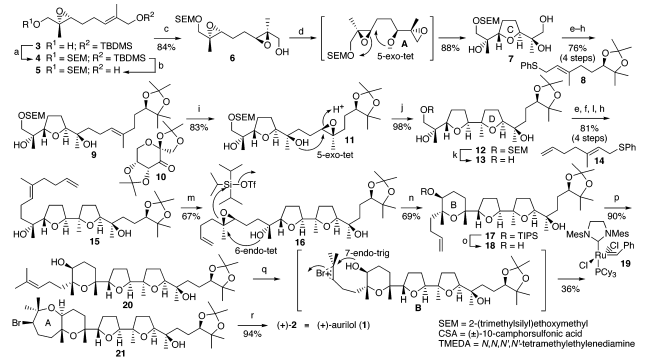


of the SEM ether in **12** afforded triol **13**, from which diene **15** was derived by the same sequence as that of **7** to **9** except for employing sulfide **14**.¹² Shi's epoxidation of the diene **15** with *ent*-**10**,¹⁰ enantiomeric to **10**, proceeded in a regioselective manner to provide monoepoxide **16** with the terminal alkene intact. The construction of the desired B ring was regioselectively carried out by treating the bishomoepoxy alcohol **16** with TIPSOTf and 2,6-lutidine in CH₃NO₂ at 0 °C for 20 min. The unprecedented 6-endo-tet cyclization promoted by a silyl triflate against Baldwin's rule⁵ would be very stimulating¹⁴ because Brønsted acid-catalyzed cyclizations for these types of bishomoepoxy alcohols, such as **11**, generally afford 5-exo-tet regioselectivity, regardless of the relative configurations of the tertiary alcohol—epoxide substrates.¹¹

After removal of the TIPS group in **17**, cross metathesis of olefin **18** with 2-methyl-2-butene using Grubbs' catalyst **19**¹⁵ produced alkene **20** in 90% yield.¹⁶ After many attempted experimentations,¹⁷ it has been found that (CF₃)₂CHOH with high polarity and non-nucleophilicity¹⁸ is the solvent of choice to successfully form the A ring. The regio- and stereoselective 7-endo-trig bromoetherification of **20** was brought about under the optimal condition to give the A,B-ring system **21**,¹⁹ wherein removal of the acetonide, finally furnished the target molecule **2**. The spectral characteristics (¹H and ¹³C NMR) of the synthetic **2**, $[\alpha]^{19}_{\text{D}} + 4.5$ (*c* 0.21, CHCl₃), were identical to those reported for the natural product, $[\alpha]^{30}_{\text{D}} + 4.6$ (*c* 0.41, CHCl₃).¹ Thus, it has been found that the hitherto unknown relative configuration of (+)-aurilol (**1**) is *erythro* as indicated in **2**.

In conclusion, we have accomplished the first asymmetric total synthesis of a cytotoxic bromotriterpene polyether (+)-aurilol (4.74% overall yield in 21 steps from 3) featuring the highly regio- and stereocontrolled biogenetic-like A–D ether ring formations. The total synthesis has realized the total assignment of the incomplete stereostructure of aurilol (1), which is difficult to determine the stereochemistry otherwise. Application of this synthetic strategy to other bromotriterpene polyethers is in progress.

Scheme 1. Total Synthesis of Target Molecule 2^a



^a Reaction conditions: (a) SEMCl, *i*-Pr₂NEt, CH₂Cl₂, 0 °C to room temperature, 12 h, 99%; (b) Bu₄NF, THF, 0 °C, 1 h, 100%; (c) *t*-BuO₂H, Ti(O*i*-Pr)₄, L-(+)-DET, MS 4A, CH₂Cl₂, -25 °C, 16 h (>20:1); (d) 1 M aq NaOH, 1,4-dioxane, reflux, 5 h; (e) MsCl, Py, CH₂Cl₂, 0 °C to room temperature, 1 h; (f) K2CO3, MeOH, rt, 15 min; (g) 8, BuLi, TMEDA, THF, -78 °C, 1 h; (h) Na, i-PrOH, THF, reflux, 4 h; (i) 10, Oxone, (MeO)2CH2/CH3CN/H2O, pH 10.5, 0 °C, 2 h (>15:1); (j) CSA, CH₂Cl₂, rt, 10 min; (k) Bu₄NF, THF, reflux, 30 h, 92%; (l) 14, BuLi, TMEDA, THF, -78 °C, 1 h; (m) ent-10, Oxone, (MeO)₂CH₂/CH₃CN/H₂O, pH 10.5, 0 °C, 2 h (>10:1); (n) TIPSOTf, 2,6-lutidine, CH₃NO₂, 0 °C, 20 min; (o) Bu₄NF, THF, 0 °C to room temperature, 15 h, 100%; (p) 19, 2-methyl-2-butene, reflux, 12 h; (q) 2.5 equiv of NBS, MS 4A, (CF₃)₂CHOH, 0 °C, 10 min (>10:1); (r) 80% aq AcOH, rt, 14 h.

Acknowledgment. This research was financially supported by the Novartis Foundation (Japan) for the Promotion of Science.

Supporting Information Available: Characterization data for 2–7, 9, 11-13, 15-18, 20, and 21, and experimental procedures for synthesis of 2 (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

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0 °C, 15 h, 80% (98% ee); (2) Me2C(OMe)2, CSA, CH2Cl2, 0 °C, 2 h, 97%; (3) K₂CO₃, MeOH, rt, 4 h, 98%; (4) (PhS)₂, Bu₃P, THF, rt, 4 h, 96%

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JA050123P