

Total Synthesis of Brevetoxin B. 1. CDEFG Framework

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With its imposing structure, brevetoxin B (1), produced by *Gymnodinium breve* Davis, stood as a formidable challenge to synthetic chemists since its discovery and structural elucidation in 1981.¹ Brevetoxin's beautifully arranged molecular assembly includes 11 *trans*-fused rings, each containing an oxygen atom, with each fusion consisting of a C–C bond separating two adjacent ring oxygens and with all adjacent substituents flanking the oxygens placed *syn* to each other except on ring K. Its unprecedented architecture, its association with the "red tide" catastrophes,² and its potent neurotoxicity and interference with the function of sodium channels attracted serious attention from chemists³ and biologists⁴ alike. We now wish to announce, in this and the following communication,⁵ the total synthesis of brevetoxin B (1) in its naturally occurring form.

Figure 1 outlines the strategic bond disconnections and retrosynthetic analysis of 1. The adopted strategy benefited from convergency (oxocene disconnections) and synthetic technologies developed in these laboratories specifically for constructing oxocene⁶ and tetrahydropyran⁷ systems.

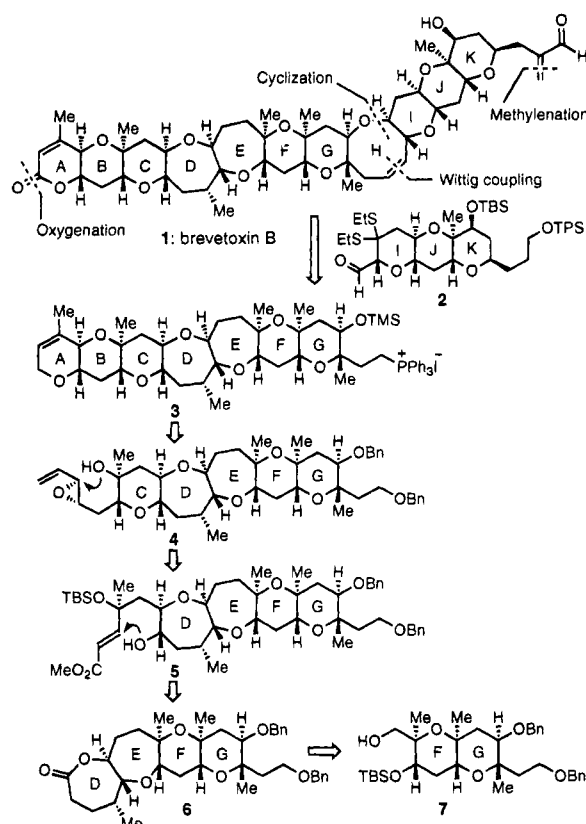


Figure 1. Strategic bond disconnections and retrosynthetic analysis of brevetoxin B (1).

The construction of the CDEFG framework 4 described herein began with the previously reported intermediate 7 (Scheme 1).⁸ Swern oxidation of 7 followed by a Wittig reaction with the appropriate reagent furnished, in 99% overall yield, compound 9 via aldehyde 8. Hydrogenation of 9 and selective, acid-induced monodesilylation gave alcohol 11 via 10 in 97% overall yield. Oxidation of 11 in a sequential fashion using Swern and NaClO₂ conditions resulted in carboxylic acid 12 (97%), which upon desilylation with TBAF led to 13 (91%). Lactonization of hydroxy acid 13 by the Yamaguchi method⁹ and enol triflate formation gave 15 via 14 in 84% overall yield. Generation of the higher order cuprate derived from the lithio derivative of iodide 17a¹⁰ and 17b followed by coupling¹¹ with triflate 15 and partial acid-induced orthoester hydrolysis resulted in formation of 18 via 16 (84% yield over two steps, *ca.* 2.4:1 ratio at C* in favor of the desired isomer, *vide infra*). Regio- and stereoselective hydroboration of 18 followed by oxidative workup and alkaline hydrolysis furnished hydroxy acid 19 in 73% overall yield. Finally, lactonization⁹ of 19 and separation of the C* epimers afforded pure lactone 6 (60% yield, plus 25% of its C* methyl epimer), whose structure was determined by X-ray crystallographic analysis (see ORTEP drawing of a derivative¹⁰ of 6, Figure 2).

The fusion of the remaining three rings onto the DEFG system 6 to afford the targeted polycyclic framework 4 proceeded as depicted in Scheme 2. Thus, conversion of lactone 6 to its enol triflate (97%) followed by Cr/Ni-mediated coupling¹² with

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Scheme 1. Construction of DEFG Ring System 6^a