

Biomimetically Inspired Total Synthesis and Structure Activity Relationships of 1-O-Methyllateriflorone. 6π **Electrocyclizations in Organic Synthesis**

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Abstract: The total synthesis of 1-O-methyllateriflorone (2) is described. The construction of the cage-like domain of the molecule involved a biomimetic Claisen/Diels-Alder cascade, whereas the novel spiroxalactone framework was generated by an intramolecular Michael reaction within precursor 16a involving the carboxylate residue as the nucleophile. This finding might bear on the biosynthetic pathway by which nature forms lateriflorone. Described herein is also an interesting cascade sequence involving facile 6π electrocyclizations which leads to complex benzopyran systems. The biological evaluation of a small library of lateriflorone analogues and related systems establishing the first SAR within this class of compounds is also included. Among the most active compounds against tumor cells are 2, 16b, 56, 58, and 59.

Introduction

The continuing phytochemical studies of Garcinia species (belonging to the Guttiferae family of tropical plants) led to the isolation of several xanthone-derived natural products, which possess unusual molecular architectures and diverse biological properties. Thus, some of these compounds exhibit noteworthy cytotoxic and antibacterial properties, and the plant sources from this family have been used by natives as folk medicines for centuries.¹ The signature structural motif of many of these compounds is an intriguing 4-oxatricyclo[4.3.1.0]decan-2-one scaffold attached onto a common xanthonoid ring system. Figure 1 displays a number of representative members of this class of cage-like xanthonoids, including lateriflorone (1),² morellin (3),³ desoxymorellin (4),3 morellic acid (5),3 bractatin (6),4 1-Omethylbractatin (7),⁴ 1-O-methylneobractatin (8),⁴ forbesione (9),⁵ gaudichaudione H (10),⁶ scortechinone A (11),⁷ and

scortechinone B (12).7 Among all, lateriflorone (1), isolated from the stem bark of Garcinia Lateriflora Bl (Guttiferae),² is the most interesting in that it possesses the additional complexity of a unique spiroxalactone moiety, making its total synthesis the ultimate challenge within this group of natural products.

Based on some scattered speculations, the biogenetic pathway8 for the formation of the xanthonoid ring system of these compounds might involve an intramolecular oxidative coupling of benzephenone or benzophenone-like intermediates, which, in turn, are formed by the condensation of shikimate and acetatederived moieties. In case of lateriflorone (1), it is hypothesized² that the unique spiroxalactone could be formed either by an oxidative cyclization of 13 (Scheme 1) or by the spiroketalization between the two fragments 14 and 15. In the later case, the C7-keto functionality of 14 could react with the C7-hydroxyl group of 15 to form a hemiketal, which could subsequently undergo lactonization in a Michael fashion to generate the novel spiroxalactone system. Lateriflorone (1) exhibits potent cytotoxicity against the P388 cancer cells (ED₅₀ 5.4 µg/mL). Like with all other xanthonoids of this family, the mechanism of action of this compound is not known, although it is assumed that the intriguing cage domain of the molecule plays a role in their biological action. In this Article, we describe full details of our investigations in this area which culminated to the total synthesis of 1-O-methyllateriflorone (2), the application of 6π electrocyclizations in the synthesis of novel benzopyran systems, and the synthesis of several lateriflorone analogues and precursors and their biological evaluation.

(2) Kosela, S.; Cao, S.-G.; Wu, X.-H.; Vittal, J. J.; Sukri, T.; Masdianto; Goh,

Synop. 1996, 392-393.

 (6) (a) Cao, S.-G.; Sng, V. H. L.; Wu, X.-H.; Sim, K.-Y.; Tan, B. H. K.; Pereira, J. T.; Goh, S. H. *Tetrahedron* 1998, 54, 10915–10924. For related gaudichaudiones, see: (b) Cao, S.-G.; Wu, X.-H.; Sim, K.-Y.; Tan, B. K. H.; Pereira, J. T.; Wong, W. H.; Hew, N. F.; Goh, S. H. *Tetrahedron Lett.* **1998**, *39*, 3353–3356. (c) Wu, X.-H.; Tan, B. K. H.; Cao, S.-G.; Sim, K.-Y.; Goh, S. H. *Nat. Prod. Lett.* **2000**, *14*, 453–458.

⁽¹⁾ Perry, L. M.; Metzger, J. Medicinal Plants of East and Southeast Asia: Attributed Properties and Uses; The MIT Press: Cambridge, MA, 1980;

⁽²⁾ Roseia, S., Cao, S.-G., Wu, A.-H., Vilda, J. J., Sukil, I., Masdianto, Golf, S.-H.; Sim, K.-Y. Tetrahedron Lett. **1999**, 40, 157–160.
(3) (a) Rao, B. S. J. Chem. Soc. C **1937**, 853–857. (b) Kartha, G.; Ramachandran, G. N.; Bhat, H. B.; Nair, P. M.; Raghavan, V. K. V.; Venkataraman, K. Tetrahedron Lett. **1963**, 4, 459–472. (c) Ollis, W. D.; P. M. (1978). Venkataraman, K. *Tetrahedron Lett.* 1965, 4, 459–472. (c) Ollis, W. D.;
Ramsay, M. V. J.; Sutherland, I. O.; Mongkolsuk, S. *Tetrahedron* 1965, 21, 1453. (d) Karanjgaonkar, C. G.; Nair, P. M.; Venkataraman, K. *Tetrahedron Lett.* 1966, 7, 687–691. (e) Bhat, H. B.; Nair, P. M.; Venkataraman, K. *Indian J. Chem.* 1964, 2, 402.
Thoison, O.; Fahy, J.; Dumontet, V.; Chiaroni, A.; Riche, C.; Tri, M. V.; Sevenet, T. *J. Nat. Prod.* 2000, 63, 441–446.
Leong, Y.-W.; Harrison, L. J.; Bennett, G. J.; Tan, H. T.-W. *J. Chem. Res.*, 5, 2023, 2023, 2023.

Rukachaisirikul, V.; Kaewnok, W.; Koysomboon, S.; Phongpaichit, S.; Taylor, W. C. Tetrahedron 2000, 56, 8539-8543.

⁽a) Bennett, G. J.; Lee, H.-H. *J. Chem. Soc., Chem. Commun.* **1988**, 619–620. (b) Roberts, J. C. *Chem. Rev.* **1961**, *61*, 591–605. (c) Carpenter, I.; Locksley, H. D.; Scheinmann, F. *Phytochemistry* **1969**, *8*, 2013–2026.

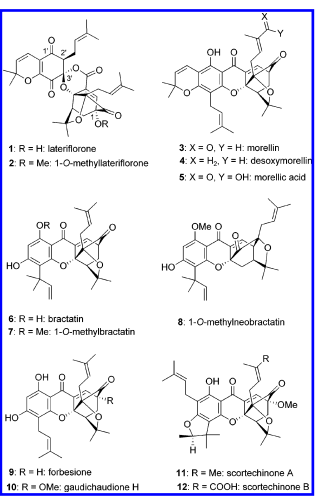
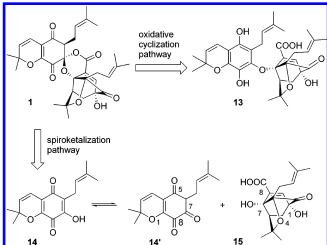


Figure 1. Selected natural products from the Garcinia family of plants.

Scheme 1. Proposed Biosynthesis of Lateriflorone (1)



Results and Discussion

1. Retrosynthetic Analysis. Inspection of the structure of lateriflorone (1)⁹ leads to the identification of the spirolactone-ketal moiety as the most appropriate strategic site for retrosynthetic disconnection. Not only are the two C-O bonds among

Scheme 2. Retrosynthetic Analysis of Lateriflorone (1)

the easiest to form in the molecule, but, most significantly, such disconnections would lead to a convergent strategy in the synthetic direction. Scheme 2 depicts the logic of the initial retrosynthetic analysis of 1. Thus, rupturing bond a generates quinone-ether carboxylic acid 16a. Intramolecular conjugate addition (path a) of the carboxyl group onto the closest enone carbon atom may be expected to lead to 1 - granted with an unknown stereochemical outcome. Alternatively, cleavage of the ether C-O bond of the spiroketal system leads to quinoneester 16b, whose similar intramolecular collapse (path b) may form the target molecule, again with the uncertainty of the stereochemical outcome at the two generated centers looming over the ring closure. In contemplating the choice between the two paths (a and b), the ease of construction of the required precursor (16a or 16b) favored path b, because forming an ester bond was considered so much easier than casting an ether linkage as needed for path a. Having defined 16b as the next target, its quinone moiety was then retrosynthetically transformed into the benzenoid system 17 whose disconnection at the ester bond led to fragments 18 and 19a. Our previous study with forbesione $(9)^{10}$ pointed to the cascade approach to the cage intermediate 19a from the aromatic system 20a via one Claisen rearrangements and one Diels-Alder reaction (intramolecular) as shown in Scheme 2.

 ^{(9) (}a) For a preliminary communication, see: Nicolaou, K. C.; Sasmal, P. K.; Xu, H.; Namoto, K.; Ritzen, A. Angew. Chem., Int. Ed. 2003, 42, 4225–4229.
 (b) For the synthesis of seco-lateriflorone, see: Tisdale, E. J.; Vong, B. J.; Li, H.; Kim, S. H.; Chowdhury, C.; Theodorakis, E. A. Tetrahedron 2003, 59, 6873–6887.

⁽¹⁰⁾ For the total synthesis of 1-O-methyl forbesione, see: Nicolaou, K. C.; Li, J. Angew. Chem., Int. Ed. 2001, 40, 4264–4268.

2. Construction of the Benzopyran Domain. The synthesis of the required prenylated 2,2'-dimethybenzopyran fragment 18 is summarized in Scheme 3. Thus, the known compound 21¹¹ was smoothly and selectively brominated at the para position with bromine in the presence of NaOAc to afford p-bromophenol 22 (92% yield) whose exposure to MOMCl-Et₃N led to MOM ether 23 (93% yield). Dakin oxidation (m-CPBA) of aldehyde 23 followed by in situ cleavage of the intermediate formate ester with NaHCO3 furnished the new bromophenol 24 in 71% overall yield, which was protected as a TIPS ether by treatment with TIPSCl in the presence of imidazole, leading to compound 25 in 96% yield. Combining bromide 25 with B(O'Pr)₃ in ether followed by sequential addition of 'BuLi at -78 °C and NaOH-H₂O₂ at 0 °C furnished phenol **26** (86% yield) via the corresponding borate derivative. ¹² The phenol (26) was then methylated (K₂CO₃-MeI) to afford methoxy compound 27 in 88% yield. Subsequent hydrogenolysis at the benzyl group in 27 (H₂, 10% Pd(OH)₂/C) afforded phenolic compound 28 in quantitative yield. Propargylation¹³ of 28 using in situ generated HC≡CC(Me)₂OCOCF₃ in the presence of DBU and catalytic amounts of CuCl2 afforded propargylic ether 29 in 76% yield (90% yield based on 84% conversion). Compound 29 underwent Claisen rearrangement¹⁴ in refluxing xylene to furnish benzopyran system 30. Removal of the solvent followed by treatment of the crude product with TBAF in THF gave desilylated product 31 in 93% overall yield from 29. Finally, a second propargylation employing the same reaction conditions as mentioned above afforded benzopyran system 32 in 80% yield (91% based on 88% conversion). Selective reduction (H₂) of the acetylenic moiety in 32 in the presence of Lindlar catalyst generated the corresponding olefin (33) in 95% yield. A second Claisen rearrangement under thermal conditions (DMF, 120 °C) converted 33 to the targeted benzopyran fragment 18 in 70% yield.

3. Construction of the Caged Domain and Coupling of the Two Fragments. The sequence devised and executed for the synthesis of the initially designed precursor 20a is shown in Scheme 4. Thus, commercially available 2,3,4-trihydroxybenzaldehyde (34) was perbenzylated with BnBr in the presence of K₂CO₃ and catalytic amounts of KI in DMF, generating the known aldehyde 3515 in 85% yield. Selective debenzylation of the phenolic group adjacent to the aldehyde moiety was achieved by the action of MgBr₂•Et₂O in ether, leading to o-hydroxybenzaldehyde derivative 36 in 83% yield. Bromination with molecular bromine in acetic acid of the later compound (36) resulted in the formation of p-bromophenol 37 in 89% yield. NaBH₄ reduction of aldehyde 37 afforded diol 38 (91% yield), which was subjected to acetonide formation, furnishing ketal **39** in 95% yield. This bromo compound (**39**) was then converted to phenolic substrate 40 in 90% overall yield following the standard protocol mentioned above via the borate intermediate. Thus, lithium-halogen exchange ("BuLi), borate formation

Scheme 3. Construction of Benzopyran Fragment 18a

^a (a) Br₂ (1.1 equiv), NaOAc (1.2 equiv), AcOH, 25 °C, 2 h, 92%; (b) Et₃N (5.0 equiv), 4-DMAP (0.1 equiv), MOMCl (3.0 equiv), 25 °C, 16 h, 93%; (c) m-CPBA (1.1 equiv), $0 \rightarrow 25$ °C, CH₂Cl₂, 6 h; then saturated aqueous NaHCO3, 16 h, 71%; (d) TIPSCl (1.5 equiv), imid (2.0 equiv), DMF, 25 °C, 2 h, 96%; (e) premix 25 and B(O'Pr)₃ (2.2 equiv) in ether; then 'BuLi (2.2 equiv) at -78 °C, 2 h, $-78 \rightarrow 0$ °C, 1 h; then MeOH, 10% aqueous NaOH (4.8 equiv), H2O2 (5.0 equiv), 0 °C, 0.5 h, 86%; (f) K2CO3 (10 equiv), MeI (10 equiv), acetone, 25 °C, 16 h, 88%; (g) 10% Pd(OH)₂/C (10 wt %), H₂ (1 atm), EtOAc/EtOH (1:1), 25 °C, 0.5 h, 100%; (h) propargyl alcohol (1.3 equiv), DBU (1.5 equiv), TFAA (1.2 equiv), MeCN, 0 °C, 0.5 h; DBU (1.35 equiv), CuCl₂ (0.01 equiv), 10 min; then TFA propargyl ester was added, 4 h, 0 °C, 76% (90% based on 84% conversion); (i) xylene, 140 °C, 0.5 h; (j) TBAF (1.5 equiv), THF, 0 °C, 5 min, 93% for two steps; (k) propargyl alcohol (1.3 equiv), DBU (1.5 equiv), TFAA (1.2 equiv), MeCN, 0 °C, 0.5 h; DBU (1.35 equiv), CuCl₂ (0.01 equiv), 10 min; then TFA propargyl ester was added, 4 h, 0 °C, 80% (91% based on 88% conversion); (1) Lindlar catalyst (10 wt %), H2 (1 atm), quinoline (3.0 equiv), EtOAc, 25 °C, 2 h, 95%; (m) DMF, 120 °C, 1 h, 70%. Bn = benzyl, 4-DMAP = 4-(dimethylamino)pyridine, MOM = methoxymethyl, m-CPBA = m-chloroperbenzoic acid, TIPS = triisopropylsilyl, imid = imidazole, DBU = 1,8-diazabicyclo[5.4.0]undec-7-ene, TFAA = trifluoracetic anhydride, TBAF = tetra-n-butylammonium fluoride.

[B(OMe)₃], and oxidation with alkaline (NaOH) hydrogen peroxide resulted in 40 whose phenolic group was protected as

⁽¹¹⁾ Parker, K. A.; Georges, A. T. Org. Lett. 2000, 2, 497–499.
(12) Luszniak, M. C.; Topiwala, U. P.; Whiting, D. A. J. Chem. Res., Miniprint **1998**, 7, 1401-1417

⁽¹³⁾ Godfrey, J. D., Jr.; Mueller, R. H.; Sedergran, T. C.; Soundararajan, N.; Colandrea, V. J. *Tetraheron Lett.* **1994**, *35*, 6405–6408.

^{(14) (}a) Quillinan, A. J.; Scheinmann, F. J. Chem. Soc., Perkin Trans. 1 1972, -1387. (b) Subramanian, R. S.; Balasubramanian, K. K. Tetrahedron Lett. 1988, 29, 6797—6800. (c) Joshi, S. C.; Trivedi, K. N. Tetrahedron 1992, 48, 563—570. (d) Yamaguchi, S.; Ishibashi, M.; Akasaka, K.; Yokoyama, H.; Miyazawa, M.; Hirai, Y. Tetrahedron Lett. 2001, 42, 1091-

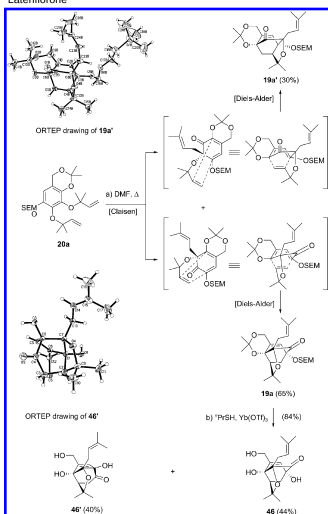
⁽¹⁵⁾ Hegedues, B.; Krasso, A. F. Helv. Chim. Acta 1970, 53, 959-963.

Scheme 4. Construction of Intermediate 20°

^a (a) K₂CO₃ (5.0 equiv), BnBr (5.0 equiv), KI (0.1 equiv), DMF, 25 °C, 24 h, 85%; (b) MgBr₂•OEt₂ (1.1 equiv), ether, 25 °C, 10 h, 83%; (c) Br₂ (1.1 equiv), NaOAc (1.15 equiv), AcOH, 25 °C, 2 h, 89%; (d) NaBH₄ (1.5 equiv), EtOH, $0 \rightarrow 25$ °C, 0.5 h, 91%; (e) 2,2-dimethoxypropane (5.0 equiv), p-TsOH (0.01 equiv), CH₂Cl₂, 1 h, 95%; (f) "BuLi (1.1 equiv), ether, -78 °C, 2 h; then B(OMe)₃ (3.0 equiv), 1 h, $-78 \rightarrow 0$ °C; then 10% aqueous NaOH (4.8 equiv), H₂O₂ (5.0 equiv), 0 °C, 0.5 h, 90%; (g) SEMCl (1.5 equiv), $({}^{1}\text{Pr})_{2}$ NEt (2.0 equiv), $\text{CH}_{2}\text{Cl}_{2}$, $0 \rightarrow 25$ °C, 2h, 70%; (h) 10% Pd/C (10 wt %), H₂ (1 atm), EtOAc, 25 °C, 45 min, 95%; (i) 'BuOK (2.2 equiv), THF, 0 °C; then concentrated and suspended in MeCN; then 18-Crown-6 (2.2 equiv), 15 min, bromoisobutyraldehyde (5.0 equiv), $0 \rightarrow 25$ °C, 1 h, 70%; (j) CH₃P⁺Ph₃Br⁻ (3.0 equiv), NaHMDS (3.0 equiv), THF, 0 °C, 1 h, 75%; (k) 'BuOK (1.1 equiv), THF, 0 °C; then concentrated and suspended in MeCN; then 18-Crown-6 (1.1 equiv), 15 min, bromoisobutyraldehyde $(5.0 \text{ equiv}), 0 \rightarrow 25 \text{ °C}, 1 \text{ h}, 75\%; (1) \text{ CH}_3\text{P}^+\text{Ph}_3\text{Br}^- (2.0 \text{ equiv}), NaHMDS}$ (2.0 equiv), THF, 0 °C, 1 h, 75%. Ts = p-toluenesulfonyl, HMDS = hexamethyldisilazane.

a SEM ether (SEMCl, Et₃N) to afford **41** (70% yield). Subsequent hydrogenolysis of the benzyl groups (H₂, 10% Pd/C, 95% yield) in **41** led to the formation of dihydroxy compound **42**. The di-potassium salt of **42**, generated by addition of KO'Bu, was then suspended in acetonitrile and reacted with bromoisobutyraldehyde¹⁶ in the presense of 18-Crown-6 to afford an unseparable mixture of regioisomeric lactols (**43a**: **43b**, ca. 1.7:1 ratio,70% yield) which reacted with methylene phosphorane (generated from MeP+Ph₃Br⁻ and NaHMDS), leading to olefins **44a**:**44b** (ca. 1.7:1 ratio of the regioisomers,

Scheme 5. First Attempt To Prepare the Cage-like Domain of Lateriflorone^a



^a (a) DMF, 120 °C, 2 h, **19a** (65%) and **19a**' (30%); (b) "PrSH (9.0 equiv), Yb(OTf)₃ (0.1 equiv), CH₂Cl₂, 25 °C, 1.5 h, 84%.

75% yield). The phenolic group within **44** was alkylated with bromoisobutyraldehyde under the same reaction conditions as mentioned above to afford regioisomeric aldehydes **45a** and **45b** in 75% combined yield. A second Wittig reaction on the latter mixture then furnished the required di-olefin **20a** in 75% yield.

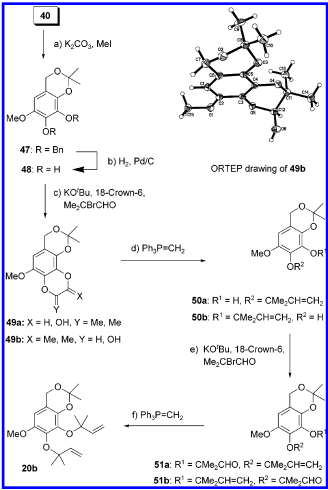
Having constructed the aromatic precursor **20a**, we then proceeded to the next stage which aimed at the rearrangement of this substrate to the desired cage-like intermediate via the projected Claisen/Diels—Alder cascade. ^{10,17} Thus, heating **20a** in DMF at 120 °C for 2 h led to the expected product **19a** (65% yield) and its regioisomer **19a**′ (30% yield), presumably via the shown intermediates (see Scheme 5). The structures of these products were based on both spectroscopic and X-ray crystallographic techniques (see ORTEP drawing ¹⁸ of **19a**′, Scheme 5). Attempts to selectively remove the acetonide group from **19a**, however, were unsuccessful; instead, the various conditions tried led to unwanted products, including triol **46** (44% yield) and rearranged triol **46**′ (40% yield), both of which were

⁽¹⁶⁾ Beckwith, A. L. J.; Thomas, C. B. J. Chem. Soc., Perkin Trans. 2 1972, 861.

^{(17) (}a) Quillinan, A. J.; Scheinmann, F. Chem. Commun. 1971, 966–967. (b) Tisdale, E. J.; Chowdhury, C. Vong, B. G.; Li, H.; Theodorakis, E. A. Org. Lett. 2002, 4, 909–912.

⁽¹⁸⁾ See the Supporting Information for crystallographic data of compounds 19a', 46', 49b, 19b', 17, 58, and 2.

Scheme 6. Construction of Key Building Block 20ba



 a (a) K_2CO_3 (5.0 equiv), MeI (10 equiv), DMF, 25 °C, 16 h, 99%; (b) 10% Pd/C (10 wt %), H_2 (1 atm), EtOAc, 25 °C, 45 min, 98%; (c) ′BuOK (2.2 equiv), THF, 0 °C; then reaction mixture concentrated and suspended in MeCN; then 18-Crown-6 (2.2 equiv), 15 min, bromoisobutyraldehyde (5.0 equiv), $0 \rightarrow 25$ °C, 1 h, 70%; (d) $CH_3P^+Ph_3Br^-$ (3.0 equiv), NaHMDS (3.0 equiv), THF, 0 °C, 1 h, 75%; (e) ′BuOK (1.1 equiv), THF, 0 °C; then reaction mixture concentrated and suspended in MeCN; then 18-Crown-6 (1.1 equiv), 15 min, bromoisobutyraldehyde (5.0 equiv), $0 \rightarrow 25$ °C, 1 h, 75%; (f) $CH_3P^+Ph_3Br^-$ (2.0 equiv), NaHMDS (2.0 equiv), THF, 0 °C, 1 h, 80%.

obtained upon exposure to "PrSH—Yb(OTf)₃. ¹⁹ Apparently, the rearranged keto-triol **46**′ is formed by the Lewis acid-induced α-ketol rearrangement²⁰ of the parent triol **46**. Although these two triols were chromatographically unseparable, the rearranged compound (**46**′) crystallized preferentially from an ether/hexane solution of the mixture, thus enabling its X-ray crystallographic analysis ¹⁸ (see ORTEP drawing of **46**′, Scheme 5).

In the face of this rather unexpected circumstance, we decided to target lateriflorone's 1-O-methyl derivative (2) to avoid the complications arising from the deacetonization step. We, therefore, adopted the methoxy derivative 20b (Scheme 6) as the substrate for the Claisen/Diels—Alder cascade sequence. Its construction proceeded along lines similar to those already described above for compound 20a and is summarized in Scheme 6. Thus, phenolic compound 40 was methylated (K₂-CO₃, MeI) to afford methoxy derivative 47 (99% yield), and

the latter compound was debenzylated (H₂, 10% Pd/C, 98% yield), leading to bis-phenol **48**. This compound was found to be quite sensitive on standing and was, therefore, taken immediately to the next step which involved generation of the di-potassium salt (KO/Bu-18-Crown-6) followed by quenching with bromoisobutyraldehyde to afford a mixture of regioisomeric lactols (**49a**:**49b**, ca. 1:1 ratio, 70% yield). These two lactols were separated by chromatography, allowing the X-ray crystallographic analysis¹⁸ of the one that crystallized (**49b**) from its ether—hexane solution (see ORTEP drawing of **49b**, Scheme 6). Each of the two lactols (**49a** and **49b**) was subjected to Wittig olefination (Ph₃P=CH₂) to afford the corresponding phenolic olefin (**50a** and **50b**) in 70% yield. Reiteration of the last two-step sequence furnished the targeted di-olefin **20b**, via **51a** and **51b** (60% overall yield).

Upon heating in DMF at 120 °C for 1 h, the methoxy derivative 20b entered into the expected Claisen Diels-Alder cascade channel, leading to the indicated intermediate products 19b and 19b' in 47% and 42% yields, respectively (see Scheme 7). An X-ray crystallographic analysis 18 of 19b' revealed its structure, and by extension that of 19b. Both structures 19b and 19b' were also supported by nOe studies. Selective removal of the acetonide group from 19b was achieved by treatment of **20b** with catalytic amounts of p-TsOH in methanol, furnishing diol 52 in 98% yield. A two-step oxidation protocol (DMP;²¹ NaClO₂²²) then was employed to convert diol **52** to the required hydroxy carboxylic acid 54 in 93% overall yield via intermediate aldehyde 53. The crucial coupling of carboxylic acid 54 with phenol 18 (see Scheme 3) was brought about by EDC and 4-DMAP, leading to advanced intermediate ester 17 in 64% yield. Crystalline 17 yielded to X-ray crystallographic analysis (see ORTEP drawing of 17, Scheme 7).

4. Final Stages of the Synthesis. Having constructed the ester bridge between the two domains, the next task called for oxidation of the molecule's aromatic nucleus to a quinone and ring closure to lateriflorone's spirolactone skeleton. To this end, compound **17** was exposed to the action of 0.25 N HCl in MeOH:Et₂O (1:1) solution, leading to a spontaneously equilibrating mixture²³ of phenolic esters (**55a:55b**, ca. 1:1, 96% yield) (see Scheme 8). Careful separation of the two components of this mixture by HPLC revealed the rapid equilibration of each back to the original ca. 1:1 composition. It was, however, decided to proceed to the next step in the hopes that at least some of the desired oxidation product, or even the targeted lateriflorone derivative, might result. Oxidation of this mixture (**55a:55b**) under a variety of conditions²⁴ did not lead, however, to the desired outcome, but rather to an array of other products,

⁽¹⁹⁾ Nicolaou, K. C.; Veale, C. A.; Hwang, C.-K.; Hutchinson, J.; Prasad, C. V. C.; Ogilvie, W. W. Angew. Chem., Int. Ed. Engl. 1991, 30, 299-303.
(20) Paquette, L. A.; Hofferberth, J. E. Org. React. 2003, 62, 477-567.

^{(21) (}a) Dess, D. R.; Martin, J. C. J. Org. Chem. 1983, 48, 4155–4156. (b) Meyer, S. D.; Schreiber, S. L. J. Org. Chem. 1994, 59, 7549–7552.

^{(22) (}a) Lindgren, B. O.; Nilsson, T. Acta Chem. Scand. 1973, 27, 888. (b) Kraus, G. A.; Taschner, M. J. Org. Chem. 1980, 45, 1175–1176. (c) Kraus, G. A.; Roth, B. J. Org. Chem. 1980, 45, 4825–4830.

 ^{(23) (}a) Ihara, M.; Nakajima, S.; Hisaka, A.; Tsuchiya, Y.; Sakuma, Y.; Suzuki, H.; Kitani, K.; Yano, M. J. Pharm. Sci. 1990, 79, 703-708. (b) Sidelmann, U. G.; Hansen, S. H.; Gavaghan, C.; Carless, H. A. J.; Lindon, J. C.; Farrant, R. D.; Wilson, I. D.; Nicholson, J. K. Anal. Chem. 1996, 68, 2564-2572.

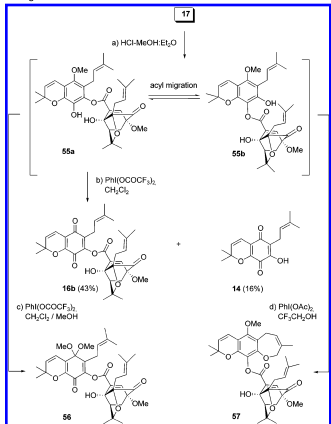
⁽²⁴⁾ For related applications of hypervalent iodine reagents to dearomatize benzenoid systems, see: (a) Varvoglis, A. Tetrahedron 1997, 53, 1179–1255. (b) Kita, Y.; Tohma, H.; Kikuchi, K.; Inagaki, M.; Yahura, T. J. Org. Chem. 1991, 56, 435–438. (c) Scheffler, G.; Seike, H.; Sorensen, E. J. Angew. Chem., Int. Ed. 2000, 39, 4593–4596. (d) Pelter, A.; Ward, R. S. Tetrahedron 2001, 57, 273–282. (e) Tohma, H.; Morioka, H.; Takizawa, S.; Arisawa, M.; Kita, Y. Tetrahedron 2001, 57, 345–352. (f) Quideau, S.; Pouységu, L.; Oxoby, M.; Looney, M. A. Tetrahedron 2001, 57, 319–329. (g) Canesi, S.; Belmont, P.; Bouchu, D.; Rousset, L.; Ciufolini, M. A. Tetrahedron Lett. 2002, 43, 5193–5195.

Scheme 7. Synthesis of Advanced Intermediate 17a

^a (a) DMF, 120 °C, 1 h, **19b** (47%) and **19b'** (42%); (b) *p*-TsOH (20 mol %), MeOH, 25 °C, 16 h, 98%; (c) DMP (2.0 equiv), NaHcO₃ (2.0 equiv), CH₂Cl₂, 25 °C, 0.5 h, 93%; (d) NaClO₂ (6.0 equiv), NaH₂PO₄ (6.0 equiv), 2-methyl-2-butene (75.0 equiv), THF:'BuOH:H₂O (2:4:1), 0.5 h, 100%; (e) EDC (1.5 equiv), 4-DMAP (1.5 equiv), **18** (2.0 equiv), CH₂Cl₂, 0 to 25 °C, 16 h, 64%. DMP = Dess−Martin periodinane, EDC = 1-ethyl-(3-dimethylaminopropyl)carbodiimide hydrochloride.

among which the most interesting are those shown in Scheme 8. Thus, exposure to PhI(OAc)₂ in CF₃CH₂OH afforded a complex mixture from which was isolated novel benzo-oxepene ring system **57** (15% yield), presumably formed from **55b** via radical chemistry.²⁵ On the other hand, reaction of **55a:55b** with PhI(OCOCF₃)₂ in MeOH:CH₂Cl₂ (1:1) led to benzoquinone monoketal **56** (60% yield) through the participation of a molecule of methanol.²⁶ Hydroxyquinones **16b** (43% yield) and

Scheme 8. Oxidation of PhenoIs **55ab** with Hypervalent Iodine Reagents^a



 a (a) 0.25 M HCl in MeOH:ether (1:1), 0 → 25 °C, 1 h, 96%; (b) PhI(OCOCF₃)₂ (1.2 equiv), CH₂Cl₂, -78 → 25 °C, 2 h, **16b** (43%) and **14** (16%); (c) PhI(OCOCF₃)₂ (1.2 equiv), CH₂Cl₂:MeOH (1:1), -78 → 25 °C, 2 h, 60%; (d) PhI(OAc)₂ (1.2 equiv), pyridine (cat.), CF₃CH₂OH, -20 → 0 °C, 30 min, 15%.

 14^{9b} (16% yield) were obtained as the only characterizable products from 55a:55b upon treatment with PhI(OCOCF₃)₂ in CH₂Cl₂ (see Scheme 8).

The failure to obtain the lateriflorone structure directly from the oxidation of **55a:55b** did not end the chase for lateriflorone, because the newly formed hydroxyquinone **16b** held considerable promise as a potential precursor to the coveted architecture. Toward this goal, several attempts were made, including protocols involving acidic conditions (e.g., PPTS, *p*-TsOH, TFA, amberlyst 15), basic conditions (e.g., Et₃N, DBU, LiHMDS, NaH), sealed tube high temperature (180 °C), and high pressure (100 psi) treatment in xylene under microwave irradiation, as well as silica supported (both acidic and basic) microwave irradiation. In all cases, decomposition and/or recovery of starting material was observed with no evidence of spiroxalactone formation.

At this juncture, we reasoned that the poor nucleophilicity of the tertiary hydroxyl group was responsible for the failure to obtain the desired spiroxalactone moiety from hydroxyquinone **16b**, and we proceeded to design a more electrophilic Michael acceptor to enhance the chances for ring closure involving the same tertiary hydroxyl group. Toward this purpose, hydroxydimethyl ketal **56** was treated with PPTS in refluxing benzene, and, according to our expectation, spiroxalactone compound **58**

⁽²⁵⁾ For a precedent of radical generation with PhI(OAc)₂, see: Dorta, R. L.; Francisco, C. G.; Hernández, R.; Salazar, J. A.; Suárez, E. J. Chem. Res., Synop. 1990, 240–241.

⁽²⁶⁾ Corey, E. J.; Wu, L. I. J. Am. Chem. Soc. 1993, 115, 9327-9328.

Scheme 9. Synthesis of 1-O-Methyllateriflorone (2)a

 a (a) PPTS (1.0 equiv), benzene, reflux, 4 h, **58** (69%) and **16b** (11%); (b) 1 N aqueous HCl, THF, 0 → 25 °C, 16 h, 80%; (c) excess CH₂N₂, ether, 0 °C, 0.5 h, 81%; (d) PPTS (1.0 equiv), benzene, reflux, 4 h, 83%. PPTS = pyridinium p-toluene sulfonate.

was formed in 69% yield, together with quinone **16b** (11% yield) (see Scheme 9). The latter compound might be formed by simple rupture of the ketal moiety prior to cyclization. Compound **58** crystallized in beautiful yellow crystals, whose X-ray crystallographic analysis¹⁸ revealed its lateriflorone-like molecular architecture, including the correct stereochemistry at C-3' (see ORTEP drawing, Scheme 9). From compound **58**, 1-*O*-methyllateriflorone (**2**) was in sight, the two compounds separated only by ketal hydrolysis. While mild acidic conditions employing PPTS or TFA left **58** unchanged, an attempt to hydrolyze the enol methyl ether within **58** employing aqueous

HCl in THF at room temperature led to red quinone ether carboxylic acid **16a** in 80% yield. This acid was derivatized and characterized as its methyl ester **59**, a red colored substance, obtained in 81% yield upon methylation with diazomethane.

Compound **16a** presented yet another opportunity to effect the long-sought ring closure to the lateriflorone framework, the expectation being that acid treatment would initiate the required conjugate addition. Indeed, exposure of **16a** to PPTS in refluxing benzene led to the formation of a single yellow substance whose spectral data were consistent with the expected, lateriflorone-like structure **2**. Also, upon crystallizing fron an ether—hexane

solution in beautiful yellow crystals, this compound yielded to X-ray crystallographic analysis 18 (see ORTEP drawing, Scheme 9), which confirmed beyond doubt its structure (2). Thus, the quest for this unusual molecular architecture was complete. Attempts to remove the methyl group from the C-20 oxygen were not successful, presumably due to the inherent instability of the labile $\alpha\text{-ketol}$ moiety, especially under the Lewis acid conditions employed in the attempts.

The exclusively regio- and stereoselective manner by which both substrates **16a** and **56** cyclize under acid conditions to form the spirolactone framework is noteworthy. In both cases, the reaction may proceed under thermodynamic control, leading to the natural stereochemistry at C-3'. In the case of **16a**, the cyclization leads to what might actually be the thermodynamically most stable configuration at both C-2' and C-3', which also happens to be the natural arrangement at those centers.

5. Synthetic Technology for the Construction of Benzopyrans via Facile 6π Electrocyclizations. From the several attempts to accomplish the required ring closure to the spirolactone fragments, the one involving Et₃N led to the most interesting results, even though it, too, like the others failed to produce the desired outcome. Thus, exposure of 16b to Et₃N in CH₂Cl₂ at room temperature (the red solution turned immediately to yellow) followed by concentration and chromatographic purification led to isolation of compounds 63a: 63b (30%, equilibrium mixture of regioisomers due to internal acyl migration, yellow) and 63c (66%, colorless) (see Scheme 10). To explain the formation of these intriguing structures from **16b**, we propose initial conjugation of **16b** to its two geometrical isomers 60-Z and 60-E, which then follow different reaction paths, leading to the observed products via a series of bond reorganizations, including a 6π electrocyclization^{14b,27} (or a conjugate addition, $62-Z \rightarrow 63a$ and $61-E \rightarrow 63c$) as the key ring-forming process.

The mild conditions and high yield associated with this cascade sequence 28 boded well for its further exploitation to generate benzopyran-type 29 compounds from the corresponding quinonoid systems. As shown in Table 1, this entry into this series of compounds $(64 \rightarrow 67a-c; 65 \rightarrow 68a-c; and 66 \rightarrow 69a-c)$ is quite general and holds promise for the construction of a variety of natural-product-like structures.

6. Biological Investigations. The campaign toward lateriflorone produced a number of key building blocks and advanced intermediates which were considered worthy of biological evaluation. Following the lead of the natural substance, we proceeded to screen such compounds (see Table 2) as cytotoxic agents against ovarian cancer cells 1A9 (parental), A2780, and

(28) For recent reviews on cascade reactions in organic synthesis, see: (a) Nicolaou, K. C.; Snyder, S. A.; Montagnon, T.; Vassilikogiannakis, G. E. Angew. Chem., Int. Ed. 2002, 41, 1668–1698. (b) Nicolaou, K. C.; Montagnon, T.; Snyder, S. A. Chem. Commun. 2003, 551–564.

(30) Skehan, P.; Storeng, R.; Scudiero, D.; Monks, A.; McMahon, J.; Vistica, D.; Warren, J. T.; Bokesch, H.; Kenney, S.; Boyd, M. R. *J. Natl. Cancer Inst.* 1990, 82, 1107–1112.

Scheme 10. Cascade Sequence Leading from Quinone **16b** to Benzopyrans **63a**-**c**^a

^a (a) Et₃N (10 equiv), CH₂Cl₂, 25 °C, 1 h, **63ab** (30%) and **63c** (66%).

AD10 (transformed 1A9, producing drug transporter pgp). The first noteworthy observation was that these compounds are not substrates for pgp because the IC_{50} values are not so different in the two assays as shown in Table 2. Second, it was noted that compound **58** is the most potent of the series, being 4–5 times more potent than 1-O-methyllateriflorone (2). Interestingly, the benzoquinone monoketal **56**, which lacks the spiro-

^{(27) (}a) Büchi, G.; Yang, N. C. J. Am. Chem. Soc. 1957, 79, 2318–2323. (b) Becker, R. S.; Michl, J. J. Am. Chem. Soc. 1966, 88, 5931–5933. (c) Rodríguez-Otero, J. J. Org. Chem. 1999, 64, 6842–6848. (d) Li, C.; Johnson, R. P.; Porco, J. A., Jr. J. Am. Chem. Soc. 2003, 125, 5095–5106. (e) Malerich, J. P.; Trauner, D. J. Am. Chem. Soc. 2003, 125, 9554–9555.

⁽²⁹⁾ For recent advances in benzopyran synthesis, see: (a) Nicolaou, K. C.; Pfefferkorn, J. A.; Roecker, A. J.; Cao, G.-Q.; Barluenga, S.; Mitchell, H. J. J. Am. Chem. Soc. 2000, 122, 9939-9953. (b) Nicolaou, K. C.; Pfefferkorn, J. A.; Mitchell, H. J.; Roecker, A. J.; Barluenga, S.; Cao, Q.; Affleck, R. L.; Lillig, J. E. J. Am. Chem. Soc. 2000, 122, 9954-9967. (c) Nicolaou, K. C.; Pfefferkorn, J. A.; Barluenga, S.; Mitchell, H. J.; Roecker, A. J.; Cao, G.-Q. J. Am. Chem. Soc. 2000, 122, 9968-9976.

Table 1. Synthesis of Benzopyran System via 6π Electrocyclizations^a

Entry	Substrate	Product(s)	Yield (%)
1 >	OAc	OR^{2} OR^{1} 67a : $R^{1} = Ac$, $R^{2} = H$	
	64	67b : $R^1 = H$, $R^2 = Ac$	36 ^b
		OH OAc	54
2 >	OPiv OFiv	67c OR ² OR ¹ 68a: R ¹ = Piv, R ² = H 68b: R ¹ = H, R ² = Piv	43°
		OH OPiv 68c	52
з >	O OR	O OR ²	
	66	69a : $R^1 = R$, $R^2 = H$ 69b : $R^1 = H$, $R^2 = R$	41 ^d
R	= MeO'. OMe	OH OR 69c	50

 a Reactions were carried out on 1.0 mmol scale in dichloromethane with 10 equiv of Et₃N at room temperature for 1 h. b 2:1 equilibrium mixture. c 1:1 equilibrium mixture. d 10:1 equilibrium mixture.

lactone moiety, exhibited approximately 2.5 times the activity of 2. It is also of interest to note that the seco-analogues (open forms) 16a and 16b were found to be more or less equipotent to their cyclized counterpart, compound 2. Finally, the two

Table 2. Cytotoxicity of Selected Compounds against 1A9, A2780, and AD10 Human Carcinoma Cells^a

entry	compounds	1A9 (IC ₅₀ in μM)	A2780/AD10 (pgp expresser)
1	52	150	150
2	54	87	136
3	17	63	99
4	57	28	63
5	18	56	81
6	14	96	104
7	16b	17	25
8	16a	37	49
9	56	9	10
10	58	5	5
11	59	17	23
12	2	25	37

^a The antiproliferative effects of the tested compounds against the parental 1A9 and resistant clones (A2780 and AD10) were determined in a 72 h growth inhibition exposure using the SRB (sulforhodamine-B) assay.³⁰

precursor domains of lateriflorone, building blocks chromene quinone 14 and hydroxy acid 54, showed poor activity against these cell lines.

Conclusion

Described herein is a convergent strategy for the synthesis of the unprecedented and highly unusual structure of 1-Omethyllateriflorone (2). The synthetic journey to this target required several redesigns and attempts to cast the final C-O bond of the novel spirolactone moiety. The successful final fusion turned out to be that involving the carboxylate residue, and not the tertiary alcohol, a finding that may bear on the biosynthetic pathway through which nature is generating this structure. On the way to the final destination, we also uncovered a number of interesting cascade sequences to complex benzopyran systems involving facile 6π electrocyclizations which may find applications in complex molecule constructions. We were also able to produce and biologically evaluate a small library of lateriflorone analogues and related systems. These chemical biology investigations established the first structure activity relationships within this class of compounds.

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Supporting Information Available: Experimental procedures, compound characterization, and selected ¹H and ¹³C NMR spectral data (PDF and CIF). This material is available free of charge via the Internet at http://pubs.acs.org.

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