Total Synthesis of *seco*-Plakortolide E and (–)-*ent*-Plakortolide I: Absolute Configurational Revision of Natural Plakortolide I

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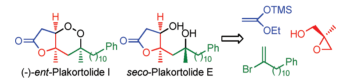
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ABSTRACT



A first total synthesis of (-)-ent-plakortolide I and seco-plakortolide E was accomplished from (S)-2-methylglycidol. The relevant key reactions involve a diastereoselective Mukaiyama aldol reaction, a regioselective hydroperoxysilylation, and elaboration of the 1,2-dioxane ring by intramolecular Michael addition of a hydroperoxide group to a butenolide. This synthesis allowed the revision of the absolute configuration of plakortolide E.

Marine sponges of genus *Plakortis* and *Plakinastrella* are a prolific source of cyclic peroxides, with the majority possessing a 1,2-dioxan ring system.¹ Plakortolides are a family of secondary metabolites found in these sponges which have in common an aromatic unit connected via a methylene chain to a 4,6-dimethyl peroxylactone ring.² They differ in absolute configuration at C-3, C-4, C-6; the substitution pattern; the level of unsaturation; and the chain length. Plakortolide **1** was isolated from the extract

of the Madagascar sponge *Plakortis aff. simplex*. The structure and relative configuration of **1** were established by NMR spectroscopic methods.^{2g}

Its absolute configuration was determined by comparison of the optical rotation with that of its natural enantiomer, with the absolute stereochemistry established by application of Mosher's method to a MTPA derivative.^{2d}

Crews and co-workers reported the isolation from Fidjian sponge *Plakortis sp.* of a new cytotoxic compound plakortolide E.^{2c} Later on, Garson and co-workers found that NMR data reported for plakortolide E were inconsistent with its structure and are likely those of the corresponding *seco*-plakortolide E **2**.²ⁱ

Only one racemic synthesis of plakortolide I has been reported.³ [4 + 2] Photocycloaddition of a singlet oxygen to a diene and iodolactonization are the keys steps for the construction of the peroxylactone framework of plakortolide I. A drawback of this strategy is the lack of stereo-chemical control at the newly formed stereogenic centers.

As a part of an ongoing project devoted to the directing effect of a double bond in the regiocontrol of intramolecular

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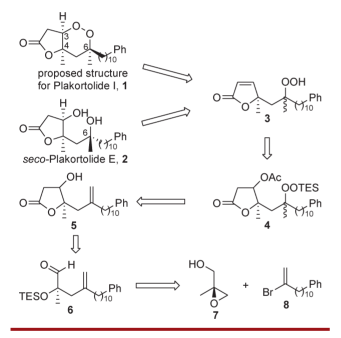
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cyclization of hydroxyl vinyl epoxides,⁴ we have recently reported a study concerning the application of this concept to forge 1,2-dioxane ring systems from β -hydroperoxy vinyl epoxides with the objective of synthesizing plakortolides. Unfortunately, the vinyl group had no directing effect on the regioselectivity of β -hydroperoxy vinyl *cis*-epoxide cyclization and exclusive formation of a 1,2-dioxolane was observed.⁵ Consequently, a new synthetic route was elaborated for the construction of the 4,6-peroxylactone ring system of plakortolides.

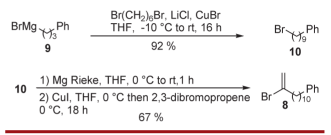
Herein we report the first total synthesis of (-)-*ent*-plakortolide I 1 and *seco*-plakortolide E 2 which differ in absolute stereochemistry only at C-6. In our retrosynthetic analysis we planned to install the 1,2-dioxane ring in the late stage of the synthesis because of its poor stability during functional group elaboration (Scheme 1).

Scheme 1. Retrosynthetic Analysis of Plakortolide I 1 and *seco*-Plakortolide E 2



We envisioned that 1 and diol 2, which is available by the O-O bond cleavage of its corresponding endoperoxide, could arise from a common intermediate 3 via an intramolecular Michael addition of a hydroperoxide group to a butenolide. As in our previous study of plakortolide synthesis,⁵ we selected the regioselective hydroperoxysilylation of alkenes to introduce the peroxide functionality at C-6.⁶ We have shown on a model butenolide bearing a 2-methylallyl substituent that the hydroperoxysilylation was not chemoselective; consequently the double bond of the butenolide moiety has to be protected during the installation of the hydroperoxide group.⁷ We selected the oxotetrahydrofuranyl acetate as a mask form of butenolide because of the inertness of the acetate group in the peroxidation reaction conditions and easy dehydroacetylation in mild basic media. The masked butenolide **4** could originate from the protected hydroxyaldehyde **6** via a Mukaiyama aldol reaction with trimethysilyl ketene acetal and functional group manipulation. We believed that the protected homoallylic alcohol **6** could be prepared by Cu(I)- catalyzed addition of an organometallic derived from the vinyl bromide **8** to the known epoxide **7**.

Scheme 2. Synthesis of the Vinyl Bromide 8 from (3-Phenylpropyl)magnesium Bromide



We first prepared the fragment **8** from (3-phenylpropyl) magnesium bromide following the route in Scheme 2. Cu(I)-Catalyzed coupling between the Grignard reagent **9** and 1,6-dibromohexane (3 equiv) gave 9-phenylnonyl bromide **10** in excellent yield after its separation from the excess of 1,6-dibromohexane by fractional distillation.⁸ A second coupling between (9-phenylnonyl)magnesium bromide⁹ and 2,3-dibromopropene in excess led to the vinyl bromide **8** in moderate yield. We next focused our attention to the addition of organometallics derived from the bromide **8** to the protected 2-methylglycidol **12**, easily obtained in highly enantiomeric purity from 2-methallyl alcohol (Scheme 3).¹⁰

In the first experiment, heating of **8** and an excess of Mg for 30 min followed by addition, at low temperature, of a catalytic amount of CuI and epoxide **12** led to no detectable coupling product (Table 1, entry 1).

Conversely, the desired homoallylic alcohol 13 was obtained in fair yield by prolonged heating of the mixture of Mg and bromide 8 and by and increase of the reaction coupling time (Table 1, entry 2). The reaction conditions reported by Alexakis and co-workers, i.e. addition of the corresponding higher order cuprate of 8 to epoxide 12, in the presence of $BF_3 \cdot Et_2O$, led to decomposition (entry 3).¹¹ Surprisingly, uncatalyzed addition of cuprate

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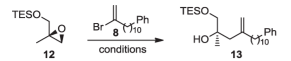
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 Table 1. Cu(I)-Catalyzed Addition of Organometallics Derived from Bromide 8 to Epoxide 12



entry	conditions	yield of $13 \ (\%)^a$
1	8 , Mg (1.5 equiv), THF, Δ, 30 min then 12 , CuI, -40 to 0 °C, 2 h	b
2	8 , Mg (1.5 equiv), THF, Δ , 3 h then 12 , CuI, -30 to 0 °C, 18 h	52
3	8 , <i>t</i> -BuLi (2.1 equiv), Et ₂ O, -78 °C, CuCN (0.5 equiv) then 12 , BF ₃ · Et ₂ O, -78° to 0 °C, 1 h	_c
4	8 , <i>t</i> -BuLi (2.1 equiv), Et ₂ O, -78 °C, CuCN (0.5 equiv) then 12 , -25 to 0 °C, 1 h	70

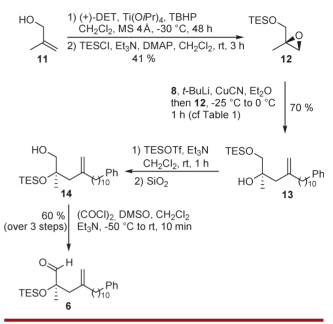
 a Isolated yield. b No metal-bromine exchange was observed. c Decomposition.

derived from **8** to **12** afforded the tertiary alcohol **13** in good yield (entry 4).

Prior to the Mukaiyama aldol reaction, we had to transform the monoprotected diol 13 to the desired α - triethylsilvloxyaldehyde 6. After protection of the tertiary alcohol within 13 as a TES ether, the resulting bis-TES ether was subjected to a selective oxidative deprotection under standard Swern conditions. Conversely to literature precedents, no reaction occurred.¹² Consequently, we adopted a stepwise method which consisted of selective deprotection of the primary TES ether with silica gel¹³ followed by Swern oxidation to provide aldehyde 6 in 60% over three steps from 13. With 6 in hand, the stage was set up for the Mukaiyama aldol addition. After some experimentation, we found that treatment of aldehyde 6 at -78 °C with [(1-ethoxyethenyl)oxy]trimethylsilane 15, in the presence of $TiCl_2(OiPr)_2$,¹⁴ led to the exclusive formation of the syn-adduct 16¹⁵ (Scheme 4).

Exposure of 16 to an excess of TBAF effected TMS and TES ether deprotection and subsequent lactonization. Acetylation of the resulting β -hydroxylactone furnished the acetate 17 in 59% yield over three steps to form 6.

Scheme 3. Preparation of Key Aldehyde 6 from 2-Methallyl Alcohol



Regioselective hydroperoxysilylation of the disubstituted olefin within **17** accomplished by using $Co(thd)_2$,¹⁶ in the presence of oxygen and triethylsilane, afforded the protected hydroperoxide in almost quantitative yield. Finally treatment of **4** at low temperature with DBU gave, via a three-step sequence, compounds **1** and **20** (58%) along with the intermediate **19** and the epoxyalcohol **18**. The formation of epoxide byproduct in the intramolecular Michael addition of the hydroperoxide groups is wellprecedented.¹⁷

Exposure of **4** to DBU at 0 °C which effected β -elimination followed by successive addition of trifluoroethanol and TBAF minimized the epoxide formation providing **1** and **20** in 72% yield, separated by preparative TLC. Structures of **1** and **20** and their relative configurations were confirmed by 2D NMR experiments (HSQC, NOESY).

The synthetic sample of **1** was found to be identical in all respects to the natural product except for the sign of the optical rotation (Lit. values: $[\alpha]_D^{20} + 8 (c \ 0.017, CHCl_3);^{2g}$ for the enantiomer: $[\alpha]_D^{20} - 8 (c \ 0.05, CHCl_3);^{2d}$ observed: $[\alpha]_D^{20} - 9 (c \ 0.7, CHCl_3)$. Since our synthetic route from (*S*)-2-methylglycidol was unambiguous, Kashman and co-workers misassigned the absolute configuration of plakortolide I. Thus, the revised absolute configuration of plakortolide I is 3*S*, 4*S*, 6*R*.

In order to confirm Garson's assumption that the structure of "plakortolide E" was in fact that of its corresponding

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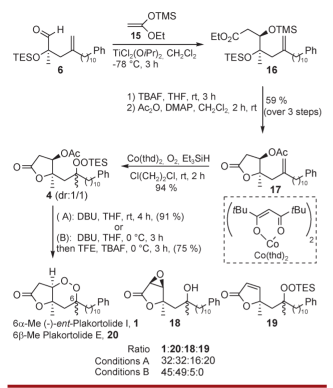
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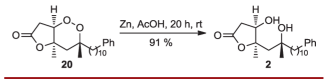
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Scheme 4. Completion of the Synthesis of Plakortolides I and E from Aldehyde 6



seco-plakortolide $\mathbf{2}$, the peroxy ring of $\mathbf{20}$ was reductively cleaved, using Zn/AcOH, to afford $\mathbf{2}$ in excellent yield (Scheme 5).

Scheme 5. Reductive Cleavage of the Peroxy Ring of 20



NMR data of synthetic **2** were in perfect agreement with those of "plakortolide E" reported by Crews and coworkers as well as its specific rotation ($[\alpha]_D^{20}$ +7.7 (*c* 0.15, CHCl₃); Lit.^{2c} $[\alpha]_D^{20}$ +10 (*c* 0.09, CHCl₃)).

In conclusion, we have achieved the first total synthesis of the natural *ent*-plakortolide I **1** and *seco*-plakortolide E **2** leading to the structural revision for plakortolide E to structure **2** and revision of the absolute configuration of **1**. The use of the regioselective hydroperoxysilylation of a *gem*-disubstituted olefin coupled to a high-yielding Mukaiyama aldol reaction allowed rapid construction of the peroxylactone core of plakortolides. Synthesis of **1** and **2** comprises respectively 11 and 12 steps starting from 2-methallyl alcohol with 3.5% and 3.3% overall yields.

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Supporting Information Available. Experimental procedures and full spectroscopic data for all new compounds. This material is available free of charge via the Internet at http://pubs.acs.org.