

244. A Short and Efficient Synthesis of ($\pm$)-Modhephene by a Stereoelectronically-Controlled Ene-Reaction

Preliminary Communication¹⁾

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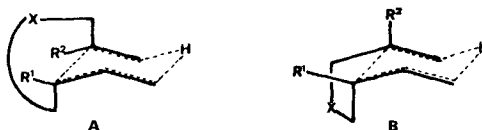
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Summary

($\pm$)-Modhephene (**6**) has been synthesized from the easily available trimethylpentalenone **1** in 6 steps in 26% overall yield (*Scheme 2*). The remarkably smooth 1,4-addition/enolate trapping **1** $\rightarrow$ **2** and subsequent selenoxide elimination after oxidation furnished the key intermediate **3** which underwent an expedient and highly stereoselective intramolecular *ene*-reaction to give the propellane **4**, readily convertible to ($\pm$)-**6**.

In conjunction with a general program exploring the scope and limits of intramolecular *ene*-reactions [1] we have observed previously that 1,6-dienes with the H-donor site *cis* relative to the enophile cyclize thermally to give exclusively 5-membered rings containing *cis*-related H-donor and acceptor sites [2]. These results could be rationalized by inspection of the alternative transition states **A** and **B**; whereas **A** shows a severe angle strain, **B** is perfectly attainable (*Scheme 1*).

Scheme 1



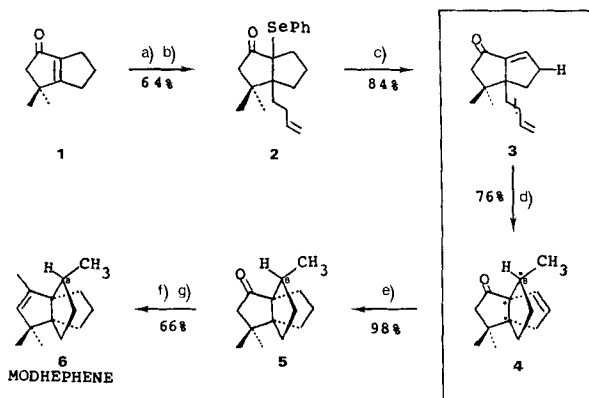
As published in a recent short communication we have exploited these findings for a stereoselective synthesis of the naturally occurring propellane modhephene (**6**) [3]. Modhephene, isolated by Zalkow *et al.* from rayless goldenrod (*Isocoma wrightii*) [4] and by Bohlmann *et al.* from South African *Berkheya* species [5] has attracted considerable attention as a target for synthesis. However all synthetic

¹⁾ Presented by one of us (*W.O.*) at the First International Conference on Chemistry and Biotechnology of Biologically Active Natural Products, Varna, Bulgaria, September 1981.

approaches to **6** so far are either non-stereoselective (requiring GC.-separation of intermediates) [6] [7] or lengthy [3] [8]²⁾.

In contrast, we report here a direct and highly stereocontrolled synthesis of ($\pm$)-modhephene (**6**) (Scheme 2).

Scheme 2



a) $\text{CH}_2=\text{CH}(\text{CH}_2)_2\text{MgBr}$ (30 mmol), THF (43 ml), $\text{CuBr} \cdot \text{DMS}$ (7.5 mmol), DMS (20 ml), -78° ; addition of **1** (15 mmol) in THF (15 ml) over 8 h at -78° , then $\rightarrow \text{RT.}$ over 16 h. b) PhSeBr (20 mmol), THF/HMPA 3:1 (10 ml), 0° , 10 min, RT. , 30 min. c) 5% aq. H_2O_2 -solution (10 mol-equiv.), Py (3 mol-equiv.), CH_2Cl_2 , RT. , 1.5 h. d) 1% solution of **3** in toluene at 250° , 16 h, sealed pyrex tube. e) H_2 (3 atm.), 10% Pd/C, EtOH, RT. , 12 h. f) $\text{H}_2\text{C}=\text{P}(\text{Ph})_3$ (6 mol-equiv.) THF, 90° , 6 h. g) TsOH , CH_2Cl_2 , 25° , 6 h.

The pentalenone **1**, readily available from cyclopentene (60% yield)³⁾ underwent a surprisingly smooth conjugate addition of butenylmagnesium bromide in the presence of freshly crystallized $\text{CuBr}/\text{Me}_2\text{S}$ -complex⁴⁾.

Subsequent enolate-trapping with freshly prepared phenylselenenyl bromide [11] furnished the crystalline selenide **2**⁵⁾ (m.p. $64-70^\circ$) in 64% yield. Oxidation of **2** with H_2O_2 -solution/pyridine led to a selenoxide elimination giving the key precursor **3**⁵⁾ in 84% yield.

The stage was now set for the decisive *ene*-reaction. Heating a solution of **3** in toluene at 250° for 16 h using a sealed, pre-dried Pyrex tube gave the desired [3.3.3]-

²⁾ Thus, our previous synthesis of ($\pm$)-**6** suffers from the need to introduce the *gem*-dimethyl group at an advanced stage [3].

³⁾ The *Friedel-Crafts* acylation of cyclopentene with 3,3-dimethylacryloyl chloride [9] (1 mol-equiv. of AlCl_3 , CH_2Cl_2) was greatly improved when carried out at -78° . Heating the crude reaction mixture with NEt_3 furnished 1-(1-cyclopentenyl)-3-methyl-2-buten-1-one (79% yield) which on *Nazarov* cyclization (SnCl_4) [10] furnished the pentalenone **1**.

⁴⁾ Cuprate addition to the hindered $\text{C}(\beta)$ of **1** previously required promotion by borontrifluoride etherate [8]. Furthermore, numerous attempts have failed to achieve the conjugate addition of butenylcuprates to 4,4-dimethyl-3-[2-(1,3-dioxolan-2-yl)ethyl]-2-cyclopentenone [3].

⁵⁾ All new compounds were characterized by IR., $^1\text{H-NMR}$. (360 MHz) and mass spectroscopy. Their purity was confirmed by GC. analysis.

propellane **4**⁵) in 76% yield. According to our prediction the relative configuration of C(8) with the methyl group pointing towards the cyclopentene bridge has been nicely controlled in the cyclization step⁶).

Catalytic hydrogenation of **4** furnished the ketone **5**⁵) (98% yield), identified by comparison (IR., ¹H-NMR. and MS.) with a sample of **5** prepared previously by Karpf & Dreiding [6]. Modifying the published procedure [6] ketone **5** was successively treated with methyldiene triphenylphosphorane and *p*-toluenesulfonic acid in CH₂Cl₂ at 25° to give (±)-modhephene (**6**) showing IR., ¹H-NMR. and mass spectra identical to those of the natural sesquiterpene.

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⁶) The configuration of **4** was established by its conversion to **5** and **6**. Two not yet fully characterized minor compounds (10% and 8% yield, respectively) were separated from **4** by simple chromatography (SiO₂, hexane/ether 40:1).