

Total Synthesis of Albolic Acid and Ceroplastol II, 5–8–5-Membered Tricyclic Insect Sesterterpenoids, via a Lactol-Regulated Silyloxy–Cope Rearrangement

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Optically active albolic acid and ceroplastol II, 5–8–5-membered tricyclic sesterterpenoids, were stereoselectively synthesised from two C₁₀ synthons (iridoids) via CrCl₂-condensation, lactol-regulated silyloxy–Cope rearrangement with a normally disfavoured boat transition geometry, TiCl₂-ring closure, and C₅-homologation.

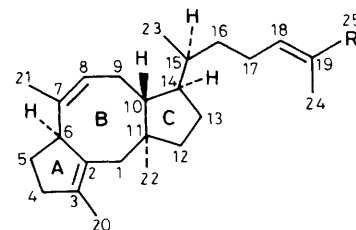
Albolic acid (**1**)¹ and ceroplastol II (**2**)² are sesterterpenoids isolated from the wax secretion of *Ceroplastes albolineatus*, a scale insect. The total syntheses of (**1**) and (**2**) are described herein.³

The carbon framework of (**1**) can be constructed by the CrCl₂-condensation of two iridoid synthons,^{4–6} a Cope rearrangement, and intramolecular eight-membered ring formation; the required stereochemistry of (**1**) at C-6 and C-14 can be transferred from (3*R*)-irida-1,8-dien-7-al (**3**) and (3*S*,8*R*)-9-benzyloxy-7-chloro-1-iridene (**4**) to the expected condensate (**5**).[†]

However, the usual silyloxy–Cope rearrangement of compounds like (**5**) is known to give products having the opposite configuration at the C-11 methyl to that required, through a chair transition state.⁴ Therefore, it is essential to perform the

Cope rearrangement via the boat transition state, and this has been achieved by means of a 'lactol-regulated Cope rearrangement'⁷ where a lactol formed in the 1,5-diene system precludes formation of the *Z*-type enol ether via the chair transition state.

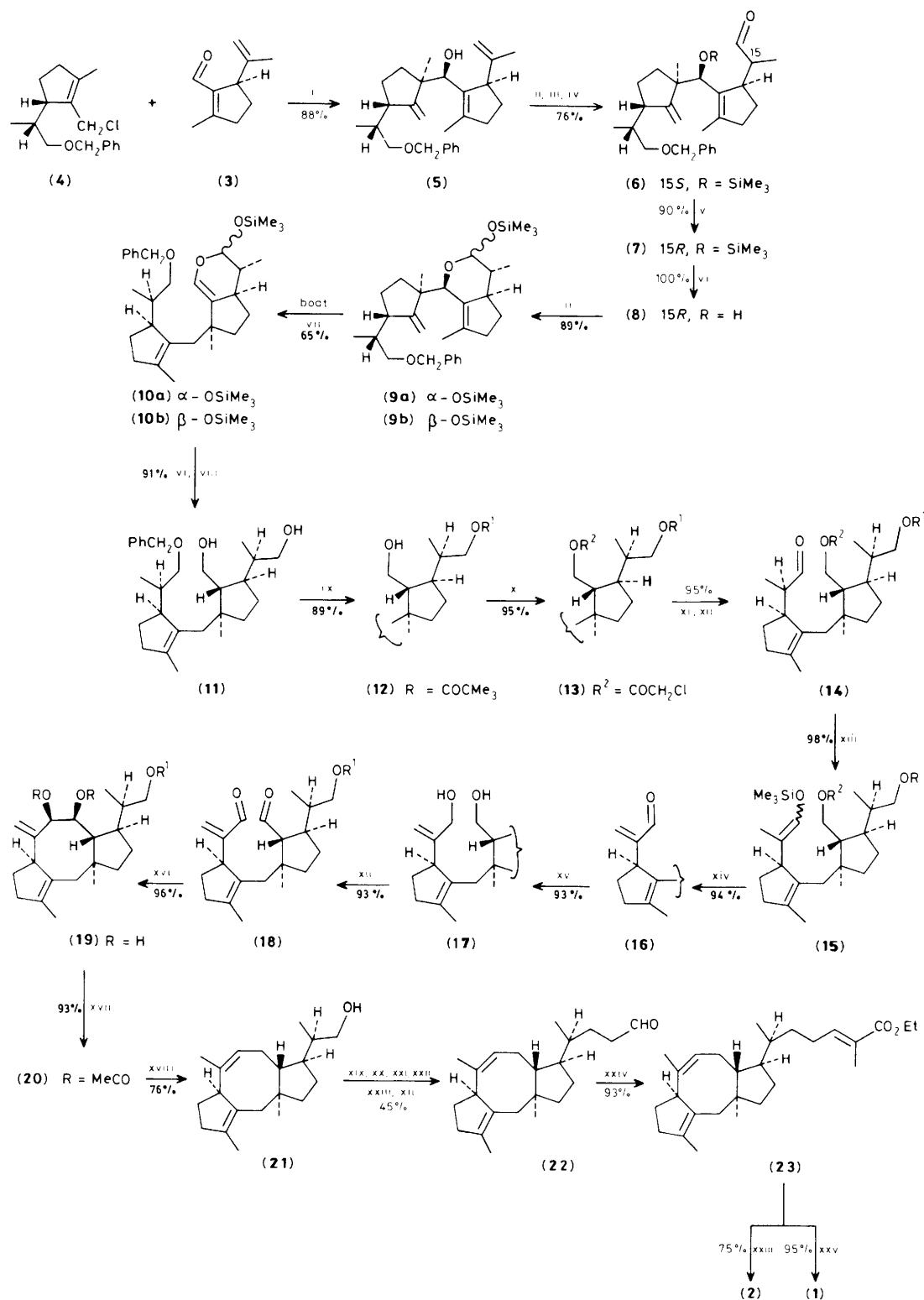
Compound (**5**) was obtained from (**3**) and (**4**) in 88% yield by a CrCl₂-condensation in the presence of PrⁱOH, the yield of



(**1**) R = CO₂H

(**2**) R = CH₂OH

[†] The new compounds described in this paper showed satisfactory elemental analyses, together with structure-consistent spectroscopic data.



Scheme 1. Reagents and conditions: i, CrCl_2 , tetrahydrofuran (THF), dimethylformamide (DMF), Pr^iOH ; ii, Me_3SiCl /pyridine (Py); iii, $[\text{Me}_2\text{CHCH}(\text{Me})_2\text{BH}]_2\text{H}_2\text{O}_2/\text{OH}^-$; iv, pyridinium chlorochromate/ CH_2Cl_2 ; v, KF-Florisil/ MeOH ; vi, pyridinium toluene-*p*-sulphonate/aq. THF; vii, 190°C , C_7H_8 ; viii, $\text{NaBH}_4/\text{aq. NaHCO}_3$, MeOH ; ix, Bu^iCOCl , Py; x, ClCH_2COCl /Py; xi, H_2 /Pd/C; xii, $(\text{COCl})_2$ /dimethyl sulphoxide (DMSO), Et_3N , CH_2Cl_2 ; xiii, $\text{CF}_3\text{SO}_3\text{SiMe}_3$, Et_3N , CH_2Cl_2 ; xiv, $\text{Pd}(\text{OAc})_2/\text{MeCN}$; xv, NaBH_4 , CeCl_3 , MeOH ; xvi, TiCl_4 , Zn/THF ; xvii, $\text{Ac}_2\text{O}/\text{Py}$; xviii, $\text{Li}/\text{liq. NH}_3$, EtOH ; xix, tosyl chloride/Py; xx, $\text{NaCH}(\text{CO}_2\text{Me})_2/\text{DMF}$; xxi, NaCN/DMSO ; xxii, CH_2N_2 ; xxiii, lithium aluminium hydride/THF; xxiv, $\text{Ph}_3\text{P}=\text{C}(\text{Me})-\text{CO}_2\text{Et}$ /benzene; xxv, 3 M NaOH/MeOH .

the epimeric alcohol was only 3%. Compound (**5**) was regio- and stereo-specifically converted to the silyloxy aldehyde (**6**). The unnatural configuration at C-15⁸ was isomerised using KF-on-Florasil to give the aldehyde (**7**) which gave the hydroxy aldehyde (**8**) on further hydrolysis.

Upon heating at 190 °C (**9a**) and (**9b**), derived from (**8**), underwent a Cope rearrangement without cleavage of the lactol ring system to the thermolysates (**10a**) and (**10b**).[‡] However, heating the lactol trimethyl silyl (TMS) ethers derived from (**6**) under similar conditions failed; this is thought to be due to a severe steric interaction between the A-ring and the axial C-15-Me. Hydrolysis and subsequent NaBH₄-reduction of (**10a**) and (**10b**) produced a single compound (**11**). The ¹³C n.m.r. data of (**11**) were used to ascertain the correct C-ring configuration for (**1**) or (**2**) by comparison with related compounds.^{5,7}

Thus, the Cope rearrangement of (**9**) has proceeded *via* a normally disfavoured boat transition geometry to give the desired product. See Scheme 1 for detailed synthesis leading to (**21**).^{9–11}

The final step of the synthesis was a conventional C₅-elongation of the side chain; *i.e.*, (**21**) was converted to the aldehyde (**22**) and a Wittig reaction of (**22**) with ethyl (2-triphenylphosphoranylidene)propionate gave (*E*)-(**23**), which corresponds to ethyl albolate, as the major product (93% *vs.* 4% of *Z*-isomer); a portion of (**23**) was saponified to the acid (**1**), and the remainder was reduced to the alcohol (**2**). The ¹H n.m.r. spectra of (**1**) and (**2**) measured in CCl₄ were identical with those reported. In addition, the 3,5-dinitroben-

zoate of (**2**) was identical with the sample prepared from the 3,5-dinitrobenzoate of ceroplastol I¹² in every respect.

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- 8 Hydroboration of irida-1,8-diene derivatives always occurs stereoselectively to give (3*S**, 8*R**)-irid-1-en-9-ols without exception. See S. Yamasaki, T. Hatsui, and H. Takeshita, *Kyushu Daigaku Seisan Kagaku Kenkyusho Hokoku*, 1986, **81**, 7. The chemical shifts of Me adjacent to the α-carbon of the aldehydes, δ 13.0 for (**6**) and 7.4 for (**7**), parallel those for dehydroiridodial (δ 10.7) chrysomelidial (δ 7.8).⁶
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[‡] (**10a**) gave another thermolysate in 32% yield which must be a chair transition product judging from the appearance of silylenol ether and an aldehydic proton signals. A 1,3-silyl migration probably occurred prior to Cope rearrangement.