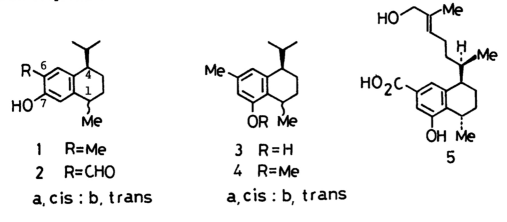
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STEREO- AND REGIOSELECTIVE SYNTHESIS OF HYDROXYCALAMENENES VIA (ARENE)TRICARBONYL CHROMIUM COMPLEXES

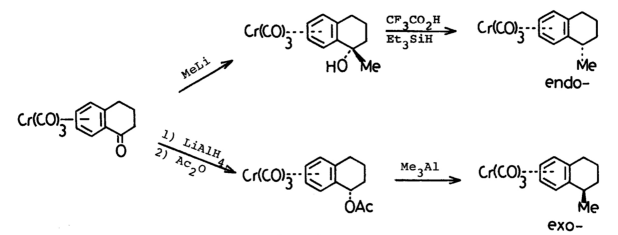
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Stereoisomers of 7- and 8-hydroxycalamenenes were synthesized stereo- and regioselectively via (arene)tricarbonyl chromium complexes.

Phenolic cadinane-type sesquiterpenoids, 7-hydroxy-*cis*-calamenene $(\underline{1a})$,¹⁾ 7hydroxy-*cis*-calamenal $(\underline{2a})$,²⁾ and 8-hydroxy and methoxy-*trans*-calamenene ($\underline{3b}$ and $\underline{4b}$),³⁾ and a prenylated analog, dihydroxyserrulatic acid $(\underline{5})$,⁴⁾ have been isolated from plant origin, and a remarkable fish-poison activity has been shown for the compounds <u>la</u> and <u>3b</u>. For the general synthesis of these terpenoids, it is necessary to introduce the benzylic substituents stereoselectively, and the aromatic substituent regioselectively at the proper position in the terpenoid skeleton. We now wish to report the stereo- and regioselective synthesis of the hydroxycalamenenes by employing stereoselective alkylation at the benzylic position and regioselective functionalization at aromatic part of (arene)tricarbonyl chromium complexes.



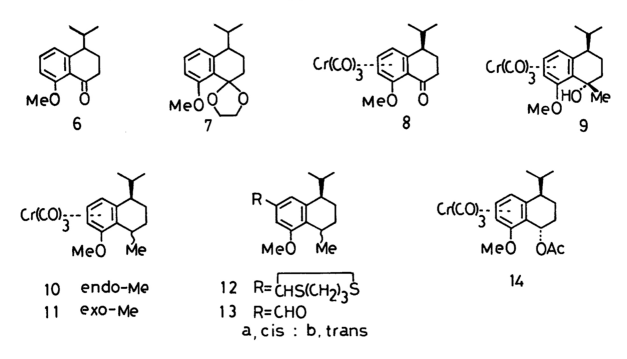
(Arene)tricarbonyl chromium complexes have some characteristic properties due to the strong electron-withdrawing ability and steric bulkiness of $Cr(CO)_3$ group. One consequence is a three dimensional structure of the planar aromatic moiety, which controls stereochemistry on nucleophilic or electrophilic attack at the reactive center of an alicyclic ring. Another one is remarkable stability of the benzylic carbocations.⁵⁾ As a synthetic application of the above two concepts, both isomers of tricarbonyl(1-methyltetralin)chromium were selectively prepared from tricarbonyl(α -tetralone)chromium as follows (Scheme 1), giving the key reactions for the synthesis of these terpenoids.



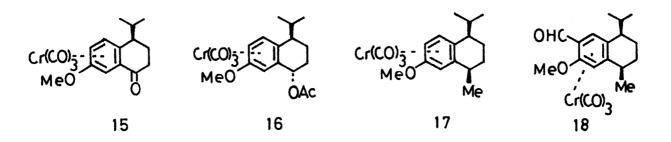


Since attempts to the direct complexation of 4-isopropyl-8-methoxy-1tetralone ($\underline{6}$), obtained from m-methoxybenzaldehyde by several steps, with Cr(CO)₆ gave a complex mixture⁶) under various conditions, the tetralone <u>6</u> was converted into the ethylene acetal derivative <u>7</u>. The acetal <u>7</u> smoothly provided the corresponding (n^6 -arene)tricarbonyl chromium complex under usual thermal conditions and subsequent acid treatment gave the desired (4-*exo*-isopropyl-8-methoxy-1tetralone)Cr(CO)₃ <u>8</u>, (mp 102-103 °C) in 82% overall yield, along with a trace of the *endo*-isomer. This reaction proceeded selectively so that the isopropyl group at the chiral benzylic position was oriented far from the bulky Cr(CO)₃ group, and afforded the *exo*-methylated product <u>9</u>, (mp 138 °C) in 61% yield. Ionic hydrogenolysis⁷ of the benzylic alcohol in the complex <u>9</u> with an excess of triethylsilane and trifloroacetic acid resulted in a stereoselective hydride displacement, giving the (1-*endo*-methyl-4-*exo*-isopropyl-8-methoxytetralin)Cr(CO)₃ (<u>10</u>, mp 120 °C) in 75% yield without stereoisomeric contamination. Nucleophilic addition⁸⁾ of 2lithio-1,3-dithian to the complex <u>10</u> and subsequent oxidative decomplexation of the anionic chromium intermediate with iodine gave 6-dithianylated compound <u>12b</u>, (25% yield) and 6-formylated compound <u>13b</u> (36% yield). trans-8-Methoxycalamenene (<u>4b</u>) was obtained from the compound <u>12b</u> by desulfurization with Raney Ni, and also from <u>13b</u> by hydrogenation with Pd/C, in good yields. Demethylation of the compound <u>4b</u> with BBr₃ afforded trans-8-hydroxycalamenene (<u>3b</u>).

The endo-acetate complex <u>14</u>, prepared from <u>8</u> by reduction and subsequent acetylation, was converted into (1-exo-methyl-4-exo-isopropyl-8-methoxytetralin) $Cr(CO)_3$ (<u>11</u>, mp 106 °C) via stereoselective exo-methyl displacement at the benzylic position by Me₃Al treatment⁹) in 99% yield. The complex <u>11</u> was converted into *cis-8-hydroxycalamenene* (<u>3a</u>) through 6-substituted products <u>12a</u> and <u>13a</u> under the same reaction sequence as described above.



Similarly, the *cis*-7-hydroxycalamenene (<u>la</u>) was synthesized as follows. (4-*exo*-Isopropyl-7-methoxy-1-tetralone)Cr(CO)₃ (<u>15</u>), major cyclization product of [4-(p-methoxyphenyl)-5-methylhexanoyl chloride]Cr(CO)₃ with AlCl₃, was converted into (1-*exo*-methyl-4-*exo*-isopropyl-7-methoxytetralin)Cr(CO)₃ (<u>17</u>, mp 125 °C) through an *endo*-acetate complex <u>16</u> (mp 155 °C) by reduction, acetylation and methylation with Me₃Al in 80% overall yield. Directed lithiation¹⁰⁾ (BuLi, TMEDA, THF, -78 °C, 2 h) of the complex 17 and subsequent quenching with dimethylformamide gave 6-formylated complex <u>18</u> (mp 164 °C) in 83% yield without formation of regioisomeric products. Decomplexation of <u>18</u> (exposure to sunlight), hydrogenolysis with Pd/C and subsequent demethylation with BBr₃ afforded *cis*-7-hydroxycalamenene (<u>1a</u>) in good yield. The stereoselective synthesis of *trans*-7-hydroxycalamenene (<u>1b</u>) from the complex <u>15</u> was already reported.¹¹



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