

170. Two New Sesquiterpenoid Ketones from Cassia Oil

by Alan F. Thomas

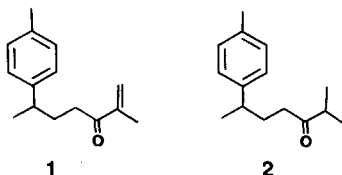
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Summary

The synthesis of 2-methyl-6-(4'-methylphenyl)-1-hepten-3-one (**1**) and 2-methyl-6-(4'-methylphenyl)-3-heptanone (**2**), constituents of the oil of *Cinnamomum cassia*, is described.

Cassia, or Chinese cinnamon oil is a steam distillate of the bark, leaves, and twigs of the tree *Cinnamomum cassia*. The most recent study of the constituents is that of *ter Heide* [1]. We have examined a sample of the commercial oil from the People's Republic of China, and in this paper we report the synthesis of the two sesquiterpene components **1** and **2** which we identified uniquely from their mass spectra¹⁾.

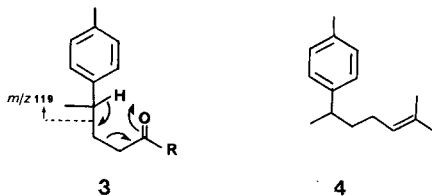


The fraction containing the heavier ketones was isolated from the crude oil by distillation and column chromatography. The fraction with b.p. 115–128°/0.001 Torr (15% of the oil) consisted mainly of (*E*)-2-methoxycinnamaldehyde, a well known constituent; this crystallized, and the mother liquors were then chromatographed on silica gel when a fraction weighing 1.0 g was eluted just after the hydrocarbons. This was a very complex mixture, but gas chromatographic (GLC.) separation of various zones, followed by ¹H-NMR.- and GLC./mass spectrometry (MS.) enabled a number of components to be identified. In zone A (with shortest retention time) issuing from an *SP 1000* GLC. column, tetradecanal and very probably 4,8,12-trimethyltridecanal were identified²⁾. Shortly after, a

- ¹⁾ Cassia oil has a reputation for being regularly adulterated [2]. Our sample contained a fairly high amount of (*Z*)-cinnamaldehyde, and the sesquiterpenes copaene, epiaromadendrene, and spathulenol. These seem unlikely materials for adulteration.
- ²⁾ Chemical Abstracts reports the presence of 4,8,12-trimethyltridecanal in cold-pressed lemon oil [3], but the original paper to which this refers [4] does not mention it. The same aldehyde, together with the ketone 6,10,14-trimethylpentadecan-2-one, is reported to arise during a chemical degradation of α -tocopherol [5], and the ketone has also been found in sediments [6].

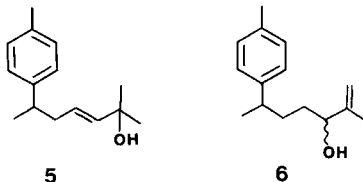
complex zone B contained ketone **2**, together with 1-phenylheptanone. A zone C followed, which contained ketone **1**, followed by a somewhat larger amount of a ketone probably identified as 6, 10, 14-trimethylpentadecan-2-one²).

The ketones **1** and **2** were identified on the basis of their molecular ions, and the fact that both had their most important fragment at m/z 132, corresponding to a *McLafferty* γ -hydrogen shift (**3**). Furthermore, loss of the side chain from the C-atom *a* to the benzene ring leaves the characteristic fragment at m/z 119, the second most important in the MS. of both **1** and **2**.



Ketone **2** has already been prepared in 40% yield by *Watanabe et al.* [7] by a *Friedel-Crafts* reaction between toluene and 2-methyl-6-hepten-3-one, but neither **1** nor **2** has been reported to be naturally occurring.

Our synthesis of **1** and **2** started from ($\pm$)- α -curcumene (**4**), prepared from 4-methylphenylmagnesium bromide and 6-methyl-5-hepten-2-one followed by reduction with sodium in liquid ammonia [8]. Sensitized photooxygenation of **4** yielded the tertiary alcohol ($\pm$)-**5**, and a mixture of the *threo*- and *erythro*-alcohols **6**. Chromatography on silica gel enabled the separation of **5** and **6**, although we found no conditions that separated the two isomers of **6**. The mixture **6** was oxidized with manganese dioxide, and the resulting ketone was identical (MS. and retention time on capillary GLC.) with ketone **1**. Catalytic hydrogenation of **1** gave a ketone identical (MS. and capillary GLC.) with **2**.



It is not often that substances can be identified only by mass spectra, but this work shows that in simple cases it is possible.

Thanks are extended to Mr. *Christian Starkemann* for the workup of the natural oil, and to *Roberto Di Giorgio* for assistance with the synthesis.

Experimental Part

General remarks. NMR. spectra were recorded in CDCl_3 on a *Hitachi-Perkin-Elmer* R 20B (60 MHz), a *Bruker* HX-90 (90 MHz), or a *Bruker* VH-360 (360 MHz) instrument. Chemical shifts are given in ppm downfield from tetramethylsilane ($=0$ ppm), coupling constants J in Hz. Mass spectra were measured either on an *Atlas* CH-4 mass spectrometer using an inlet temperature of ca. 150° , or on a *MAT* 112 instrument coupled with a capillary gas chromatograph, both spectrometers

using electrons of 70 eV. Results are quoted as m/z (% most important fragment), and generally the ten most important fragments are given. Column chromatography was carried out on *Merck H* (Type 60) silica gel using a *Jobin-Yvon* medium pressure preparative chromatograph.

Workup of the oil of Cinnamomum cassia. The commercial oil (2.025 kg) was distilled, first at 0.01 Torr up to 110° through a 20-cm *Vigreux* column, thereby obtaining 1.675 kg of distillate. Further distillation with a 10-cm *Vigreux* column yielded 312 g (15%) of material with b.p. 115–128°/0.001 Torr. A large proportion of this fraction crystallized as (*E*)-2-methoxycinnamaldehyde. The mother liquors (115 g) were chromatographed on 800 g of silica gel in hexane/ether 7:3, thereby separating 28 g of less polar constituents from more 2-methoxycinnamaldehyde eluted just after. Rechromatography of these less polar constituents on 500 g silica gel in hexane/ether 95:5 gave the following fractions: (1) 1.5 g of hydrocarbons, principally *s-guaiazulene*, *rimuene*, and other *diterpenes* $C_{20}H_{32}$; (2) 1.2 g of a complex mixture of *ketones* and *esters*; (3) 11.6 g of (*E*)-cinnamyl acetate; (4) 5.0 g of (*E*)-2-methoxycinnamyl acetate; (5) 5.2 g of a mixture of cinnamaldehyde and (*Z*)-2-methoxycinnamaldehyde [the latter was identified by means of its 1H -NMR. (90 MHz): 3.88 (*s*, 3 H, CH_3O); 6.16 ($d \times d$, $J=8$ and 11, 1 H, $CH=CHCHO$, (*Z*)); 7.76 (d , $J=11$, 1 H, $CH=CHCHO$, (*Z*)); 9.85 (d , $J=8$, 1 H, CHO); an (*E*)-configuration of the double bond implies a coupling constant of 16 Hz, and the aldehyde proton signal at higher field [9]]; (6) 2.5 g of (*E*)-2-methoxycinnamaldehyde.

Fraction (2) was separated into zones by GLC. on an *SP 1000* column, each zone being subsequently rechromatographed on an *OV 17* column. From the first zone (from the *SP 1000* column) were thus identified *tetradecanal* (MS. and 1H -NMR.), and 4,8,12-trimethyltridecanal. - 1H -NMR. (360 MHz): 0.84 (*d*, $J=6$, 3 H), 0.86 (*d*, $J=6$, 6 H), and 0.88 (*d*, $J=6$, 3 H, together 4 CH_3); 1.0–1.6 (*m*, 15 H); 1.66 (*m*, 2 H); 2.42 (*m*, 2 H) and 9.78 (*t*, $J=2$, 1 H, CH_2CH_2CHO). Addition of $Eu(fod)_3$ shifted the *d* at 0.88, and double irradiation clearly established the presence of the $>CHCH_2CH_2CHO$ group. - MS.: 240 (M^+ , trace), 196 ($M^+ - 44$, 5), 85 (48), 81 (40), 71 (73), 57 (100), 69 (60), 56 (70), 55 (58), 43 (98), 41 (58).

The next, highly complex zone from the *SP 1000* column contained 1-phenylheptanone (identified by MS. and retention time of an authentic sample), and ketone 2.

The third zone from the *SP 1000* column contained ketone 1 and many other substances, the main component of the mixture being 6,10,14-trimethylpentadecan-2-one. - 1H -NMR. (90 MHz): 0.87 (*d*, $J=6$, 12 H, 4 CH_3); 1.0–1.8 (*m* of very similar shape to that in the 1H -NMR. of 4,8,12-trimethyltridecanal, *ca.* 17 H); 2.13 (*s*, 3 H, 3 $H-C(1)$); 2.42 (*t*, $J=7$, 2 H, 2 $H-C(3)$). - MS.: 250 ($M^+ - 18$, 5), 85 (22), 71 (47), 69 (25), 59 (43), 58 (90), 57 (43), 55 (30), 43 (100), 41 (25).

The main constituent of fraction (2) was eluted in a later zone from the *SP 1000* column, and was identified as benzyl benzoate (1H -NMR., MS.).

Sensitized photooxygenation of α -curcumen. A solution of 35 g of α -curcumen (4) [8] in 400 ml of methanol containing rose bengale was irradiated while oxygen was bubbled through. After 7 h, 4.4 l of O_2 had been absorbed (theory: 4.33 l at 20°/730 Torr), and the solution was added dropwise to 2 l of ice-cooled aqueous (15%) Na_2SO_3 -solution. After the addition was completed, the mixture was stirred for 15 h, then extracted with pentane. Usual workup gave 35.8 g of crude material, showing approximately equal amounts of two components by GLC. (*Apiezon L*). Of this, 30.2 g were chromatographed in ether/hexane 2:3 over 1 kg of silica gel. There were thus obtained, first, 14.7 g of ($\pm$)-(3*E*)-2-methyl-6-(4'-methylphenyl)-3-hepten-2-ol (5) which was practically pure. For analysis it was collected by GLC. - 1H -NMR. (60 MHz): 1.20 (*s* and *d*, $J=6.5$, 9 H); 2.29 (*s*, superimposed on *m*, 3 H and 2 H); 2.68 (5 bands, 1 H, $H-C(arom.)$); 5.52 (*s*, superimposed on *m*, 2 H); 7.04 (*s*, 4 H). - MS.: 203 (2), 200 (4), 188 (16), 131 (29), 119 (100), 118 (21), 105 (12), 91 (15).

$C_{15}H_{22}O$ (218.36) Calc. C 82.51 H 10.16% Found C 82.38 H 10.20%

The next substance eluted from the silica gel was a mixture of threo- and erythro-($\pm$)-2-methyl-6-(4'-methylphenyl)-1-hepten-3-ol (6) which was collected from GLC. (*Apiezon L*). - 1H -NMR. (60 MHz): 1.22 (*d*, $J=7$, 3 H); 1.63 (br.) superimposed on 1.3–1.8 (*m*, 7 H in all); 2.29 (*s*, 3 H); 2.64 (1H); 4.00 (br., 1 H, $H-C(3)$); 4.75–4.95 (2 H, 2 $H-C(1)$); 7.04 (*s*, 4 H). - MS.: 218 (M^+ , 0.5), 200 (9), 132 (85), 120 (11), 119 (100), 117 (11), 105 (13), 91 (15), 41 (11).

$C_{15}H_{22}O$ (218.36) Calc. C 82.51 H 10.16% Found C 82.40 H 10.15%

Synthesis of ($\pm$)-2-methyl-6-(4'-methylphenyl)-1-hepten-3-one (1). A suspension of manganese dioxide (*Merck*, activated 2 h at 120°), was shaken with 9.3 g of 6 in 500 ml of petroleum ether

(b.p. 50–70°) for 3 h. After filtration, the solution was concentrated and the residue distilled to obtain 6.2 g (68%) of **1**, b.p. 71–74°/0.01 Torr. For analysis the material was purified by GLC. (Apiezon L). – ¹H-NMR. (60 MHz): 1.24 (d, *J* = 6.5, 3 H); 1.83 (d, *J* = 1, 3 H); 2.30 (s, 3 H); 5.67 and 5.78 (each 1 H, 2 H–C(1)); 7.04 (s, 4 H). – MS.: 216 (*M*⁺, 12), 133 (12), 132 (100), 131 (8), 119 (60), 117 (15), 105 (8), 91 (12), 41 (14). The pure substance polymerized within a few h at room temp.

Synthesis of (±)-2-methyl-6-(4'-methylphenyl)-3-heptanone (2). The preceding ketone (**1**, 1.5 g) in 150 ml of ethanol was hydrogenated in the presence of 0.2 g of 10% Pd/C. After 35 min, the theoretical amount of hydrogen had been absorbed. The catalyst was removed by filtration, and the residue was distilled in a bulb tube, bath temp. 90°/0.01 Torr, yield 1.3 g. For analysis, the substance was collected by GLC. (Apiezon L). – ¹H-NMR. (60 MHz): 1.00 (d, *J* = 6, 6 H); 1.22 (d, *J* = 6.5, 3 H); 2.30 (s, 3 H); 7.04 (s, 4 H). – MS.: 218 (*M*⁺, 9), 133 (13), 132 (100), 119 (38), 117 (8), 105 (9), 43 (9).

C₁₅H₂₂O (218.36) Calc. C 82.51 H 10.16% Found C 82.76 H 10.11%

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