Analysis Of The Enantiomers Ratio Of Citronellal From Indonesian Citronella Oil Using Enantioselective Gas Chromatography

Edy Cahyono1*, Harno Dwi Pranowo2, Muchalal2, Tryyono2

1Department of Chemistry, Universitas Negeri Semarang (Unnes), Semarang, Indonesia, 2Department of Chemistry, University of Gadjah Mada, Yogyakarta, Indonesia

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ABSTRACT

Citronellal 97.3% has been isolated from Java citronella oil (Cymbopogon winterianus) from Yogyakarta Indonesia by fractional distillation under reduced pressure (5 cmHg, 110-120 °C). Citronellal has two optical isomerics that can be separated by capillary column of chiral GC phase. Enantioselective capillary GC with heptakis(2,3-di-O-acetyl-6-O-tert-butyldimethylsilyl)β-cyclodextrin (β-DEX-225) as stationary phase has been used for analysis of the enantiomers ratio of citronellal. The analysis of enantiomer ratio showed that citronellal contain of 88.21% ee of (R)-(+) -citronellal. Physical properties of isolated citronellal showed that the compound was (+) enantiomer. Structure identification of citronellal was carried out by GC-MS, IR, and 1H NMR, resulted identical fragment and spectra with standard citronellal. Theoretical study with semiempirical-AM1 method showed that energy of (R)-(+) -citronellal on the β-DEX 225 was lower than its (S)-(−)-citronellal.

1. INTRODUCTION

Citronellal is one of monoterpenes in citronella oil. The citronellal content of high quality oil is 28-45% [1]. Citronellal has two optical isomerics with molecular weight of 154.25. Citronellal is important monoterpenic aldehyde that produces some of the most intense aromas. These it become key aroma impact compounds in many essential oils (EOs). It causes the characteristic lemon-like odor. Citronellal (3,7-dimethyl-6-octenal) occurs as two oils (EOs). It presents at high concentrations (70–80%) in Citronellal is important monoterpene aldehyde which the enantiomerically pure (>97.7%) genuine and 2 commercial lemon balm oils contained the samples. The results showed only 4 samples out of 28 reported in the literature. The most marked difference was observed for lower-concentration genuine samples, for which the (R)-(+) -citronellal content varied from 96.5 to 83.7%. Citronellal in Java-type citronella oil (Cymbopogon winterianus Jowitt) could be biosynthesized either by direct conversion of geraniol to citronellal, via enol formation, or by conversion of geraniol to citronellol and then oxidation of citronellol to citronellal. Obviously, the investigation on the enantiomer distribution of some compounds in authentic and commercial citronella EOs proved that citronellal in these EOs was produced by oxidation of citronellol-amounts of (R)-(+) -citronellol and of (R)-(+) -citronellol were homogeneous (means of 86.7% and 80.1% respectively) [4,5].

Cyclodextrins are chiral cyclic oligomers composed of six or more D-glucose units bonded through α-(1-4) linkage. The cavities have different diameters dependent on the number of glucose units as showed in the Table 1 (empty diameters between anomeric oxygen atoms given in the Fig. 1). The side rim depth is the same for all three (at about 0.8 nm) [6]. The β-DEX 225 capillary column is a chiral GC phase designed for separating enantiomers and other isomers. The β-DEX 225 column provides good separation of many enantiomers that are not separable or poorly separable on other chiral columns. β-Cyclodextrin consists of seven glucose residues. The mouth of the torus-shaped cyclodextrin molecule has a larger circumference than the base. Secondary hydroxyl groups at C2 and C3 atoms of the glucose units are located around the mouth.
2. EXPERIMENTAL

2.1 Materials, Method and Instruments

Citronella oil (*Cymbopogon winterianus*) obtained from Sari Warna Yogyakarta, and standards citronellal (+)/(−)-citronellal (*Fluka*), (+)/(−)-citronellal (*E.Merck*), and (R)/(−)-citronellal (*Sigma Aldrich*) then called as citronellal A, B, and C respectively.

2.2 Isolation of Citronellal

Citronellal was isolated from citronella oil that contain more than 35% of citronellal (GC-MS) by fractional distillation under reduced pressure (5 cmHg). To obtain higher concentration of citronellal, distillate was redistilled. Isolated citronellal was analyzed by GC, FTIR spectrophotometer, and ¹H-NMR spectrometer. Chromatogram and spectra citronellal isolated compared to its standard citronellal.

2.3 Enantioselective Gas Chromatography

GC conditions were adapted from Nhu-Trang et al. method [3]: carrier gas: hydrogen, 9.25 psig, flow 1.5 mL min⁻¹; injector and FID temperatures 250 °C. The temperature was programmed from 60 °C (held for 10 min)
at 1° min⁻¹ to 110 °C and then at 6° min⁻¹ to 175 °C which was held for 2 min.

### 2.4 Molecular Modeling

The stabilities of citronellal enantiomers on the β-DEX-225 chiral column were evaluated by quantum mechanical calculations. Geometry of citronellal enantiomers were optimized by Semiempirical AM1 method. Because of molecule of heptakis(2,3-di-O-acetyl-6-O-tert-butylidimethylsilyl)-β-cyclodextrin is too large, thus optimization was done by Molecular Mechanic MM+ and then followed optimization of citronellal enantiomer in the β-DEX-225 cavity structure by Semiempirical AM1 method.

### 3. RESULTS & DISCUSSION

#### 3.1 Isolation of Citronellal

The results of physical properties determination of citronella oil are followed: specific rotation $[\alpha]_{D}^{27}$-2,30°, specific gravity $[\rho]_{27}$ 0.8856 g/cm³, and refractive index $[\eta]_{20}$ 1.4684. The results of GC-MS analysis showed that citronella oil contain of 42.12% citronellal.

Fractional distillation of 250 mL citronella oil obtained two fractions. The amount of first fraction (80-110 °C) is 18.0 mL distillate, and the second fraction (110-120 °C) is 99.5 mL distillate. Redistillation of second fraction was obtained 68.0 mL distillate with higher citronellal contain (97.30%).

The results of physical properties determination of isolated citronellal are followed: specific rotation $[\alpha]_{D}^{27}$ +8.35°, specific mass $[\rho]_{27}$ 0.8570 g/mL, and refractive index $[\eta]_{20}$ 1.4478, while physical properties of standard citronellal B are followed: $[\alpha]_{D}^{27}$ +3.80°, $[\rho]_{27}$ 0.8469 g/cm³, and $[\eta]_{20}$ 1.4373. Based on physical properties and spectrum data, we assumed that main contain of isolated citronellal is (R)-(+)citronellal.

Structural analysis of citronellal by FTIR spectrophotometer showed the identical spectrum with IR spectrum of standard citronellal C. Figure 3 (a) showed spectrum with $\nu_{\max}$: 1724.2 (s, C=O aldehyde), 2870.08 and 2715.77 (w, C-H aldehyde), 2924.09 (s, C–H sp³), 1643.35 (w, C=C), 1450.47 (m, –CH₂–), 1381.03 (m, -CH₃) cm⁻¹.

$^1$H-NMR spectrum of isolated citronellal appropriate to $^1$H-NMR spectrum of standard citronellal C.

Figure 4 (a) showed spectrum (60 MHz, TMS) that consist of peak A ($\delta$=9.75 ppm, t, 1H, aldehyde proton), peak B ($\delta$=5.08 ppm, t, 1H, proton C=C olefin), peak C ($\delta$=2.36 ppm, t, 1H, Hc), peak D ($\delta$=1.88-2.30 ppm, m, 2H, –CH₂– bonded to C=C olefin), peak E ($\delta$=1.68 ppm, s, 6H, isopropylidene=C(CH₃)₂ proton), peak F ($\delta$=1.32 ppm, m, 1H, chiral C proton), and peak G ($\delta$=1.06 ppm, d, -CH₃ proton).
3.2 Enantioselective Gas Chromatography

The results of samples analysis by enantioselective gas chromatography showed in Figure 5.

Isolated citronellal contained 88.21% ee, commercial enantiomerically pure (standard citronellal C) contained 91.34% ee, and commercial racemic citronellal (standard citronellal C) contained 51.18% ee of (R)-(+-)-citronellal. The retention time of (R)-(+-)-citronellal is longer than (S)-(+-)-citronellal. It is means that (R)-(+-)-citronellal stronger adsorbed on the stationary phase and approximated relative boiling point of (R)-(+-)-citronellal is higher than its enantiomer.

3.3. Molecular Modeling

Energy of complex (R)-(+-)-citronellal in the β-DEX-225 cavity is lower than its enantiomer (Table 2). This means that (R)-(+-)-citronellal more stable and stronger bonded in the β-DEX-225 cavity than (S)-(+-)-citronellal. The calculation results of total energy and heat formation parameters were supported the experiment data that showed the retention time of (R)-(+-)-citronellal was longer than its enantiomer. Fig. 6 showed citronellal molecule possible to pass the β-DEX-225 cavity if its position parallel to cavity. We assumed that the enantioselectivity not only determined of polarity of citronellal enantiomer molecule but also the other parameters like geometry and volume of molecule.

| No | Parameter                      | (R)-(+-)-citronellal in β-DEX-225 | (S)-(+-)-citronellal in β-DEX-225 |
|----|--------------------------------|----------------------------------|----------------------------------|
| 1  | Total energy, kkal/mol          | -42651.9144                      | -42651.8401                      |
| 2  | Binding Energy, kkal/mol        | -2770.4235                       | -2770.4235                       |
| 3  | Heat of formation, kkal/mol     | -64,2028                         | -64,1285                         |
| 4  | Total dypole, Debyes            | 2,646                            | 2,789                            |

Table 2. The results of geometry optimization (R)-(+-)-citronellal and (S)-(+-)-citronellal in the β-DEX-225 cavity by Semiempirical-AM1 method

4. CONCLUSION

Citronellal 97.3% has been isolated from citronella oil (Cymbopogon winterianus-Java type) by fractional distillation under reduced pressure at 5 cmHg and 110-120 °C. Enantioselective GC analysis on a Supelco β-DEX 225 column showed that citronellal contain of 88.21% ee of (R)-(+-)-citronellal.

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