Development of a Rapid and Simple Non-Derivatization Method to Determine Constituents and Antioxidative Capacity of Camellia Oils by HPTLC

Guang-Ping Lv1*, Meijun Aoli1*, Bin Zhou2, Jing Zhao1#

1State Key Laboratory of Quality Research in Chinese Medicine, Institute of Chinese Medical Sciences, University of Macau, Macao, China; 2School of Pharmacy, Jiangxi Science and Technology Normal University, Nanchang, China.

Email: zhaojing.cpu@163.com

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ABSTRACT

Camellia oil is an edible vegetable oil with high value of nutrition and health protection function such as antioxidant and adjusting blood fat. In this study, a simple, rapid and effective HPTLC method was developed for analyzing the composition and antioxidant constituents of camellia oil. The HPTLC was performed on G60 plate with n-hexane-diethyl ether-acetic acid (6:4:0.1, v/v/v) as mobile phase combined with two coloration methods (ethanol containing 10% phosphomolybdic acid, ethanol containing 0.03% DPPH) and scanning densitometry technique. The unsaturated fatty glyceride, free fatty acids, sterols and lipids including triolein, oleic acid, ergosterin, β-sitosterol, tocopherol and phospholipids in camellia oils were determined and performed densitometrically at $\lambda_{s1} = 620$ nm and $\lambda_{s2} = 517$ nm. The results show that the main components of different samples of camellia oil are similar, however the contents are diverse. The antioxidative test shows that camellia oil has obvious antioxidant capability as olive oil, especially the pressed virgin oil. Therefore, this non-derivatization HPTLC method can be used for composition and antioxidative capacity determination of camellia oils.

Keywords: Antioxidant; Camellia oleifera Abel; Camellia Oil; HPTLC; Non-Derivatization

1. Introduction

Food provides not only essential nutrients required for life, but also bioactive compounds useful to maintain good health and disease prevention. Camellia oil is an edible vegetable oil that can be obtained from the mature seeds of Camellia oleifera Abel which mainly distributed in southern China, such as Guangxi, Hunan, and Jiangxi provinces [1]. It is an important woody oil crop [2] and also grown as an ornamental plant in Western countries. Camellia oil can be used both for medicine and food dual purposes such as directly given in the form of injection [3] or used as dissolvent of liquid soluble drug or ointment base [4], especially it listed as medicinal oil in the Chinese pharmacopoeia [5]. Simultaneously, camellia oil is high quality natural cooking oil, because of its high quality and healthful properties. The monounsaturated fatty acid contains the best natural vegetable oil, which is hailed as olive oil of eastern world [4,6-10]. Besides, it also has been used in cosmetics [11-13].

Camellia oil was traditionally applied as a health caring medicine to prevent coronary heart disease, hypertension, cerebrovascular disease [14], arteriosclerosis, stomachache [7] as well as burning injury in China [15]. Recent evidence concerning an association between bioactive phytochemicals in the diet and health has imparted impetus to the utilization of camellia oil for most healthy edible oil, keeping its bioactive phytochemicals intact. Camellia oil is predominantly composed of glyceride of unsaturated fatty acid such as oleic acid (75% - 80%) and linoleic acid, while the saturated fats are presented in low amounts [13]. Camellia oil also contains many other natural antioxidants with various biological activities, such as phytosterols, tocopherol, free fatty acids (FFAs) [16]. The camellia oil is helpful in protecting liver against carbon tetrachloride toxicity [17]. It has been observed to suppress cholesterol content in the body and provide resistance to oxidative stress [18]. Antioxidants are essen-
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2. Materials and Methods

2.1. Materials

Acetic acid, n-hexane, diethyl ether, petroleum ether (60°C - 90°C), acetone, ethyl acetate, toluene-methanol and chloroform were purchased from UNI-CHEM d.o.o. (Belgrade, Serbia and Montenegro). All chemicals reagents used were analytical grade. Triolein (98%, HPLC), oleic acid (99%, HPLC), beta-sitosterol (98%, HPLC), ergosterol (98%, HPLC), tocopherol (98%, HPLC) and 2, 2-diphenyl-1-picrylhydrazyl (DPPH) were purchased from Sigma Company (USA). Refined camellia oils were collected from Fangchenggang of Guangxi Province; Refined camellia oil (SM07) was collected from Yichun of Jiangxi Province; Pressed virgin oil (SM08) was collected from Jiangxi Province; Virgin oil (SM09) was made in the Institute of Chinese Medical Sciences, University of Macau, Macao, China; Refined olive oil (SM10) was collected from Italy. All refined oils are commercially available. Voucher specimens of these samples were deposited at the Institute of Chinese Medical Sciences, University of Macau, Macao SAR, China.

2.2. Preparation of Oil Samples and Standard Solutions

The seeds of Camellia oleifera Abel were powdered by a laboratory grinder after discarding the shells and dried to constant weight at 65°C. The prepared sample (5 g) was macerated with 50 mL solvent (petroleum ether, 60°C - 90°C) at 80°C for 6 h by using Soxhlet apparatus. Finally, the solvent was dried under nitrogen protection. The residual is the crude oil samples (extracted), marked as SM09.

A stock solution containing 0.25 mg/mL of phospholipid, tocopherol, ergosterin and β-sitosterol, 0.5 mg/mL of oleic acid and 1 mg/mL of triolein was prepared in methanol, vortex for 2 min. The solution was stored at 4°C freezer for use.

The oil sample (25 μL, accurately weighed) was dissolved in 1 mL of n-hexane and vortex for 2 min; the sample (2.5%) was collected for analysis.

2.3. HPTLC Methods

Chromatography was performed on silica G₆₀ TLC plate (20 cm × 10 cm, Merck, Darmstadt, Germany), and a HPTLC system (Desaga GmbH, Germany) including AS30 HPTLC Applicator, CD 60 HPTLC densitometer with Pro Quant Windows software. The plate was heated at 105°C for 30 min before use. The plate was allowed to cool to room temperature before spotting the samples. Mixed standards (3 μL) and samples (3 μL) of camellia oil were spotted on the plate as bands 4 mm wide, 18 mm apart and 10 mm from the bottom edge. The plate was developed to a distance of 95 mm with n-hexane-diethyl ether-acetic acid (6:4:0.1, v/v/v) in a Desaga 20 cm × 10 cm glass flat-bottom chamber after equilibration with mobile phase vapor for 30 min. The developed plate was colorized with ethanol containing 10% phosphomolybdic acid solution and heated at 95°C on a YOKO-XR plate heater (Wuhan YOKO technology Ltd., China) for 5 min to make spots colored clearly. The plate was then scanned at λ₟₄₃ (scan wavelength) = 620 nm in positive signal mode by use of the densitometer.

The chromatography for evaluating antioxidant capa-

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bility of camellia oil, was performed on silica G₅₀ TLC plate and a HPTLC system, the procedure was the same as mentioned above, the developed plate was colorized with ethanol containing 0.03% DPPH solution and heated at 40°C on a YOKO-XR plate heater (Wuhan YOKO technology Ltd., China) for 30 min and at room temperature for 12 h to make spots colored clearly. The treated plate scanned at \( \lambda_{254} \) (scan wavelength) = 517 nm in negative signal mode by use of the densitometer.

2.4. Validation of the Method

The precision was expressed as instrumental precision, identical and different plate precision. Instrumental precision was performed by scanning the same spot of the investigated compound of SM08 sample (3 \( \mu L \)) six times, and variations were expressed by the relative standard deviations (RSD). Identical plate precision was determined by analyzing six spots of the SM08 sample (3 \( \mu L \)) on one plate, while different plate precision was tested by determining one spot of the SM08 sample (3 \( \mu L \)) per plate on six TLC plates. The RSDs of the investigated compounds were calculated.

The repeatability was evaluated by preparing and analyzing six solutions of the SM08 sample (25 \( \mu L \) each). One spot of each solution was analyzed on the same plate, and RSDs of the investigated compounds were calculated.

The stability of six investigated compounds were also determined by injecting SM08 sample at 0, 4, 8, 12, 16, 20 and 24 h, and scanning each band respectively. The RSDs of the investigated compounds were calculated.

3. Result and Discussion

3.1. Optimization of Method

Mobile phases and developing mode were optimized to obtain good separation. Petroleum ether-acetone (4:1 or 3:1, v/v), n-hexane-ethyl acetate (5:1 or 4:1, v/v), toluene-methanol-acetic acid (90:4:4, v/v/v), chloroform-acetone-acetic acid (85:15:1, v/v/v), n-hexane-acetone-acetic acid (90:10:1 or 80:20:1, v/v/v), were tested and demonstrated not available for simultaneous separation of low- and high-polarity compounds in camellia oil in one run. The optimum development reagents are: n-hexane-diethyl ether-acetic acid (6:4:0.1, v/v/v). As the results, five main investigated compounds and the components in ten oil samples were well separated in once development (Figure 1).

The saturation time (10 min, 20 min, 30 min, 40 min) and the spray reagents (5% vanillin-H₂SO₄, iodine vapor, ethanol containing 10% phosphomolybdic acid) were also optimized. The results show that saturating the plate for 30 min and using the ethanol containing 10% phosphomolybdic acid as spray reagents show good separation for HPTLC analysis.

3.2. Method Validation

Instrumental precision (RSD, %) was less than 1.47% (n = 6). For identical plates, and the different plates precision were also determined. The overall RSDs of identical and different plate were less than 2.09% (n = 6) and 3.01% (n = 6), respectively. The repeatability for all the analysis was less than 3.71%. The stability results prove it was stable within 24 h, which RSDs was less than

Figure 1. HPTLC of mixed standards and oil samples developed on silica G₅₀ TLC plate (20 cm × 10 cm) and colored with ethanol containing 10% phosphomolybdic acid, viewed at white light. ST, mix standards; X, phospholipid; A, tocopherol; B, ergosterin; C, β-sitosterol; D, oleic acid; E, triolein. SM02 to SM07 are refined camellia oil; SM08 is pressed virgin camellia oil; SM09 is extracted virgin camellia oil; SM10 is refined olive oil.

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2.0%. The established method showed good stability, precision and repetitiveness.

### 3.3. Determination of Main Components in Oil Samples

The proposed method was applied to the determination of oil samples. The representative chromatograms of the standard solutions and the oil samples were shown in Figure 1. The results show that, the components could be simultaneously separated in one run. The main components of camellia oil and olive oil samples are similar: tocopherol with Rf values of 0.23, ergosterin with Rf values of 0.41, \(\beta\)-sitosterol with Rf values of 0.46, oleic acid with Rf values of 0.67, triolein with Rf values of 0.88 and one unknown compound with Rf values of 0.94, respectively. However, the results show that pressed virgin oil is different from others, which contains more free fatty acid. Therefore, TLC could rapidly and easily analyze free fatty acid in vegetable oil and discriminate the pressed virgin oil [25,31].

Maximum detection wavelengths which are set in scan detector are important for the detection sensitivity. Spots of tocopherol, ergosterin, \(\beta\)-sitosterol, oleic acid and triolein were scanned from 400 to 900 nm so as to record their spectrum and to obtain their wavelengths of maximum absorption. Densitograms were recorded at the wavelength of maximum absorption (620 nm) of primary components. The detection wavelengths of the investigated compounds were selected based on their absorption spectra.

The contents of components in ten oil samples were determined using TLC scanning (TLCs). Their TLCs profiles were shown in Figure 2. Generally, the components are showing high similarities of ten oil samples in tocopherol, ergosterin, \(\beta\)-sitosterol, triolein. However, there are two obvious discrimination: 1) Virgin oil samples (SM08, SM09) have high content of polarity compounds which relatively more ingredients were retained in the origin shown in Figure 1. From the band (Rf-0) which corresponding with standard, it may be the phospholipids for this kind of components was reported as common component in virgin camellia oil [32]. 2) Pressed virgin oil (SM08) has more FFAs which was shown in Figure 1 bond D. FFAs in edible oils are undesirable, it may results in lower flavour quality and stability of the oil, moreover high levels of FFAs will result in rancidity of the oil and usually removed from crude oil by refining in industrial production [33].

### 3.4. Antioxidative Capacity of Oil Samples

The quality of edible oil is determined by various factors such as flavour, free fatty acids (FFAs) content, oxidation, etc. The free radical scavenging activity of the oil samples was done using DPPH. The TLC chromatogram of camellia oil and olive oil samples was colored with ethanol containing 0.03% DPPH solution. Figure 3 shows the results, which revealed sterols and unsaturated fatty acid esters in camellia oil have high antioxidant capacity [18].

The contents of antioxidative components in ten oil samples were determined using TLC scanning (TLCs). For the DPPH have maximum absorption at 517 nm. The plate colored with ethanol containing 0.03% DPPH, scanned at \(\lambda_{\text{abs}}\) (scan wavelength) = 517 nm in negative signal mode by use of the densitometer.

The densitogram results for oil samples in Figure 4 provide a visible method of the antioxidative capacity between virgin oil and refined oil. It also proves that pressed virgin oils can significantly reduce the superoxide radicals (Figure 4), because it has more FFAs content. Camellia virgin oil with more antioxidative components than other refined oil. The vegetable oil sold in markets has lost parts of antioxidative components during the refining procedures in order to increase its stability and flavour. The camellia oil sold in markets has similar components and antioxidative capability comparing with imported olive oil, which has high price. Therefore, camellia oil has potentiality to be used as excellent healthy edible oil.

### 4. Conclusion

In this study, a simple and rapid method to simultaneous determination of unsaturated fatty glyceride, free fatty acids, sterols and other components of camellia oil was set up. This research breaks through the traditional derivatization analysis method and exhibits a powerful potential for the analysis of FFAs, phytosterols, etc. from edible oils and evaluates the antioxidative capacity of camellia oil.

Figure 2. TLC densitograms (at 620 nm) of ten oil samples. SM01 to SM07 are refined camellia oil; SM08 is pressed camellia oil; SM09 is extra camellia oil; SM10 is refined olive oil.
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Figure 3. Chromatograms of oil samples’ antioxidative capability. Oil samples developed on silica G60 TLC plate (20 cm × 10 cm) and colored with ethanol containing 0.03% DPPH, viewed at white light. SM01 to SM07 are refined camellia oil; SM08 is pressed virgin camellia oil; SM09 is extra virgin camellia oil; SM10 is refined olive oil.

Figure 4. Typical TLC densitograms of ten oil samples 1), refined camellia oil 2), pressed virgin oil 3), extracted virgin camellia oil 4). SM01 to SM07 are refined camellia oil; SM08 is pressed camellia oil; SM09 is extracted camellia oil; SM10 is refined olive oil.

vegetable oil directly. The developed non-derivatization TLC method can be used as an economical alternative method for routine quality control of vegetable oil.

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REFERENCES


[29] A. Zeb, “Triacylglycerols Composition, Oxidation and...


