Jae B. Park

Diet, Genomics, and Immunology Laboratory, BHNRC, ARS, USDA, Bldg. 307C, Rm. 131, Beltsville, MD 20705, USA

Correspondence should be addressed to Jae B. Park; jae.park@ars.usda.gov

Received 19 April 2017; Revised 10 July 2017; Accepted 25 July 2017; Published 23 August 2017

Academic Editor: Günther K. Bonn

Copyright © 2017 Jae B. Park. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Javamide-I/javamide-II are phenolic amides found in coffee. Recent reports suggested that they may contain several biological activities related to human health. Therefore, there is emergent interest about their quantities in coffee-related products. Green coffee extract is a powder extract made of unroasted green coffee beans, available as a dietary supplement. However, there is little information about the amounts of javamide-I/javamide-II in green coffee extract products in the market. Therefore, in this paper, javamide-I/javamide-II were extracted from green coffee extract products and their identifications were confirmed by NMR. After that, the amounts of javamide-I/javamide-II were individually quantified from seven different green coffee extract samples using the HPLC method coupled to an electrochemical detector. The HPLC method provided accurate and reliable measurement of javamide-I/javamide-II with excellent peak resolution and low detection limit. In all seven green coffee extract samples, javamide-II was found to be between 0.28 and 2.96 mg/g, but javamide-I was detected in only five samples in the concentration levels of 0.15–0.52 mg/g, suggesting that green coffee extract products contain different amounts of javamide-I/javamide-II. In summary, javamide-I/javamide-II can be found in green coffee extract products sold in the market, but their amounts are likely to be comparatively different in between green coffee extract brands.

1. Introduction

Coffee plants consist of several species such as Coffea arabica, Coffea canephora, Coffea liberica, Coffea excelsa, and Coffea stenophylla [1, 2]. Particularly, Coffea arabica (Arabica coffee) and Coffea canephora (Robusta coffee) are two most commercially important coffee species [2]. Green coffee extract is a powder extract made of unroasted green coffee beans with potential health effects [3–7]. Particularly, green coffee extract is often sold as a weight loss supplement, although more thorough studies are needed to determine its effectiveness as a weight loss supplement [7]. In general, green coffee extract can be made of any unroasted green coffee beans, but most of green coffee extract products found in the market are acclaimed to be made of Arabica beans. Javamide-I (N-coumaroyltryptophan) and javamide-II (N-caffeoyltryptophan) are safflomide-type phenolic amides found in roasted and unroasted coffee beans [3, 8]. Recently, they were reported to contain several biological activities related to human health [8, 9]. Thus, there is a great deal of interest about how much javamide-I/javamide-II can be found in coffee-related products, particularly green coffee extract found in the market. Therefore, in this paper, javamide-I/javamide-II in the seven green coffee extract products purchased from the market were extracted and their identities were confirmed using NMR spectroscopic methods. Then, their amounts in the seven extract products were determined using the HPLC method with excellent and reliable peak resolution and low detection limit.

2. Materials and Methods

2.1. Materials. Seven green coffee extract products were purchased from online stores. Javamide-I/javamide-II standards were prepared as reported previously [8, 9]. All other reagents and chemicals were purchased from Sigma Chemical Co. (St. Louis, MO).
2.2. HPLC Analysis of Javamide-I/Javamide-II Standards. HPLC analysis was performed as described previously with slight modification [8]. Briefly, a 150 mm × 2.1 mm i.d., 4 μm, Nova-Pak C18 (Waters, Milford, MA) was used as the stationary phase to analyze the standards and green coffee samples. The samples were separated using a gradient condition; buffer A (10 mM phosphate, pH 5) for 0–5 min, a linear gradient from buffer A to buffer B (40% acetonitrile) for 5–45 min, and buffer B for 10 min (1mL/min). The samples were injected by an autosampler into Alliance 2690 HPLC system (Waters, Milford, MA), and were monitored by CoulArray electrochemical detector with four electrode channels (ESA, Chelmsford, MA).

2.3. Extraction of Green Coffee Extract Samples. Green coffee extract samples were extracted in two different ways: collectively (for the NMR analysis and extraction recovery of javamide-I/javamide-II) and separately (for the quantification of javamide-I/javamide-II). For the NMR confirmation, the equally mixed green coffee extract sample was extracted with 70% methanol (50 mL per 1 g green coffee) with gentle shaking at room temperatures for 24 hr. Then, the samples were extracted twice, providing 1st and 2nd extraction samples. The 1st and 2nd extraction samples were utilized to isolate javamide-I/javamide-II for NMR analysis and determine the extraction recovery. Meanwhile, for the quantification of javamide-I/javamide-II in each green coffee extract sample, seven green coffee samples were individually extracted using the same extraction method.

2.4. NMR Analysis. The peaks of javamide-I/javamide-II were isolated from the green coffee extract using the method described previously. The chemical structures of isolated javamide-I/javamide-II were verified using NMR spectroscopic methods. For NMR experiments, the samples were prepared by dissolving javamide-I/javamide-II (20 mg) in d6-DMSO (0.75 mL). 1H and 13C spectra were acquired at ambient temperature on the JEOL BCX-400 NMR spectrometer operating 400 MHz for 1H and 100 MHz for 13C.

2.5. Quantification of Javamide-I/Javamide-II in Green Coffee Extract. Seven green coffee extract samples were individually analyzed using the HPLC method as described previously. Since the extraction was performed twice (1st and 2nd), the 1st and 2nd extraction samples were independently analyzed using the HPLC method to determine the amounts of javamide-I/javamide-II in each extraction. The amounts of javamide-I/javamide-II were quantified using CoulArray electrochemical detector with four electrode channels (ESA, Chelmsford, MA).

2.6. Statistical Analysis. All statistical analyses were performed with the SigmaPlot II.0 (Chicago, IL). Data points in all figures were represented as the mean ± SD of 5 samples.

3. Results and Discussion

3.1. HPLC Analysis of Javamide-I/Javamide-II Standards and Their Standard Curves. Javamide-I/javamide-II standards were separated and detected using the HPLC method as described in Materials and Methods (Figure 1). The HPLC condition was examined to improve the separation and resolution of the compounds using a gradient condition described in Materials and Methods. A HPLC condition was selected for optimizing the detection, separation, resolution, and reproducibility of the peaks of javamide-I/javamide-II in the chromatography. Total HPLC running time for the assay was 50 min, and javamide-I/javamide-II were detected at a retention time of 28.8 and 31.3 min, respectively. The peak areas of javamide standards (1, 10, 50, 100, 200, and 300 pmol) are utilized to produce their standard curves, which provided satisfactory linear responses at the concentrations between 1 and 300 pmol (for javamide-I correlation coefficient (R) = 0.99 (f(x) = 0.0204), and for javamide-II, correlation coefficient (R) = 0.99 (f(x) = 0.0414)). Also, the implemented limit of detection (LOD) and limit of quantification (LOQ) were 5/5.5 pmol, respectively (Figure 1).

3.2. Extraction and HPLC Analysis of Green Coffee Extract Mixture Sample. Since all seven green coffee extract samples purchased from the market are fine powders, the sample was prepared by mixing equal amounts of the seven extract powders and was extracted with 70% methanol as described in Materials and Methods. After that, the HPLC analysis of javamide-I/javamide-II was performed as described previously. After several preparations of green coffee mixture samples, the 1 g green coffee per 50 mL 70% methanol was found optimal for extraction and HPLC separation of the javamide in the samples. The HPLC method provided excellent separation and resolution of the peaks of javamide-I/javamide-II in the green coffee samples (Figure 2). Like the chromatograms of javamide-I/javamide-II standards, javamide-I/javamide-II in the green coffee extract samples were detected at the same retention time (28.9 and 31.2 min).

3.3. NMR Confirmation of Javamide-I/Javamide-II. For NMR confirmation, each peak of javamide-I/javamide-II was purified from the green coffee extract using the method described previously. The purified javamide-I was analyzed using NMR spectroscopic methods (Figures 3 and 4). The NMR data were following: IH NMR (d6-DMSO, 400 MHz) δ 7.58 ((IH, d, J) 8.2 Hz, H-18), 7.45 ((IH, d, J) 8.2 Hz, H-1/H-5), 7.37 ((IH, d, J) 15.6 Hz, H-7), 7.33 ((IH, d, J) 7.8 Hz, H-15), 7.20 (IH, s, H-13), 7.06 ((IH, t, J) 7.3 Hz, H-16), 6.98 ((IH, t, J) 7.3 Hz, H-17), 6.59 ((IH, d, J) 8.2 Hz, H-2/H-4), 6.46 ((IH, d, J) 15.6 Hz, H-8), 4.72 ((IH, t, J) 7.3 Hz, H-10), 3.24 ((IH, dt, J) 6.0, 6.9 Hz, H-11), 9.68 (IH, br s, OH-a), 12.89 (IH, br s, OH-a’), 10.79 (IH, br s, NH-beta), 8.38 (IH, br s, NH-alpha); 13C NMR (d6-DMSO, 100 MHz) 174.7 (C, C-20), 166.8 (C, C-9), 157.5 (C, C-3), 141.7 (C, C-7), 136.4 (C, C-14), 130.6 (C, C-1), 130.6 (C, C-5), 127.9 (C, C-6), 127.4 (C, C-19), 124.0 (C, C-8), 123.0 (C, C-13), 121.7 (C, C-16), 119.8 (C, C-17), 118.8 (C, C-18), 115.7 (C, C-2), 115.7 (C, C-4), 115.5 (C, C-15), 109.3 (C, C-12), 60.3 (C, C-10), 28.2 (C, C-11). Based on the NMR data, the structure of the isolated compound was determined as being coumaroyl-L-tryptophan (N-coumaroyltryptophan; javamide-I). Likewise, the purified javamide-II was analyzed using NMR spectroscopic methods. The NMR data were following:
IH NMR (d6-DMSO, 400 MHz) δ 7.58 ((IH, d, J) 8.2 Hz, H-18), 7.33 ((IH, d, J) 7.8 Hz, H-15), 7.32 ((IH, d, J) 15.6 Hz, H-7), 7.20 (IH, s, H-13), 7.06 ((IH, t, J) 7.3 Hz, H-16), 7.06 (IH, s, H-5), 6.98 ((IH, t, J) 7.3 Hz, H-17), 6.82 ((IH, d, J) 8.7 Hz, H-2), 6.67 ((IH, dd, J) 8.2, 1.4 Hz, H-1), 6.46 ((IH, d, J) 15.6 Hz, H-8), 4.72 ((IH, t, J) 7.3 Hz, H-10), 3.24 ((IH, dt, J) 6.0, 6.9 Hz, H-11), 9.48 (IH, br s, OH-a), 12.89 (IH, br s, OH-a'), 10.79 (IH, br s, NH-beta), 8.38 (IH, br s, NH-alpha); 13C NMR (d6-DMSO, 100 MHz) 174.7 (C, C-20), 166.8 (C, C-9), 146.5 (C, C-3), 145.9 (C, C-4), 141.7 (C, C-7), 136.4 (C, C-14), 128.0 (C, C-6), 127.4 (C, C-19), 124.0 (C, C-8), 123.2 (C, C-1), 123.0 (C, C-13), 121.7 (C, C-16), 119.8 (C, C-17), 118.8 (C, C-18), 117.2 (C, C-2), 115.2 (C, C-5), 111.1 (C, C-15), 109.7 (C, C-12), 60.3 (C, C-10), 28.2 (C, C-11). Based on the NMR data, the structure of the isolated compound was determined as being (E)-(3-(3,4-dihydroxyphenyl)acryloyl)tryptophan (N-caffeoyltryptophan; javamide-II).

3.4. Extraction Recovery of Javamide-I/Javamide-II in Green Coffee Extract Samples. In order to determine the extraction recovery, the extraction was performed twice. The amounts of javamide-I in the 1st extraction were 95% higher than those
of the 2nd extraction, high enough for no subsequent 3rd extraction (Table I(a)). The recovery was determined by measuring the amounts following the addition of javamide-I standard (0.2 mg) and found to be more than 98% (Table I(b)). Likewise, the amounts of javamide-II in the 1st extraction were 96% higher than those of the 2nd extraction (Table 2(a)) and the recovery was found to be more than 98% (Table 2(b)).

3.5. HPLC Quantification of Javamide-I/Javamide-II. For the quantification of javamide-I/javamide-II, seven green coffee extract samples were individually extracted and prepared for HPLC analyses using the same extraction method described in Materials and Methods. Javamide-I was detected in five samples at the range of 0.15–0.52 mg per g (Table 3). However, javamide-II was detected in all seven green coffee samples between 0.28 and 2.96 mg per g (Table 4), suggesting that javamide-II could be present in higher amounts than javamide-I in the green coffee extract samples, which was also confirmed in roasted coffee beans (data not shown here). Altogether, this study clearly suggests that javamide-I/javamide-II can be found in green coffee extract products sold in the market, but their amounts are likely to be significantly different depending on green coffee extract brands.

4. Conclusion
In this paper, javamide-I/javamide-II were extracted from seven green coffee extract products sold in the market and
Table 2: (a) The amounts of javamide-II in 1st and 2nd extractions. The mixture of seven green coffee extract samples (S1–S7) as described in Table 1 was extracted twice with 70% MeOH (50 mL), providing the 1st and 2nd extracts. The amounts of javamide-II were presented as mg per 1 g green coffee sample (n = 5). (b) The recovery yield of javamide-II. The purified javamide-II (1 mg) was added to the mixture of the seven green coffee extract sample containing 1 mg javamide-II and extracted with 70% MeOH. The recovery was determined using 1st and 2nd extracts (n = 5).

| Sample | 1st (mg) | 2nd (mg) | 1st and 2nd (mg) | 1st/2nd (%) |
|--------|----------|----------|------------------|-------------|
| S1–7   | 1.01 ± 0.02 | 0.03 ± 0.001 | 0.63 ± 0.02 | 97.1 |

| Sample | Initial (mg) | J-II added (mg) | 1st (mg) | 2nd (mg) | Yield (%) |
|--------|--------------|-----------------|----------|----------|-----------|
| S1–7   | 1            | 1               | 1.89 ± 0.05 | 0.07 ± 0.005 | 98.0 |

Table 3: The amounts of javamide-I in seven green coffee samples. Seven green coffee extract samples (S1–S7) as described in Table 1 were individually extracted twice with 70% MeOH extracts (50 mL), providing the 1st and 2nd extracts. The amounts of javamide-I were presented as mg per 1 g green coffee sample (n = 5).

| Sample | 1st (mg) | 2nd (mg) | 1st and 2nd (mg) | 1st/2nd ratio (%) |
|--------|----------|----------|------------------|-------------------|
| S1     | 0.41 ± 0.02 | 0.01 ± 0.001 | 0.42 ± 0.02 | 97.0 |
| S2     | 0.51 ± 0.02 | 0.01 ± 0.001 | 0.52 ± 0.02 | 97.6 |
| S3     | 0.15 ± 0.01 | ND        | 0.15 ± 0.01 | —     |
| S4     | ND         | ND        | —           | —     |
| S5     | 0.15 ± 0.01 | ND        | 0.15 ± 0.01 | —     |
| S6     | ND         | ND        | —           | —     |
| S7     | 0.17 ± 0.02 | ND        | 0.17 ± 0.02 | —     |

Table 4: (a) The amounts of javamide-II in seven green coffee samples. Seven green coffee extract samples (S1–S7) were individually extracted twice with 70% MeOH extracts (50 mL) as described in Table 1, providing the 1st and 2nd extracts. (b) The amounts of javamide-I/javamide-II in the samples. The amounts were presented as mg per 1 g green coffee sample (n = 5).

| Sample | 1st (mg) | 2nd (mg) | 1st and 2nd (mg) | 1st/2nd ratio (%) |
|--------|----------|----------|------------------|-------------------|
| S1     | 1.68 ± 0.05 | 0.07 ± 0.005 | 1.75 ± 0.05 | 95.6 |
| S2     | 2.88 ± 0.06 | 0.08 ± 0.005 | 2.96 ± 0.06 | 97.0 |
| S3     | 0.68 ± 0.02 | 0.02 ± 0.001 | 0.70 ± 0.02 | 96.4 |
| S4     | 0.27 ± 0.01 | 0.01 ± 0.001 | 0.28 ± 0.01 | 95.2 |
| S5     | 0.65 ± 0.02 | 0.02 ± 0.001 | 0.67 ± 0.02 | 96.2 |
| S6     | 0.27 ± 0.01 | 0.01 ± 0.001 | 0.28 ± 0.01 | 95.2 |
| S7     | 0.76 ± 0.02 | 0.02 ± 0.001 | 0.78 ± 0.02 | 96.8 |

| Sample | Javamide-I (mg) | Javamide-II (mg) |
|--------|-----------------|------------------|
| S1     | 0.42 ± 0.02     | 1.75 ± 0.05      |
| S2     | 0.52 ± 0.02     | 2.96 ± 0.06      |
| S3     | 0.15 ± 0.01     | 0.70 ± 0.02      |
| S4     | ND               | 0.28 ± 0.01      |
| S5     | 0.15 ± 0.01     | 0.67 ± 0.02      |
| S6     | ND               | 0.28 ± 0.01      |
| S7     | 0.17 ± 0.02     | 0.78 ± 0.02      |

Their amounts were quantified using the HPLC method which provided accurate and reliable measurement of javamide-I/javamide-II with excellent peak resolution and low detection limit. Javamide-II was detected in all seven green coffee extract samples at the range between 0.28 and 2.96 mg per g green coffee, but javamide-I was detected only in five samples at the range of 0.15–0.52 mg per g green coffee, indicating that the amount of javamide-II may be found higher than that of javamide-I in green coffee extract products. This study suggests that different amounts of javamide-I/javamide-II
can be found in green coffee extract products and their amounts are likely to be very different depending on green coffee extract brands.

**Disclosure**

Mention of trade names or commercial products in this publication is solely for the purpose of providing specific information and does not imply recommendation or endorsement by the U. S. Department of Agriculture.

**Conflicts of Interest**

There is nothing to disclose regarding financial, consulting, and personal matters related to this paper.

**Acknowledgments**

This study was funded by USDA (Project no. 1235-51000-054-00D).

**References**

[1] A. C. Charrier and J. Berthaund, “Botanical classification of coffee,” in *Coffee: Botany, Biochemistry and Production of Beans and Beverage*, M. N. Clifford and K. C. Willson, Eds., pp. 13–47, Croom Helm, London, UK, 1985.

[2] A. Farah and T. F. Dos Santo, “The coffee plant and beans,” in *Coffee in Health and Disease Prevention*, V. Victor Freedy, Ed., pp. 5–10, Academic Press, New York, NY, USA, 2015.

[3] O. Babova, A. Occhipinti, and M. E. Maffei, “Chemical partitioning and antioxidant capacity of green coffee (Coffea arabica and Coffea canephora) of different geographical origin,” *Phytochemistry*, vol. 123, pp. 33–39, 2016.

[4] R. Revuelta-Iniesta and E. A. S. Al-Dujaili, “Consumption of green coffee reduces blood pressure and body composition by influencing 11β-HSD1 enzyme activity in healthy individuals: a pilot crossover study using green and black coffee,” *BioMed Research International*, vol. 2014, Article ID 482704, 9 pages, 2014.

[5] S. J. Song, S. Choi, and T. Park, “Decaffeinated green coffee bean extract attenuates diet-induced obesity and insulin resistance in mice,” *Evidence-Based Complementary and Alternative Medicine*, vol. 2014, Article ID 718379, 14 pages, 2014.

[6] B.-K. Choi, S.-B. Park, D.-R. Lee et al., “Green coffee bean extract improves obesity by decreasing body fat in high-fat diet-induced obese mice,” *Asian Pacific Journal of Tropical Medicine*, vol. 9, no. 7, pp. 635–643, 2016.

[7] I. Onakpoya, R. Terry, and E. Ernst, “The use of green coffee extract as a weight loss supplement: a systematic review and meta-analysis of randomised clinical trials,” *Gastroenterology Research and Practice*, vol. 2011, Article ID 382852, 6 pages, 2011.

[8] J. B. Park, “Finding potent Sirt inhibitor in coffee: Isolation, confirmation and synthesis of Javamide-II (N-caffeoyltryptophan) as Sirt1/2 inhibitor,” *PLoS ONE*, vol. 11, no. 3, Article ID e0150392, 2016.

[9] J. B. Park, “Javamide-I-O-methyl ester increases p33 acetylation and induces cell death via activating caspase 3/7 in monocytic THP-1 cells,” *PhytoMedicine*, vol. 23, no. 13, pp. 1647–1652, 2016.