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Bioactivity-Guided Extract Optimization of *Osmanthus fragrans* var. *aurantiacus* Leaves and Anti-Inflammatory Activities of Phillyrin

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Abstract: The aim of this study was to identify the optimal extraction conditions for leaves of *Osmanthus fragrans* var. *aurantiacus*. Inhibitory effects of various extracts on NO production were compared. Antioxidant evaluations for total phenol and flavonoid contents were carried out using various extracts of *O. fragrans* var. *aurantiacus* leaves obtained under optimal extraction conditions that showed the greatest effect on NO production. The optimal method for extracting *O. fragrans* var. *aurantiacus* leaves resulted in an extract named OP OFLE. OP OFLE showed DPPH and ABTS radical scavenging activities in a concentration-dependent manner. Phillyrin (PH) was isolated as a major compound from OP OFLE by HPLC/DAD analysis. OP OFLE and PH reduced inducible nitric oxide (iNOS) and cyclooxygenase (COX)-2 protein expression and downregulated proinflammatory cytokines such as interleukin (IL)-1β, IL-6, IL-8, and tumor necrosis factor (TNF)-α in LPS-stimulated RAW 264.7 and HT-29 cells. To determine the signal pathway involved in the inhibition of NO production, a Western blot analysis was performed. Results showed that OP OFLE decreased phosphorylation of extracellular regulated kinase (pERK) 1/2 and the expression of nuclear factor-kappa B (NF-κB). Our results suggest that extracts of *O. fragrans* var. *aurantiacus* leaves and its major components have biological activities such as antioxidative and anti-inflammatory properties.

Keywords: *Osmanthus fragrans* var. *aurantiacus*; phillyrin; RAW 264.7 cells; antioxidant; optimal extraction; anti-inflammation; cytokine

1. Introduction

Inflammation is necessary for the human body’s defensive response to injury, lipid peroxide, and infection. However, prolonged inflammatory processes can exacerbate existing diseases or lead to various new diseases [1]. Various risks factors, including diabetes and high blood pressure, are increased by the injurious effects of low density lipoprotein cholesterol, initiating a chronic inflammatory reaction and resulting in vulnerable plaques, rupture, and thrombosis, and inflammation generally increases the incidence of neurodegenerative diseases, cardiovascular diseases, obesity, and cancer [2]. Inflammation can also induce acute and chronic inflammatory responses in the brain, heart, intestine, kidney, liver, lung, pancreas, and the reproductive system, potentially causing tissue impairment or disease. Both infectious and non-infectious agents and cell damage can activate inflammatory cells and trigger inflammatory signaling pathways [3].

Lipopolysaccharide (LPS)-stimulated macrophages have generally been used for evaluating the anti-inflammatory effects of various materials [4]. LPS can also activate Toll-like receptors (TLR) to induce phosphorylation and activation of MAP kinases (MAPKs) that are important for inducible nitric oxide synthase (iNOS) [5]. Activated macrophages can transcriptionally express iNOS, which can catalyze oxidative deamination of L-arginine...
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to produce nitric oxide (NO) [6]. Once activated by LPS stimulation, macrophages play a decisive role in the inflammatory response by releasing various factors such as NO, prostaglandin mediators, and proinflammatory cytokines such as tumor necrosis factor (TNF)-α, interleukin (IL)-1β, and IL-6 [7]. In addition, under inflammatory conditions, macrophages can significantly increase the production of both NO and superoxide anion (O2), which can rapidly react with each other to form peroxynitrite anion (ONOO), thus activating COX-1 and COX-2 as constitutive and inducible forms of cyclooxygenase, respectively [8]. Activation of nuclear factor-κB (NF-κB), an important transcription factor that regulates the transcription of TNF-α, IL-1β, and genes encoding inducible enzymes such as COX-2 and iNOS, is involved in immune and inflammatory responses [9]. Among mitogen-activated protein kinases (MAPKs), extracellular signal-regulated kinase (ERK) and p38 MAPK are thought to play a key role in the regulation of pro-inflammatory cytokines for cellular responses [10]. Intestinal epithelium is a crucial gut mucosal barrier that participates in innate immunity [11]. The IL-8 production can be induced by other inflammatory signals such as LPS, IL-1, and TNF [12].

Natural antioxidants are of increasing interest as medicines and food additives [13]. The use of antioxidants to prevent oxidative damage caused by excessive free radicals is significant for the cosmetics, medical, and food industries, among others [14]. Oxidative stress is associated with cancer, Alzheimer’s disease, diabetes, aging, obesity, Parkinson’s disease, inflammation, and other diseases [15]. Reactive oxygen species can induce protein oxidation, lipid oxidation, base modification, DNA strand break, and modulation of gene expression [16].

The *Osmanthus fragrans* var. *aurantiacus*, as an evergreen tree, belongs to the Oleaceae family. It is a rich source of iridoid, secoiridoid, phenylpropanoid, and lignan glycosides [17]. In a previous study, we determined the anti-inflammatory activities of fractions and triterpenoids from *O. fragrans* var. *aurantiacus* leaves [18]. Natural products are affected not only by the species and tissues used for extraction, but also by the extraction process such as temperature, pretreatment time, and extraction time [19]. The purpose of this study was to discover phytochemicals with anti-inflammatory activities in *O. fragrans* var. *aurantiacus* leaves extract (OFLE) using optimal extraction methods. Various extraction conditions were tested to find the optimal one for preparing an extract with potent anti-inflammatory properties. To determine the antioxidant activity of OFLE, radical scavenging activity was assessed using 1,1-diphenyl-2-picrylhydrazyl (DPPH) and 2,2-azino-bis-30ethylbenzthiazoline-6-sulphonic acid (ABTS) assays. In addition, total phenolic acid and flavonoid contents in the OP OFLE were analyzed. Finally, the anti-inflammatory activities of OP OFLE and its major components were evaluated using LPS-stimulated RAW 264.7 and HT-29 cells.

2. Results

2.1. Effects of OFLEs Extracted at Various Solvent Ratios on LPS-Induced NO Production and Cytotoxicity

RAW 264.7 cells were pretreated with eleven OFLEs (extracted with 0, 10, 20, 30, 40, 50, 60, 70, 80, 90, and 100% EtOH for leaves harvested in June) at various concentrations (10, 25, 50, and 100 µg/mL) for 1 h and then induced with LPS (1 µg/mL) for 20 h. The control group was not treated with either LPS or OFLEs. The supernatant (100 µL) was harvested and then NO production was measured with a Griess reagent. Viabilities of cells treated with the OFLEs were measured by MTT assay. Effects of eleven OFLEs (0, 10, 20, 30, 40, 50, 60, 70, 80, 90, 100% EtOH extracts) on NO production and cell viability were then analyzed. Results showed that all OFLEs inhibited LPS-induced NO production in a concentration-dependent manner. They showed no cytotoxicity (Figure 1A). Such inhibitory effects of OFLEs on NO production were not attributable to their effects on cell viability. Inhibitory activities of OFLEs extracted with 60, 70, and 80% EtOH on NO production (71.3, 61.6, and 41.1% inhibition, respectively) were significantly higher than those of other OFLEs at a concentration of 100 µg/mL (Figure 1B).
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Inhibitory activities of OFLEs extracted with 60, 70, and 80% EtOH on NO production (71.3, 61.6, and 41.1% inhibition, respectively) were significantly higher than those of other OFLEs at a concentration of 100 μg/mL (Figure 1B).

Figure 1. Effects of OFLEs obtained with various solvent ratios on cell viability and NO production. RAW 264.7 cells were cultured in the presence of eleven OFLEs (harvested in June) at various concentrations (10, 25, 50, and 100 μg/mL) for 1 h and then induced with LPS (1 μg/mL) for 20 h. Cell viability (A) and NO production (B) were measured using MTT assay and Griess reagent, respectively. Nitrite concentrations of non-treated and LPS-treated controls were 0.9 ± 0.1 μM and 17.6 ± 0.2 μM, respectively. Each experiment was performed in triplicates. Data are presented as mean ± SD. * p < 0.05, ** p < 0.01, *** p < 0.001 vs. LPS-treated group.

2.2. Effects of OFLEs Extracted with Various Solvent Ratios for Leaves Harvested at Different Time on LPS-Induced NO Production and Cell Viability

After the first experiment, nine OFLEs (extracted with 60, 70, and 80% EtOH for leaves harvested in March, June, and September) at different concentrations were also evaluated for their effects on NO production and cell viability. Inhibitory effects of OFLEs on NO production were not attributable to their effects on cell viability (Figure 2A). For leaves harvested in June, the OFLE extracted with 60% EtOH showed a significantly higher inhibitory activity (69.1% inhibition) on NO production than other OFLEs at a concentration of 100 μg/mL (Figure 2B).
leaves harvested in June, the OFLE extracted with 60% EtOH showed a significantly higher inhibitory activity (69.1% inhibition) on NO production than other OFLEs at a concentration of 100 μg/mL (Figure 2B).

Figure 2. Effects of OFLEs obtained from leaves with different harvest time and various solvent ratios on cell viability and NO production. Inhibitory effects of different harvest time OFLE (10, 25, 50, and 100 μg/mL) by solvent ratio in LPS-induced RAW 264.7 cells. Cells were treated with each extract for 1 h and then stimulated with LPS (1 μg/mL) for 20 h. Cytotoxicity (A) was determined by MTT assay. Culture media were evaluated for NO production (B) using Griess reagent. Each determination was conducted in triplicates. Data are represented as mean ± SD. * p < 0.05, ** p < 0.01, *** p < 0.001 vs. LPS-treated group.

2.3. Effects of OFLEs Obtained with Various Extraction Time on LPS-Induced NO Production and Cell Viability

After the second experiment, five OFLEs obtained with a sonication time of 60, 90, 120, 150, and 180 min at different concentrations were evaluated for their effects on NO production and cell viability. Results showed that their inhibitory effects on NO production were not attributable to their effects on cell viability (Figure 3A). The OFLE obtained from leaves harvested in June and extracted with 60% EtOH for 120 min showed significantly higher inhibitory activity (69.8% inhibition) on NO production than other OFLEs at a concentration of 100 μg/mL (Figure 3B). Therefore, extracting leaves harvested in June...
with 60% EtOH condition for 120 min was derived as the optimal extraction method for OFLE. Such extract was named OP OFLE.

Figure 3. Effects of OFLEs obtained with various extraction time on cell viability and NO production. Effect of OFLEs (10, 25, 50, and 100 µg/mL) obtained with different extract time (60, 90, 120, 150, and 180 min) on LPS-stimulated cell viability (A) and NO production (B) in mouse macrophages. Cells were cultured in the presence of OFLE at various concentrations for 1 h before stimulation with 100 µg/mL LPS. At 20 h, the culture supernatant was assayed for cell viability and NO production. Values are presented as mean percentage ± SD of five samples from three different experiments compared with the corresponding LPS-induced control (100%). *p < 0.05, **p < 0.01, ***p < 0.001 vs. LPS-treated group.

2.4. Effects of OP OFLE and OFLE Extracted with 60% MeOH on LPS-Induced NO Production and Cell Viability

In the last experiment, we evaluated the effects of OP OFLE and OFLE extracted with 60% MeOH on NO production in LPS-stimulated mouse macrophages. OP OFLE and the OFLE extracted with 60% MeOH suppressed the production of NO at concentrations of 25, 50, and 100 µg/mL. In addition, they showed no cytotoxicity (Figure 4A). When these two extracts were compared, OP OFLE showed a higher inhibition effect on NO production.
than the methanol extract at all concentrations. In particular, OP OFLE at 100 µg/mL showed the highest NO inhibitory activity, resulting in an inhibition of 67.6% (Figure 4B).

![Figure 4](image-url)

**Figure 4.** Effects of OP OFLE and MeOH OFLE on cell viability and NO production. Inhibitory effects of OP OFLE and MeOH OFLE (10, 25, 50, and 100 µg/mL) on cell viability and NO production of LPS-stimulated RAW 264.7 cells were determined. Cells were pretreated with each extract at different concentrations for 20 h. Cytotoxicity (A) was determined by MTT assay. Culture media were used to determine NO production (B) with Griess reagent. Data are represented as mean ± SD. * p < 0.05, ** p < 0.01 vs. LPS-treated group.

2.5. DPPH and ABTS Radical Scavenging Activities of OP OFLE

DPPH and ABTS assays were performed to investigate the antioxidant activities of OP OFLE at 50 and 100 µg/mL. In this experiment, ascorbic acid (100 μM) was used as a positive control. OP OFLE exhibited a concentration-dependent radical scavenging activity. It showed a strong antioxidant activity in both DPPH and ABTS experiments. In the DPPH assay, it showed 45.4% radical scavenging activity at 100 µg/mL. In the ABTS assay, it showed 42.6% radical scavenging activity (Figure 5).
Figure 5. DPPH radical scavenging activity (A) and ABTS radical scavenging activity (B) of OP OFLE (50 and 100 µg/mL) are shown. Data are presented as mean ± SD (n = 3) of three individual experiments. *p < 0.05, **p < 0.01, ***p < 0.001.

2.6. Determination of Total Phenolic Content (TPC)

TPC indicates the phenolic content in a sample. Phenolic compounds present in plants have redox properties. Such properties allow them to act as antioxidants. Results were derived from a calibration curve \( y = 0.0077x - 0.0224, R^2 = 0.9925 \) of gallic acid (Sigma Aldrich, St. Louis, MO, USA) and expressed in gallic acid equivalents (GAE) per gram of dry OP OFLE weight. The TPC in OP OFLE was 31.5 mg GAE/g. High phenolic content is responsible for the bioactivity of OP OFLE. Therefore, this extract is expected to reveal good antioxidant and antibacterial activities.

2.7. Determination of Total Flavonoid Content (TFC)

TFC indicates the content of flavonoids in a sample. Results were derived from a calibration curve \( y = 0.0013x - 0.0554, R^2 = 0.9966 \) of quercetin (Sigma Aldrich, St. Louis, MO, USA) and expressed in quercetin equivalents (QE) per gram of dry extract weight. The TFC in OP OFLE was 18.5 mg QE/g. High flavonoid content is responsible for the bioactivity of OP OFLE. Therefore, this extract is expected to show good antioxidant and antibacterial activities.
2.8. Isolation of PH, Lignan Glycode from OP OFLE

OP OFLE was subjected to column chromatography and assigned a major compound, which was identified as lignan glycoside, PH, by analyzing its spectral data, including nuclear magnetic resonance (NMR) and liquid chromatography-mass spectroscopy (LC-MS) data with those reported studies (Figure 6) [20].

\[ O.\ fragrans\ var.\ aurantiacus\ leaves\]

Extracted with DW/EtOH, respectively
Evaporated in vacuo

\[ \text{<LC condition>}\]
- Column: Shiseido C₁₈ UG120 (250 x 4.6 mm)
- Mobile phase: (A) Distilled water (B) Acetonitrile
- Temperature: 30 °C
- Flow rate: 1 mL/min

\[ \text{OP OFLE}\]
Separated using multi-gradient RP HPLC system

\[ \text{Phillyrin}\]

\[ \text{<HPLC chromatogram>}\]

Figure 6. Isolation scheme and structure of PH isolation from OP OFLE.

2.9. Effects of OP OFLE and PH Treatment on iNOS and COX-2 Expression

RAW 264.7 cells were used to determine the effects of OP OFLE (50, 100 µg/mL) and PH (100, 200 µM) on protein expression. As shown in Figure 7, iNOS and COX-2 protein levels were increased significantly in the group induced with LPS without any pretreatment. However, pretreatment with OP OFLE or PH resulted in dose-dependent decreases of these protein expression levels induced by LPS. Compared to the control group without any pretreatment, OP OFLE at 100 µg/mL and PH at 200 µM reduced iNOS protein expression by 91.7% and 92.5%, respectively. Additionally, we assessed effects of OP OFLE and PH on COX-2 protein through Western blot analysis. Both OP OFLE and PH inhibited its expression. Pretreatment with OP OFLE at 100 µg/mL showed the highest inhibitory effect on COX-2 protein expression, resulting in a 41.8% reduction. PH at 100 and 200 µM also attenuated the expression of COX-2 protein upregulated by LPS stimulation by 54.9% and 91.8%, respectively. The expression of LPS-stimulated COX-2 protein was inhibited by 41.8% and 91.8% after treatment with OP OFLE (100 µg/mL) and PH (200 µM), respectively (Figure 7).
2.10. Effects of OP OFLE and PH Treatment on NF-κB Activation and pERK1/2 Expression

It has been reported that NF-κB activation was induced by IL-1β, IL-6, TNF-α, and iNOS in RAW 264.7 cells [21]. To investigate the anti-inflammatory effects of OP OFLE and PH, Western blotting was carried out after RAW 264.7 macrophage cells were treated with OP OFLE (50, 100 µg/mL) or PH (100, 200 µM) for 2 h prior to LPS stimulation to determine their inhibitory effects on NF-κB and pERK1/2. Results show that OP OFLE and PH downregulated the expression of NF-κB and pERK1/2 in RAW 264.7 cells in a concentration-dependent manner, with OP OFLE at 100 µg/mL and PH at 200 µM showing high inhibitory effects. These results verified that OP OFLE and PH could suppress NF-κB and pERK1/2 protein expression levels (Figure 8).

![Western blot images showing downregulation of NF-κB and pERK1/2](image)

**Figure 8.** Western blot images showing downregulation of NF-κB and pERK1/2 in RAW 264.7 cells treated with OP OFLE (50, 100 µg/mL) or PH (100, 200 µM) for 2 h prior to LPS stimulation.

| Sample (µM) | OP OFLE (µg/mL) | PH (µM) |
|-------------|-----------------|---------|
| LPS (1 µg/mL) | - | 50, 100 | 100, 200 |
| iNOS | - | + | +, + |
| COX-2 | - | + | +, + |
| β-actin | - | - | - |

2.11. Effects of OP OFLE and PH on Expression Levels of Pro-Inflammatory Cytokines IL-1β, IL-6, IL-8, and TNF-α

Activated HT-29 cells and RAW 264.7 cells can produce pro-inflammatory cytokines such as IL-1β and IL-6 [22]. The inhibitory effects of OP OFLE and PH on pro-inflammatory mediators were evaluated using ELISA kits. Stimulation with LPS significantly increased...
levels of pro-inflammatory cytokines. However, pretreatment with OP OFLE or PH decreased levels of pro-inflammatory cytokines in a concentration-dependent manner. Particularly, IL-1β production was significantly inhibited by OP OFLE at 100 µg/mL and PH at 200 µM. In addition, OP OFLE and PH down-regulated IL-8 production in LPS-stimulated HT-29 cells. These results showed that OP OFLE and PH could suppress pro-inflammatory cytokines (Figure 9).
Figure 8. Effects of OP OFLE and PH on pERK1/2 activation and NF-κB expression. Inhibition of the OP OFLE (50 and 100 μg/mL) and PH (100 and 200 μM) on LPS-induced pERK1/2 (A) and NF-κB (B) proteins in RAW264.7 macrophages. Cells were pretreated with OP OFLE or PH at different concentrations for 2 h and then stimulated with LPS for 1 h. Expression of pERK1/2 and NF-κB proteins were detected by Western blot using specific pERK1/2 and NF-κB antibodies. β-Actin was used as a loading control. Relative density of NF-κB compared to that of β-actin was calculated. Blots shown are representatives of three independent experiments with similar results. * p < 0.05, ** p < 0.01, *** p < 0.001 vs. LPS-stimulated group.

Figure 9. Effects of OP OFLE and PH on expression levels of IL-1β, IL-6, TNF-α, and IL-8. Inhibitory effects of OP OFLE (50 and 100 μg/mL) and PH (100 and 200 μM) on IL-1β (A), IL-6 (B), TNF-α (C), and IL-8 (D) expression levels in LPS-induced RAW 264.7 cell and HT-29 colonic epithelial cells were determined. These four proinflammatory cytokines were assessed with ELISA kits. Data are expressed as mean ± SD (n = 3) of three individual experiments. * p < 0.05; ** p < 0.01; and *** p < 0.001 vs. LPS-stimulated group.
3. Discussion

Nowadays, there is an increasing recognition that inflammation is associated with the pathogenesis of diverse human diseases ranging from infection to immune-mediated disorders, diabetes, cancer, cardiovascular pathology, neurodegeneration, metabolic syndrome, and aging itself [23]. Inflammation is a normal immune process that responds to tissue damage, microbial pathogen infection, and chemical irritation. However, prolonged inflammation can cause various chronic diseases [24]. The necessity of discovering new anti-inflammatory drugs becomes important. Great effort has been made to develop drugs for treating inflammation [25].

During our search for candidates of botanical drugs to control inflammation, *O. fragrans* var. *aurantiacus* leaves have been found to possess anti-inflammatory phytochemicals. In this study, we focused on extracts of *O. fragrans* var. *aurantiacus* leaves showing the highest bioactivity. In addition, representative active ingredients were determined. To find the optimal extraction method for OFLE, different solvent ratios, harvest time of leaves, and extraction time were used. Each OFLE was then tested for its inhibitory effect on NO production. It was observed that leaves harvested in June and extracted with 60% EtOH for 120 min showed the highest inhibitory effect on NO production. We also evaluated the effect of OFLE extracted with MeOH on NO production. Results revealed that OP OFLE had a stronger inhibitory effect on NO production than MeOH OFLE. OP OFLE was then subjected to HPLC analysis to elucidate its major active constituents, yielding PH.

A previous study revealed that phenylpropanoids and lignans, such as phenylpropanoid dimers, are the main ingredients in *Osmanthus* species that possess biological activities such as antioxidant, antifungal, and antibacterial effects [26]. In particular, it has been reported that lignans including phillygenin and phylliryn from *O. fragrans* var. *aurantiacus* flowers have inhibitory activities on NO production [13]. This study evaluated the potential of OP OFLE and lignans isolated from OFLE to treat inflammation. In macrophages, LPS as a well-known endotoxin can induce the production of inflammatory mediators such as NO and prostaglandin E2 (PGF2) synthesized by iNOS and COX-2 and inflammatory cytokines such as IL-1β and TNF-α [27]. We examined the cell viability of macrophages and inhibitory effects of these extracts on NO production. Results showed that the extract suppressed NO production without showing any cytotoxicity OP OFLE and PH lignan showed strong NO production inhibitory effects. Their treatment decreased iNOS and COX-2 levels in LPS-induced RAW 264.7 cells.

MAPKs (including JNK kinase and p38MAPK) and NF-κB can activate some proinflammation cytokines [28]. NF-κB commonly regulates the promoter region of many proinflammatory cytokines. In activated macrophages, NF-κB in synergy with other transcriptional activators plays a central role in coordinating the expression of genes encoding iNOS, IL-1, IL-6, and TNF-α [29]. OP OFLE and PH decreased the phosphorylation levels of pNF-κB and pERK1/2 (Figure 8). We evaluated levels of IL-1β, IL-6, and TNF-α proteins upon treatment of cells with OP OFLE and PH and revealed that the levels of these LPS-stimulated pro-inflammatory cytokines were suppressed in macrophage and colonic epithelial cells (Figure 9). These results showed that OP OFLE and PH could potentially be considered therapeutic agents for treating inflammatory diseases by down-regulating inflammatory mediators.

Phenolic compounds including flavonoids, phenylpropanoids, and lignans are secondary metabolites of plants [30]. They have attracted increasing attention in traditional medicine for cancer prevention or treatment due to their ability to act as antioxidant agents [31]. In addition, flavonoids that are readily consumed by humans appear to exhibit anti-allergic, anti-cancer, and anti-inflammatory activities [32]. Flavonoids can exert antioxidant activities by donating hydrogen atoms to free radicals to suppress the generation of free radicals excessively produced by oxidative stress in vivo [33]. Lignans exhibit numerous biological activities, including antiproliferative and antioxidant effects. In addition, they have protective functions against cardiovascular diseases and other degenerative pathologies associated with oxidative stress [34]. As a result of evaluating TPC and TFC,
O. fragrans var. aurantiacus leaves were found to contain high levels of phenolic compounds and flavonoids. O. fragrans var. aurantiacus leaves possess high amounts of phenolic compounds, including lignan and flavonoids with potent antioxidant activities. Therefore, they have disease fighting properties and provide various defenses. Additionally, the antioxidant effects of OP OFLE were assessed with DPPH and ABTS assays by measuring radical scavenging capacity [35]. Results of DPPH and ABTS assays showed that OP OFLE at 100 µg/mL concentration exhibited high radical scavenging activities (DPPH and ATBS: 45.4% and 42.6%, respectively).

In summary, we determined the optimal extraction method by evaluating the anti-inflammatory effects of OFLEs on NO production in LPS-induced macrophages. OP OFLEs exhibited potent DPPH and ABTS radical scavenging activities and inhibited NO production. Compounds 1–3 isolated from leaves of O. fragrans var. aurantiacus showed no cytotoxicity. OP OFLE and PH isolated as a major compound from it downregulated the iNOS, COX-2, pNF-κB, and p-ERK levels. In addition, levels of pro-inflammatory cytokines such as IL-1β, IL-6, IL-8, and TNF-α were decreased by treatment with OP OFLE and PH.

We found that the anti-inflammatory activity of PH isolated from O. fragrans var. aurantiacus leaves is due to different mechanisms of action. On the basis of previous studies and our results, OP OFLE could be used as a potential antioxidant with anti-inflammatory properties and applied to cosmetics, health functional foods, and pharmaceutical compositions.

4. Materials and Methods

4.1. Plant Materials

Leaves of O. fragrans var. aurantiacus were collected from Seoul National University Forests, Gwangyang, Korea in March, June, and September of 2018. A voucher specimen (SCNUP 21) was deposited at the laboratory of Pharmacognosy, College of Pharmacy, Sunchon National University, Suncheon, Jeonnam, Korea.

4.2. Preparation of Extracts

To prepare ethanol extracts, 1 g of ground O. fragrans var. aurantiacus leaves was mixed with 10 mL of 0, 10, 20, 30, 40, 50, 60, 70, 80, 90, or 100% EtOH and extracted by sonication at room temperature (RT) (2 h × 2 cycles). Extracts were filtered through No. 2 Whatman filter paper (Whatman, Pleasanton, CA, USA) and evaporated in vacuo at 39 °C using a rotary evaporator (Eyela; Tokyo, Japan). Finally, concentrated extracts were kept in dark at 4 °C. After the first experiment, three candidates (extracts with 60, 70, and 80% EtOH) were selected. Leaves were then extracted by sonication according to their harvest time (March, June, and September). After the second experiment, the harvest time and the extraction solvent mixture ratio condition (June, 60% EtOH) were selected and sonication extraction time was performed at five conditions (60, 90, 120, 150, 180 min). After the first, second, and third experiments, the optimum extraction condition for leaves of O. fragrans var. aurantiacus was derived as extraction of leaves harvested in June with 60% EtOH for 120 min. In the final experiment, the OP OFLE was compared with the extract obtained with 60% methanol.

4.3. Extraction and Isolation

The OP OFLE was additionally obtained by extraction 10 g of leaves and then dissolved with 100% MeOH and separated by column chromatography. Compounds (3.4 mg, tR 23.38 min) were obtained after isolation using a multi-gradient reverse phase (RP) HPLC system (Shiseido, C18, 250 × 10 mm, H2O-CH3CN = 95:5 → 30:70, 1 mL/min) and were then subjected to proton NMR (1H-NMR), carbon NMR (13C-NMR), and LC-MS analyses.

Phillyrin (PH): C27H34O11; white amorphous powder; 1H-NMR (400 MHz, C5D5N): δH 6.85–7.05 (2H, m, H-2, 2′), 6.85–7.05 (2H, m, H-5, 5′), 6.85–7.05 (2H, m, H-6, 6′), 4.87 (1H, d, J = 6.4 Hz, Glc-1), 4.10 (1H, d, J = 9.6 Hz, H-9α), 3.75 (3H, s, 3-OCH3), 3.75 (3H, s, 3′-OCH3), 3.75 (3H, s, 4-OCH3), 3.36 (2H, d, J = 6.8 Hz, H-7, 7′), 3.05–3.71 (4H, m, H-8), 2.84 (1H, m, H-8); 13C NMR (100 MHz, C5D5N): δC
148.9 (C-3, 3'), 145.9 (C-4, 4'), 135.3 (C-1, 1'), 118.2 (C-6, 6'), 115.2 (C-5, 5'), 110.4 (C-2, 2'), 100.1 (Glc-1), 86.7 (C-7, 7'), 77.0 (Glc-5), 76.9 (Glc-3), 73.2 (Glc-2), 70.3 (C-9, 9'), 69.7 (Glc-4), 60.7 (Glc-6), 55.7 (3-OCH₃), 55.3 (3'-OCH₃), 55.5 (4-OCH₃), 54.0 (C-8, 8'); ESIMS m/z 535.32 [M + H]+.

4.4. Cell Culture

Human colon epithelial cells, HT-29 cells, and mouse macrophages, RAW 264.7 cells, were purchased from Korean Cell bank (Seoul, Korea). These cell lines were cultured in an incubator at 37 °C with a humidified atmosphere and 4.5% CO₂. These cell lines were grown in Dulbecco’s modified Eagle’s medium (DMEM) (Hyclone, Logan, UT, USA) supplemented with 100 µg/mL streptomycin solution (Hyclone, Logan, UT, USA), 10% fetal bovine serum (FBS) (HyClone, Logan, UT, USA), and 100 IU/mL penicillin.

4.5. Cell Viability Assay

RAW 264.7 cells were plated in a 96-well plate at a density of 1 × 10⁵ cells/well in DMEM containing 10% FBS and 1% penicillin for 24 h. Cells were treated with different concentrations of samples for 1 h and then stimulated with LPS at 1 µg/mL for 20 h. Cell viability was measured by 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl tetrazolium bromide (MTT) assay. Cultured cells were incubated with MTT (0.05 mg/mL) at 37 °C for 4 h. After removing the media, 100 µL DMSO was added. After shaking for 5 min, absorbance was measured at 570 nm using a microplate reader (Bio Tek Instruments, Winooski, VT, USA).

4.6. Measurement of NO Production

The supernatant of NO culture produced by RAW 264.7 cells were assessed by Griess reagent. The RAW 264.7 cells were planted at 1 × 10⁵ cells/well density in a 96-well plate and cultured in an incubator for 24 h. Cells were then treated with samples at different concentrations for 1 h and then stimulated with LPS for 20 h. The supernatant was then harvested and the same amount of the Griess reagent was added to the supernatant. The mixture was reacted in dark conditions at RT for 15 min and the absorbance was then measured at a wavelength of 550 nm using a microplate reader.

4.7. Measurement of Cytokine Production

Levels of IL-1β, IL-6, and TNF-α in RAW 264.7 cells and the level of IL-8 in HT-29 cells were measured using ELISA kits (BD OptEIA™, San Diego, CA, USA) according to the manufacturer’s instructions.

4.8. Western Blot Analysis

RAW 264.7 cells were seeded in 6-well plates at a density of 1 × 10⁵ cells/well. After treatment with OP OFLE (50, 100 µg/mL) and PH (100, 200 µM), cells were stimulated with LPS. Subsequently, protein was extracted using proprep solution (iNtRON, Biotechnology, Daejeon, Korea) for each treatment group. Protein concentrations were quantified using BCA solution through Bradford assay. The same amount of protein was separated by electrophoresis using 10% sodium dodecyl sulfate-poly acrylamide gel. Separated proteins were transferred to PVDF membranes using a transfer device (Thermo Fisher Scientific, Waltham, MA, USA). Membranes were then blocked with 5% skim milk power in platin buffer [4 M NaCl and 1 M Tris-HCl (PH 7.5)] for one hour at RT. Primary antibodies against iNOS, COX-2, NF-κB, ERK1/2, and pERK1/2 were incubated with the membrane for 24 h at 4 °C. Membranes were then washed with wash buffer (4 M NaCl, Tween-20, 1 M Tris-HCl (PH 7.5)) in DW and then incubated with a second antibody (1:2000 dilution) at RT for 2 h. Protein signals were detected with a bio image system (Microchemi 4.2 Chemilumineszenz-System, Neve Yamin, Israel) using ECL solution (Thermo Fisher Scientific, Waltham, MA, USA) [36].
4.9. Total Phenolic Content

Total phenolics content (TPC) of each extract was determined using the Folin–Denis method [37]. Briefly, 160 \( \mu \text{L} \) of sample was dissolved in EtOH (5, 10 mg/mL) and mixed thoroughly with 160 \( \mu \text{L} \) of Folin–Denis reagent (Sigma-Aldrich Co., St. Louis, MO, USA) for 3 min at RT. After adding 160 \( \mu \text{L} \) of 10% \((\text{w/v})\) sodium carbonate, the mixture was allowed to stand at RT for a further 60 min in the dark. After centrifugation at 10,000 rpm for 10 min, the supernatant (100 \( \mu \text{L} \)) was collected and its absorbance was measured at 700 nm. A standard curve was prepared using gallic acid (Sigma-Aldrich, St. Louis, MO, USA) with a concentration range from 0 to 500 \( \mu \text{g/mL} \). Total phenolic content was expressed as mg gallic acid equivalents (GAE)/g of sample.

4.10. Total Flavonoid Content

Total flavonoid content of sample was determined by the Moreno method [38]. Briefly, 10 \( \mu \text{L} \) of each extract (5, 10 mg/mL EtOH) was mixed with 4 \( \mu \text{L} \) of 10% aluminum nitrate (Sigma-Aldrich Co.), 4 \( \mu \text{L} \) of 1 M potassium acetate (Sigma-Aldrich Co.), and 82 \( \mu \text{L} \) of MeOH. The mixture was allowed to stand at RT for a further 40 min in the dark. Its absorbance was then measured at 415 nm. A standard curve was prepared using quercetin (Sigma-Aldrich Co.) with a concentration range from 0 to 500 \( \mu \text{g/mL} \). Total flavonoid content was expressed as mg quercetin equivalents (QE)/g of sample.

4.11. DPPH Radical Scavenging Activity Assay

The electron donating ability (EDA) was measured by modifying Amarowicz’s method [39] to determine DPPH radical inhibitory activity. Briefly, 0.1 mL of each sample solution (dissolved in EtOH) was mixed with 0.1 mL of 0.2 mM 2,2-diphenyl-1-picrylhydrazyl (DPPH, Sigma-Aldrich Co.) and allowed to stand at RT for 30 min. Its absorbance was then measured at 517 nm. The electron donating ability was expressed as the absorbance reduction rate of the sample solution added group compared to the no addition group using the following formula: DPPH radical scavenging activity (%) = (1 – absorbance of sample addition group/absorbance of no addition group) \times 100. As a positive control, 2,6-di-tert-butyl-4-methylphenol (BHT, Sigma-Aldrich Co.), a commercially available antioxidant, was used [40].

4.12. ABTS Radical Scavenging Activity Assay

ABTS radical inhibitory activity was measured by mixing 0.1 mL of each sample solution (dissolved in EtOH) and 0.1 mL of ABTS solution (7.4 mM 2,2-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid); ABTS) mixed with 2.6 mM potassium persulfate). After incubating at RT for 6 min, the absorbance of the mixture was measured at 734 nm. The electron donating ability was expressed as the absorbance reduction rate of the sample solution added group compared to the no addition group [40].

4.13. Statistical Analysis

All data are expressed as means ± standard deviation (SD) of at least three independent experiments. Comparison of data was achieved using Duncan’s multiple range tests with the software package SPSS statistics version 20. Statistical significance was indicated by asterisks (* \( p < 0.05 \); ** \( p < 0.01 \); and *** \( p < 0.001 \)).

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