Antioxidative and Anti-Aging Effects of *Pinus Rigida Mill*. Ethyl Acetate Extract on the Human Dermal Fibroblast Cell Line CCD-986sk Damaged by Ultraviolet B Radiation

Young Ah Jang¹, Bo Ae Kim² and Jin Tae Lee*³

¹Gennolab Co., LTD. Gyeongbuk 38540, Republic of Korea
²College of Sciences & Technology, Division of Biomedical & Cosmetics, Mokwon University, Daejeon 302-729, Korea
³Department of Cosmeceutical Science, Daegu Haany University, Gyeongbuk 38540, Republic of Korea

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*Corresponding author: Jin Tae Lee, Department of Cosmeceutical Science, Republic of Korea

Abstract

We analyzed the antioxidant and anti-wrinkle effects of *Pinus rigida Mill*. (pitch pine) to evaluate its use as a cosmetic material. 3-[4,5-Dimethylthiazol-2-yl]-2,5-diphenyl-tetrazoliumbromide assay was performed to evaluate the toxicity of the sample. To evaluate the anti-wrinkle activity of *P. rigida Mill.*, the amount of type I procollagen and TIMP-1 synthesized was measured, and Western blot analysis and RT-PCR were performed to determine the expression levels of wrinkle-related proteins and genes. To investigate the inhibitory effect of ROS of *P. rigida Mill.*, we conducted experiments to induce oxidative damage by UVB radiation using zebrafish embryos. Cytotoxicity to EtOAc extract (PRE) was measured using the human dermal fibroblast cell line CCD-986sk, and cytotoxicity was not observed below the concentration of 20 μg/mL. PRE has been shown to increase type I procollagen synthesis and promote the expression of its regulator, TIMP-1. In addition, *P. rigida Mill.* reduced the expression of MMP-1, -2, and -3 in CCD-986sk cells stimulated with UVB among the matrix metalloproteinases found in the dermal layers, which cause collagen reduction. After treatment with *P. rigida Mill.* extract at various concentrations, fluorescence spectra of ROS levels in the zebrafish embryos induced by UVB showed that fluorescence intensity decreased in a concentration-dependent manner compared with the intensity of the positive control group. Based on these findings, it is considered that *P. rigida Mill.* has value as a cosmetic material for antioxidant and anti-wrinkle benefits.

Keywords: Matrix Metalloproteinases; *Pinus Rigida Mill.*; Photo Aging; Reactive Oxygen Species; Zebra Fish

Introduction

Based on the characteristics of aging, it can be divided into endogenous and exogenous aging. Endogenous aging is caused by cell and tissue damage over time, abnormal immune function, and imbalance of epidermal homeostasis, whereas exogenous aging is caused by ultraviolet (UV) radiation, free radicals, toxicity, allergens, and physical stimuli [1]. A typical risk factor for exogenous aging is UV, and constant exposure of the skin to it triggers fibroblasts to produce reactive oxygen species (ROS), including superoxide anion, peroxides, and singlet oxygen, which are considered harmful to humans. This reaction causes structural changes in the extracellular matrix, such as changes in collagen, elastin, proteoglycan, and fibronectin [2, 3]. Excessively produced ROS can trigger DNA denaturation, which causes mutations in elastic fiber proteins and reduces collagen, leading to wrinkle formation and skin laxity [4]. Collagen reduction is caused by matrix metalloproteinase (MMP) in the dermal layers, which is activated by UV light [5, 6]. MMPs, such as zinc-dependent peptides involved in the reconstruction of the extracellular matrix (endopeptidases), are related to skin cell death, skin ulcers, tumor invasion, and metastasis [7]. Among MMPs, MMP-1 degrades type I and type III collagen, which comprise the largest portion of the extracellular matrix [5, 6]. MMP-3, also known as stromelysin 1, activates pro-MMP-1 by degrading proteoglycan, laminin, fibronectin, gelatin, type III collagen, and type IV collagen in the basement membrane [8, 9]. MMP-2 is a gelatinase that degrades collagen fragments
degraded by MMP-1 and degrades type IV collagen, playing a major role in skin aging [12]. Homeostasis in MMPs produced in the body is maintained via modulation of TIMP proteins (tissue inhibitors of MMPs and TIMPs) and is regulated by the action of their proteolytic enzyme system. TIMP-1 is a general inhibitor associated with the MMP family and has various functions, including regulating cell proliferation and apoptosis and controlling angiogenesis and inflammatory response [13]. Collagen degradation by MMP activation is caused by the signal transduction pathway of the transcription factor activator protein-1 (AP-1) via the activation of the mitogen-activated protein kinase pathway by oxidative stress such as UV light [14, 15]. Thus, suppression of the activity of MMP-1 and increase of collagen synthesis can maintain the elasticity of the skin tissue and prevent wrinkle formation and skin laxity. Recently, to reduce skin irritation caused by various synthetic substances, attention has been paid to functional cosmetic materials possessing antioxidant, anti-wrinkle, and whitening properties such as vitamin E, flavonoids, phenolic acid, and carotenoid obtained from natural plants. Pinus rigida Mill. (pitch pine), native to the eastern part of North America, is an evergreen coniferous tree belonging to the pine family. Among the coniferous trees found in Korea, P. rigida Mill. has the third largest distribution following pine and larch [16]. However, because its adhesiveness and paintability are poor, its use as timber is considered difficult. To date, various studies have been conducted to explore its increased efficient and high value-added applications [17]. Pine trees comprise leaves, pine cones, pollens, resin, and husks, whereas pine leaves that have been most actively studied contain essential oil components such as α-pinene, β-pinene, and camphene; flavonoids such as quercetin and kaempferol; and resins. Further, it has been reported to be effective in antioxidative and antibacterial and microorganism growth inhibition [18-20]. Pine bark contains approximately 20%-30% phenolic compounds. Dihydroquercetin-3'-O-β-galactoside catechin, epicatechin-(4β-6)-catechin, catechin-(4α-8)-catechin, catechin-(4α-6)-catechin, and epicatechin-(4β-6)-catechin are known compounds of proanthocyanidin [21, 22]. In this study, rigid pine bark was applied to UVB-damaged human fibroblasts, and the effect of photosaging through the expression of ROS and inhibition of MMPs was examined to verify the functionality of the bark as a cosmetic material.

Materials and Methods

Plant material

The inner bark of P. rigida Mill. used in this experiment was collected in November 2016 from P. rigida stand in Gyeongsan City (Gyeongbuk-do) by removing the outer bark. The specimens and extracts have been stored in Skin Immunity Pharmacological Laboratory of Department of Cosmeceutical Science, Daegu Haany University.

Sample Extraction

Ethanol extract of ligand was loaded with 10-times the amount of acetone and water in a ratio of 7:3(v/v). Following 3-day extraction, the solution was filtered using a filter paper (Whatman No. 2, Tokyo, Japan), and the obtained acetone crude extract was concentrated under reduced pressure. Then, the crude extract was suspended in water. Fraction extraction was then conducted thrice with the same amount of CHCl₃, and the water layer was extracted thrice with same amount of EtOAc. As a result, EtOAc and water extracts were obtained. In this study, EtOAc extract (PRE) obtained from P. rigida Mill. was used.

Cell Strain Culture and Cytotoxicity Measurement

The human fibroblast cell line CCD-986sk used for cytotoxicity measurement was purchased from American Culture Collection (ATCC; Manassas, VA, USA). Dulbecco’s modified Eagle’s medium (DMEM), fetal bovine serum (FBS), penicillin/streptomycin, and 0.4% trypan blue stain were purchased from Gibco BRL Co. (Rockville, NE, USA) for cytotoxicity measurement and culture, and 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyl-tetrazoliumbromide (MTT) was purchased from Sigma Chemical Co. (St. Louis, MO, USA). Primary and secondary antibodies to MMP-1, MMP-2, MMP-3, and β-actin were procured from Santa Cruz (Santa Cruz, CA, USA). Each cell was cultured using DMEM culture medium supplemented with 10% FBS and 1% penicillin/streptomycin (100 U/mL) and subcultured in an incubator at 37 °C and 5% CO₂.

MTT Assay Cytotoxicity Measurement for Safety Evaluation

CCD-986sk cells were uniformly dispensed in a 96-well plate (0.18mL) at a density of 5 × 10⁴ cells/well. PRE was prepared at concentrations of 5, 10, 20, 40, 60, 80, and 100μg/mL and 0.02 mL of the solutions was added. Then, the cells were cultured in a 5% CO₂ incubator at 37°C for 24h. Further, 0.02mL MTT solution prepared at a concentration of 5mg/mL was added, and the solution was cultured for 4h. Following completion, the culture solution was removed, and 0.15mL DMSO was added to each well. After incubation at room temperature for 15min, absorbance was measured at 540nm with an ELISA reader. Cytotoxicity was measured by the absorbance reduction rate of two groups: sample solution addition group and non-addition group.

Type I Procollagen Synthesis Measurement

To measure the amount of procollagen type I synthesized when the CCD-986sk cells were treated at different concentrations by PRE, the cells were dispensed in a 24-well plate at a density of 1 × 10⁵ cells/well and then stabilized for 24h. Next, the cultured medium was removed, 1mL PBS was added; the cells were irradiated with UVB (20mJ/cm²) and treated with PRE at different concentrations, and the cells were cultured for 48h. The supernatant from each well was then recovered and added to each well of procollagen Type I C-Peptide ELIA kit (Takara-Bio Inc., Shiga, Japan), and the total amount of procollagen type I was measured according to the manufacturer’s protocol.

Measurement of TIMP-1 Synthesis

After the CCD-986sk cells were cultured, the cells were inoculated in a 6-well plate at a density of 1 × 10⁴ cells/well and cultured for 24h. Following the 24h culture, the medium was...
removed and 1mL PBS was added; after UVB irradiation (20mJ/cm²), PBS was removed. After DMEM medium not containing 10% FBS and 1% penicillin streptomycin was substituted by diluting it with 5, 10, and 20μg/mL PRE, it was cultured for 48h. The supernatant was collected, TIMP-1 (R&D system Inc., MN, USA) was tested according to the kit protocol, and the absorbance was measured at 450nm with a microplate reader.

**Measurement of Protein Expression Using Western Blot**

CCD-986sk cells were uniformly dispensed in a 6-well plate at a density of 6 × 10⁴ cells/well and cultured for 24h. Then, PRE was irradiated with UVB at a dose of 20mJ/cm² and treated with the control, i.e., 25μg/mL epigallocatechin gallate (EGCG), at concentrations of 5, 10, and 20μg/mL and cultured for 48h. The cells were lysed with RIPA buffer (Pierce, IL, USA) and centrifuged (12,000 rpm, 4°C, 30min), and the cell membrane components were then removed to obtain the supernatant. Next, proteins were quantified by Bradford assay. It was then transferred with tris buffer nonspecific binding of the antibody. It was then transferred at a voltage of 60V for more than 2h. After blocking with tris buffer electrophoresed using 10% sodium dodecyl sulfate polyacrylamide gel electrophoresis and transferred to a PVDF membrane to inhibit the control, i.e., 25μg/mL epigallocatechin gallate (EGCG), whereas the control EGCG was treated at a concentration of 25μg/mL for 48h. Total RNA was then washed thrice with PBS and collected, and the trizol reagent was isolated using Invitrogen (Carlsbad, CA, USA). Total RNA (2μg) and PCR primer octonucleotide were mixed with the RT-PCR mixer, and RT-PCR was performed. (Table 1) summarizes the primers used. CCD-986sk cells were lysed with lysis buffer and centrifuged at 12,000 rpm at 4°C for 20min. cDNA was prepared by incubation at 42°C for 1h under RT conditions, and reverse transcriptase was inactivated by incubation at 94°C for 5min. The subsequent PCR conditions were based on 30 cycles of denaturation at 94°C for 30s, annealing at 50°C for 1min, and extension at 72°C for 90s, and the optimal conditions were adjusted according to the type of target cDNA. The amplified cDNA was separated by electrophoresis using 1.2% agarose gel and confirmed by EZ-Capture MG.

**Table 1:** The sequences of the primers of MMP-1, MMP-2, MMP-3 and GAPDH.

| Gene      | Primer   | Sequence (5' → 3')                           |
|-----------|----------|---------------------------------------------|
| MMP-1     | Forward  | AGC GTC TGA CAC TAA GCT AA                  |
| MMP-1     | Reverse  | GTT TTC CTC AGA AAG AGC AGC AT              |
| MMP-2     | Forward  | TGG CCA TCC TCT TCA AAG TGG TAG G           |
| MMP-2     | Reverse  | CAC TGT CCA CCC CTC AGA GC                  |
| MMP-3     | Forward  | TTG TCC TTT GAT GCA GTC AGC                 |
| MMP-3     | Reverse  | GAT GTC GCC CAA AAG TGC                    |
| GAPDH     | Forward  | CACCAGTACAGGGTACAAC                        |
| GAPDH     | Reverse  | CATGGTGCTTTCGTCGCT                         |

**Zebrafish Breeding Condition and Embryo Acquisition**

Zebrafish (Danio rerio) was obtained from the Zebra Fish Bank of Chungnam National University and was used for the extension of the breeding period. All experimental protocols were approved and conducted according to the guidelines and regulations of the Animal Ethics Committee of Chungnam National University. Wild-type zebrafish were provided by the Zebrafish Center for Disease Modeling, Korea. The rearing condition was kept constant at 28°C ± 1°C with a 16-h/8-h light/dark cycle (Rotifer continuous culture system, GDBC, Korea). They were fed brine shrimp (Ocean Star International, Utah, USA). Male and female fish with at least five spawning experiences were fed adequate feed the day prior to spawning; 24 h later, irradiation of light was used to produce embryos. The produced embryos were incubated in an incubator at 28°C ± 1°C for 2 days in pre-manufactured Egg water (Sea salt solution, Aquarium systems, France) to avoid exceeding a maximum of 50 eggs in a 100-mm petri dish, and the Egg water was replaced periodically and used in the experiment.

**Oxidative Damage Induction Using**

UVB Spawned zebrafish embryos were dispensed into each 60-mm petri dish (5 embryos/dish), followed by 2-day culturing in an incubator (TempMini H2200-H, China) to obtain young fish. PRE was dissolved in the Egg water at concentrations of 5, 10, and 20μg/mL and treated with 5mL of each solution. The negative control group Egg water was treated with 5mL of each solution. After pretreatment of the sample, the sample was washed with the Egg water and irradiated for 1min at a dose of 50mJ/cm² UVB using a UV irradiator (UV-X000, LAB24, Korea). After UVB irradiation, 2'-7'- dichlorofluorescein diacetate (DCFH-DA, Sigma-Aldrich, USA) was applied and the solution was incubated for 1h in an incubator. Later, the zebrafish larvae were washed with the Egg water and observed with a glaucoma microscope (Dino-Lite, Taipei).
Statistical Analysis

Statistical analysis was performed using SPSS 10.0, and significant differences were verified at statistical significance level of p < 0.05, 0.01 using t-test using ANOVA (Analysis of Variance) program.

Results and Discussion

Cell Viability of PRE in UVB-Stimulated Fibroblasts

To ensure the synthesis of type I procollagen and expression of MMPs in the fibroblasts of PRE, cytotoxicity was first examined. The cells treated with PRE at concentrations of 5, 10, 20, 40, 60, 80, and 100μg/mL showed cell viability of above 90% at concentrations below 20μg/mL. It was considered that there was no toxicity, and the experiment was conducted at a concentration of 20μg/mL or less (Figure 1).

PRE Type I Procollagen and TIMP-1 Expression Rates

The dermis of the skin occupies most of the complete skin surface and reflects the characteristics of the skin such as flexibility, elasticity, and tension. The biosynthesis and degradation of the dermal collagen are important in skin aging and skin regeneration processes in wound healing [23]. Collagen, which accounts for 90% of the dermis, is divided into collagen types I-V and is synthesized as a procollagen precursor in the cell; then, it is secreted out of the cell and polymerized into collagen fiber [24]. Approximately, 80% of the collagen is type I collagen and is downregulated by UV light, and protein MMPs are involved in its degradation [25]. Procollagen is a precursor of collagen and has a peptidase called a propeptide. Propeptide is involved in collagen formation in the endoplasmic reticulum; therefore, the amount of collagen biosynthesis can be estimated by measuring the amount of propeptide to antagonize the effect of biosynthesis [26]. The results of the estimation of the amount of procollagen biosynthesis of PRE are shown in (Figure 2A). The control had an expression level of 100.0% ± 4.5% and the group irradiated with UVB (20mJ/cm^2) of 38% ± 2.9%; the levels noted for 10ppm PRE, 20ppm PRE, and EGCG were 48% ± 2.8%, 59% ± 1.9%, and 72% ± 3.5%, respectively.

Thus, it was confirmed that the expression level of procollagen increased by the addition PRE in a concentration-dependent manner. The action of MMPs that degrade collagen is regulated by inhibitors such as 2-macroglobulin or TIMP-1 [27,28], which are secreted to maintain cellular homeostasis. To date, four types of TIMP have been identified, and TIMP inhibits the function of MMP by forming strong non-covalent bonds with inactivated MMP at a ratio of 1:1 [29]. According to (Figure 2B), the control had an expression level of 100.0% ± 1.9%, whereas the irradiated group had a level of 43% ± 1.1%; the levels noted for 10ppm PRE, 20ppm PRE, and EGCG were 49% ± 2.3%, 51 ± 2.7%, and 62 ± 3.5%. Thus, it was confirmed that PRE increased the expression level of TIMP-1.
Protein synthesis and mRNA expression measurement of MMP-1, -2, and -3

MMPs are enzymes that degrade the extracellular matrix and are induced by AP-1 activated by UV light; they are secreted by many cells including human keratinocytes and fibroblasts, and their expression increases after UV irradiation in the normal epidermis [10]. Among MMPs, MMP-1 is known as collagenase 1 and involves type I and III collagen as substrates; MMP-2 is known as gelatinase and degrades type IV and type VII collagen; and MMP-3, also known as stromelysin 1, degrades laminin (an adhesive protein in the basement membrane) and breaks down type IV collagen, producing pro-MMP-1 (a zymogen) [11]. Reduction of total collagen production in the extracellular matrix of the skin following UV irradiation reduces elasticity in photoaged skin and causes skin laxity [30]. Therefore, various plant-derived natural materials capable of reducing the production of MMPs and increasing procollagen production can reduce photoaging. Western blot analysis was conducted to determine the effect of PRE extracts on MMP-1,-2, and -3 expressions by UVB irradiation of fibroblasts. The results are shown in (Figure 3). It was confirmed that as the concentration of PRE extract increased, MMP-1 decreased significantly compared with the untreated group.

![Figure 3: Expressions of MMP-1, -2 and MMP-3 evaluated by protein level in CCD-986sk fibroblasts after PRE treatment. CCD-986sk cells (1x10^5) were grown and exposed to UVB for 1 min; the cells were then subject to 5, 10, 20 μg/mL of PRE and 25 μg/mL of EGCG treatment. The protein levels were normalized using EZ-Capture MG. The results were expressed as the average of triplicate. *p<0.05, **p<0.01 compared with control.](image)

The expression level of MMP-1 was found to be 16% ± 2.5% and 35% ± 1.9% for 10 and 20 μg/mL PRE, respectively, and 28% ± 1.8% for 25 μg/mL EGCG. The expression level of MMP-2 was 31% ± 2.9% and 58% ± 2.4% for 10 and 20 μg/mL PRE, respectively, and 58% ± 2.2% for 25 μg/mL EGCG. The expression level of MMP-3 was 50% ± 2.9% and 49% ± 2.4% for 10 and 20 μg/mL PRE, respectively, and 58% ± 2.2% for 25 μg/mL EGCG, highlighting its efficacy as a MMPs inhibitor. RT-PCR was performed to observe the expression of mRNA according to the treatment with PRE extract, and the result of the measurement based on the expression of GAPDH as a housekeeping gene is shown in (Figure 4). The expression levels of MMPs increased in the UVB-irradiated groups. As PRE concentration increased, MMP-1 showed a significant reduction of more than 50% for 10 and 20 μg/mL PRE and MMP-2 decreased in a concentration-dependent manner. Therefore, PRE is believed to promote the expression of TIMP-1 in fibroblasts damaged by photoaging and to reduce the degradation of extracellular matrix proteins by active MMPs, resulting in the suppression of wrinkle formation.

![Figure 4: Expressions of MMP-1, -2 and MMP-3 evaluated by mRNA level in CCD-986sk fibroblasts after PRE treatment. CCD-986sk cells (1x10^5) were grown and exposed to UVB for 1 min; the cells were then subject to 5, 10, 20 μg/mL of PRE and 25 μg/mL of EGCG. The mRNA levels were normalized using EZ-Capture MG. The results were expressed as the average of triplicate. *p<0.05, **p<0.01 compared with control.](image)
Protective Effect of PRE in Zebrafish that Induced ROS with UVB

Zebrafish is readily available as a vertebrate with human-like genetic information and organ systems and exhibits a short interval between generations (Figure 5). Further, because zebrafish embryos remain transparent for several days following birth, their organs can be observed. Thus, it has recently been used as an animal model for various human diseases [31]. ROS measurement of UVB-induced zebrafish employs the property that DCFH-DA reacts with ROS in the form of DCFH, which is liberated from the acetyl group, to produce a fluorescent material [32]. After pretreatment of PRE extracts at concentrations of 5, 10, and 20μg/mL, the ROS levels in the zebrafish embryos induced by UVB damage were examined based on fluorescence intensity. As a result, it was confirmed that the fluorescence intensity decreased in a concentration-dependent manner compared with the positive control group only irradiated with UVB. In particular, the ROS level in the group treated with 20μg/mL PRE reduced by 19.6% compared with the level observed in the positive control group.

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