Apigenin inhibits UVA-induced cytotoxicity in vitro and prevents signs of skin aging in vivo

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Abstract. Apigenin (4',5,7-trihydroxyflavone) is a flavone that has been reported to have anti-inflammatory, antioxidant and anti-carcinogenic properties. In this study, we investigated the protective effects of apigenin on skin and found that, in experiments using cells, apigenin restored the viability of normal human dermal fibroblasts (nHDFs), which had been decreased by exposure to ultraviolet (UV) radiation in the UVA range. Using a senescence-associated (SA)-β-gal assay, we also demonstrate that apigenin protects against the UVA-induced senescence of nHDFs. Furthermore, we found that apigenin decreased the expression of the collagenase, matrix metalloproteinase (MMP)-1, in UVA-irradiated nHDFs. UVA, which has been previously identified as a photoaging-inducing factor, has been shown to induce MMP-1 expression. The elevated expression of MMP-1 impairs the collagen matrix, leading to the loss of elasticity and skin dryness. Therefore, we examined the clinical efficacy of apigenin on aged skin, using an apigenin-containing cream for clinical application. Specifically, we measured dermal density, skin elasticity and the length of fine wrinkles in subjects treated with apigenin cream or the control cream without apigenin. Additionally, we investigated the effects of the apigenin-containing cream on skin texture, moisture and transepidermal water loss (TEWL). From these experiments, we found that the apigenin-containing cream increased dermal density and elasticity, and reduced fine wrinkle length. It also improved skin evenness, moisture content and TEWL. These results clearly demonstrate the biological effects of apigenin, demonstrating both its cellular and clinical efficacy, and suggest that this compound holds promise as an anti-aging cosmetic ingredient.

Introduction

Aging is defined as the progressive accumulation of damage over time, leading to the disruption of functions at the cellular, tissue and organ levels. Eventually, disease and death are induced by a complex, multifactorial process, involving genetic, endogenous and environmental factors (1). The consequences of human aging are mostly visible in the skin, manifesting as increased wrinkling, sagging, uneven texture and decreased elasticity (2). It has been suggested that aged skin has a disrupted barrier function and altered permeability, resulting in a dry appearance and an enhanced risk of skin disorders (3,4). A number of processes are also impaired in aged skin, including angiogenesis, lipid and sweat production, immune function and vitamin D synthesis. This often results in impaired wound healing, atrophy, vulnerability to external stimuli and in the development of several benign and malignant diseases (3). Aging of the skin is induced by both intrinsic and extrinsic factors (5,6), all of which lead to a reduced structural integrity and the loss of physiological function (6). Critically, understanding the mechanisms of skin aging is necessary for developing improved skin care products that delay this process and reduce the hazardous effects of aging-inducing factors (5,7).

Apigenin (4',5,7-trihydroxyflavone), a member of the flavone subclass of flavonoids, is widely found in herbs, fruits and vegetables, and thus, is a substantial component of the human diet. It has been shown to possess a variety of biological characteristics, including antioxidant (8) and anti-inflammatory properties (9). It has also been shown to exert tumor-inhibitory effects (10) and to promote neurogenesis (11). Apigenin has also been shown to enhance wound healing and tissue repair in the skin of diabetic rats (12). During the process of wound healing, fibroblasts secrete collagen, and the formation of collagen-rich granulation tissue is vital for the pathophysiological mechanisms of wound closure (13).

Ultraviolet (UV) radiation that reaches the earth's surface is comprised of wavelength ranges referred to as UVB (280-315 nm) and UVA (315-400 nm); UV radiation within the UVC wave-
band (100-280 nm) is absorbed entirely within the atmosphere. UV irradiation is the main culprit implicated in premature skin aging, which is referred to as photoaging. A number of studies have explored the relative contributions of UVA and UVB to the aging phenotype (14-16). Notably, while UVA photons are, on average, 1,000 times less energetic than UVB photons, they are capable of inducing aging-related changes even in the dermis, partly due to their greater average depth of skin penetration than UVB photons (17).

Photoaging is characterized by the macro- and micro-structural deterioration of the skin, which includes damage to collagen fibers, the excessive deposition of abnormal elastic fibers and increased levels of glycosaminoglycans (18-20). Among these factors, matrix metalloproteinases (MMPs) are thought to play a major role in mediating UV-induced skin aging (17). The MMPs are a family of structurally related molecules, including collagenase-1 (MMP-1), collagenase-3 (MMP-13), gelatinases A and B (MMP-2 and MMP-9, respectively), stromelysin-1 (MMP-3), membrane-type MMPs, and others, all of which are capable of degrading components of the extracellular matrix, such as collagen, elastin, fibronectin, proteoglycans and laminin (21). In particular, MMP-1 initiates the degradation of type I and III fibrillar collagen, MMP-9 (gelatinase B) further degrades the collagen fragments produced, and MMP-3 (stromelysin-1) degrades type IV collagen and activates pro-MMP-1 (22-24).

In the present study, we examined the protective effects of apigenin on skin aging and demonstrated that apigenin induces anti-aging effects in skin by improving its barrier function and reducing UVA-induced damage.

Materials and methods

Cell culture. Normal human dermal fibroblasts (nHDFs; Lonza, Basel, Switzerland) were cultured in Dulbecco's modified Eagle's medium (DMEM; Gibco Life Technologies, Carlsbad, CA, USA), supplemented with 10% fetal bovine serum (FBS; Sigma-Aldrich, St. Louis, MO, USA) and 1% penicillin/streptomycin (Gibco Life Technologies) at 37°C in an atmosphere of 5% CO₂. Apigenin was purchased from Sigma-Aldrich and dissolved in dimethyl sulfoxide (DMSO).

UVA irradiation. The nHDFs (1x10⁵/well) were seeded into 6-well plates and cultured until 70-80% confluent. Prior to irradiation, the cells were washed twice with phosphate-buffered saline (PBS). Fresh PBS was then added, and the cells were irradiated with UVA light (25 J/cm²; UVA; UVA lamp; UVP, Inc., Upland, CA, USA). The radiation intensity was monitored by a fiber optic spectrometer system USB2000 (Ocean Optics, Dunedin, FL, USA). The control cells were treated identically, except for the exposure to UV light. Following irradiation, various concentrations (0-100 µM) of the treatment agent (apigenin) in fresh medium were added to cells at 37°C for 24 h.

Cell viability assay. The nHDFs were seeded at a density of 2x10⁵ cells/well in 96-well plates and incubated for 24 h. These cells were washed with PBS and fixed by treatment with 0.5 ml fixing solution/well (4% formaldehyde, 0.5% glutaraldehyde in PBS buffer, pH 7.2) for 1 h. The fixed cells were stained in staining solution mix (staining solution, 470 µl; staining supplement, 5 µl; 20 mg/ml X-Gal in dimethyl-formamide, 25 µl) for 24 h, at 37°C. After 1 day, 70% glycerol (1 ml/well) was added, and images were captured using an Olympus IX51 microscope (Olympus, Tokyo, Japan).

Isolation of total RNA and quantitative PCR. Total RNA was isolated using TRIzol reagent (Invitrogen Life Technologies, Carlsbad, CA, USA) according to the manufacturer's instructions. The purity and concentration of the RNA were evaluated using a MaestroNano®, a microvolume spectrophotometer (Maestrogen, Las Vegas, NV, USA), and cDNAs were synthesized using the miScript II RT kit (Qiagen, Hilden, Germany) according to the manufacturer's instructions. In order to evaluate the expression of MMP-1, quantitative PCR was performed using the following primers: forward, 5’-TCT GACGTGATCCTCCAGAGACGC-3’ and reverse, 5’-CATGG TGACACCAGTGACTGCAC-3’ using EvaGreen dye (Solis Dynex, Tartu, Estonia) with Line-Gene K software (Bioer Technology Co., Ltd., Hangzhou, China). The Ct value for each gene was normalized to β-actin using the following primers: forward, 5’-GGATTCCTATGTGGGCGACGA-3’ and reverse, 5’-CGCTCGGTAGGATCTTCAATGC-3’. The relative expression levels of each gene were calculated using the 2^-ΔΔCt method, as previously described (25).

Senescence-associated (SA)-β-galactosidase assay. The expression of galactosidase as a marker for senescent nHDFs was determined using the SA-β-galactosidase staining kit (BioVision, Inc., Milpitas, CA, USA) following the manufacturer's instructions. The nHDFs were seeded at a density of 2x10⁵ cells/well in 60 mm cell culture plates and incubated at 37°C until they were 90% confluent. The cells were then pre-treated with apigenin, irradiated with UVA, and incubated for 24 h. These cells were washed with PBS and fixed by

Human subjects and clinical evaluation. All clinical evaluations were approved by the Ethics Committee of the Korea Institute for Skin and Clinical Sciences and performed in accordance with the Declaration of Helsinki Principles. We enrolled 40 women, aged over 30 years, in a randomized and double-blinded clinical trial. The subjects were selected based on age and were not pregnant or nursing. All subjects were informed about the objective of the study, signed an informed consent, and agreed to use only our products for skin care during the study duration. Factors for dropping out of the trial included itching, erythema, or hindrance to evaluation by excessive drinking or smoking. Th subjects were divided into the control and experimental groups, each containing 20 subjects (control group, 44.40±5.97 years; experimental group, 45.30±6.29 years). All subjects were subjected to the same conditions, apart from the experimental group which

water-soluble tetrazolium salt (WST-1) assay. WST-1 solution was added to the cultured cells at a volume equal to 10% of the culture medium, and the cells were then incubated at 37°C for 1 h. Cell viability was evaluated by measuring the absorbance at 450 nm using an iMark microplate reader (Bio-Rad, Hercules, CA, USA).
was administered the test treatment. The study duration was 4 weeks, and no participants dropped out. Biometric parameters were measured 3 times: before application, and then at 2 and 4 weeks after application. An investigator also questioned the subjects about their condition and performed visual evaluations for skin disorders, such as erythema, itching, scaling, edema, tingling and burning sensations, at each visitation.

**Experimental procedures.** To investigate the effects of apigenin on dermal density, skin elasticity, skin texture, moisture, transepidermal water loss (TEWL) and fine wrinkles around the eyes (also known as crow’s feet), the subjects were instructed to apply 2 g of the test treatment to the face, including the eye rim, every morning and night for 4 weeks. The subjects and investigators were blinded to the test and control treatments. At each visit, all subjects washed with the cleanser provided and lay quietly in a room with a constant temperature (22±1°C) and humidity (45±5%), so that they would all be evaluated under the same conditions. The cream provided to the experimental group contained 1% (wt%) apigenin; whereas, the cream provided to the control group was prepared using the same volume of water in the place of apigenin.

**Measurement of skin elasticity.** To evaluate the improvement in skin elasticity, a DermaLab USB elasticity probe (Cortex Technology Inc., Hadsund, Denmark) was applied to the skin, and the results were analyzed using the associated application software, version 1.09. The measurement was performed by applying a single fixed elasticity probe on the left cheek of a subject. To analyze the measured value (in MPa), Young's modulus (E) was used, and the detected value is dependent on skin elasticity. To evaluate improvement, measurements were taken 3 times, before treatment and both at 2 and 4 weeks after application.

**Measurement of dermal density.** To evaluate dermal density, a DUB® SkinScanner (taberna pro medicum, Luneburg, Germany) was utilized. Dermal density was measured (in µm) 3 cm beside the left eye, applying a couplant for ultrasonic examination. The analysis range was limited to the region between the dermis and the upper panniculus. To evaluate improvement, measurements were performed three times, before treatment and both 2 and 4 weeks after first application.

**Measurement of length of crow's feet.** To evaluate the improvement of wrinkles, particularly crow’s feet, a Robo skin analyzer CSS50 (Inforward Inc., Tokyo, Japan) was used. All facial images were captured under the same position and with equal lighting. The capturing was performed 3 times at each evaluation, on the front, left and right sides of the face. To evaluate improvement, measurements were performed 3 times, before treatment and both at 2 and 4 weeks after application. We analyzed the captured images matching the facial feature points to reenact accurately, and the measurement unit was in mm.

**Evaluation of skin moisture.** To evaluate improvement in skin moisture, a DermaLab USB moisture probe (Cortex Technology Inc.) was applied to the skin, and the data were analyzed using the associated application software, version 1.09. All subjects were evaluated on the same region of the right cheek, 5 times consecutively, and we calculated the mean value, excluding the maximum and minimum values. To evaluate improvement, measurements were performed 3 times, before treatment and both at 2 and 4 weeks after application. The probe measures skin conductance in micro Siemens (µS), and the numerical value is dependent on skin moisture.

**Measurements of TEWL.** To evaluate improvements in TEWL, a DermaLab USB TEWL probe (Cortex Technology, Inc.) was applied to the skin, and the data were analyzed using the associated application software, version 1.09. The measurement was performed 5 times consecutively, on the right cheek of the subjects, and we calculated the mean value, excluding the maximum and minimum values. To evaluate improvement, measurements were performed 3 times, before treatment and both at 2 and 4 weeks after application.

**Measurements of facial evenness.** To evaluate improvements in facial evenness, a PRIMOS Lite system (field of view 45x30; GFMesstechnik GmbH, Teltow, Germany) was used, and the captured clinical images were analyzed using the associated imaging software, PRIMOS Lite version 5.6E. The images were captured 3 times consecutively, on the left side of the forehead of the subjects. We analyzed facial evenness by calculating surface roughness, Ra (average of all heights and Depths to the reference plane) value. The Ra value, which is the most well-used measurement for facial evenness, is the arithmetic mean of the absolute values within the total measurement range. To evaluate improvement, measurements were performed 3 times, before treatment and both at 2 and 4 weeks after application.

**Statistical analysis.** For cellular efficacy tests, all results are presented as the mean percentage ± standard deviation (SD) of 3 independent experiments. Differences with a P-value <0.05, as determined by the Student's t-test, were considered statistically significant. For clinical efficacy tests, statistical analyses were conducted using SPSS software (SPSS, version 17.0 for Windows; IBM SPSS, Armonk, NY, USA). Paired Student's t-tests were performed in the cases of repeated measurements on the same subject. To analyze subject questionnaires, the mean values, standard deviations and percentages were calculated. The formula used to measure the percentage change for each skin parameter was ‘Percentage change = [(A - B)/B] x100’, where A is defined as the individual value of any parameter at the 2-and 4-week visits, and B represents the zero hour of the assessed parameter.

**Results**

**Cytotoxicity of apigenin and UVA in human dermal fibroblasts.** To determine whether apigenin affects nHDF viability, the cells were exposed to apigenin at concentrations ranging from 0-100 µM for 24 h. As shown in Fig. 1, apigenin reduced cell viability by 1.23% at 1 µM, 3.63% at 5 µM, and 12.61% at 20 µM. The apigenin-induced cytotoxicity increased significantly at concentrations >50 µM. Therefore, we used the concentration of 20 µM as the maximum concentration in all subsequent experiments.

To evaluate the effects of apigenin on the viability of damaged cells, the nHDFs were irradiated with 25 J/cm² UVA,
and these cells were then treated with apigenin at various concentrations. As shown in Fig. 2, this dose of UVA reduced cell viability by 34.60%; however, treatment with 10 and 20 µM apigenin increased viability back to 90.00 and 98.92%, respectively, suggesting that apigenin protects cells from UVA-induced cytotoxicity.

Senescent cell detection assay. We then investigated the ability of apigenin to inhibit senescence, using a SA-β-galactosidase assay. When the cells were irradiated with 25 J/cm² UVA, the percentage of senescent cells was found to be as high as 61.29%. This number decreased in a dose-dependent manner to 50.49, 32.03 and 17.34% when cells were post-treated with 5, 10 and 20 µM apigenin, respectively (Fig. 3). These results indicate that UVA acts as a stimulator of senescence, and that apigenin can inhibit UVA-induced cellular senescence.

Analysis of MMP-1 mRNA expression. UVA radiation corresponds to 90-95% of solar UV radiation (26) and is mainly responsible for the high production of reactive oxygen species (ROS) in skin, leading to oxidative stress (27). ROS are able to induce several disruptive cellular processes, such as senescence, DNA cleavage, lipid peroxidation and cell death (28). In addition, UVA induces the expression of MMP-1 in dermal fibroblasts in vivo and stimulates the expression of MMP-1, MMP-2 and MMP-3 in cell culture, all of which are induced during wrinkling and skin aging (29,30). MMP-1, when generated from fibroblasts, has been reported to ultimately promote a decrease in collagen (23,31,32). Collagen is the most abundant protein in the dermis, and type-1 collagen, in particular, provides structure to skin and composes >90% of collagen in the body (33). We found that the mRNA expression of MMP-1 in the UVA-irradiated HDFs increased up to 2.91-fold as compared with the non-irradiated cells. However, treatment with 5, 10 and 20 µM apigenin reduced MMP-1 expression 2.43-, 1.85- and 1.31-fold, respectively (Fig. 4), indicating that apigenin inhibits the UVA-induced induction of MMP-1 expression.

Evaluation of dermal density. To evaluate the effects of apigenin on skin aging in vivo, we measured the density of the
dermis in subjects treated with a cream containing 1% apigenin. Using a DUB SkinScanner, we found that the subjects using the non-apigenin-containing control cream displayed a mean density of 50.41 µm before use, and densities of 50.02 and 50.05 µm after 2 and 4 weeks, respectively (Fig. 5). However, subjects using the apigenin-containing cream displayed a mean density of 49.96 µm before use, and densities of 55.31 and 62.32 after 2 and 4 weeks of application, respectively (Fig. 5). Dermal density measurement, represented as a mathematical value, is proportional to density, and these experimental data were statistically significant (P<0.001). To compare the results from the treatment and control groups, we calculated the improvement as a percentage based on the density values before and after application. Using this metric, the dermal density improvement was found to be -0.77 and -0.72% after 2 and 4 weeks of application, respectively. These results suggest that the topical application of apigenin enhances dermal thickness.

Evaluation of crow's feet length. We then measured the length of crow's feet in the subjects treated with the apigenin-containing cream and the controls. In the control group, the mean length was found to be 63.10 mm before application, and 63.95 and 64.25 mm after 2 and 4 weeks of application, respectively (Fig. 6). In the experimental group, the mean length was 65.05 mm prior to application, and 56.60 and 48.65 mm after 2 and 4 weeks of application, respectively (Fig. 6). The measured values for the experimental group were statistically significant (P<0.001). To compare the results from the control and experimental groups, we calculated the improvement as a percentage based on the values before and after application. For the control group, the percentage improvement was calculated as -1.35 and -1.82% after 2 and 4 weeks of application, respectively. Whereas for the experimental group, the percentage improvement was found to be 12.99 and 25.21% after 2 and 4 weeks of application, respectively. These data suggest that the topical application of apigenin can lead to a reduction in wrinkle length.

Evaluation of skin elasticity. The dermis is composed of an extracellular matrix consisting of fibrous proteins, such as collagen and elastin, and is involved in the regulation of skin elasticity. Factors such as ROS, UV, or age can cause skin damage, wrinkle formation and a reduction in elasticity through the structural denaturation of collagen and elastin (34). In order to examine the effects of apigenin on skin elasticity, we measured elasticity in subjects treated with apigenin-containing cream or the control cream using a DermaLab USB elasticity probe. In the control group, elasticity was found to be 7.38 MPa before application, and 7.33 and 7.27 MPa after 2 and 4 weeks of application, respectively. In the experimental group, elasticity was 7.40 MPa before application, and 9.14 and 10.60 MPa after 2 and 4 weeks of application, respectively (Fig. 7). The experimental group values were statistically significant (P<0.001). To compare the results from the control and experimental groups, we...
calculated the improvement as a percentage based on the values before and after application. For the control group, the percentage improvement was calculated to be -0.68 and -1.42% after 2 and 4 weeks of application, respectively. However, the experimental group showed an improvement of 23.60 and 43.34% after 2 and 4 weeks of treatment, respectively. These results suggest that the topical application of apigenin increases skin elasticity.

Apigenin-containing cream improves skin hydration. Keratinocyte moisture content is pivotal for maintaining moisture in the skin. Normal keratinocytes maintain 10-30% moisture; however, when the moisture content drops below 10%, keratinocytes are unable to maintain the skin’s barrier function, and skin becomes dry, acquires an uneven texture and produces wrinkles, accelerating senescence (4). To evaluate the effect of apigenin treatment on skin moisture, we analyzed the skin moisture content in our study subjects using the DermaLab USB moisture probe. We found that the moisture content in the control group was 391.25 µS before use, and 393.30 and 397.87 µS after 2 and 4 weeks of application, respectively (Fig. 8). By contrast, the moisture content in the experimental group, which used the apigenin-containing cream, was 399.48 µS before use, and 528.75 and 604.74 µS after 2 and 4 weeks of application, respectively (Fig. 8). To compare the results from the control and experimental groups, we calculated the degree of improvement as a percentage based on the values before application. In the control group, moisture was increased by 0.52 and 1.69% after 2 and 4 weeks, respectively. These changes were not statistically significant (P>0.05), indicating that the control cream had no measurable effect on moisture content. Conversely, the use of the apigenin-containing cream significantly improved skin moisture; however, when the moisture content drops below 10%, keratinocytes are unable to maintain the skin's barrier function, and skin becomes dry, acquires an uneven texture and produces wrinkles, accelerating senescence (4). To investigate the effects of the apigenin-containing cream on skin texture, facial skin evenness was measured using a PRIMOS Lite system. Evenness in the control group was 19.71 Ra before use, and 19.43 and 19.30 Ra after 2 and 4 weeks of application, respectively (Fig. 9). By contrast, skin evenness in the experimental group was 20.05 Ra before use, and 18.41 and 16.11 Ra after 2 and 4 weeks of application, respectively (p<0.001). Through these experiments, we identified an improvement in TEWL as an outcome of using the apigenin-containing cream.

Use of apigenin-containing cream improves the evenness of skin texture. The thickness of the stratum corneum changes depending on its moisture content, and insufficient moisture in this layer gradually roughens skin texture (35). To investigate the effects of the apigenin-containing cream on skin texture, facial skin evenness was measured using a PRIMOS Lite system. Evenness in the control group was 19.71 Ra before use, and 19.43 and 19.30 Ra after 2 and 4 weeks of application, respectively (Fig. 10). By contrast, skin evenness in the experimental group was 20.05 Ra before use, and 18.41 and 16.11 Ra after 2 and 4 weeks of application, respectively (Fig. 10). To compare the results from the control and experimental groups, we calculated the improvement as a percentage based on the value before application. Consequently, we found that skin texture in the control group improved by 1.42 and 2.06%, after 2 and 4 weeks of use, respectively. These data were not statistically significant (P>0.05), indicating that the control cream had no measurable effect on skin texture. However, the use of the apigenin-containing cream significantly improved...
Itching 0  Prickling 0  Scaling (epidermis) 0  Tightness 0  Swelling (edema) 0  Burning 0  Erythema 0

Table I. Adverse skin reactions reported by the subjects.

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<td>Erythema</td>
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*0, none; 1, mild; 2, severe; 3, very severe.

the evenness of skin texture by 8.20 and 19.65% after 2 and 4 weeks of application (P<0.001), respectively, suggesting that the use of the apigenin-containing cream can improve skin texture.

Analysis of adverse effects of apigenin-containing cream. In this study, investigators questioned the subjects individually about the condition of their skin and performed a visual evaluation of skin reactions, including erythema, itching, scaling, tingling, tightness, pricking and burning sensations at each visit. No extraordinary reactions were reported based on either visual evaluation or the questionnaire (Table I).

Discussion

Apigenin has been reported to have various biological activities in various cell types, such as antioxidant, anti-inflammatory, anti-mutagenic and anti-tumorigenic properties (36-39). However, it has not been determined whether apigenin can affect the aging process. In this study, we examined the effects of apigenin, particularly with regard to skin aging and wrinkling, using cellular and clinical efficacy experiments.

We first evaluated the viability of nHDFs that were irradiated with 25 J/cm² UVA and found that the post-treatment of UVA-irradiated nHDFs with apigenin significantly reduced cell cytotoxicity. This suggests that apigenin can reduce and/or mitigate UVA-induced cellular damage. We then evaluated the effects of apigenin on cellular senescence using a SA-β-galactosidase assay and found that while the percentage of senescent nHDFs increased in response to UVA irradiation, treatment with apigenin reduced the percentage of senescent cells in a dose-dependent manner. Furthermore, under the same conditions, the mRNA expression of MMP-1 collagenase-1, which is a reported initiator for the degradation of type I and III fibrillar collagen and is induced in response to UVA irradiation, was reduced by apigenin in a dose-dependent manner (17). Thus, we demonstrated the cellular efficacy of apigenin in inhibiting UVA-induced growth arrest, cellular senescence and the expression of MMP-1 in nHDFs.

UVA, a component of the UV spectrum, has been reported to have a greater average skin penetration than other UV types. Based on previous studies (17,29), UV irradiation, including UVA, is the main cause of skin aging, referred to as photoaging. It has been reported that collagen fibers, which are dermal components that maintain skin elasticity, are fragmented in photoaging skin (40). Furthermore, UV irradiation was found to decrease collagen synthesis and increase collagenolytic MMP synthesis in dermal fibroblasts (22,23,41), which are believed to contribute to the furrows observed in photoaging skin. MMPs are capable of degrading all components of the extracellular matrix and are upregulated by UV.

Based on the results of our cellular experiments demonstrating an apigenin-mediated protection from UVA-induced toxicity and an inhibition of MMP-1 upregulation in nHDFs, we constructed an apigenin-containing cream and a non-apigenin-containing control and enrolled 40 women (≥30 years old) in a randomized and double-blinded clinical trial to examine the effects of apigenin on aging skin in vivo. We then evaluated a number of features associated with aging in the skin of subjects who used either the apigenin-containing cream or the control. Of note, we detected a significant improvement in dermal density and skin elasticity, and a reduction in the length of fine wrinkles, particularly crow's feet, after 2 and 4 weeks of application in the subjects using the apigenin-containing cream. These results indicate that the topical application of apigenin reduces aging phenomena and its aging-associated clinical signs.

The skin is an important organ that separates the human body from the external environment. It has been previously reported that both the barrier function and the water-holding capacity of human skin are decreased by solar UV exposure (42). Therefore, the dryness of photoaging skin cannot be explained only under the direct influence of UV irradiation to skin cells. Therefore, we also measured skin moisture content, TEWL, and skin texture in subjects treated with apigenin-containing cream or the control. We found that all these parameters were significantly improved after 2 and 4 weeks of use in those who applied the apigenin-containing cream. These results strongly suggest that the topical application of the apigenin cream can improve aging skin, by enhancing the skin's barrier function.

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