Anti Cutaneous Aging Effect of Red Djulis
(Chenopodium formosanum) Extract on Gene Expression of Human Dermal Fibroblast


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** Equal contribution

Abstract: Red djulis (Chenopodium formosanum) is a native cereal plant in Taiwan; it contains abundant polyphenols, betalain and dietary fiber. The appearance of red djulis is bright red. Therefore, it is also called the “ruby of cereals”. The antioxidative activity of red djulis extract is well-understood. However, the antiaging function still remains unclear. This study examined the potential of red djulis extract for enhancing collagen secretion and preventing cutaneous aging using red djulis extracts. The red djulis extracts are comprised of an abundant active component that can effectively enhance the ability of collagen secretion of dermal fibroblasts, prevent the glycation of collagen and resist the damage of ultraviolet light exposure. After fibroblast treatment with red djulis extracts, TGM1, KRT1, KRT10 and SOD2 genes were up-regulated significantly by 2.3, 4.3, 4.4 and 27.3 times, respectively, compared to those of the control group. Additionally, it can increase COL1A2 gene expression by 43% and decrease MMP9 gene expression 33%. Therefore, it was demonstrated that red djulis extracts affect gene expressions related to the skin barrier, antioxidation and collagen. Moreover, we found positive effects on skin barrier integrity, endogenous antioxidant activity and skin collagen-preservation. The preparation of the red djulis extracts is environmental friendly and can promote the economic value of Chenopodium formosanum; thus, the proposed extract is suitable for applications in the development of food products, especially beverages, skin care and cosmetic products.

Keywords: Chenopodium formosanum; human dermal fibroblast; UV exposure; antioxidant activity; anti-aging; red djulis

1. Introduction

Collagen, one of the primary extracellular matrix (ECM) components of the dermis, is the most abundant protein in mammals, comprising about 30% of total protein[1-4]. Collagens co-polymerize to form extended mechanically stiff fibrils which confer tensile strength to the tissue providing the elasticity of skin[5]. However, the overall collagen content per unit area of the skin surface is known to diminish approximately 1% per year leading to gradual loss of skin elasticity (sagging) with age (intrinsic aging)[6, 7]. Additionally, chronic sun exposure leads to marked degradation of skin collagen structure resulting in wrinkle formation in photo aged skin (extrinsic aging)[8]. Therefore, in addition to sun exposure prevention, novel and effective anti-aging ingredients for skin care have drawn a lot of attention[9, 10].

Red djulis is a native cereal plant in Taiwan. The whole plant is colorful and traditionally called the “ruby of cereals” for its bright red grain color (Figure 1). Besides the high content of dietary fiber and starch, red djulis also possesses high protein levels and abundant essential amino acids, which make it a nutritious food. Djulis is particularly rich in lysine, which increases calcium absorption to help build collagen, and acts as a building block for the fibrils and fibers of collagen[5, 8, 11]. What is more, the betalain pigment in djulis is considered to possess not only good coloring potential but also...
positive physiological benefits for human health, such as melanoma cell growth inhibition and boosting of antioxidant activity[11-14], through antioxidant molecules such as betacyanin, mesembryanthin, and other polyphenols and flavonoids[11, 12]. Chyau et al., revealed the role of the ethanolic extracts of red djulis (Chenopodium formosanum) and its bioactive compounds in preventing adipogenesis in 3T3-L1 adipocytes[15]. Specifically, treatment with red djulis extracts under UVB light exposure can induce an accumulation of betacyanins and flavonoids in the immortal keratinocyte cell line (HaCat) for protection against UV-induced damage[14]. In addition, red djulis extracts have been reported to have antioxidant and moisturizing effects, and can stimulate the proliferation of fibroblasts and keratinocytes in the course of wound healing and regeneration[14, 16]. Therefore, djulis extracts contain an abundance of antioxidants and may act as an active ingredient for skin care product[17, 18]. Therefore, this in vitro study using genetic analysis aims to determine the anti-Cutaneous aging effect of red djulis extract and exploring the feasibility of application in the biomedical field.

2. Materials and Methods

2.1. Preparation and antioxidant analysis of red djulis extract

The red djulis used in the present study (Chenopodium formosanum) was purchased from Pingtung County, Taiwan. The whole grain red djulis (unhulled) stored at 4 °C prior to use. To achieve optimal extraction conditions, before carrying out the extraction step of the present study, 2.5g of grounded red djulis was combined was 25mL of water to form a mixture, and extracted with water at different temperatures for 30 minutes. The ratio of grounded red djulis to water was 1:10 (w/v), and the temperatures used for extraction were 25°C, 50 °C, and 70 °C, respectively. The mixture was then centrifuged at 4600 X g for 20 minutes to separate solid substances. The supernatant was collected after centrifugation and final filtration before subjecting the red djulis extract to analysis. Further, sonication at 40 kHz was applied to facilitate red djulis extraction. Finally, the red djulis extract was disinfected and sterilized at 121°C for 15 minutes, stored in a sealed container at 4 °C and kept out of light.

The antioxidant analysis of red djulis extract was applied using total phenol content by the Folin-Ciocalteu colorimetric method. Four mL of 2% sodium carbonate water solution and 5mL of Folin-Ciocalteu reagent was added to red tubes with 1 mL djulis extract or standard solution. The standard solution was gallic acid at a concentration of 200 to 1000 mg/mL. (7.5% in water) in order to create basic conditions (pH ~10) for the redox reaction between phenolic compounds and the Folin-Ciocalteu reagent. After incubation for 90 min at room temperature, the absorbance was read at 750 nm by a microplate reader (BioTek Instruments, Winooski, VT), against the blank. Further, incubation at 25 ° C for 30 minutes, absorbance was measured as 750 nm by the microplate reader (BioTek Instruments, Winooski, VT), against the blank. Total phenolic content was expressed as the equivalent of gallic acid.

The total flavonoid content of crude extract was determined by the aluminium chloride colorimetric method[19]. Fifty μL of red djulis extract was increased to 1 mL with methanol, mixed with 4 mL of distilled water and then 0.3 mL of 5% NaNO₂ solution; 0.3 mL of 10% AlCl₃ solution was added after 5 min of incubation, and the mixture was reacted for 6 min. Two mL of 1 mol/L NaOH solution were added, and the final volume of the mixture was brought to 10 mL with double-distilled water. The mixture was reacted for 15 min; absorbance was measured at 500 nm. The total flavonoid content was calculated from a calibration curve, and the result was expressed as mg rutin equivalent per g dry weight.

2.2. Chemical characterization

Varian 400 NMR instrument was used to record ¹H NMR and ¹³C NMR spectrums. The chemical shifts of spectroscopic data are given in δ (ppm) and coupling constants in hertz (Hz). Bruker amaZon SL mass spectrometer equipped with an ESI ionization source (Bruker, Bremen, Germany) was used for acquiring mass data. The HPLC system was composed of Hitachi L-2310 series pump (Hitachi,
Tokyo, Japan), L-2420 UV-VIS detector (Hitachi, Tokyo, Japan), and an ODS column (5 µm, 250 × 10 mm, Discovery® HS C18, Supelco Inc, Tokyo, Japan). The Medium pressure liquid chromatography (MPLC) was performed on a CombiFlash® RF (Teledyne ISCO, Lincoln, USA). Sephadex LH-20 (Amersham Pharmacia Biotech AB, Sweden) was used for separation. LiChrospher® Si 60 (5 µm, 250–10, Merck, Darmstadt, Germany) and LiChrospher® 100 RP-18e (5 µm, 250–10, Merck, Darmstadt, Germany) were used for NP-HPLC and RP-HPLC (Merck, Darmstadt, Germany), respectively.

2.3. Cell lines and chemicals

The human skin fibroblast cell line CCD-966SK (CRL-1881) was derived from the American Type Culture Collection (Manassas, VA, USA). All cell culture media and reagents including MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] were purchased from Sigma-Aldrich (St. Louis, MO, USA) or Gibco (Thermo Fisher Scientific, Inc., Carlsbad, CA, USA) and were of reagent grade or cell-culture grade. Cells were cultured in MEM with 10% fetal bovine serum, and incubated at 37 °C in a humidified atmosphere of 5% CO2.

2.4. Cell viability

Briefly, cells were seeded into 96-well plates at a density of 5 × 10^3 cells/well. After 24 h of culturing, the medium was removed and replaced with a fresh medium containing various concentrations (0.25, 0.5, 1 and 2 mg/mL) of red djulis extract. Cell viability was assessed through MTT assay. Briefly, 15 µl of 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) (Sigma; 4 mg/ml) was added and the cells were incubated for an additional 4 hours. The medium was removed and 50 µl /well of DMSO was added to resolve formazan crystal. The plate was placed on a shaker and incubated for 10 min and the absorbance was measured at 570 nm. Cell viability in response to treatment was calculated as: Cell viability (%) = (OD_sample / OD_control) × 100%.

2.5. Quantification of gene expressions by real-time PCR

The treated CCD-966SK fibroblasts or human primary epidermal keratinocytes were harvested, and total RNA was isolated from cells using an RNA purification kit (Geneaid, Taiwan). DNA-free total RNA was reversely transcribed to cDNA using a SuperScript™ Reverse Transcriptase kit (Invitrogen, Life Technologies Co., CA, USA). Quantitative real-time PCR was conducted using an ABI StepOnePlus™ Real-Time PCR System (Thermo Fisher Scientific, Inc., CA, USA) and the SYBR Green Master Mix (KAPA Biosystems, MA, USA) for was used transcript measurements. The reaction mixture was cycled as follows: One cycle at 95 °C for 20 s, followed by 40 cycles of 95 °C (1 s), 60 °C (20 s), and plate reading was conducted after each cycle. The melting curves of the PCR products were analyzed during the quantitative real-time PCR. The gene-specific primers used in this study are listed in Table 1. Real-time PCR reactions were performed using the ABI StepOnePlus™ system with KAPA SYBR FAST ABI Prism (KAPA). Antioxidant-related genes (SOD2 and CAT) and collagen-related genes (COL1A1 and TIMP1) were detected (Table.1). GAPDH was used as the reference gene to normalize relative expression. The GAPDH gene was used as a normalization control. Data were analyzed using the ABI StepOne™ Software v2.2.3 (Thermo Fisher Scientific, Inc., Carlsbad, CA, USA). All PCR assays were performed in duplicate three times.

2.6. Wound healing assay

In vitro wound scratch assay was performed as described in Yarrow et al.[20] with modifications. Briefly, CCD-966SK cells were grown to 80% confluence in 24-well plates. The scratching was performed by scraping with a sterile 1-ml pipette tip across the center of the well, followed by incubation with media only (mock) or with 2 mg/ml of SGE in culture media for 17 h. The wound closures were subsequently photographed under a microscope (Eclipse Ti-U, Nikon Corporation, Tokyo, Japan) using a CCD digital camera. Cell migration was analyzed by Image J (image processing and analysis in Java: https://imagej.nih.gov/ij/download.html) and expressed as percentage of wound closure.
coverage by cells moving into the scratched wound area. Cell migration data were expressed as the percentage of wound coverage.

2.7. Antiglycosylation analysis

AGE formation was determined by using collagen and fructose as glycation reactants. The modification procedure was described as in previous study by Peter-Katalinic (2005)[21]. Collagen solution (60 mg/mL) and fructose (1.5 M) were prepared in 0.2 M potassium phosphate buffer (pH 7.4, containing 0.06% sodium azide). For each reaction of glycation, 250 μL of collagen solution, 250 μL of fructose solution, and 250 μL of deionized water diluted sample or control (3mM aminoguanidine) were mixed. 100 μL of the above mixture was withdrawn immediately to a well of a 96-well plate and subjected to intensity determination by a spectrofluorometer (Paradigm microplate reader, Beckman Coulter, Fullerton, CA, USA) set at 360 nm of excitation and 460 nm of emission. The data were defined as Fluorescence sample 0hr or Fluorescence control 0hr. 450 μL of the remaining mixtures were taken into a 1.5 mL microfuge tube and incubated at 50°C for 24hr. After incubation, the reacted solutions were measured with a spectrofluorometer again and the result was defined as Fluorescence sample 24hr or Fluorescence control 24hr. The antiglycosylation activity rate of the sample was calculated as follows:

\[
\text{Antiglycosylation activity (%) = } 1 - \frac{\text{Fluorescence sample 24hr - Fluorescence sample 0hr}}{\text{Fluorescence control 24hr - Fluorescence control 0hr}} 
\times 100\%
\]

2.8. Collagen secretion assay

The soluble collagen was determined by the method described by Aramwit et al. (2009). Human skin fibroblast cells (CCD-966SK) were seeded at an initial concentration of 4 × 10^4 cells per well of 24-well plate in MEM containing 10% FBS. After 24 h, the culture medium was replaced by fresh medium. The sample extracts were diluted with various concentrations with phosphate buffer saline (PBS) and added into each well. Cells without a sample extract served as negative controls. After incubation for 48h at 37°C with 5% CO2, the supernatants were collected. The total amount of soluble collagen type I was assayed using the Sircol® Collagen Assay Kit (Bicolor Life Science Assays, Northern Ireland, UK). Briefly, 100 μL of experimental supernatant was mixed with 1 mL of dye solution at room temperature for 30 min. Then the samples were centrifuged at 15,000 × g for 10 min to form a pellet of collagen. The supernatant was removed and the soluble collagen produced was dissolved in 1 mL of alkali reagent. The resultant alkali reagent solutions were assayed by a spectrophotometer at a wavelength of 540 nm. The amount of collagen was calculated based on a standard curve of soluble collagen (bovine skin collagen type I standard from American disease free animals).

2.9. Collagen immunofluorescence staining

For immunohistochemistry, antigen retrieval was performed by heating sections of 10 mM citrate buffer (pH 6.0) in a microwave oven. Reactions with endogenous peroxidases and proteins were blocked by incubation with 3%H2O2 diluted in methanol and 10% normal goat or rabbit serum, depending on the host animal for the secondary antibody. Then incubation with primary antibodies was done overnight at 4 °C. The primary antibodies used were as follows: rabbit anti-collagen I (1:50; Abcam, Cambridge, UK), rabbit anti-collagen III (1:100; Abcam), mouse anti-collagen IV (1:50; Dako, Glostrup, Denmark), rabbit anti-collagen V (1:50; Abcam), and rabbit anti-collagen VI (1:50; Abcam).
The EnVision labeled polymer-HRP system (Dako) was used as the secondary antibody and peroxidase activity was visualized with a liquid diaminobenzidine substrate (Dako). Nuclei were counter-stained with hematoxylin. Collagen staining of the nodular lesions was classified into three categories according intensity compared with background staining (strongly positive, weakly positive, and negative).

2.10. Trosinase activity inhibition assay

Trosinase activity inhibition assay was conducted following Chan et al (2011)[22]. Briefly, 100 μl of freshly prepared L-DOPA solution, at different concentrations (2.5, 5.0 and 10.0 mM), was added for dopachrome formation. The relationship between total protein and concentration of L-DOPA for dopachrome formation was observed. The reaction mixture (200 μl/well) consisting of cell-extracted protein and L-DPA in 0.1 M sodium phosphate buffer (pH 6.8) was added into wells of a 96-well plate in triplicate. The plate was incubated at 37°C and absorbance was measured at 475 nm for a time course of up to 4 hours.

2.11. Statistical analysis

The significance of treatment was evaluated using paired t-test with the software Statistical Product and Service Solutions (SPSS® 18.0; SPSS Inc., Chicago, IL). A difference was considered statistically significant when the p value was < 0.05. All results were expressed as mean ± standard deviation (SD).

3. Result and Discussion

3.1. Optimal extraction condition of red djulis extracts

When evaluating the optimal conditions for total flavones and total phenol content from red djulis, the amount of total flavones and total phenol may be affected by the type of solvent, pH level, solvent-water ratio and extraction time[23]. To avoid the Islamic legal limitations and cost consideration, authors decided to use water extraction. The antioxidant compounds of different extraction conditions are shown in Table 2. The amount of total flavones and total phenol of the unhulled red djulis groups were higher than those of hulled red djulis and the difference reached almost three to four fold. Furthermore, the maximum amount of total flavones and total phenol extract from red djulis was obtained from unhulled red djulis at 50°C for 30 minutes. A previous study revealed that the djulis hull contains numerous antioxidant compounds and could be used to develop functional foods[16, 24]. Therefore, we decided to obtain extracts from unhulled red djulis at 50°C for 30 minutes for further experiments.

3.2. Chemical characterization

The obtained extract was partitioned with ethyl acetate (EtOAc), inclusive of n-butanol (n-BuOH) to obtain EtOAc layer (12.3 g), n-BuOH layer (22.1 g) and H2O layer (108.7 g) respectively. Following the bioassay-guided fractionation isolation (BGFI), the EtOAc layer was subjected to a Sephadex LH-20 and eluted with MeOH to yield 9 fractions (Fr. 1-Fr. 9). Later on, Fr. 2 was separated by MPLC with LiChroprep® RP-18 to afford five fractions (Fr. 2-1-Fr. 2-5) by elution with a linear gradient of mixtures of MeOH–H2O (from 10:90 to 90:10). Fr. 2-2 was separated by MPLC with a Discovery® HS C18 (250 × 10 mm, 5 μm) column and eluted by mixtures of MeOH–H2O (10:90), to yield compound 2 (3.2 mg). Fr. 3 was purified by preparative HPLC with isocratic solvent system MeOH–H2O (30:70), to afford compound 1 (5.2 mg). Fr. 4 was loaded onto a preparative HPLC system and eluted by mixtures of MeOH–H2O (20:80) to yield compound 4 (4.6 mg). Fr. 5 was treated similarly to yield compound 6 (10.0 mg). However, Fr. 6 was applied to a RP-18 column by MPLC eluted with a gradient solvent of mixtures of MeOH–H2O (from 10:90 to 90:10) and further purified by HPLC with mixture of MeOH-H2O (35:65) to obtain compound 3 (5.6 mg) and compound 5 (20.3 mg). The structures of all isolates were determined by analysis of their spectroscopic data (NMR and
MS) data shown in Figure 2 and Figure 3, the promote ability of collagen secretion will be discuss at later sections.

3.2. In Vitro Efficacy Evaluation

3.2.1. Cytotoxicity Assessment

The effect of red djulis extracts on cell viability of human dermal fibroblasts (CCD-966SK) was analyzed. The results showed that cell viabilities of red djulis extracts with concentrations of 0.1875 to 3 mg/mL were not significantly different with those of the control group (100%) (Figure 4). The results indicated that red djulis extracts are without cytotoxic compound.

3.2.2. Wound Healing Assay

The wound healing rate of red djulis extracts was 2.3 fold greater than that of the control group. After 24 h exposure to the test sample, red djulis extracts were enhanced so that the cell migrates towards the provisional gap (Figure 5). According to comprehensive cytotoxicity results, the red djulis extracts had low cytotoxicity and enhanced cell migration. These results seem to be correlated with of antioxidant compound of red djulis extracts.

Every living cell that undergoes aerobic metabolism produces reactive oxygen species (ROS) as its byproduct. The ROS is removed by antioxidative compounds of red djulis extracts, thus maintaining the healing physiological functions of fibroblast. Our findings are consistent with those of previous studies [14, 25, 26].

3.2.3. Gene Expressions of CCD-966SK Fibroblasts

Skin-barrier-related genes (TGM1, KRT1, KRT10), antioxidant-related genes (SOD2) and collagen-related genes (COL1A1 and MMP9) were selected for analysis of the mRNA expression in human dermal fibroblasts (CCD-966SK) by qPCR. The TGM1 gene provides instructions for producing an enzyme called transglutaminase 1. This enzyme is involved in the formation of the cornified cell envelope, which is a structure that surrounds skin cells and helps to form a protective barrier between the body and the environment. The KRT1 and KRT10 gene encode a member of the type I (acidic) and type II (basic or neutral) cytokeratin families, respectively. They are coexpressed during the differentiation of the epithelial tissues. The SOD2 gene encodes superoxide dismutase 2 (SOD2, mitochondrial), which is a member of the iron/manganese SOD family, catalyzing the dismutation of the superoxide radical into hydrogen peroxide or oxygen. The COL1A2 (collagen type I alpha 2) encodes the pro-alpha2 chains of type I collagen, which is a fibril-forming collagen found in most connective tissues and is abundant in dermis, bone, cornea and tendon. The MMP9 gene encodes matrix metallopeptidase 9, which is a class of enzymes involved in the degradation of the extracellular matrix (collagen) [11]. The "photo aging" or "photo-aging effect" refers to aging cells with UV light caused by factors including but not limited to UV light activated protein kinase pathway split via silk (MAP Kinase pathway) and increased phosphorylation of the dermal layer of matrix metalloproteinase (matrix metalloproteinases, MMPs) content. The matrix metalloproteinases decomposes collagen, decreasing the collagen content of the skin; will promote the UV light reactive oxygen species (reactive oxygen species, ROS), such as oxygen ions, free radicals and peroxides such as organic and inorganic produce, other cause denaturation of collagen (denature) and loss of function.

In the present study, after treatment with red djulis extracts, TGM1, KRT1, KRT10 and SOD2 genes were up-regulated significantly 2.3, 4.3, 4.4 and 27.3 times, respectively, compared to those of the control group (Figure 6). Additionally, the extract can increase COL1A2 gene expression 43% and decrease MMP9 gene expression 33% (Figure 7). Therefore, it was demonstrated that red djulis extracts affect the gene expressions related to skin barrier, antioxidation and collagen, and shows positive effects on skin barrier integrity, endogenous antioxidant activity and skin collagen-preservation.
3.4. Stimulation of Collagen Secretion

After treatment with red djulis extracts, collagen content in human dermal fibroblast obviously increased based on immunofluorescence staining compared with the control (Figure 8 a). Additionally, the collagen content of the red djulis extract group was about 1 higher than that of the control group (Figure 8 b). These results revealed that red djulis extracts can promote collagen secretion from skin fibroblast cells.

Analysis of red djulis extract used in this study show that the extract promotes the secretion of collagen, wherein the vertical axis represents the percentage of collagen secretion rates, with the control group shown as 100% for the sake of reference. According to the findings, with 0.2mg/mL of unhulled treated red djulis extract, collagen secretion in human skin fibroblasts reached 138% compared to the untreated control group. Thus, treatment with red djulis extract led to an increase of 38%; and 0. shelling red djulis extract treated 2mg/mL of collagen secretion rates of human skin fibroblasts 201.4% is reached, the control group protein secretion in human skin fibroblasts treated to extract collagen without red djulis rate more than 2 times. Furthermore, the fractions of TCI-CF-01 TCI-CF-02 and TCI-CF-03 from red djulis extract treated 20 mg/mL of collagen secretion rates of human skin fibroblasts 133.3% 122.9% and 119.8% are reached, respectively (Figure 8c). Thus, experiments confirmed that unhulled red djulis extract, hulled red djulis extract and fractions of red djulis extract can effectively promote the collagen secretion rate of cells.

3.5. Anti- Advanced Glycation End-products (AGEs)

Red djulis extracts inhibited the formation and accumulation of Amadori Product (primary glycation products), α-dicarbonyl compounds (secondary glycation products) and Advanced Glycation End-products (AGEs), with inhibition of about 58%, 53% and 80%, respectively, compared with the control (Figure 9). The results showed that red djulis extracts decreased the formation of AGEs and the glycation of collagen. The experiment indicated that red djulis extracts have the ability to protect and maintain the function of collagen in the skin. Moreover, according to our analysis, red djulis extracts inhibit the glycation of collagen. The extract reached an optimal effect at a concentration of 8% (w / w), inhibiting collagen glycation by up to 62.2%.

3.6. Red Djulis Extracts Inhibited Melanin Synthesis

The result showed that after treatment with different doses of red djulis extracts, the tyrosinase activity was significantly inhibited by 30% and 43% compared with the control group (Figure 10), and the result was dose dependent. Red djulis extracts also significantly reduced melanin synthesis by about 30% compared to the control group (Figure 8). This study indicated that red djulis extracts had a whitening effect by reducing the activity of tyrosinase, a key enzyme in melanin synthesis, and inhibiting melanin synthesis.

4. Conclusion

In summary, red djulis extracts contain abundant bioactive compounds rich in flavonoids and phenolic 15. Thus, the extract not only promotes secretion of collagen of dermal fibroblasts, but also effectively inhibits the glycation effect, and may enhance the dermal fibroblasts viability and migration rate. After fibroblasts were treated with red djulis extracts, TGM1, KRT1, KRT10 and SOD2 genes were up-regulated significantly by 2.3, 4.3, 4.4 and 27.3 times, respectively, compared to those of the control group. Additionally, the extract can increase the COL1A2 gene expression by 43% and decrease MMP9 gene expression by 33%. Therefore, it was demonstrated that red djulis extracts affect gene expressions related to the skin barrier, antioxidation and collagen. The results show positive effects on skin barrier integrity, endogenous antioxidant activity and skin collagen-preservation. Furthermore, the water soluble red djulis extracts do not harm the environment and could provide flexible formulation for safe anti-aging skin care products, cosmetic raw materials or functional beverages.
Reference

17. Sun, L.-C.; Sridha, K.; Tsai, P.-J.; Chou, C.-S., Effect of traditional thermal and high-pressure processing (HPP) methods on the color stability and antioxidant capacities of Djulis (Chenopodium formosanum Koidz.). *LWT* 2019, 109, 342-349.


27. TGM1 Available online: http://www.ncbi.nlm.nih.gov/gene/7051;


### Legends

| Table 1 | Real-time quantitative PCR primers used in this study. |
| Table 2 | Antioxidative compounds of red djulis extract. |
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| Figure 8 | Effects of red djulis extracts in fibroblasts with or without red djulis extract treatment: (a) Collagen immunofluorescence staining (b) collagen secretion of red djulis extracts treatment and (c) collagen secretion of different red djulis extract fractions treatment. |
| Figure 9 | The effect of red djulis extracts on formation of Advanced Glycation End-products (AGEs). |
| Figure 10 | Effect of red djulis extracts on tyrosinase activity. |

### Table 1.

<table>
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KRT14-F 5'- TTCTGAACGAGATGCGTGAC - 3' epidermal barrier structural genes 29
KRT14-R 5'- GCAGCTCAATCTCCAGGTTTC - 3'

SOD1-F 5' - GGTGGGCAAAGGATGAAGAG - 3' anti-oxidative gene 30
SOD1-R 5' - CCACAAGGCAAACGACTTCC - 3'

COL1A1-F ATCAACCCGAGGAATTTCCGT provides instructions for making part 31
COL1A1-R CACCAGGAGCCGCACTTCC type I collagen

MMP9-F GGGACCCAGACATCGTCATC enzymes engaged in the degradation and 32
MMP9-R TCGTCATCGTCGAAATGGGC remodeling of extracellular matrix (ECM)

Table 2

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<th>Extraction Temperature (°C)</th>
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Figure 1.
Figure 2
Figure 3
Figure 4

The bar graph shows the cell viability (%) in response to different concentrations of Red djulis extracts (mg/ml). The x-axis represents the concentration (0, 0.75, 1.5, 3 mg/ml), while the y-axis represents cell viability (%) ranging from 0 to 140. The graph indicates a trend where cell viability decreases with increasing extract concentration.
Figure 5

Red djulis extracts, 0.23mg/ml
Red djulis extracts, 0.25mg/ml
Red djulis extracts, 0.5mg/ml
Figure 6.

![Bar chart showing relative expression ratio for TGM1, KRT1, KRT10, and SOD2 between Control and Red Djulis Extract. The chart indicates significant differences with asterisks (*) and a three-star symbol.](chart.png)
**Figure 7**

The bar chart represents the relative expression ratio of COL1A2 and MMP9 between the Control and Red Djulis Extract groups. The y-axis indicates the relative expression ratio (test/control), while the x-axis separates the expression of COL1A2 and MMP9. The chart includes error bars for the Control group and an asterisk for the Red Djulis Extract group, indicating statistical significance.
(a) Collagen
Dapi (Nucleus)

Control Red Djulis Extract

(b) Collagen assay

<table>
<thead>
<tr>
<th>% of control</th>
<th>Control</th>
<th>Red Djulis Extract</th>
</tr>
</thead>
<tbody>
<tr>
<td>100.00</td>
<td>201.41</td>
<td></td>
</tr>
</tbody>
</table>
Figure 8

(c)

Collagen secretion (%) vs. TCI-CF-01 to TCI-CF-06.
Figure 9

Relative Advances Glycation End-products (AGES)

Control

Red Djulis Extract
Figure 10

Melanin content (%)

- Control
- Red djulis extracts, 0.5mg/mL
- Red djulis extracts, 1mg/mL