Effects of Sizes and Conformations of Fish-Scale Collagen Peptides on Facial Skin Qualities and Transdermal Penetration Efficiency

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Fish-scale collagen peptides (FSCPs) were prepared using a given combination of proteases to hydrolyze tilapia (Oreochromis sp.) scales. FSCPs were determined to stimulate fibroblast cells proliferation and procollagen synthesis in a time- and dose-dependent manner. The transdermal penetration capabilities of the fractionated FSCPs were evaluated using the Franz-type diffusion cell model. The heavier FSCPs, 3500 and 4500 Da, showed higher cumulative penetration capability as opposed to the lighter FSCPs, 2000 and 1300 Da. In addition, the heavier seemed to preserve favorable coiled structures comparing to the lighter that presents mainly as linear under confocal scanning laser microscopy. FSCPs, particularly the heavier, were concluded to efficiently penetrate stratum corneum to epidermis and dermis, activate fibroblasts, and accelerate collagen synthesis. The heavier outweighs the lighter in transdermal penetration likely as a result of preserving the given desired structure feature.

1. Introduction

Collagens are the major structural element of connective tissues in vertebrate, comprising 30% or so of total protein. They also exist in the interstitial tissues of virtually all parenchymal organs, wherein they stabilize organs and keep them in good shapes [1]. To date, over 20 types of collagens have been identified, and they were otherwise divided into three groups, fibrous collagen, fibril associated collagen, and basement membrane collagen [1–3]. Skin, the largest organ in human body protects the body from varied external insults. It is composed of three layers, namely the epidermis, the dermis, and the subcutaneous layer. Each layer provides some extents of physical strength and flexibility in concert with other physiological functions [4]. The dermis contains considerable amounts of extracellular matrix (ECM), such as collagens and glycosaminoglycans (GAGs) produced mainly by fibroblasts. Collagen type I [5] known to maintain the integral of the dermis is the most abundant collagen in human body. Aging skin is admittedly related to the reduction of collagen type I [6].

Collagen has been widely processed as products in food, cosmetic, biomedical, and pharmaceutical industries. Oral consumption of collagen peptides may provide some beneficial effects for the body. For examples, collagen peptides as a food supplement may improve low bone mineral density in people in malnutrition and people suffering degenerative joint diseases [7–9]. Reports also indicated that consumption of collagen can thicken hairs [10], improve nail disorders such as brittle nails [11], increase the size of collagen fibrils in Achilles tendon [12], induce fibroblast density, and enhance formation of collagen fibrils in dermis etcetera [13]. Traditionally, collagens are derived from livestock sources, such as bovine hide and bones as well as swine...
skin [14]. Because of the outbreaks of bovine spongiform encephalopathy (BSE) and foot-and-mouth disease (FMD) in recent years, collagens and collagen-derived products from such sources may have intimidated considerable users [15]. Collagens and collagen-derived products from swine sources have an additional disadvantage as religious sake in some parts of the world [16]. “Fish wastes” therefore turn out to be a good alternative. In fish processing, a large portion of wastes would generate, which typically accounts for 50%–70% of raw materials including skins, bones, scales, viscera, and heads [17]. These so-called “wastes” were difficult to handle, generally used as low-value feedstuffs or fertilizers as a whole, so that serious environmental issues arise from time to time [18]. Given 30% or so of collagen contained in these “wastes”, they have drawn great attention lately [19].

The stratum corneum (SC) has been known to minimize passive water loss from the body, to ward off absorption of chemicals in the environment, and to prevent microbial invasion [19, 20]. Topical administration of cosmetics or biomedical materials on skin therefore has to overcome the stratum corneum barrier so as to reach the fibroblast cells in the dermal layer. Raw collagens derived from whatever sources without any process for either cosmetic or biomedical purposes may not have desirable effects, for instance the type I collagen proliferation. We considered factors such as molecular volume (MV) or molecular weight (MW) of collagens may play a key role on their penetration ability when administrated topically [21], of which relevant reports remained few.

In this paper, we prepared testing collagens using scale wastes of tilapia (Oreochromis sp), which is a high value-added fish in Taiwan fishery. The prepared collagens formulated into skin essence were assessed for their physiological effects on facial skin in users. In addition, we determined the levels of fibroblasts proliferation and collagen synthesis in embryonic fibroblast cell lines after being subjected to FSCP treatments. Given molecular sizes of FSCPs on transdermal efficiencies were also explored. In fact, our results showed that the transdermal penetration efficiency of FSCPs is positively correlated to given FSCPs’ molecular sizes and structural features in a nude mice model.

2. Materials and Methods

2.1. Preparation of Fish Scale Collagen Peptides (FSCPs). Fish scale collagen peptides (FSCPs) were isolated from tilapia (Oreochromis sp., scales (Ko-Fwu Fishes Co., Taiwan) by enzymatic hydrolysis according to our patented protocol [22]. In short, fish scales were washed to rid of impurities and then heated for 15 minutes at 121°C to soften scales. The heated scales (200 g) were smashed into small pieces by disperser (Kinematica, NY, USA), then subjected to hydrolysis under 1% Protease N for 2.5 hours and 0.5% Flavourzyme (Novozymes, Chiba-shi, Japan) for another 0.5 hour at an optimal pH and temperature. Hydrosylates were stirred and heated in a boiling water bath for 10 min to inactivate enzymes. Then, the hydrolysates were centrifuged at 12,000 g for 20 min. The supernatants (FSCPs) were lyophilized and stored at −20°C for use.

2.2. Moisture Contents and Relative Elasticity Assays. FSCP-based skin essences were formulated into 5%, 7%, and 10% of total FSCPs as the common basal ingredients. Sixty-two voluntary Taiwanese women (within 23 to 60 years of age) were subjected to the FSCP-based skin essence treatment on facial skin twice a day for 30 days. These test subjects were forbidden using any other cosmetic products during the test. Moisture contents and relatively elasticity of facial skins were measured every other week (at the 0, 2nd, and 4th week) by the skin probe of Cutometer MAP 580 (KOKO, Leichlingen, Germany) at 20–22°C and at 40%–60% relative humidity. Each measurement took place at 30 min after washing face with DI water.

2.3. Cell Proliferation and Collagen Release Assays. About 2 ×10^5 Detroit 551 cells (human embryonic skin fibroblast cell line, ATCC CCL-110) and STO cells (mouse embryonic fibroblast cell line, ATCC CRL-1503) were seeded into each well of a 96-well plate and maintained in eagle’s minimum essential medium (EMEM) and Dulbecco’s modified Eagle’s medium (Sigma, St. Louis, MO, USA) plus 10% fetal bovine serum (FBS) (Invitrogen, San Diego, CA, USA), respectively. Cells were cultured overnight. The medium was refreshed with new medium containing 0, 0.4, 0.8, 1.6, 3.2, 6.3, 12.5, 25, 50, 100, and 200 mM FSCPs; cells were incubated for 48 hours before MTT and procollagen type I production assays. For MTT assay, 100 µl of the MTT solution [a mixture of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) and yolk lipoprotein (YLP) (Invitrogen, San Diego, CA, USA) in PBS] was added into wells and reacted for 3 hours at 37°C. The plates were stirred for 2 min, and corresponding absorbance was recorded at 570 nm using an ELISA reader [6]. For the procollagen type I production assay, the procollagen type I C-peptide ELISA kit (Takara Bio Inc, Otsu, Japan) was used. Experimental procedures followed the manufacturer’s instruction. In short, 20 µl of culture medium and 100 µl of the antibody-POD conjugate solution were sequentially added into microtiter plates and reacted for 3 hours at 37°C. After 4 × 5 min washing with the provided washing buffer solution (400 µl/well), 100 µl of the substrate solution was added in. After 15 min room temperature incubation, the stop solution (100 µl) was added in; corresponding absorbance was recorded at 450 nm using an ELISA reader.

2.4. Molecular Size Effects of FSCPs. The enzymatic hydrolysates from the fish scales were fractionationized into five different peptide pools, using the Millipore minianin system (Millipore, Bedford, MA) with four different molecular weight cutoff membranes (5, 3, 1, and 0.5 kDa). A 50 µl aliquot of fish scale hydrolysate was loaded onto a Superdex peptide 10/300 GL (10 mm × 300 mm × 13 µm) gel filtration column (Pharmacia, Uppsal, Sweden) and eluted with 200 mM phosphate buffer (with 0.25 M NaCl, pH 7.8) at a flow rate of 0.5 mL/min. FSCPs were calibrated against protein standards (Bio-Rad, California, USA) of cytochrome C (Mr = 12500 Da), aprotinin (Mr = 6512 Da), vitamin B12 (Mr = 1355 Da), and cytidine (Mr = 246 Da).
Average molecular weights (MW) of fractionated FSCPs were determined to be 4500, 3500, 2000, and 1300 Da, respectively; each fraction was lyophilized for use.

2.5. FSCPs Labeled with Fluorescein Isothiocyanate. FITC-labeled FSCPs were obtained by using the Fluoro Tag FITC conjugation kit (Sigma, St. Louis, MO, USA). The FITC-labeled FSCPs were prepared based on a conjugation reaction between the isothiocyanate group in FITC and the primary amino groups in collagen peptides [23]. In short, FSCPs solution (750 ppm, 1 mL) was rebuffered with a phosphate buffer solution (pH 7.4). The new FSCPs solution was mixed with 0.25 mL of 0.1 M sodium carbonate-bicarbonate buffer (pH 9.0) containing FITC, which was gently stirred for 2 hours at room temperature. The FSCPs labeled with FITC were isolated by Sephadex G-25 column (Bio-Rad, Japan). Mice were subjected to destratum corneum by a 30% w/w solution of urea and 1.25% w/w solution of sodium thioglycolate, and the treated skin tissues were observed under fluorescence microscope (BX-51, Olympus, Japan) equipped with a digital camera [24].

2.6. In Vivo Transdermal Delivery Efficiency of FITC-Labeled FSCPs. Six-week-old female C3H/HeN mice were anesthetized using acepromazine maleate (i.p.). Hairs covering the areas of abdomen skin were removed with a shaver. Residual hairs were removed by hair-remove-cream (Yanagiya, Japan). Mice were subjected to destratum corneum by treating mice with 10% alpha hydroxy acids (AHAs; BIOPEUTIC, USA) for 5 minutes and cleanup for use. An aliquot of 25 μg FITC-labeled FSCPs in sterile DI water (100 μl) was introduced into nonwoven fabrics (cosmetics mask; Widetex Biotech Co., Taiwan). The fabric covered 1 cm² of hairless dorsal back skin which then was topped with 1.5 cm² transparent dressing film (Tegaderm, Neuss, Germany). The nonwoven fabric was removed after one hour. The skin tissues treated with the FITC-labeled FSCPs were embedded in O.C.T. Embedding Medium (Sakura Finetek USA, Torrance, California) and sectioned into 10 μm thickness. The skin tissues were observed under fluorescence microscopy (BX-51, Olympus, Japan) and treated with a blocking solution (95% ethanol and 5% acetic acid). After rinsed with PBS for 20 seconds and then rinsed with 50%, 70%, 90%, and 100% ethanol for 30 seconds, the slides were observed under a confocal spectral microscope (TCS SP5, Leica, Wetzlar, Germany).

2.7. The Franz-Type Diffusion Cell Model. The procedure was modified from the method described by Kim et al. [25]. Nude mice skin was mounted on the receptor compartment of the Franz-type diffusion cell (PermeGear, HT, USA). A phosphate buffer solution (6.80 g/l potassium phosphate monobasic, 3.78 g/l sodium chloride, and 0.235 g/l sodium acid; pH 7.4) containing 750 ppm a given fraction of FSCPs was added in the tight interface, facing nude mice skin. The donor cap was filmed with a parafilm. An isotonic phosphate buffer solution (pH 7.4) was used as the receptor solution. The receptor solution was stirred by a magnetic follower rotating at 250 rpm (which would increase mixing efficiency and reduce tendency of forming a stagnant boundary layer next to membrane surface). Samples were performed at 1, 2, 3, 4, 6, and 24 hours. The amount of total collagen peptides from the receptor solution was measured by the BCA protein assay (Pierce, Rockford, USA).

The cumulative penetration amount (μg) = \[ C \times (ppm) \times \frac{Vi}{Vr} + \sum_{n-1}^{Vi} C \times Vr \] (C = concentration; Vi = initial volume; Vr = sampling volume).

The unit area cumulative penetration amount (μg/cm²) = cumulative penetration amount/area measure (0.66 cm²).

2.8. Transdermal Penetration Efficiency of FITC-Labeled FSCPs Determined by Confocal Spectral Microscope. Various sizes of FSCPs were labeled with FITC, which would penetrate into nude mice skins in the Franz-type diffusion cell model. The skins of nude mice treated with FITC alone served as a control. After one hour, the treated skin tissues were dissected out, embedded in O.C.T. compound and sectioned in 10 μm thickness. The slides were then mounted and treated with a blocking solution (95% ethanol and 5% acetic acid). After rinsed with PBS for 20 seconds and then rinsed with 50%, 70%, 90%, and 100% ethanol for 30 seconds, the slides were observed under a confocal spectral microscope (TCS SP5, Leica, Wetzlar, Germany).

2.9. Statistical Analysis. The graphs and statistical analyses were performed using SigmaPlot and SigmaStat. The statistical analyses between groups were determined by nonparametric one-way analysis of variance (Kruskal-Wallis test) and Mann-Whitney tests. Differences were considered significant if the P value was ≤.05.

3. Results

3.1. Effects of FSCPs on Human Facial Skin Moisture Contents and Relative Elasticity. While collagen peptides have been well documented to be an ideal material in cosmetic industries [26], fish scale collagen peptides (FSCPs) as a sustainable collagen source however did not draw great attention. In this study, we examined the treating effects of FSCPs as a major component in a skin essence on the facial skin moisture contents and relative elasticity by using standard professional skin probes. The FSCP-based skin essences were first formulated into 5%, 7%, and 10% of total FSCPs as the common basal ingredients. Sixty-two voluntary Taiwanese women (within 23 to 60 years of age) were subjected to the FSCP-based skin essence treatment on facial skin twice a day for 30 days. The effects with all FSCP-based skin essences on the skin moisture contents were found improving significantly but in a time- and dose-dependent manner during the testing course (P ≤ .05; Figure 1(a)). Likewise, the effects with all FSCP-based skin essences on the relative elasticity of facial skin were also found increasing in the same manner as for the skin moisture test by 24%, 33%, and 35%, respectively (P ≤ .05; Figure 1(b)). Thereby, the FSCP-based skin essences did improve facial skin qualities (moisture and elasticity) in a time- and dose-dependent manner in these tested subjects.

3.2. Cell Proliferation and Collagen Release in Embryonic Skin Fibroblast Cell Lines after FSCPs Treatments. To determine
Figure 1: Effects of FSCP-based skin essence on moisture contents and relative elasticity of facial skins. The skin moisture contents (a) and relative skin elasticity (b) were measured by the professional skin measurement probes. Bars are expressed as means ± SEM in each group for the 62 subjects. Difference letters in the same group indicate significant difference (P ≤ .05).

Figure 2: Cell proliferation and collagen release in embryonic skin fibroblast cell lines treated by FSCPs. Comparisons of the proliferation activity (a) and procollagen synthesis ability (b) on Detroit 551 cells (human embryonic skin fibroblast line) and STO cells (mouse embryonic fibroblast) treated with 0, 0.4, 0.8, 1.6, 3.2, 6.3, 12.5, 25, 50, 100, and 200 mM FSCPs. The group with no FSCPs acts as a control. After 48 hours posttreatment, the samples were analyzed by MTT and procollagen type I C-peptide ELISA assays. Bars stand as means ± standard deviation (triplicate for each group).

whether the skin quality improving was resulted from fibroblast cells proliferation and mass collagen synthesis in dermis after the FSCPs treatment, model cells Detroit 551 (human embryonic skin fibroblast) and STO (mouse embryonic fibroblast) were chosen for the purpose stated. The cells were grown in media in the presence of various concentrations of FSCPs. On the contrary, the cells grown in media without FSCPs served as controls. Cell proliferation and procollagen release in the tested cells were determined by MTT assays and procollagen type I C-peptide ELISA assays, respectively. As shown in Figure 2(a), FSCPs (0.4–200 μg/mL) can dose-dependently stimulate the cell proliferation in both fibroblast cells. Likewise, the procollagen type I synthesis was found mass produced in a concentration-dependent manner in the cells with FSCPs treatments (Figure 2(b)). The procollagen synthesis in the presence of 200 μg/mL FSCPs was found most prominent and peaked as high as 250%. FSCPs thereby were determined to be able to effectively stimulate and
induce fibroblast cell proliferation and collagen synthesis so as to improve the facial skin quality.

3.3. In Vivo Determination of Transdermal Delivery Efficiency Using FITC-Labeled FSCPs. As mentioned above, the stratum corneum is the outmost layer of the epidermis, which serves as a physical barrier, for instance, preventing pathogens from invasion and also limiting desired macromolecules to pass by. To evaluate whether the processed FSCPs are capable of overcoming the stratum corneum barrier, an animal model experiment was set to determine the transdermal efficiency of the prepared FSCPs. Six-week-old female C3H/HeN mice were first shaved and treated with hair-removal cream to remove residual hairs. The mice were then subjected to treatments with or without alpha hydroxyl acids (AHAs, for weakening the stratum corneum) before topical application with the nonwoven fabric that contained 25 μg FITC-labeled FSCPs. The mice pretreated with 10%
AHAs and topped with the nonwoven fabric containing no FITC-labeled FSCPs served as controls. These treated skins were dissected out and immediately embedded with O.C.T. in due course. Then, the samples were sectioned, mounted on glass slides, and observed under fluorescence microscopy. As shown in Figure 3, the FITC positive signals laid mainly on the area of superficial epidermis. And, the FITC positive signals in the skin pre-treated with AHA were found deeper and brighter than those without AHA treatment. The FITC positive signals were also found in the hair follicles of epidermis. Thus, the stratum corneum barrier was determined to be a key factor that influences the penetration efficiency of FSCPs in mice skin. Nonetheless, FSCPs without AHA treatment still considerably pass by the stratum corneum, which agreed with the results in the human facial skin test.

3.4. Transdermal Penetration Ability of FSCPs in Franz-Type Diffusion Cell Model. We considered factors such as molecular volume (MV) or molecular weight (MW) of collagens may play a role on their penetration capability when administrated topically. The size effect for given FSCPs was determined by employing the Franz-type diffusion cell model. Fish scale hydrolates were first subjected to gel chromatography (superdex peptide 10/300 GL columns), whereby FSCP hydrolysates were fractionated by sizes. As shown in Figure 4(a), the eluted peptides were collected into six fractions, >6.5 KDa (3.65%), 3–6.5 KDa (16.8%), 1.5–3.0 KDa (32.0%), 0.7–1.5 KDa (29.3%), 0.35–0.7 KDa (14.6%), and <0.35 KDa (3.7%). These peptide fractions were further subjected to ultrafiltration membrane refining (5, 3, 1, and 0.5 kDa cutoff) into four refined groups I–IV (Mw = 3–5, 1–3, 1–0.5, and <0.5 KDa) with average molecular
Figure 6: Peptide conformations of various sizes of FITC-labeled FSCPs in nude mice skins. Nude mice skins were treated with various molecular sizes of FITC-labeled FSCPs in the Franz-type diffusion cell model. The topical sites of tested nude mice skins were collected in one hour. The vertical cross-sections of the nude mice skins were imaged for FITC signals using cryosection and confocal spectral microscope systems. (a) 1,300 Da FSCPs; (b) 2,000 Da FSCPs; (c) 3,500 Da FSCPs; (d) 4,500 Da FSCPs; and (e) the mixture FSCPs (FSCPs without ultracentrifugation membrane treatment).

weights of 4500, 3500, 2000, and 1300 Da, respectively (Figure 4(b)). These collagen peptides were corroborated by gel chromatography (Sp-Sephadex C-25 column) against standard markers. As shown in Figure 5, all groups were found to be able to penetrate the nude mice skin in the Franz-type diffusion cell model but in various extents and in a time-dependent manner. In short, groups I and II exhibited relatively better transdermal penetration ability in the first 4 hours. The highest cumulative penetration amounts for groups I and II at 24 hours were 1825.17 ± 495.66 and 1986.43 ± 671.11 μg/cm², respectively. In contrast, the FSCPs mixture (total FSCP hydrolates without ultrafiltration membrane refining) showed the lowest transdermal penetration capability (1010 ± 583.34 μg/cm²) (Figure 5).

3.5. Transdermal Penetration Depth and Peptide Conformation in Nude Mice Skin. To probe the penetration depth and peptide conformation for given FSCPs, each FSCP group was first labeled with FITC and then applied to nude mice skins in the Franz-type diffusion cell model for one hour. The skin treated with given FITC-labeled FSCPs proceeded according to the same procedure described in the previous section. Interestingly, groups I (Figure 6(d)) and II (Figure 6(c)) exhibited a coil-like structure in epidermis as opposed to groups III (Figure 6(b)) and IV (Figure 6(a)) which showed less folded or in a linear form. In contrast, the mixed FSCPs did not have a defined structure (Figure 6(e)). The penetration depths for each FSCPs group were also measured by confocal spectral microscopy. The distances for groups IV–I and the mixture were 21.21, 21.44, 28.72, 27.73, and 14.58 μm, respectively. The heavier FSCPs penetrated deeper than the lighter. As a result, the transdermal penetration efficiency of given FSCPs is positively correlated to FSCPs’ sizes and/or conformations thereof. The folded structure seemed to help penetration, which however is an interesting issue for future study.

4. Discussion

Many reports have pointed out collagen’s versatile applications, prominently for cosmetic industry [13]. Fish scale collagens appear to be an ideal collagen source as they are cheap, rich and sustainable [25, 26]. Since collagen is the key component in maintaining the elasticity of skin, diminishing
collagen contents in skin would therefore result in wrinkled and flabby skins [27]. In this paper, we have demonstrated the formulated FCSP-based skin essence possesses such desirable effects as increasing skin moisture contents and relative elasticity (Figure 1). The FCSPs at 200 μg/mL provided the best effects by manifesting embryonic skin fibroblast cell proliferations (Detroit 551 cells and STO cells) and procollagen synthesis. These results otherwise are reinforced by the reports of Katayama et al., wherein they demonstrated pentapeptides from type I collagen promoted extracellular matrix production in fibroblasts [28, 29].

Since the stratum corneum acts as a physical barrier for multiple purposes in skins [19, 20], it does influence the penetration efficiency of the FSCP-based skin essences as they were present mainly in the surface of epidermis. However, given FITC-labeled FSCPs still e

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