Biological activity of the thrombospondin epidermal growth factor domain from banana shrimp (*Fenneropenaeus merguiensis*)

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ABSTRACT: Human epidermal growth factor (EGF) plays a role in cell proliferation and has been used to promote wound healing in patients. However, its role in cancer development via the EGF receptor (EGFR) has recently been reported. Thrombospondin (TSP) from banana shrimp (*Fenneropenaeus merguiensis*) contains an EGF domain at its carboxy-terminal end though it shares only 30% nucleotide sequence similarity with human EGF. It is challenging to investigate its biological activities. In this work, the EGF domain from banana shrimp TSP (TSP3-2) was cloned, and the recombinant His-TF-TSP3-2 was evaluated for its effects on proliferation, cytotoxicity and collagen production in a skin fibroblast cell line. Additionally, genotoxicity was determined using a hypoxanthine-guanine phosphoribosyltransferase mutation assay. Fibroblast cell proliferation increased 1.2–1.4 fold after treatment with 0.5–15 \(\mu\)g/mL His-TF-TSP3-2. The migration rate after 48 h of the His-TF-TSP3-2-treated cells (at 0.5 \(\mu\)g/mL) in a scratch assay was greater than that of the control. The collagen production was 1.7- and 1.2-fold increased after treatment with His-TF-TSP3-2 at 0.5 and 1.0 \(\mu\)g/mL, respectively. Low toxicity to the genome and no significant mutant frequency was found in the His-TF-TSP3-2-treated cells. Further investigations are required to confirm a potential use for this protein.

KEYWORDS: thrombospondin (TSP), proliferation, growth factor, collagen

INTRODUCTION

Extracellular matrix proteins are known to maintain tissue and organ structures. They have a range of functions including acting as signal modulators involved in cell adhesion, proliferation, migration, and differentiation [1–3]. Thrombospondins are glycoproteins in the extracellular matrix and have been reported to have various functions in humans [2]. For example, the thrombospondins TSP1 and TSP2 are potent endogenous inhibitors of angiogenesis [3]; TSP1 can cause an imbalance in the host immune response that ultimately leads to reduced phagocytic function [4] and potently inhibits fibroblast migration [5]. The human TSPs, including TSP-1, TSP2, TSP-3, TSP-4, and TSP-5/COMP have been classified [6]. The general structure of human TSPs consists of repeated chitin-binding domains, an EGF-like domain, repeated thrombospondin-type domains and a C-terminal domain [2] while *Penaeus monodon* TSP (*Pm*TSP) contains 4 chitin-binding type II domains, an EGF-like domain, 8 thrombospondin type III domains and a thrombospondin C-terminal domain. Additionally, *Marsupenaeus japonicas* TSP (*Mj*TSP) has 5 chitin-binding type II domains, 6 EGF-like domains, 8 thrombospondin type III domains and a thrombospondin C-terminal domain [7]. In humans, TSP transcripts have been found in organs such as the intestine, ovary, and heart. Each human TSP exhibits high expression in a specific organ. For example, TSP1, TSP2, TSP3, TSP4, and TSP5 show the highest expression in the lung, ovary, uterus, pituitary gland, and trachea, respectively [2]. Both *Pm*TSP and *Mj*TSP are TSP3 genes and are expressed at very high level in the ovary, intestine and heart [7, 8].

Some TSPs are localized to different cell types or structures within the same tissue. The expression of TSPs changes in response to physiological or pathological changes; for example, TSP1 is found at
high level during ischemia. Moreover, TSP1 appears to exert a protective effect in myocardial infarction (MI) patients, in whom the high level is present [9]. TSP1 plays a role in collagen homeostasis through interactions with matrix metalloproteinases and directly affects collagen fibril formation [10]. TSP4 has been reported to protect against heart disease after MI injury or protein misfolding [11]. TSP4 expression is induced in the heart and vasculature under pathological conditions including myocardial infarction, myocardial pressure overload, and hypertension. TSP4 is linked to remodeling processes, in which it may affect extracellular matrix protein organization [12]. TSP5 has been reported to support chondrocyte attachment through interactions with integrins [13].

In shrimp, only TSP3 has been reported, and it exhibits high expression in ovarian tissues. The structures of TSPs from _P. monodon_, _Fenneropenaeus chinensis_ and _M. japonicus_ show fewer EGF domains than human TSPs [7, 14, 15]. The corresponding biological activities have not been reported. Therefore, TSP from the ovarian tissue of banana shrimp (_F. merguiensis_) was cloned and characterized in this study.

Currently, increasing numbers of slow or non-wound healing patients are found among the aging and obese populations [16]. Bioactive compounds that enhance wound healing have been introduced such as EGF; growth hormone, and basic fibroblast growth factor [17–19]. The topical use of EGF was reported to increase epithelial cell proliferation, and reduce the healing time of skin grafts, venous ulcers, and diabetic foot ulcers, and it was found that epidermal growth factor receptor (EGFR) expression was increased [20]. EGFR was observed to play a critical role in oncogenesis and metastasis while EGFRi (EGFR inhibitor) is a promising target for the treatment of metastatic cancers. However, EGFR inhibition showed significant cutaneous side effects and limitations to long-term use [16, 21]. Therefore, an alternative EGF without a role in cancer progression or a specific EGFRi that does not affect EGFR signaling in normal cellular function is needed. And the EGF domain of TSP has this required character. Thus, the EGF domain of TSP was cloned and evaluated for its biological activities such as wound healing, collagen production, and genotoxicity.

### MATERIALS AND METHODS

#### Banana shrimp TSP sequence analysis

Total RNA was extracted from banana shrimp ovaries using the TRIzol reagent (Invitrogen, Carlsbad, CA, USA) following the manufacturer’s protocol. To convert the RNA to cDNA, 400 ng of total RNA was incubated with 200 ng of random primers at 70 °C for 5 min, followed by cooling on ice for 5 min. This mixture was supplemented with 1 × avian myeloblastosis virus (AMV) buffer, 1 mM deoxynucleotide triphosphate, and 10 U of AMV reverse transcriptase (Promega, Madison, USA) in a 25 μL reaction mixture, followed by incubation at 48 °C for 2 h. Primers for full-length TSP were designed based on the GenBank database accession number FJ644567 as follows: forward primer, 5'–CATATGATGGCGTTTCTTTCCGCAA–3', and reverse primer, 5’–CTCGAGTTATTGGTTAGACTTGCCC–3’. The obtained PCR fragment was cloned into the pGEM-T Easy vector (Promega, Madison, USA), and the nucleotide sequence was determined with bioinformatics tools. Phylogenetic analysis was analyzed by the neighbor joining (NJ) and maximum likelihood (ML) methods using MEGA software. Five hundred bootstrap replications were performed for the NJ trees to check the reproducibility of the results. TSP3-2 sequence alignment was performed by ClustalX2 and GeneDoc software. Then, the TSP3-2 fragment was cloned into the pGEM-T Easy plasmid. The primers for the TSP3-2 gene were as follows: forward primer, 5’-CATATGGAACCGATCCA-3’, and reverse primer, 5’-ACAGTATCTAACCCAAGCCAAAAGCTT-3’.

#### Preparation of banana shrimp TSP3-2 protein

The shrimp TSP3-2 gene was cloned in frame with a histidine fusion system into the _HindIII_ and _NdeI_ restriction sites in the pCold vector (Takara, Tokyo, Japan) to obtain the pCold-TSP3-2 clone. In the first step, the TSP3-2 fragment and pCold vector were digested with the _HindIII_ and _NdeI_ restriction enzymes. The fragments were observed by a 1.2% agarose gel and then purified from the gel with the QIAquick PCR purification kit (QIAGEN, USA). TSP3-2 was subsequently ligated into the pCold vector at the _HindIII_ and _NdeI_ restriction sites. The whole ligation reaction was incubated with competent _E. coli_ Top10F' cells. Colonies were grown overnight at 37 °C on 100 μg/mL ampicillin plates. The plasmid DNA was extracted to select positive clones. The pCold-TSP3-2 clone was then
transformed into *E. coli* BL21(DE3) cells.

The *E. coli* strain BL21(DE3) containing pCold-TSP3-2 was grown overnight at 37 °C in LB medium containing 100 µg/mL ampicillin. Pre-warmed medium (300 mL) was inoculated with 30 mL of an overnight culture and grown at 37 °C with vigorous shaking until a log-phase OD was reached. Protein expression was induced with 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG), and the cells were further grown at 30 °C for 16–18 h. The bacterial cells were harvested by centrifugation (4000 × g at 4 °C), resuspended in lysis buffer (50 mM NaH$_2$PO$_4$, pH 8.0, 300 mM NaCl, 6 M urea and 5 mM imidazole), lysed by sonication (20 × 10 s, 200–300 W), and centrifuged at 2000 × g for 20 min at 4 °C. The cell debris and supernatant were examined by an SDS-PAGE gel with Coomassie brilliant blue staining.

The protein in the supernatant was further purified by elution from Ni-NTA resin connected to the AKTA prime chromatography system (GE Healthcare Bio-Sciences AB, Uppsala, Sweden). The column was equilibrated with 25 mL of 50 mM phosphate buffer pH 7.4 containing 300 mM NaCl and 10 mM imidazole and washed with 300 mL of the same buffer at a flow rate of 5 mL/min. Then, 10 mL of the sample was injected and eluted with 30 mL of the same phosphate buffer but with 300 mM NaCl and 250 mM imidazole. Ten fractions (1 mL) were subsequently collected from the flowthrough. The proteins in each fraction were detected by 12% SDS-PAGE and Western blot analysis. To remove imidazole, the purified His-TSP3-2 protein was dialyzed in the same phosphate buffer with 300 mM NaCl for 24 h. The purified proteins were kept at −80 °C until use.

**Thermal stability of HIS-TF-TSP3-2**

The thermal stability of His-TF-TSP3-2 was analyzed using Thermal shift assay methods [22]. The 5000X SYPRO orange fluorescent stain for protein was diluted 50-fold in 100 mM HEPES with 150 mM NaCl, pH 7.5. The reaction consisted of 5X SYPRO orange, 100 mM buffer, 100 mM NaCl, and 0.138 µg/mL His-TSP3-2 protein. Deionized water was added instead of the test protein in the control samples. The reactions were centrifuged for 1 min at 1000 rpm. The tubes were heated in a Mx3005P Q-PCR system from 25–90 °C in increments of 0.5 °C. A graph of the relationship between temperature and the fluorescent signal was plotted and analyzed. The melting temperature was determined from the midpoint of the unfolding transition.

**Skin fibroblast proliferation and viability assay**

A human skin fibroblast cell line (Millipore’s BiogROTM Xeno-Free Human Foreskin Fibroblasts, USA) was cultured in Dulbecco’s modified Eagle’s medium (DMEM, Gibco, USA) containing 10% fetal bovine serum (FBS, Gibco, USA) and antibiotics (100 U/mL penicillin and 100 U/mL streptomycin, Gibco, USA) with 5% CO$_2$ at 37 °C. The media was changed every other day. When the cells reached confluence, they were harvested using 0.25% trypsin-EDTA (Gibco, USA), followed by the addition of fresh culture medium to generate a new single-cell suspension for further cell passaging.

Cytotoxicity was evaluated by the method described [23]. The skin fibroblast cell line was seeded at a concentration of 5 × 10$^4$ cells/mL into a 96-well plate in DMEM containing 10% FBS. After 24 h, the culture medium was replaced by fresh medium. His-TF-TSP3-2 at various concentrations (i.e., 0, 0.5, 1, 5, 7.5, 10, 15, and 30 µg/mL) in fresh medium was added to the culture medium. Cells without His-TF-TSP3-2 served as the negative control. After incubation for 24 h, a 3-(4,5-dimethyl thiazolyl-2-yl)-diphenyl-tetrazolium bromide (MTT) solution (5 mg/mL) assay was performed to evaluate cell activity. Briefly, the cells were treated with 100 µL of fresh medium along with 50 µL of MTT solution, followed by incubation at 37 °C with 5% CO$_2$ for 4 h. Thereafter, the medium containing MTT was removed, and 100 µL of DMSO was added. The absorbance was determined with a microplate reader (Biohit 830, Helsinki, Finland) at a wavelength of 570 nm. The percentage of cell proliferation was calculated and compared to the negative control.

**In vitro scratch assay**

The spreading and migration capabilities of human skin cells were assessed using a scratch assay (injury to the cell monolayer) that measured the expansion of a cell population on a surface as described [26, 27]. The skin fibroblast cell line was seeded at a concentration of 9 × 10$^5$ cells/mL in DMEM containing 10% FBS in a 6-well plate. Once a confluent monolayer was formed, a linear scratch was generated in the monolayer with a sterile pipette tip. Any cellular debris was removed and replaced with 2 mL of His-TF-TSP3-2 and His-TF at a concentration of 0.5 µg/mL in fresh medium, and DMEM without a sample served as a control. Microphotographs were taken at 10 × magnification using an Olympus IX73 inverted microscope (Bangkok, Thailand) on day 0. Then, the plates were incubated with 5% CO$_2$ at
37°C, and photographs were taken at 24, 36, and 48 h after incubation. The images acquired for each sample were further analyzed quantitatively using computing software (ImageJ 1.42q/Java 1.6.0 10). By comparing the images from 0–24, 36, and 48 h, the distance of each scratch closure was determined and the percent migration rate was calculated. The average values for the left scratch and right scratch were determined separately. The percent migration was calculated for the left scratch and then the right scratch. Samples were examined in triplicate. The percentage of migration obtained from three wells was averaged and recorded.

\[
\text{Migration rate} = \frac{\text{DAS}_0 - \text{DAS}_{ep}}{\text{DAS}_0}
\]

where DAS₀ and DASₑ are the average distance across the scratch (day 0) and across the scratch (experimental periods), respectively.

**Determination of soluble collagen production**

Soluble collagen was quantified by the method described [23]. The BJ human skin fibroblast cell line was seeded at an initial concentration of 2 × 10⁵ cells/mL in 96-well plates in DMEM supplemented with 10% FBS. After 24 h, the culture medium was replaced with fresh DMEM plus 10% FBS, and His-TF-TSP3-2 was added to the culture medium at concentrations of 0.5 and 1 μg/mL in fresh medium. After incubation for 24 h, the supernatants were collected. Then, the total amount of soluble collagen was assayed using the Sorcol collagen assay kit (Biocolor Ltd., Carrickfergus, UK). Briefly, 100 μL of the experimental cell supernatant was mixed gently with 1 mL of dye solution at room temperature for 30 min. Thereafter, the samples were centrifuged at 15000×g for 10 min to form a collagen pellet. The supernatant was removed, and the soluble collagen was dissolved in 1 mL of alkali reagent. The resultant solutions with the alkali reagent were transferred to a 96-well plate, and measurements were performed with a microplate reader at a wavelength of 540 nm. The amount of collagen was calculated based on a soluble collagen standard curve. DMEM supplemented with 10% FBS without His-TSP protein was used as a negative control.

**Determination of TSP3-2 protein genotoxicity**

CHO cells (1 × 10⁵ cells) were seeded per well in a 48 well plate and incubated for 36–40 h. The cells were exposed to a concentration of 0.5 or 1 μg/mL of His-TF-TSP3-2 protein for 24, 48, or 72 h at 37°C, after which the medium containing the His-TF-TSP3-2 protein was removed. The 1-methyl-3-nitro-1-nitrosoguanidine (MNNG) (Tokyo Chemical Industry Co., Ltd, Japan) and 7,12-dimethylbenz(a)anthracene (DMBA) (Sigma Aldrich, US) mutagens were used as positive controls while PBS and DMSO were used as negative controls. The cells were cultured in DMEM for 8 days. Mutant cell selection was performed by treatment with 5 μg/mL 6-thioguanine (6-TG) (Sigma Aldrich, US) in DMEM. After 10 days of growth, the cells were stained and counted to determine the mutant frequency.

**Statistical analysis**

Three separate samples were used for all the experiments in this study. The mean of three samples was calculated. The results were statistically compared between groups using the independent samples t-test via SPSS software at a 95% confidence level (p < 0.05).

**RESULTS**

**FmTSP sequence analysis and preparation of HIS-TF-TSP3-2**

Full-length FmTSP was amplified based on information from the GenBank database. The nucleotide sequence of the obtained PCR fragment was determined and analyzed. The full-length TSP sequence contained 2769 bp encoding 922 amino acids. FmTSP was found to consist of 3 chitin-binding repeats, an epidermal growth factor domain (TSP3-2), 8 thrombospondin-3 domains and a thrombospondin C-terminal domain (Fig. 1ab). Phylogenetic analysis of the TSP family was performed, and the results revealed that TSPs from shrimp are closely related to each other. Moreover, shrimp FmTSP was closer to TSP3 and TSP4 than TSP1 and TSP2 (Fig. 1c). The epidermal growth factor domain of FmTSP was defined as TSP3-2, and it was cloned to characterize its activity for potential applications. TSP3-2 was 462 bp in length and translated into 154 amino acids (aa); when the protein sequence of TSP3-2 was compared with EGF proteins from other organisms, it displayed 90% similarity to human EGF, and the TSP3-2 domain contained 13 phosphorylation sites including 9 serine, 1 threonine, 2 tyrosine and 6 cysteine residues, which are characteristic of epidermal growth factor
proteins. The shrimp TSP3-2 gene was cloned in frame with a histidine fusion system into the pCold vector (Takara, Tokyo, Japan) and expressed in *E. coli* BL21(DE3). The His-TF and His-TF-TSP3-2 were examined by SDS-PAGE; they were produced after induction with IPTG as shown in lanes 2 and 4 of Fig. 2a. The His-TF-TSP3-2 protein after purified by affinity chromatography was separated on 12% SDS-PAGE; a single band was observed (Fig. 2b). His-TF-TSP3-2 was expected to contain 613 aa, consisting of 459 aa of His-TF and 154 aa of TSP3-2, with a molecular
Fig. 2 TSP3 protein expression, purification and detection. (a) The proteins were separated by 12% SDS-PAGE with Coomassie blue staining of the cell-free extracts: lane 1, molecular weight markers; lane 2, non-induced pColdTF cell lysate; lane 3, induced pColdTF cell lysate; lane 4, non-induced His-TF-TSP3-2 cell lysate; and lane 5, induced His-TF-TSP3-2 cell lysate. (b) Analysis of the purified His-TF-TSP3-2 protein by 12% SDS-PAGE: lane 1 molecular weight markers and lane 2 the purified protein. (c) His-TF-TSP3-2 was detected with an anti-His monoclonal antibody after Western blotting: lane 1, molecular weight markers; lane 2, His-TF after induction; and lane 3, His-TF-TSP3-2 after induction.

Fig. 3 Thermal stability of His-TF-TSP3-2. The linear line represents the $T_m$ of His-TF-TSP3-2. Error bars are the mean ± SD (N = 3).

Biological and physical properties of HIS-TF-TSP3-2

A thermal shift assay was used to determine the thermal stability of His-TF-TSP3-2. A graph of the relationship between temperature and the fluorescence signal was plotted and analyzed. SYPRO orange can bind to unfolded protein, and aggregation of protein:dye complexes leads to quenching of the fluorescence signal. The melting temperature was identified at the midpoint of the unfolding transition. The assay showed that the melting temperature of His-TF-TSP3-2 was 47.5 °C, which means that the His-TF-TSP3-2 protein could exhibit some activity at room temperature (Fig. 3).

Cytotoxicity was evaluated by the method described [23]. His-TF-TSP3-2 at various concentrations (i.e., 0, 0.5, 1, 5, 7.5, 10, 15, and 30 µg/mL) in fresh medium was added to the skin fibroblast cell line. The percentage of cell proliferation was calculated and compared to a negative control after 24 or 48 h of incubation. Fig. 4 shows that His-TF-TSP3-2 at 0.5 µg/mL increased cell proliferation whereas the other concentrations did not show activity after 24 h of the treatment of skin fibroblasts. After 48 h of the treatment with 0.5, 1, 5, 7.5, 10, or 15 µg/mL His-TF-TSP3-2, the proliferation of the skin fibroblasts was 1.2–1.4 fold increased compared to the control without treatment. Therefore, the best concentration for cell proliferation was 0.5 µg/mL.

In addition, the activation of the wound healing of human skin after cell injury by His-TF-TSP3-2 was determined by measuring the migration of a cell population on a scratched surface of the cell monolayer [24]. Cells treated with His-TF-TSP3-2 at a concentration of 0.5 µg/mL showed percent migration rates of 24.00, 43.79, and 67.86% after treatment for 24, 48, and 72 h, respectively. The percent migration rates of the first control cells (no treatment) were 15.69, 27.39, and 55.84% at 24, 48, and 72 h, respectively.

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were 21.76, 37.56, and 44.01% at 24, 48, and 72 h, respectively (Fig. 5). Comparison of the percent migration rates of the His-TF-TSP3-2-treated cells from 0–24, 36, and 48 h with both controls showed that the percent migration rate of the 0.5 µg/mL His-TF-TSP3-2 group was much faster than those of the controls.

Soluble collagen was also measured in the culture medium of the human skin fibroblast cell line after treatment with His-TF-TSP3-2 for 24 h. The collagen production was 1.7-fold and 1.2-fold increased compared with the control after treatment with His-TF-TSP3-2 at concentrations of 0.5 and 1 µg/mL, respectively (Fig. 6).

**Genotoxicity of the HIS-TF-TSP3-2 protein**

After CHO cells were exposed to 0.5 or 1 µg/mL of His-TF-TSP3-2 protein at 37°C for 24, 48, or 72 h, the genotoxicity was evaluated; the MNNG and DMBA mutagens served as positive controls, and the PBS and DMSO were used as negative controls. Mutant cells were selected to determine the mutant frequency, and no significant mutant frequency was found in the His-TF-TSP3-2-treated cells compared to the untreated cells (Fig. 7). The mutant frequencies were 35% and 6.5% in the MNNG and DMBA mutagen-treated cells, respectively.

**DISCUSSION**

Human EGF is known for its cell proliferation activity in medical applications including wound healing and has been used since 1989 [20]. It has been shown to enhance the healing process in peripheral tissue wounds and gastrointestinal damage. A study in transgenic mice overexpressing EGF showed that this protein may not be sufficient to transform cells, and no effects of EGF were observed on either in vitro or in vivo human gastric cancer cell growth [25, 26]. In contrast, long-term EGF administration caused epithelial organ hyperplasia in a dose-dependent manner with no change to cell phenotype differentiation, or no tumor markers were found [27]. Clinically, a topical recombinant human EGF was tested in 77 patients with skin lesions and showed promising results; however, the researchers suggested that patients should have individualized formulations for efficient treatment, and large scale studies should be performed [28]. Therefore, the balance between the advantages and disadvantages of human EGF are currently under investigation.
Thus, an EGF-like domain from other sources should be evaluated to get good candidates for treatment.

The TSP3 proteins from the shrimp species *P. monodon*, *F. chinensis*, and *F. merguiensis* (examined in this study) contained an EGF-like domain whereas 6 and 3 EGF domains have been reported in *P. japonicas* and human TSP3, respectively. TSP in shrimp has been suggested to act as a protective protein in oocytes, as it is present in the cortical rod in the ovary along with SOP (shrimp ovarian peritrophin). Additionally, TSP is similar to SOP because the structure contains a chitin-binding domain that plays a role in protecting the ovary from microorganism invaders [29, 30]. However, TSP3 in shrimp was found to increase during the ovarian maturation process, which implies a role of TSP in proliferation activity in the shrimp ovary. The effect of the EGF domain on the proliferation activity of cells was shown in both the MTT and scratch assays in this study. It was also found that treatment with shrimp EGF (His-TF-TSP3-2) at 0.5 µg/mL resulted in a 1.7-fold increase in the production of soluble collagen. The role of EGF is also critical at the initial stages of wound healing.

EGF stimulates the proliferation of fibroblasts and keratinocytes around lesions that occur during the early phase of healing [31]. This growth factor has been applied to assist in wound healing. It has been reported that an effect of EGF is to induce the expression of the gene encoding phosphatidylinositol 3-kinase (PI 3-kinase) during rabbit corneal epithelial wound repair [32]. The stimulation of cell line migration by commercial EGF has been reported. EGF induces cell migration in cultured human lens epithelial cells through the ERK and PI3k/AKT pathways [15]. Therefore, the ability of our recombinant EGF (His-TF-TSP3-2) to contribute to cell migration indicated that this EGF protein has a similar function to commercial EGF. Since only 30% nucleotide sequence similarity between EGF from *Fm*TSP3 and human TSP3 was observed, shrimp EGF may present some differences from that of humans in terms of activity and specificity. Therefore, EGF (His-TF-TSP3-2) from *Fm*TSP was evaluated for genotoxicity, and no significant mutant frequency was found in His-TF-TSP3-2-treated cells compared to untreated cells.

Moreover, our recombinant EGF protein increased the collagen level of the skin fibroblast cell lines. Several studies have shown that EGF affects collagen biosynthesis in cultured cells [19]. EGF increases the contents of acidic glycosaminoglycans (AGAGs), which are thought to be closely related to the formation of collagen fibers [33]. AIMP1-derived peptide (AdP, INCI name: sh-oligopeptide-5/sh-oligopeptide SP) has been proposed as an alternative product to EGF because of its high sensitivity to the EGF concentrations applied [34]. Although the recombinant His-TF-TSP3-2 had quite a high melting temperature and low sensitivity in humans, further pharmaceutical testing of this TSP3-2 protein is required to confirm its benefit as an anti-wrinkle or a skin renewal agent.

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