Title: MASS PRODUCING METHOD OF GROWTH FACTOR USING ADIPOSE DERIVED ADULT STEM CELLS

Abstract: The present invention relates to a method for producing large amounts of human growth factors from human adipose-derived stem cells. More specifically, the invention provides a method capable of synthesizing human growth factors in significantly large amounts by culturing adipose-derived stem cells extracted from human adipose cells in suitable media and conditions. Also, stem cell culture media produced according to the method of the invention, and human growth factors isolated from the culture media, can be advantageously used as raw materials for drugs and cosmetics.
For two-letter codes and other abbreviations, refer to the “Guidance Notes on Codes and Abbreviations” appearing at the beginning of each regular issue of the PCT Gazette.
Invention Title

MASS PRODUCING METHOD OF GROWTH FACTOR USING ADIPOSE DERIVED ADULT STEM CELLS

[Technical Field]

The present invention relates to a method for producing large amounts of growth factors from human adipose-derived stem cells. More specifically, the present invention provides a method which comprises culturing adipose-derived stem cells extracted from human fat cells in suitable media and conditions, such that human growth factors, for example, an acidic fibroblast growth factor (acidic FGF), a basic fibroblast growth factor (basic FGF), an insulin-like growth factor-1 (IGF-1), an insulin-like growth factor-2 (IGF-2), a keratinocyte growth factor, a platelet-derived growth factor (PDGF), a human transforming growth factor-alpha (TGF-\(\alpha\)), a human transforming growth factor-beta (TGF-\(\beta\)), a vascular endothelial growth factor (VEGF), an epidermal growth factor (EGF) and a nerve growth factor, can be synthesized in amounts significantly larger than the case of use of existing stem cells.

[Background Art]

As used herein, the term "stem cells" refers to cells, which remain undifferentiated into particular cells, and, if necessary, has the potential to differentiate into all kinds of cells constituting the body, including nerve, blood, cartilage and the like. Methods capable of producing such stem cells are broadly classified into two categories: (1) a method of producing stem cells from embryos developed from fertilized eggs (embryonic stem cells); and (2) a method of recovering stem cells stored in each part of the adult human body (adult stem cells). Although embryonic stem cells and adult stem cells are functionally different from each other, they all have a characteristic in that they can differentiate into various cells.

The embryonic stem cells have advantages of very good differentiation potential and long telomerizes, but have ethical problems and a shortcoming in that they are difficult to obtain in large amounts. In comparison with this,
the adult stem cells can be large amounts, but have shortcomings in that, when these cells are transplanted into other persons, they have the risk of infection and a relatively low differentiation potential.

Despite the above-described shortcomings, the adult stem cells have a characteristic in that they are considerably safe for use in medical applications. Specifically, these cells do not cause cancer even when they are transplanted into the body for organ regeneration, and they do not cause immune rejection reactions because they have been originated from the body of oneself. Thus, these adult stem cells can be used for autologous transplantation.

Also, the adult stem cells have site-specific differentiation potential to differentiate according to the characteristics of the peripheral tissues, and do not cause cancer even when they are injected in an undifferentiated state. Thus, these adult stem cells have the potential to produce cells immediately after transplantation and show also the self-renewal potential to create and store undifferentiated stem cells, if necessary.

Due to the above-described advantages, the importance of adult stem cells has recently been highlighted, and various studies have been conducted to obtain adult stem cells in vivo.

Adipose tissue plays an important role in normal growth and physiological action in vivo, but the importance thereof has been missed till now. The most general type of fat is white adipose tissue, which is located below the skin (subcutaneous fat) and located in the abdominal cavity (visceral fat) or located around the reproductive organ (gonadal fat). A slightly less general form in an adult is brown adipose tissue, which plays an important role in the production of heat during the infant stage (Gimble, New Biol. 2(4): 304-12, (1990)).

However, in fact, reproductive capability and mature stage are closely associated with adipose tissue storage in individuals. Female and male adolescence is closely associated with the production and secretion of
adipose tissue-derived hormones and are closely associated with body fat composition. Also, adipose tissue plays an important role in glucose metabolism and energy balance.

For a few years, there has been a significant advancement in the biomaterial field. On the basis of this, many materials are currently developed and used. Despite this advancement, many studies have, in fact, not been conducted on the use of human adipose tissue. However, since it was recently reported that adult stem eel's are present in adipose tissue (Zuk PA, et al., Molecular Biology of Cell, 13: 4279-4295 (2002); Rodriguez AM, et al., Biochimie, 87: 125-128 (2005)), various studies on the use of adipose-derived cells have been conducted.

Also, on the basis of the development of biochemistry and molecular biology, small amounts of signaling substances (growth factors) have recently been found in the human body, and on the basis of these findings, theories on in vivo aging have been re-established (Stanley Cohen, Nobel Lecture 1986, Dec, 8). Moreover, it has been found that the signaling substances (growth factors) decrease as age increases, and this decrease in the growth factors is closely associated with the aging of the human body (Sporn M and Roberts A, Handbook of Experimental Pharmacology, Vol. 1, Vol. 95/1, 1990, Springer-Verlag, DE, Berlin., pp. 667-698).

Thus, it has been reported that, when external growth factors are supplied into the body, the aging of the body can be inhibited, and studies on special effects of these substances have been conducted (GE Pierce and TA Mustoe, Annu Rev Med, 46. 467-481 (1995)). Specifically, studies on the structure and synthesis of the signaling substances (growth factors) have been conducted, but the most of the growth factors have protein structures, which are three-dimensionally complex, thus causing various problems in chemical synthesis and significantly increasing costs for the synthesis.

Accordingly, the present inventors have conducted studies on a method of obtaining active human growth factors at low cost while maintaining the activity thereof in the human body and, as a result, paid attention to the
fact that adipose-derived adult stem cells secrete growth factors (Rehman, J. et al., Circulation, 109: 1292 - 1298 (2004)).

However, studies on adipose-derived adult stem cells mostly relate to the use or differentiation of the cells themselves, and there have been almost no studies on a method of synthesizing growth factors from these cells.


It is considered that the reason why studies on the synthesis of growth factors using adipose-derived adult stem cells are insufficient is that the methods of using adult stem cells are focused on the use thereof through differentiation into other cells and that studies on the cytological differences between stem cells are insignificant.

Also, as described above, methods for synthesizing growth factors have been studied only on methods of producing growth factors using genetic recombination techniques, as disclosed in Korean Patent Registration No. 101436, entitled "Method for producing recombined human endothelial cell

Accordingly, the present inventors have studied on a method for producing large amounts of growth factors using adipose-derived adult stem cells and, as a result, found that adipose-derived adult stem cells obtained through the establishment of suitable culture conditions and physical and/or chemical stimulation synthesize and secrete growth factors in significantly effective amounts compared to adipose-derived adult stem cells to which specific stimulation is not applied.

[Disclosure]

[Technical Problem]

It is an object of the present invention to produce large amounts of human growth factors from adipose-derived adult stem cells, the produced human growth factors having excellent in vivo activity compared to growth factors synthesized by recombinant or chemical methods.

Another object of the present invention is to provide safe and effective drugs or cosmetics containing either human growth factors produced in large amounts from adipose-derived stem cells, or culture media of the growth factors.

[Technical Solution]

To achieve the above objects, in one aspect, the present invention provides a method for producing large amounts of human growth factors from adipose-derived stem cells, the method comprising the steps of: (i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells; (ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium! (iii) applying, to the adipose-derived stem cells, at least one physical stimulation selected from among low-oxygen culture, UV irradiation, nutrient deficiency and mechanical friction; and
(iv) optionally adding, to the culture media, at one or more vitamins selected from among vitamin A, vitamin B, vitamin C and vitamin D, wherein the step (iii) and the step (iv) are performed in conditions where the highest production of the human growth factors occurs.

In another aspect, the present invention provides a functional cosmetic composition containing human growth factors produced by said method.

In still another aspect, the present invention provides a functional cosmetic composition containing an adipose-derived adult stem cell culture medium obtained by said method.

[Advantageous Effects]

According to the present invention, it is possible to produce human growth factors in large amounts from adipose-derived adult stem cells, and it was found that the growth factors produced using the inventive production method had excellent safety and activity compared to those of growth factors produced according to existing production methods, and could act in the same fashion as the existing growth factors of the human body. Also, it is expected that a culture medium of adipose-derived stem cells and growth factors separated from the culture medium can be advantageously applied in drugs, quasi drugs and cosmetics for anti-wrinkle, wound healing, and scar removing.

[Description of Drawings]

FIG. 1 is an optical microscopic photograph of stem cells isolated from adipose tissue;

FIG. 2 shows the results of flow cytometry of originally isolated PLA eel Is;

FIG. 3 is a graphic diagram showing the results of flow cytometry after subculture;

FIG. 4 is an electrophoresis photograph of TGF β-1, bFGF and VEGF, confirmed through RT-PCR from adipose-derived stem cells;

FIG. 5 is a graphic diagram showing the concentrations of bFGF and VEGF secreted in culture media during the culture of adipose-derived stem
FIG. 6 is a graphic diagram showing the measured collagen synthesis of fibroblasts in mixed culture of fibroblasts and adipose-derived stem cells;

FIG. 7 is a graphic diagram showing the number of fibroblast cells in varying concentrations of adipose-derived stem cell culture media;

FIG. 8 is a graphic diagram showing the comparison of cell proliferation potential between adipose-derived stem cell culture media and recombinant growth factor-containing media;

FIG. 9 is a graphic diagram showing the comparison of secretion of a basic fibroblast growth factor (bFGF) between adipose-derived stem cells cultured in physical and chemical conditions according to the present invention;

FIG. 10 is a graphic diagram showing the comparison of secretion of a vascular epithelial growth factor (VEGF);

FIG. 11 is a graphic diagram showing the secretion of transforming growth factor-beta (TGF-β);

FIGS. 12, 13, 14 and 15 are photographs showing the wrinkle-reducing activity of a composition containing growth factors isolated from autologous adipose-derived adult stem cells; and

FIG. 16 is a photograph showing the results of Western blot analysis for the collagen synthesis of fibroblasts, caused by adipose-derived stem cell culture media.

[Mode for Invention]

The first aspect of the present invention relates to a method for producing large amounts of growth factors using adipose-derived stem cells.

A growth factor contained in adipose-derived stem cells is selected from the group consisting of an acidic fibroblast growth factor (acidic FGF), a basic fibroblast growth factor (basic FGF), an insulin-like growth factor-1 (IGF-I), an insulin-like growth factor-2 (IGF-2), a keratinocyte growth factor (KGF), a platelet-derived growth factor (PDGF), a human transforming growth factor-alpha (TGF-α), a human transforming growth factor-beta (TGF-
β), a vascular endothelial growth factor (VEGF), an epithelial growth factor (EGF), a nerve growth factor (NGF), and mixtures thereof.

In the present invention, adipose-derived stem cells were used to specifically stimulate the synthesis of the following growth factors.

1) Basic fibroblast growth factor (hereinafter, referred to as "bFGF")

bFGF or heparin-binding growth factor 2 (HBGF-2) includes seven kinds of factors having a homology of about 30-50% in the amino acid level (Burgess, W.H and Maciag, T., Annu. Rev. Biochem., 58: 575-606 (1989); Baird, A. and Klagsbrun, M., Cancer Cells, 3(6): 239-43 (1991)). bFGF is isolated from nerve tissue, the pituitary body, the adrenal cortex, corpora leutea, and the placenta.


2) Vascular endothelial growth factor (hereinafter, referred to as "VEGF")

VEGF (Ferrara, N. and Henzel, W.J. Biochem Biophys Res Commun, 161(2): 851-8 (1989)) is known as a vascular permeable factor (Senger, D.R. et al.,
Science, 219(4587): 983-5 (1983)), which is a homodimer, has a molecular weight of 34-42 kDa and is a heparin-bound glycoprotein. This has an angiogenesis factor, and thus can promote the mitosis of epithelial cells and improve vascular permeability.

VEGF is expressed as a primary structure known as a limited base sequence and has a homology with the A and B chains of a platelet-derived growth factor (PDGF). Such growth factors have eight conserved cystein residues and are included in chains inside and outside disulfide bonds. cDNA encoded by the VEGF protein has an amino acid homology of 53% to the platelet-derived growth factor (PDGF), and VEGF is isolated from a cDNA library of the human placenta (Maglione, D. et al., PNAS, 88: 9267 (1991)). This protein is called "placenta growth factor" (PGF), and is currently recognized as a member of the VEGF family. Based on homology with VEGF, the placenta growth factor (PGF) is suggested as an angiogenesis factor.

Genes for the human VEGFs are combined in eight exons. Alternative splicing yields base sequences of 121, 165, 189 and 206 amino acids, which encode four monomeric VEGFs. Each of the base sequences has 26 signal peptide amino acid residues, and thus can be detected. VEGF121 and VEGF165 are diffusible proteins exposed to extracellular matrixes, and VEGF189 and VEGF206 have high affinity for heparin, and thus form proteoglycan bonds with heparin in extracellular lipids. VEGF contains an N-Linked glycosylation site, which is originally a glycoprotein.

The VEGF proteins produced by general recombinant or chemical methods are produced on the basis of the diffusible proteins VEGF121 and VEGF165 sites. In the prior art, it was reported that the expression of recombinant human VEGF in E. coli was not different from original VEGF in vitro with respect to biological functions (Connolly, D.T. J. Cell. Biochem, 47(3): 219-23 (1991); Schott, R.J. and Morrow, L.A. Cardiovasc, Res., 27(7): 1155-61 (1993); Neufeld, G. et al., Prog. Growth Factor Res., 5(1): 89-97 (1994); Senger, D.R. et al., Cancer and Metastasis Reviews, 12(3-4): 303-24 (1993)).

3) Transforming growth factor- β (hereinafter, referred to as "TGF-β")
A human transforming growth factor (TGF), which is a factor that stimulates the transformation of fibroblasts differentiating into a tumor-like phenotype, consists of a mixture of two proteins TGF-α and TGF-β and is found with tumor-inhibitory factors rather than a tumor-stimulating factors (Lawrence, D.A. Eur. Cytokine Netw. 7(3): 363-74 (1996); Cox, D.A. and Maurer, T., Clin. Immunol. Immunopathol. 83(1): 25-30 (1997); Alevizopoulos, A. and Mermod, N., Bioassays. 19(7): 581-91 (1997)).

The two molecules are members of the super-family including TGF-β1 having five kinds of bone morphogenic proteins as active and inactive substances (Kingsley, D.M., Genes Dev, 8: 133 (1994)). It is known that human TGF-β1 has a molecular weight of 25 kDa, consists of bisulfide bonds and bi-glycosyl homodimers, and has a 100% conserved gene sequence between almost all mammalian species. TGF-β1 initiates intracellular signaling as a bisulfide bond is transferred from a precursor by a protease similar to subtilisin (Dubois, C.M. et al., J. Biol. Chem., 270: 10618 (1995)), and it is generally secreted as an inactive material or a composite of two (Gleizes, P-E. et al., Stem Cells, 15: 190-197 (1997)). A TGF-β1 signaling process includes two receptors (Ten Dijke, P. Curr. Opin. Cell Biol, 8(2): 139-45 (1996); Derynck, R. and Feng X.H., Biochim. Biophys. Acta 1333(2): F105-50 (1997); Padgett, R.W. et al., Bioassays, 20(5): 382-90 (1998)), and a TGF-β RII dimmer as a 75-kDa ligand-binding protein has an intracellular serine-threonine kinase enzyme, which is continually activated, and TGF-β RII phosphorylates 53-kDa signaling dimmer TGF-β RI through binding to TGF-β1. The phosphorylated TGF-β RI activates protein kinase and induces initiating a downstream signal via intracellular protein SMADS.

The TGF receptor involved in a signaling process is found in all cells and affects almost all physiological actions. The systematic and cell-specific activation thereof is a very complex mechanism, but shows three basic activities. TGF-β1 generally regulates the proliferation of cells such as inhibitory factors, promotes the precipitation of protein hydrolysates in addition to cell membranes by repeating the inhibition of
protein degradation and the synthesis of proteins, and stimulates immune
inhibitory reactions through various mechanisms.

A TGF-β1 protein produced by prior recombinant or chemical synthesis
is an active protein having a size of 25 kDa, and thus shows biological
activity similar to the original TGF-β1 protein in vitro, but it contains
only some of the inherent properties of the TGF-β1 protein.

The method according to the first aspect of the present invention
comprises the steps of: (i) isolating adipose-derived adult stem cells
extracted from mammalian adipose cells; (ii) selectively culturing the stem
cells in a serum medium and then subculturing the stem cells in a serum-free
medium! (iii) applying, to the adipose-derived stem cells, at least one
physical stimulation selected from among low-oxygen culture, UV irradiation,
nutrient deficiency and mechanical friction; and (iv) selectively adding, to
the culture media, at one or more vitamins selected from among vitamin A,
vitamin B, vitamin C and vitamin D, wherein the step (iii) and the step (iv)
are performed in conditions where the highest production of the human growth
factors occurs.

Collection of stem cells

Adipose-derived adult stem cells according to the present invention
can be collected through a purification process from cells present in adipose
tissue. Preferably, human adipose-derived adult stem cells are collected,
and for this purpose, adipose-derived stem cells are separated from human
adipose tissue.

Adipose tissue is brown or white adipose tissue derived from
subcutaneous, network membrane, intestinal, breast genital or other adipose
tissue sites, and subcutaneous white adipose tissue can be conveniently
obtained using liposuction.

As adipose tissue, it is possible to use adipose tissue wasted in a
liposuction process, which is generally performed. That is, the utility of
the adipose tissue is increased, because the need to perform invasive surgery
is eliminated. The separated liposuction material is washed, and only
adipose tissue is separated from the washed material. The extracellular matrix of the adipose tissue is treated with collagenase, and then centrifuged to collect a stromal vascular fraction containing a high density of stem cells. The pellets thus obtained are washed, and then passed through a cell filter to other tissues. Then, monocytes and cell fragments containing red blood cells are isolated from the remaining tissue using a monocyte isolation solution. The isolated monocyte cells are cultured in non-inducing media, and nonadhesive cells are removed.

**Culture of stem cells**

In the present invention, special culture media were established for in vitro culture of the adipose-derived stem cells obtained in the above-described process.

Before the start of cell culture, biological samples extracted from a supply source can be repeatedly washed with a wash medium containing general antibiotics to minimize the possibility of contamination in subsequent culture.

In the present invention, culture media are optimized so as to the maximum synthesis and secretion of growth factors in the adipose-derived stem cells.

Specifically, the in vitro culture of the adipose-derived stem cells is performed by culturing the cells in a serum medium and then subculturing the cells in a serum-free medium, such that the synthesis of growth factors is maximized.

The serum-containing medium for the initial cell culture is preferably a medium suitable for maintaining and storing cell types such as adipose-derived stem cells.

In the present invention, the medium is based on Dulbecco's Modified Eagle's Medium (DMEM), which is generally used in cell culture in the art, and it contains serum, which is generally used in cell culture.

Herein, the initial medium may also be a frozen medium obtained by adding 7-10% of dimethyl sulfoxide (DMSO) thereto, and thus the stem cells
can be frozen, and then, if necessary, can be thawed before use.

As the serum, 0.1-20% fetal bovine serum (FBS) is preferably added, and antibiotic agents, antifungal agents and reagents for preventing the growth of micoplasma causing contamination are more preferably added to the medium.

As the antibiotic agents, it is possible to use all antibiotic agents used in general cell culture, including penicillin-streptomycin. As the antifungal agent, it is preferable to use amphotericin B, and as the micoplasma inhibitory agent, it is preferably to use tylosin. In addition, the micoplasma contamination can be prevented with gentamicin, ciprofloxacin, azithromycin and the like. If necessary, oxidation nutrients such as glutamine, and energy metabolites such as sodium pyruvate, can further be added to the medium.

A more preferred medium contains 1-2 mM glutamine, 0.5-1 mM sodium pyruvate, 0.1-10% FBS, 1% antibiotic (100 IU/ml)-supplemented glucose and DMEM, and is called "complete serum medium". Herein, the concentration of glucose is approximately 1 g/L to 4.5g/L. The complete serum medium provides the storage and maintenance of adipose-derived stem cells and stable basic culture conditions in vitro and shows effective cell stabilization.

With respect to general culture conditions for the initial culture, the most suitable conditions for cell culture are applied, and thus the cell culture is performed in an incubator at humidity of 90-95% and a temperature of 35-39 °C under a condition of 5-10% C(V). When the cell culture is carried out in a condition of 5-10% CO₂, a carbon source such as sodium bicarbonate is added to a final concentration of 0.17-0.22%.

During the initial culture stage, the tissue fragments are preferably kept attached to the bottom of the culture flask, and the growth of the cells can be promoted through short stimulation caused by treatment with trypsin-EDTA according to standard cell culture techniques.

Cumulative population doubling time is maintained until the cells
being cultured in the flask reach a confluence of 75-85%. Preferably, the cells are collected at a confluence of 80% and subcultured in a serum-free medium for late-stage culture.

The present invention provides a serum-free medium for the differentiation of growth factors, which stimulates the differentiation of growth factors in adipose-derived stem cells.

The subculture process using the serum-free medium for the differentiation of growth factors is preferably performed by washing the flask, from which the culture medium has been removed, with phosphate buffer saline, suspending the cells with trypsin-EDTA, centrifuging the cell suspension, and then washing the resulting pellets with buffer solution, wherein the washing process is repeated 2-3 times. The washed pellets are suspended in the serum-free medium developed in the present invention and are subcultured about 3 times in a cell culture flask.

The serum-free medium developed in the present invention is based on DMEM not containing a pH indicator such as phenol red, and a Ham's F-12 nutrient mixture (SIGMA, Cancer Research Vol 47, Issue 1, 275-280) is added thereto at a ratio of approximately 1: 0.5-2. Herein, it is possible to add oxidation nutrients such as L-glutamine, energy metabolites such as sodium pyruvate, and carbon sources such as sodium bicarbonate. In addition, it is possible to add not only growth factors other than the growth factors targeted in the present invention, but also growth hormones.

The inherent characteristic of the serum-free medium developed in the present invention can be seen in the Ham's F-12 nutrient mixture. In this mixture, various inorganic substances and amino acids, which help to maintain the growth and homeostasis of the cells and are involved in increasing the safety and maintenance of the cells in the late-stage culture following the initial-stage culture of the adipose-derived stem cells, vitamin nutrients that can stimulate the higher production of growth factors selected from the adipose-derived stem cells, and other factors, are mixed with each other at a given ratio. The serum-free medium containing the Ham's F-12 mixture,
established in the present invention, do not show any reduction or negative effect on the culture of the adipose-derived stem cells and the production of growth factor compared to the conditions of general serum media containing animal serum. Also, some of growth factors show a higher productivity in the serum-free medium, all the components and contents of the culture medium can be determined, unlike serum media having unknown components caused by serum.

This suggests that various variables, which can result from animal serum contained in serum media, can be minimized, and the effects of the present invention can be achieved at a low cost of about 50% compared to the prior art.

Table 1 below indicates the components and contents of the Ham's F-12 nutrient mixture contained in the serum-free medium established in the present invention.
It is obvious that the components and contents of amino acids, vitamins and inorganic salts in the serum-free medium can be modified by one skilled in the art, as long as this modification does not deteriorate the object of the present invention.

To achieve the object of the present invention, it is possible to promote the synthesis of targeted growth factors by activating the adipose-
derived adult stem cells cultured according to the above method, through specific stimulation. Herein, the stimulation can be performed in conditions divided into physical conditions and chemical conditions.

Examples of physical stimulations may include UV rays, nutrients, and oxygen, and examples of chemical stimulations may include vitamins and other active compounds in a medium composition for cell culture.

In order to obtain growth factors in amounts significantly larger than those in existing methods for culturing adipose-derived adult stem cells, it is apply to physical stimulations, including low-oxygen reactions (Circulation. 2004 Mar 16; 109(10): 1292-8.), UV irradiation (FASEB J. 2003 Mar; 17(3):446-8), nutrient deficiency (Blood. 2004 Nov 1; 104(9): 2886-92. Epub 2004 Jun 24), and mechanical friction. Such physical stimulations can selectively or collectively increase growth factors targeted in the present invention.

In order to examine whether physical stimulations promote the synthesis and secretion of growth factors, in the present invention, the adipose-derived stem cells were cultured in vitro using the serum medium and then using the serum-free medium as described above, the cells were then collected and the culture media were completely removed from the cells. In this state, the cells were subjected to each of low-oxygen culture, UV irradiation, nutrient deficiency and mechanical friction, and then were normally cultured, and the concentrations of growth factors secreted in the culture medium of the adipose-derived stem cells were measured.

Specifically, the low-oxygen culture is preferably performed in conditions of about 5% carbon dioxide and 1-5% oxygen for 36-48 hours for the highest synthesis of growth factors. The UV irradiation is preferably performed by irradiating ultraviolet B having a wavelength of 280-320 nm at an energy dose of 80-120 mJ/cuf. The nutrient deficiency is preferably performed by culturing the cells in a Mg and Ca-containing Dulbecco's phosphate buffered saline for up to a maximum of 4 hours immediately before
the cells precipitated, and then normally culturing the cells in a medium obtained by adding an Ham's F-12 nutrient mixture to DMEM not containing a pH indicator such as phenol red, at a ratio of about 1:1. The mechanical friction is preferably performed by applying scratch stimulation in a lattice to the eel1 medium.

As a result, bFGF was increased 1.74 times in the case of the low-oxygen stimulation and was increased 2.71 times in the case of the UV stimulation. Also, when the low-oxygen stimulation and the UV stimulation among the physical stimulations were performed in combination, the synergic effect thereof was shown (see FIG. 9).

VEGF was increased 2.53 times in the case of the low-oxygen stimulation, 1.36 times in the case of the UV stimulation, and 1.30 times in the case of the scratch stimulation using mechanical friction. Also, when the low-oxygen stimulation, the UV stimulation and the scratch stimulation among the physical stimulations were performed in combination, the synergic effect thereof was shown (see FIG. 10).

TGF-1 was increased 1.64 times in the case of the low-oxygen stimulation, 1.75 times in the case of the UV stimulation, 2.13 times in the case of the scratch stimulation using mechanical friction, and 2.01 times in the case of the nutrient deficiency stimulation. Also, when the low-oxygen stimulation and the UV stimulation among the physical stimulations were performed in combination, the synergic effect thereof was shown (see FIG. 11).

Chemical stimulations

With respect to chemical conditions, it is preferable to expose a variety of generally widely known activating compounds directly or indirectly to individuals.

Activating compounds, which can be added to the adipose-derived stem cells according to the present invention, include cell aging-related retinoic acid, vinpocetine as a precursor thereof, picamillon serving as an assistant in this cycle, and quinic acid and quinate, which serve as protein kinase.
In addition, carbohydrate synthesis factors, such as adenine dinucleotide and acetyl-L-carnitine, which are involved in metabolism, perform an important role in cell nutrition, and such activating compounds include other additives, such as dimethylaminoethanol, which acts to stop apoptosis, L-lipoic acid and L-hydroxy acid, which are involved in cell proliferation, and coenzyme Q-10, which is involved in amino acid production. As described above, these activating compounds can be added simultaneously or individually during the culture of the adipose-derived stem cells.

The present invention focused on the stimulation effects of vitamin series among various activating compounds.

Generally, vitamin A is known to be involved in primary immune reactions, cell development processes related thereto, and a series of apoptosis reactions, and then a typical example thereof is retinoic acid, which binds to a retinoic acid receptor (RAR) to regulate and activate metabolic processes associated therewith. Among them, complexes bound to RAR-alpha, RXR-alpha and RXR-beta were reported to promote or inhibit the expression of about 128 genes, which are involved in the development of T lymphocytes as major immune cells. In particular, it was reported that bcl2 family genes known as typical anti-apoptotic proteins are definitely increased in apoptosis mechanisms (Rasooly, R. et al., J. Immunol., 175: 7916 - 7929 (2005); Spilianakis, CG. et al., Eur. J. Immunol., 35(12): 3400-4 (2005); Evans T, Exp. Hematol., 33(9): 1055-61 (2005)). This suggests that vitamin A can show the effect of inhibiting apoptosis reactions.

Vitamin B is generally known as riboflavin and performs an important role in maintaining human health. One research team in Sweden reported that treatment with this substance shows an effect on neutrophil migration, thus causing an increased primary immune response (Verdrengh, M. and Tarkowski, A., Inflamm. Res., 54(9): 390-3 (2005)). It is expected that this substance can increase initial immune responses, caused by the migration of primary immune cells.

Vitamin C has important intracellular functions of promoting the
synthesis of collagen and fibroblasts, and typical examples thereof include ascorbic acid. When it is used in combination with other cytokines TGF- and IFN-Y the effect thereof is further increased (Chung, J.H. et al., J. Dermatol. Sci., 15(3): 188-200 (1997)). Vitamin D3 has been frequently used, because it is known to influence cell development and differentiation, unlike other vitamins. Vitamin D3 mediates an important signaling system in cell growth and development processes, particularly formation processes of epidermal keratinocytes and osteogenic cells such as osteoblasts and osteoclasts. Also, it has an inhibitory effect against various cytokines, such as IL-1α, IL-6 and IL-8d, which are involved in inflammatory reactions (Alper, G. et al., Endocr. Rev., 23: 763 (2002)).

In the present invention, any one or mixture of vitamin A, vitamin C and vitamin D as chemical stimulation conditions was added to culture media in an effective amount that causes no cytotoxicity.

In order to examine whether chemical stimulation using vitamin promotes the synthesis and secretion of growth factors, in the present invention, adipose-derived stem cells were cultured in vitro using serum media and then using serum-free media as described above. Then, the cells were collected and washed with phosphate buffer saline to completely remove the media, and the cells were cultured in a medium obtained by adding an Ham's F-12 nutrient mixture to DMEM not containing a pH indicator such as phenol red, at a ratio of about 1:1, and adding thereto at least one selected from among oxidation nutrients such as L-glutamine, energy metabolites such as sodium pyruvate, and carbon sources such as sodium bicarbonate. A suitable amount at least one selected from vitamin A, vitamin B, vitamin C and vitamin D was added to the culture medium, and then the concentrations of growth factors secreted in the culture medium of the adipose-derived stem cells were measured.

Specifically, the optimal concentration of vitamin A is 2-5 µM, the optimal concentration of vitamin B2 is 50-100 µM, the optimal concentration of vitamin C is 10-100 µM, and the optimal concentration of vitamin D is 5-
10 μM. Preferably, after vitamin is added to the culture medium, the culture is performed for more than 48 hours. When a mixture of vitamins is used, the optimal concentrations of vitamins are the same as above.

As a result, it was seen that bFGF was increased 1.62 times in the case of vitamin A, 1.33 times in the case of vitamin B, 2.33 times in the case of vitamin C, and 2.80 times in the case of vitamin D.

Also, it was observed that VEGF was increased 1.59 times in the case of vitamin A, 1.68 times in the case of vitamin B, 1.68 times in the case of vitamin C, and 1.30 times in the case of vitamin D.

Moreover, it was observed that TGFβ-1 was increased 2.35 times, when UV light was used, vitamin C case, and vitamin D. Also, when vitamin A, vitamin B, vitamin C and vitamin D were added to the culture medium at the above-described optimal concentrations, bFGF was increased 3.62 times, VEGF was increased 2.03 times, and TGFβ-1 was increased 1.68.

Combination of stimulations

With respect to the case where physical stimulation was used in combination with chemical stimulation, UV light was irradiated into a culture medium, and then immediately the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of vitamins A, B, C and D, and the medium was cultured in a condition of low-oxygen stimulation for the optimal culture time. In this case, bFGF was increased 4.11 times.

Also, VEGF was increased 3.92 times, when UV light was irradiated into a culture medium followed by applying scratch stimulation, and then immediately the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of vitamins A, B, C and D, and the medium was cultured in a condition of low-oxygen stimulation for the optimal culture time.

Moreover, TGFβ-1 was increased 2.35 times, when UV light was
irradiated into a culture medium followed by applying scratch stimulation and nutrient deficiency, and then immediately the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of vitamins A, B, C and D, and the medium was cultured in a condition of low-oxygen stimulation for the optimal culture time.

Furthermore, in order to bFGF, VEGF and TGF β-1, it is most preferable to use a combination of low-oxygen stimulation, UV light stimulation and vitamins A, B, C and D. Specifically, after UV light is irradiated into a culture medium, the medium is replaced with a medium containing Ham's F-12 nutrient mixture and optimized concentrations of vitamins A, B, C and D, and the medium is cultured in a condition of low-oxygen stimulation for the optimal culture time. In this case, bFGF is increased 4.11 times, VEGF is increased 3.8 times, and TGF β-1 is increased 1.9 times.

The second aspect of the present invention provides novel uses of a culture medium obtained using the method according to the first aspect, or human growth factors purified from the culture medium.

Specifically, adipose-derived stem cell culture media or human growth factors obtained according to the present invention can be used in drugs, quasi drugs, drug supplements, and cosmetics, which are used for anti-wrinkle, wound healing, and scar removing.

The adipose-derived stem cell culture media obtained according to the present invention include all media obtained through the following cases: i) a case in which adipose-derived stem cells are cultured in serum-free media; ii) a case in which adipose-derived stem cells were stabilized in serum media, and then cultured in serum-free media; and iii) adipose-derived stem cells were activated through physical or chemical stimulation during a culture process. Also, the human growth factors according to the present invention include all human growth factors obtained by purifying the cells or culture media obtained through the above-described culture method.

It is preferable either to use culture media obtained by culturing cells in serum media and then in serum-free media through the optimized
method according to the first aspect of the present invention, or to use human growth factors purified from the said culture media.

It is more preferable either to culture media obtained by culturing cells in serum media and then in serum-free media through the optimized method according to the first aspect of the present invention and activating the cells through physical or chemical stimulation during the culture process, or to use human growth factors purified from the said culture media.

The growth factors produced from adipose-derived adult stem cells are distinguished from growth factors synthesized through existing methods, that is, growth factors synthesized from amino acids through chemical synthesis methods, and growth factors synthesized through genetic recombinant methods. In comparison with such growth factors synthesized by the genetic recombinant methods or the chemical synthesis methods, the growth factors produced according to the present invention have advantages in that they are structurally similar to the original growth factors of the human body, and thus have excellent skin compatibility and secured safety.

In terms of functions, the growth factors according to the present invention do not show isomers or three-dimensional stereospecificity and have the same form as the growth factors of the body. Thus, the growth factors according to the present invention have excellent activity compared to the growth factors produced by the recombinant or chemical synthesis methods.

Specifically, fibroblast proliferation potential was compared between the growth factors of the present invention and the growth factors produced by the recombinant or chemical synthesis methods. As a result, it was seen that the growth factors produced from adipose-derived stem cells according to the present invention had excellent activity compared to the growth factors by the existing synthesis methods (see Table 5).

Also, either the adipose-derived stem cell culture media according to the present invention or human growth factors purified from the culture media have activities of increasing intracellular collagen synthesis, promoting fibroblast proliferation, inhibiting keratinocyte proliferation caused by UV
light, mitigating skin hyperkeratization and removing wrinkles (see Tables 4, 6, 7 and 8, and FIGS. 6, 12, 13, 14, 15 and 16).

Thus, the adipose-derived stem cell culture media according to the present invention or the human growth factors purified from the culture media can be advantageously used as raw materials for drugs, quasi drugs, drug supplements and cosmetics, which are used for anti-wrinkle, wound healing, and scar removing. In particular, according to the present invention, it is possible to produce human growth factors in large amounts and to solve the prior problem in that it is almost impossible to produce industrially effective amounts of growth factors from adipose-derived adult stem cells without other operations.

Hereinafter, the present invention will be described in further detail with reference to examples. It will however be obvious to those skilled in the art that various modifications, additions and substitutions are possible, without departing from the scope and spirit of the invention as disclosed in the accompanying claims.

Example 1

(i-1) Isolation of adipose-derived stem cells

Human liposuction material collected from Leaders Clinic, Seoul, Korea was washed with the same volume of phosphate buffer saline, and only adipose tissue was separated from the liposuction material.

The extracellular matrix of the adipose tissue was enzymatically treated with 0.075% collagenase in a 5% CO₂ incubator at 37 °C for 45 minutes, and the enzymatically treated adipose tissue was centrifuged at 1200 g for 5 minutes to collect a stromal vascular fraction containing a high density of stem cells. The pellets were washed with phosphate buffer saline and passed through a 70-μm nylon cell filter to remove other tissues, and only monocyte cells and cell fragments including red blood cells were separated from the remaining material using Histopaque-1077 (SIGMA).

The separated monocytes were cultured in a non-inducing medium containing Dulbecco's Modified Eagle's Medium (DMEM), 10% fetal bovine serum
Culture of adipose-derived stem cells

The initial stage culture of the stem cells isolated from the adipose tissue was performed using DMEM containing 10% FBS. Also, 1% penicillin-streptomycin (100 IU/ml, GIBCO) was added as an antibiotic agent, amphotericin E (0.5 µg/ml, Amresco) was added as an antifungal agent, tylosin (10 µg/ml, Serva, Heidelberg) was added as a micoplasma inhibitor, and 2 mM glutamine and 1 mM sodium pyruvate were further added. The culture was conducted in a 5% CO₂ incubator at a humidity of 95% and a temperature of 37°C. During the 5% CO₂ culture, sodium bicarbonate was added at a final concentration of 0.17%.

The stem cells isolated in the above section (1-1) were suspended at a density of 10^4 cells/ml, and 10 ml of the cell suspension was transferred and cultured in a T25 flask (area: 25 cm²; volume: 50 ml) in the above-described conditions.

Cumulative doubling time was maintained until the cells being cultured in the flask reached a confluence of 80%, and the cells were subcultured at a confluence of 80%.

The flask from which the medium was washed with PBS, the cells were suspended with 0.25% Trypsin-EDTA (GIBCO), and the cell suspension was then centrifuged. Then, the number and viability of the cells were measured, and the cells were subcultured to three times. A serum-free medium used in the subculture process was based on DMEM not containing a pH indicator such as phenol red, and a Ham’s F-12 nutrient mixture (SIGMA) was added thereto at a ratio of 1:1. Also, 2 mM L-glutamine and 1 mM sodium pyruvate were added to the medium, and then sodium bicarbonate was added at a concentration of 0.1 wt%. The subculture process was repeated three times.
The cell number measurement and viability examination were performed by mixing 0.1 ml of the cell suspension with the same amount of 0.2% trypan blue (SIGMA), counting stained dyed cells and non-stained cells using a hemocytometer under a microscope, and calculating the percentages of the stained and non-stained cells relative to the total number of the cells.

Identification of stem cells

Adipose-derived stem cells express a number of adhesion and surface proteins. As these proteins, SH-2, SH-3 and SH-4 monoclonal antibodies recognize the surface epitope of human mesenchymal stem cells. As a result of the analysis of peptide base sequences and absorbance, SH-3 and SH-4 are identified to be CD73 (ecto-5'-nucleotidase), and SH-2 is identified to be CD105 (endoglin). Such cell surface markers are shared by adipose-derived stem cells (Barry, F. et al., Biochem Biophys Res Commun., 289(2): 519-24 (2001)).

The confirmation of stem cells was performed by carrying out the flow cytometry of the adipose tissue-derived stem cells, cultured in the initial culture, first subculture and second subculture, using a fluorescence activated cell sorter (Beckman Coulter). Specifically, the cells were collected with 0.25% trypsin-EDTA, washed with PBS and adjusted to a cell density of 10 cells/ml. Then, the cells were allowed to react with mesenchymal stem cell-specific markers CD73-PE and CD105-FITC (BD science) antibodies, and analyzed at 488 nm with an argon laser.

In the results of the flow cytometry of the PLA cells isolated from the adipose tissue, the initially isolated stromal vascular fraction showed a homology of 5.27% to stem cells (see FIG. 2), and after the second subculture, the cells showed a homology of 92.32% to stem cells for CD73-PE and a homology of 90.67% to stem cells for CD105-FITC (see FIG. 3).

Identification of activated growth factors of adipose-derived stem cells

In order to examine whether adipose-derived stem cells synthesize...
growth factors, adipose-derived stem cells were cultured in the same culture medium and culture conditions as in Example (1-2), and then RNA in the stem cells was analyzed through a reverse transcription-polymerase chain reaction. Specifically, the total RNA of the adipose-derived stem cells was extracted with a RNeasy Plus Mini kit (QIAGEN Corp., Valencia, California), and the RNA was amplified by PCR using a MMLV-reverse transferase (Promega Corp., U.S.A) at 37 °C for 45. Then, the MMLV-reverse transferase was inactivated at 65 °C for 45 minutes. The PCR reaction solution had a total volume of 50 µl and contained 1.5 mM MgCl₂, 0.25 mM dNTP, and 2.5 unit Taq polymerase (QIAGEN). The PCR reaction was performed in TGRADIENT (BIOMETRA) in the following conditions: a cDNA denaturation of 3 min at 94 °C, and then 30 cycles each consisting of 30 sec at 94 °C (DNA denaturation), 30 sec at 60 °C (annealing) and 30 sec at 72 °C (extension), followed by a final extension of 5 min at 72 °C. The reverse transcription was performed as described above to synthesize cDNA, which was then subjected to reverse transcription-PCR using primers of VEGF-β, bFGF and TGF β-1 (see Table 2).

<table>
<thead>
<tr>
<th>Base sequences of primers used in RT-PCR</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>VEGF 5-TACCTCCACCATGCGAGTG-3; SEQ ID NO: 1</td>
<td>5-TGATGTCTGACCCTCTCC-3; SEQ ID NO: 2</td>
</tr>
<tr>
<td>bFGF 5-GACGCGCAGTGAAGCG-3; SEQ ID NO: 3</td>
<td>5-CTCTCTCAGTGTCAGAGTG-3; SEQ ID NO: 4</td>
</tr>
<tr>
<td>TGF β-1 5-GCTGAGCCGTTTCTGATGCT-3; SEQ ID NO: 5</td>
<td>5-CGAGTGTGCTGAGTACG-3; SEQ ID NO: 6</td>
</tr>
</tbody>
</table>

As a result, a 482-bp bFGF product (1522 bp-2003 bp) corresponding to SEQ ID NO: 7, a 343-bp VEGF product (1080 bp-1422 bp) corresponding to SEQ ID NO: 8, and a 212-bp TGF β-1 product (2119 bp-2330 bp) corresponding to SEQ ID NO: 9, were identified to be growth factors in the adipose-derived stem cells (see FIG. 4).

Also, in order to examine whether growth factors are present in a culture medium, an enzyme-linked immunosorbent assay was performed.
Specifically, 1 ml of the culture medium obtained by culturing adipose-derived stem cells to passage 3 in the conditions of Example 1-2 was centrifuged at 1200 g for 5 minutes, and then filtered through a 0.22 mm filter to remove cell residue. The filtrate was serially diluted to determine optimal conditions in which a non-specific reaction did not occur. Then, the concentration of each of growth factors VEGF, bFGF and TGF β-1 secreted in the culture medium was measured using a sandwich ELISA kit (Quantikine Human FGF basic Immunoassay, R&D systems).

To examine the concentrations of optimally diluted growth factors, the culture medium was serially diluted at the same concentration ratio with a calibration dilution in the kit, the concentration at the absorbance included in the range of the standard curve was determined, and the values included in the range of the standard curve were plotted as data. 100 ml of each of growth factor-containing culture media and standard solutions of growth factors was placed in each well coated with an antibody (Quantikine Human FGF basic Immunoassay, R&D systems) to each of the growth factors and was cultured at room temperature for 2 hours. After completion of the antigen-antibody binding reaction, each well was washed four times with wash buffer, and 200 ml of a solution (Quantikine Human FGF basic Immunoassay, R&D systems), to which a secondary antibody to each of the growth factors was bound, was placed in each well, followed by culture at room temperature for 2 hours. After completion of the second antibody bonding reaction, each well was washed four times with wash buffer, 200 ml of a color-developing reagent (tetramethylbenzidine, R&D systems) was added to each well, and then the absorbance of the medium was measured at 450 nm with an ELISA reader.

As a result, bFGF showed an increase of 54.96 pg/ml after 12 hours and an increase of 76.393 pg/ml after 24 hours, and VEGF showed an increase of 87.021 pg/ml after 12 hours and an increase of 163.52 pg/ml after 24 hours (see FIG. 5).

Example 3

Activity of adipose-derived stem cells on collagen production of
One of the causes of formation of skin wrinkles is the lack of collagen. Collagen is a major protein of skin dermis and serves to maintain the skin structure and firmness. It is known that the production of collagen is decreased as age increases, and the degradation thereof is also increased to induce the collapse of the dermal layer, thus producing skin wrinkles. Thus, the effect of a substance for anti-wrinkles can be proven by testing the production and degradation of collagen.

(3-1) Coculture of adipose-derived stem cells and fibroblasts

The effect of adipose-derived stem cells on the collagen production of fibroblasts was estimated through the coculture of adipose-derived stem cells and fibroblasts. In the test, an increase in the production of collagen in fibroblasts, when adipose-derived stem cells were cocultured with fibroblasts in a transwell insert (Costar, Corning), was compared with the production of collagen in the single culture of fibroblasts.

a) Coculture

Fibroblasts (primary cell line) were obtained by finely cutting a portion of human skin tissue (Department of Laboratory Medicine, Hangang Sungsim Hospital; 9 years old; circumcision fragment) suspended in PBS, stirring the cut tissue with 50-100 ml of trypsin for 20 minutes, centrifuging the stirred tissue, and then filtering the tissue through a 7-mm nylon filter.

The filter cell suspension was seeded on the bottom of a culture dish and was added to a DMEM medium containing penicillin (100 IU/mL), streptomycin (100 g/mL) and 10% FBS. Then, the cells were cultured in an incubator containing 5% carbon dioxide at 37 °C. The fibroblasts, which reached a confluence of 80%, were dispensed into a 6-well plate at a density of 5 x 10⁴/well and cultured for 24 hours in the same culture conditions as in the subculture process.

The adipose-derived stem cells isolated in Example (1-1) were cultured in a transwell insert for 24 hours in the same conditions as the case of the
fibroblasts, and then the transwell insert was inserted into the 6-well plate in which the fibroblasts were being cultured. The stem cells and the fibroblasts were cocultured in the 6-well plate. At this time, the medium below the transwell insert was discarded, the 6-well plate was washed with PBS, a fresh medium was added thereto, and the fibroblasts and the adipose-derived stem cells were cocultured.

1 ml of the cocultured medium was taken at 24 hours and 72 hours after the culture, and the amount of collagen in the culture medium was measured.

In a control group test, fibroblasts, which reached confluence, were dispensed into a 6-well plate at a density of 5 x 10⁴ cells/well, and then cultured for 24 hours. A transwell insert having no adipose-derived stem cell seeded therein was inserted into the 6-well plate.

b) Analysis of activity for collagen production

1) Measurement of amount of collagen by enzyme-linked immunosorbent assay (ELISA)

The measurement of the amount of collagen was performed using a Procollagen type I peptide EIA kit (TAKARA BIOMEDICAL Co.). 100 μl of an antibody-PoD conjugate solution was placed in each well. Then, 20 μl of fibroblasts, which were optimally diluted with a calibration dilution so as to be included in a collagen standard curve, and 20 μl of fibroblasts cocultured with adipose-derived stem cells, were placed in each well and left to stand at 37°C for 3 hours. Then, each well was washed four times with wash buffer, and 100 μl of a color-developing reagent was added to each well, followed by culture at room temperature for 15 minutes. Then, the absorbance of the culture medium was measured at 450 nm with an ELISA reader.

2) Measurement of amount of collagen in cells by semi-quantity PCR

The total RNA of fibroblasts cocultured with stem cells was extracted using a RNeasy plus mini kit (QIAGEN), and 3 μg of the extracted RNA was amplified by PCR in a reaction solution containing 200 unit MMLV-reverse transcriptase (Promega), 50 mM Tris-HCl (pH 8.3), 75 mM KCl, 3 mM MgCl₂, 10 mM
DTT, 10 mM dNTP, 25 unit RNase inhibitor and 20 pmole Oligo-dT. The PCR reaction was performed in T-GRADIENT (BIOMETRA) at 37 °C for 45 minutes, and then the MMLV-reverse transcriptase was inactivated at 65 °C for 15 minutes. Specific base sequences for the detection of collagen I type (GenBank No. NM-000089) were constructed based on base sequences recorded in the NCBI Genbank and are shown in Table 3 below.

<table>
<thead>
<tr>
<th>Collagen type I</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>5'-CCCTCAAGTTTCAAGGAC-3': SEQ</td>
<td>ID NO: 10</td>
<td>5'-ACCAGGTCCACTTCACAC-3': SEQ</td>
</tr>
</tbody>
</table>

As a result, as shown in FIG. 6, the amount of intracellular collagen synthesis was increased 2.17 times in 1-day culture and 1.27 times in 3-day culture, compared to the control group, and the amount of total collagen synthesis was increased 1.44 times in 1-day culture and 1.14 times in 3-day culture.

Collagen synthesis of fibroblasts by adipose-derived stem cell culture medium

In order to examine the effect of a culture medium (conditioned medium) of adipose-derived stem cells on the collagen synthesis of fibroblasts, western blot analysis (specific protein detection) was performed.

To collect a culture medium of adipose-derived stem cells, 5 x 10⁵ adipose-derived stem cells were seeded into a T75 flask after subculture. Herein, serum-free DMEM was used as medium. The cells were cultured in a 5% CO₂ incubator at 37 °C for 3 days, and then the medium was harvested and filtered through a 0.22-um syringe filter. The filtrate was used as a serum-free conditioned medium in the following test.

Fibroblasts were subcultured, and then seeded into a 0.1% FBS-containing DMEM in a 6-well plate at a density of 5 x 10⁴ cells/well. After the cells were cultured for 24 hours, the medium was replaced with the above-
described serum-free conditioned medium. As control group, serum-free DMEM
was used. After 12 hours, serum was adjusted to 2%, and after 30 hours of
culture, the culture medium was harvested and subjected to western blot
analysis.

The electrophoresis of the culture medium was performed according to
the SDS-PAGE method using an electrophoresis kit (Bio-rad). In the
electrophoresis, 8% polyacrylamide gel was used and transferred to a PVDF
membrane (Bio-rad). Then, the blot was blocked with a 5% nonfat dry milk-
containing TBST (50 mM Tris, pH8.0, 138 mM NaCl, 2.7 mM KCl, and 0.1% (w/v)
Tween 20). The medium was allowed with react with collagen type I
(Santacruz) as a primary antibody overnight and allowed to react with
peroxidase-Rabbit anti-goat IgG (Zymed) as a secondary antibody for 30
minutes. Finally, the medium was allowed to react with an antibody detection
reagent (ECL, Milipore) for 1 minute, and the test results were observed.

As a result, as shown in FIG. 16, the adipose-derived stem cell
culture medium treated with fibroblasts showed an increase of more than 2
times in the amount of collagen compared to the medium untreated with
fibroblasts (quantified with a gene tool software; 2.16-fold increased).
This demonstrates that the culture medium of adipose-derived stem cells
increases the collagen synthesis of fibroblasts, and thus suggests that the
culture medium of adipose-derived stem cells can be used as an anti-wrinkle
substance for preventing skin aging.

Example 4

Stimulation of fibroblast proliferation by adipose-derived stem cells

Stem cells isolated from adipose tissue were cultured to passage 3 as
described in Example (1-2), and then 10 cells of the stem cells were seeded
into a T175 flask (area: 175 cm²; and volume: 500 ml) and cultured for 3 days.
The stem cell culture medium was collected, and added to concentrations of
10%, 25%, 50% and 100% to a 6-well plate, in which passage-4 fibroblast cells
from the skin tissue of 9 years old boy (Department of Laboratory Medicine,
Hangang Sungsim Hospital: 9 years old: circumcision fragment) were dispensed
at a density of 25,000 cells/well.

After 3 days, the proliferation of the fibroblasts was measured with a cell viability measurement kit (cell counting kit-8, Dojindo Molecular Technologies, Inc.), and then the absorbance of the cells was measured at 450 nm with an ELISA reader.

As a result, as shown in Table 4, it was demonstrated that the growth factors secreted from the adipose-derived stem cells stimulated the proliferation of fibroblasts. Also, it was observed that the proliferation of fibroblasts was stimulated as the concentration of growth factors secreted from adipose-derived stem cells was increased (see FIG. 7).

<table>
<thead>
<tr>
<th>Classification</th>
<th>Cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal media (control group)</td>
<td>35000</td>
</tr>
<tr>
<td>Treated with 10% ADSC 3-passage 3-day cultured</td>
<td>45000</td>
</tr>
<tr>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>Treated with 25% ADSC -3-passage 3-day cultured</td>
<td>93000</td>
</tr>
<tr>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>Treated with 50% ADSC 3-passage 3-day cultured</td>
<td>180000</td>
</tr>
<tr>
<td>medium</td>
<td></td>
</tr>
<tr>
<td>Treated with 100% ADSC 3-passage 3-day cultured</td>
<td>240000</td>
</tr>
<tr>
<td>cultured medium</td>
<td></td>
</tr>
</tbody>
</table>

Example 5

Comparison of growth factors secreted from adipose-derived stem cells with recombinant growth factors expressed in E.coli

Stem cells isolated from adipose tissue were cultured to passage-3 as described in Example (1-2), and then 10 cells of the stem cells were seeded into a T175 flask (area: 175cm²; volume: 500 ml) and cultured for 3 days. The culture medium was collected, and added at a concentration of 100% to a 6-well plate in which passage-4 fibroblast cells from the skin tissue of a 9 years old man were dispensed at a density of 5000 cells/well. As a control
group, a medium containing the same concentrations of recombinant growth factors VEGF and bFGF (Santa Cruz.) expressed in E. coli was used. Cell proliferation potential was compared between the adipose-derived stem cell culture medium and the recombinant growth factor-containing culture medium using a cell viability measurement kit.

As a result, when the growth factors obtained by the overexpression and isolation of growth factor genes in E. coli using the genetic recombinant method were compared with the growth factors synthesized from the adult stem cells, it was demonstrated that the growth factors synthesized from the adipose-derived adult stem cells had excellent effects compared to the growth factors obtained by the existing synthesis method. The analysis results are shown in Table 5 below.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal media + rFGF (50ng)</td>
<td>8215.3</td>
</tr>
<tr>
<td>Normal media + rVEGF (500ng)</td>
<td>4408.9</td>
</tr>
<tr>
<td>Normal media + rFGF+rVEGF</td>
<td>6727.1</td>
</tr>
<tr>
<td>Treated with 100% ADSC 3-passage 3-day cultured medium</td>
<td>17215</td>
</tr>
</tbody>
</table>

Example 6

**Culture method of adipose-derived stem cells having increased growth factor secretion**

(6-1) Physical stimulation

Adipose tissue-derived stem cells were cultured to passage 3 as described in Example (1-2), and when the cultured cells reached a confluence of 80%, the cells were collected with trypsin/EDTA. Then, the cells were accurately dispensed into a 6-well plate at a density of 25,000 cells/well.

24 hours after the dispense, when the cells were all attached, the culture medium was completely removed, and the cells were a multigas incubator containing 5% carbon dioxide and 1% oxygen in physical conditions, and was irradiated with an energy of 90 mJ/cuf at a wavelength of 280-320 nm
(model BEX-800; Ultra-Lum, Inc.) corresponding to UV B. In a nutrient
deficiency reaction, the cells were cultured in Dulbecco's phosphate buffered
saline containing Mg and Ca for a maximum of 4 hours immediately until the
cells were precipitated, and then, the medium was replaced with the serum-
free medium of Example (1-2). In scratch stimulation using mechanical
friction, the cell medium, which was being cultured attached to the plate,
was scratched with a blade in the form of a lattice having a size of 1 mm x 1
mm.

(6-2) Chemical conditions

To the serum-free medium of Example (1-2), prepared by adding a Ham's
F-12 nutrient mixture to DMEM not containing a pH indicator such as phenol
red, at a ratio of 1:1, and adding thereto 2 mM L-glutamine, 1 mM sodium
pyruvate and 0.17% sodium bicarbonate, vitamins A, B, C and D were added at
effective levels that did not cause cytotoxicity. Cells were cultured in
this medium.

The cells were cultured in the above medium in a 5% carbon dioxide
incubator at 37 °C for more than 48 hours, and then 200 µl of the culture
medium supernatant was centrifuged at 3,000 rpm and filtered through 0.22-µm
filter paper. Then, the concentrations of bFGF, VEGF and TGF β-1 in the
filtrate were measured by the enzyme-linked immunosorbent assay (ELISA) of
Example 2.

In the analysis results, the optimal concentration of vitamin A was 2-
10 µM, the optimal concentration of vitamin B2 was 50-100 µM, the optimal
concentration of vitamin C was 10-100 µM, and the optimal concentration of
vitamin D was 5-10 µM. It was seen that, at such concentrations, no
cytotoxicity occurred as a result of an MTT assay, and the synthesis of each
of growth factors targeted in the present invention was increased.

(6-3) Culture conditions of control group

As a control group, a normal medium, which was not subjected to
physical stimulation and chemical stimulation, was used. The increase in
growth factors was compared between the control group and the culture medium
subjected to physical stimulation and chemical stimulation.

In a culture process, cells were added to a serum-free medium prepared by adding a Ham's F-12 nutrient mixture to DMEM not containing a pH indicator such as phenol red, at a ratio of 1:1, and adding thereto 2 mM L-glutamine, 1 mM sodium pyruvate and 0.17% sodium bicarbonate. Then, the cells were cultured in a 5% carbon dioxide incubator at 37 °C for more than 48 hours. Then, 200 µl of the culture medium supernatant was centrifuged at 3,000 rpm, and then filtered through 0.22-µm filter paper. Then, the concentrations of bFGF, VEGF and TGF β-1 were measured by the enzyme-linked immunosorbent assay (ELISA) of Example 2.

(6-4) Combination of stimulations

The case where physical stimulation is used in combination with chemical stimulation will now be described. For bFGF, UV light was irradiated into a culture medium, and then immediately the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of 2 µM vitamin A, 50 µM vitamin B, 10 µM vitamin C and 10 µM vitamin D. The medium was cultured low-oxygen stimulation conditions of 1% oxygen and 5% carbon dioxide for 48 hours.

For VEGF, a medium was irradiated with UV light and then subjected to scratch stimulation. Immediately after this, the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of 2 µM vitamin A, 50 µM vitamin B, 10 µM vitamin C and 10 µM vitamin D. The medium was cultured low-oxygen stimulation conditions of 1% oxygen and 5% carbon dioxide for 48 hours.

For TGF β-1, a medium was irradiated with UV light and then subjected to scratch stimulation and nutrient deficiency stimulation. Then, the medium was replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of 2 µM vitamin A, 50 µM vitamin B, 10 µM vitamin C and 10 µM vitamin D. The medium was cultured low-oxygen stimulation conditions of 1% oxygen and 5% carbon dioxide for 48 hours.

To obtain the highest total amount of bFGF, VEGF and TGF β-1, a medium
was irradiated with UV light, and then replaced with a DMEM medium containing a Ham's F-12 nutrient mixture and optimized concentrations of 2 \( \mu \)M vitamin A, 50 \( \mu \)M vitamin B, 10 \( \mu \)M vitamin C and 10 \( \mu \)M vitamin D. The medium was cultured low-oxygen stimulation conditions of 1\% oxygen and 5\% carbon dioxide for 48 hours.

(6-5) Results

As shown in FIG. 9, in comparison with the control group, bFGF was increased 1.74 times in the case of low-oxygen stimulation, and 2.71 times in the case of UV light stimulation. In the case of chemical stimulation, bFGF was increased 1.62 times for vitamin A, 1.33 times for vitamin B, 2.33 times for vitamin C and 2.80 times for vitamin D.

When low-oxygen stimulation and UV light stimulation, having excellent effects among the above stimulations, were performed in combination, the synergic effect thereof was shown. In the case of chemical stimulation, bFGF in a medium containing optimized concentrations of vitamins A, B, C and D was increased 3.62 times. In case where physical stimulation was applied in combination with chemical stimulation, bFGF was increased 4.11 times, when low-oxygen stimulation was optimized, UV light was irradiated and the concentrations of vitamins A, B, C and D were optimized.

As shown in FIG. 10, in comparison with the control group, vascular endothelial growth factor (VEGF) was increased 2.53 times in the case of low-oxygen stimulation, 1.36 times in the case of UV light stimulation, and 1.33 times in the case of scratch stimulation caused by mechanical friction. In the case of chemical stimulation, VEGF was increased 1.59 times for vitamin A, 1.56 times for vitamin B, 1.68 times for vitamin C, and 1.30 times for vitamin D.

When low-oxygen stimulation and UV light stimulation, having excellent effects, among the above stimulations, were performed in combination, the synergic effect thereof was shown. In the case of chemical stimulation, VEGF in a medium containing optimized concentrations of vitamins A, B, C and D was increased 2.03 times.
In case where physical stimulation was applied in combination with chemical stimulation, VEGE could be increased 3.92 times, when low-oxygen stimulation was optimized, UV light stimulation and scratch stimulation were used and the concentrations of vitamins A, B, C and D were optimized.

As shown in FIG. 11, in comparison with the control group, transforming growth factor beta-1 (TGFβ-1) was increased 1.64 times in the case of low-oxygen stimulation, 1.75 times in the case of UV light stimulation, 2.13 times in the case of scratch stimulation caused by mechanical friction, and 2.01 times in the case of nutrient deficiency stimulation. In the case of chemical stimulation, TGFβ-1 was increased 1.20 times for vitamin A, 1.56 times for vitamin B, 1.20 times for vitamin C, and 1.16 times for vitamin D.

When low-oxygen stimulation and UV light stimulation, having excellent effects, among the above stimulations, were performed in combination, the synergic effect thereof was shown. In the case of chemical stimulation, TGFβ-I in a medium containing optimized concentrations of vitamins A, B, C and D was increased 1.68 times.

In case where physical stimulation was applied in combination with chemical stimulation, TGFβ-1 could be increased 2.35 times, when low-oxygen stimulation was optimized, UV light stimulation, scratch stimulation and low-oxygen stimulation were used and the concentrations of vitamins A, B, C and D were optimized.

Also, in order to the highest total amount of bFGF, VEGF and TGFβ-1, it is most preferable to subject a medium to low-oxygen stimulation and UV light stimulation and add vitamins A, B, C and D to the medium. Specifically, a medium is irradiated with UV light, and then replaced with a medium containing a Ham's F-12 nutrient mixture and optimized concentrations of vitamins A, B, C and D, and the medium is cultured in a condition of low-oxygen stimulation for the optimized culture time. In this case, bFGF is increased 4.11 times compared to the control group, VEGF is increased 3.8 times, and TGFβ-1 is increased 1.9 times.
Example 7

Keratinocytes defensive effect of adipose-derived stem cells against UV light

The keratinocytes defensive effect of growth factors secreted from adipose-derived stem cells against UV light was tested in the following manner.

Keratinocytes were dispensed into a 6-well plate at a suitable cell concentration and added to a KGM (keratinocyte growth medium; Clonetics.) in a 5% carbon dioxide incubator at 37°C. To the medium, 2.5 cc of 100% ADSC 3-passage 3-day cultured medium was added to adjust the KGM to 5 cc. Also, 2μM retinol was added to 5 cc of KGM. Also, as a negative control group, 5 cc of KGM was administered alone to keratinocytes. Each of the test samples was stabilized for 4 hours, such that the raw materials were sufficiently dispersed. After that, each of the test samples was irradiated with UV light using a 40W double lamp for 8 minutes in an aseptic laboratory. After 24 hours, the UV defense function of the test sample was measured by determining the viability of the cells in comparison with the negative control group, which was untreated with the culture medium or retinol and irradiated with UV light for 8 minutes.

As a result, as shown in Table 6 below, the cell group, which was cultured as described in Example (1-2) and contained VEGF, bFGF and TGFβ-1, showed a defensive effect of 67%, the cell group administered with retinol showed a defensive effect of 54%, and the negative control group showed a defensive effect of 55%. This indicates that retinol has a low defensive effect against UV lights, whereas the growth factors secreted from adipose-derived stem cells has an increased defensive effect against UV light.

Table 6
Measurement results of anti-aging effects
Effects of adipose-derived stem cell growth factors on skin photoaging caused by UV light irradiation in nude mice.

In order to assess the activity of a culture medium, which was cultured as described in Example (1-2) and contained VEGF, bFGF and TGFβ-1, the following test was performed using thirty 15-20-week-old nude mice.

Irradiating the back of nude mice with UV light at a dose of 2 mJ/cm² using an UV simulator was performed two times a week for 4 weeks, thus inducing the abnormal hyperkeratinization of the skin. Then, growth factors synthesized by a chemical synthesis method, growth factors produced by a genetic recombination method, and growth factors synthesized from adipose-derived adult stem cells, were applied on one side of the back of the nude mice in an amount of 1 cc two times a day for 2 weeks, while the other side was untreated for comparison. After 2 weeks, the anti-aging effects of the growth factors were observed visually and assessed (Jin Ho Chung et al. Archives of Dermatology, 137-8 (2001)). The measurement results are shown in Table 7 below.

<table>
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<th>Tested materials</th>
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<tr>
<td>100% ADSC 3-passage 3-day cultured medium</td>
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<tr>
<td>Retinol</td>
<td>54%</td>
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<tr>
<td>Negative control group</td>
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When each of the growth factors was applied to the nude mice whose aging was artificially induced, the growth factors synthesized from adipose-
derived adult stem cells showed the effect of reducing abnormal skin hyperkeratinization in all of the 30 animals, and this effect was excellent compared to that of the growth factors synthesized by the chemical synthesis method or the genetic recombination method.

Example 9

Anti-wrinkle effect of growth factors secreted from autologous adipose-derived stem cells in human beings

In order to verify the wrinkle-reducing effect of growth factors synthesized from adipose-derived adult stem cells on 30-50-years old women, 20 persons per test group were selected and the following test was conducted.

1 cc of a adipose-derived stem cell culture medium purified to have a VEGF content of 30 ng, a bFGF content of 30 ng and a TGF-1 content of 70 ng was applied on the brow or around one eye of each of the women subjects two times a day for 8 weeks, and a negative control group (1 cc of physiological saline as a basic solution) was applied around the other eye for comparison. After 4 weeks and 8 weeks, the wrinkle-reducing effect of the test sample was visually observed and the measurement results are summarized in Table 8 below.

<table>
<thead>
<tr>
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<tr>
<td>Growth factor-containing culture medium</td>
<td>8</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Negative control group</td>
<td>2</td>
<td>4</td>
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As can be seen from the results of Table 8 and FIGS. 12 to 15, the application of the growth factors synthesized from adult stem cells showed an excellent wrinkle-reducing effect.

[Industrial Applicability]
The human growth factors produced according to the method of the present invention are substances having secured stability and physiological activity, thus can be used to develop drugs, quasi drugs, cosmetics and the like for anti-wrinkles, wound healing, and scar removing.

Also, the adipose-derived stem cell culture medium according to the present invention contains various growth factors in large amounts, and thus can be used by themselves in drugs, quasi drugs, cosmetics and the like for anti-wrinkles, wound healing, and scar removing.

[Sequence List Text]

SEQ ID NOS: 1-6 and 10-11 according to the present invention are amplification primer pairs for the detection of specific proteins, and SEQ ID NOS: 7-9 are amplification products encoding for human growth factors.
[CLAIMS]

[Claim 1]

A method for producing large amounts of human growth factors from adipose-derived stem cells, the method comprising the steps of:

(i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells;

(ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium;

(iii) applying, to the adipose-derived stem cells, at least one physical stimulation selected from among low-oxygen culture, UV irradiation, nutrient deficiency and mechanical friction; and

(iv) optionally adding, to the culture media, at one or more vitamins selected from among vitamin A, vitamin B, vitamin C and vitamin D,

wherein the step (iii) and the step (iv) are performed in conditions in which the highest production of the human growth factors occurs.

[Claim 2]

The method of Claim 1, wherein the human growth factors are a basic fibroblast growth factor (bFGF), a vascular endothelial growth factor (VEGF) and/or a transforming growth factor-beta (TGF-β).

[Claim 3]

A method for the highest culture of a basic fibroblast growth factor (bFGF) from adipose-derived stem cells, the method comprising the steps of:

(i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells;

(ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium; and

(iii) irradiating the stem cells with UV light, replacing the medium with a medium containing optimized concentrations of vitamins A, B, C and D, and then culturing the stem cells in the replaced medium in a condition of low-oxygen stimulation.

[Claim 4]
A method for the highest culture of a vascular endothelial growth factor (VEGF) from adipose-derived stem cells, the method comprising the steps of:

(i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells;

(ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium; and

(iii) irradiating the stem cells with UV light, applying scratch stimulation to the medium, replacing the medium with a medium containing optimized concentrations of vitamins A, B, C and D, and then culturing the stem cells in the replaced medium in a condition of low-oxygen stimulation.

[Claim 5]

A method for the highest culture of a transforming growth factor-beta (TGF-β) from adipose-derived stem cells, the method comprising the steps of:

(i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells;

(ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium; and

(iii) irradiating the stem cells with UV light, applying scratch stimulation, applying nutrient deficiency stimulation, replacing the medium with a medium containing optimized concentrations of vitamins A, B, C and D, and culturing the stem cells in the replaced medium in a condition of low-oxygen stimulation.

[Claim 6]

A method for producing large amounts of human growth factors from adipose-derived stem cells, the method comprising the steps of:

(i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells;

(ii) optionally culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium; and

(iii) irradiating the stem cells with UV light, replacing the medium
with a medium containing optimized concentrations of vitamins A, B, C and D, and then culturing the stem cells in the replaced medium in a condition of low-oxygen stimulation.

[Claim 7]
<262> The method of any one of Claims 1 to 6, wherein the serum medium contains 0.1-20% of serum.

[Claim 8]
<263> The method of any one of Claims 1 to 6, wherein the serum-free medium contains a Ham's F-12 nutrient mixture.

[Claim 9]
<264> The method of any one of Claims 1 to 6, wherein the low-oxygen culture is performed in conditions of about 5% carbon and 1-5% oxygen.

[Claim 10]
<265> The method of any one of Claims 1 to 6, wherein the UV irradiation is performed by irradiating UV light having a wavelength of 280-320 nm at an energy dose of 80-120 mJ.

[Claim 11]
<266> The method of Claim 1 or 5, wherein the nutrient deficiency is performed in a buffer solution containing Mg and Ca.

[Claim 12]
<267> The method of Claim 1, wherein the mechanical friction is performed by applying scratch stimulation to the cell medium.

[Claim 13]
<268> The method of any one of Claims 1 to 6, wherein the vitamin A is added in an amount of 2-5 µM.

[Claim 14]
<269> The method of any one of Claims 1 to 6, wherein the vitamin B is added in an amount of 50-100 µM.

[Claim 15]
<270> The method of any one of Claims 1 to 6, wherein the vitamin C is added in an amount of 10-100 µM.
[Claim 16]

<271> The method of any one of Claims 1 to 6, wherein the vitamin D is added in an amount of 5-10 μM.

[Claim 17]

<272> A human basic fibroblast growth factor (bFGF) obtained according to Claim 1.

[Claim 18]

<273> A human vascular endothelial growth factor (VEGF) obtained according to Claim 1.

[Claim 19]

<274> A human transforming growth factor-beta (TGF-β) obtained according to Claim 1.

[Claim 20]

<275> A functional cosmetic composition containing an adipose-derived adult stem cell culture medium obtained according to Claim 1.

[Claim 21]

<276> A functional cosmetic composition containing human growth factors obtained according to Claim 1.

[Claim 22]

<277> The cosmetic composition of Claim 20 or 21, wherein the functionality is an anti-wrinkle or anti-aging activity.

[Claim 23]

<278> A functional cosmetic composition for anti-wrinkle and/or anti-aging, which comprises a culture medium of stem cells, in which the stem cells are obtained by a method comprising the steps of:

<279> (i) isolating adipose-derived adult stem cells extracted from mammalian adipose cells; and

<280> (ii) culturing the stem cells in a serum medium, and then subculturing the stem cells in a serum-free medium.
[Figure 5]

12hr 24hr
average value of three repeated tests

12hr 24hr
average value of three repeated tests

[Figure 6]

intracellular collagen synthesis  total collagen synthesis

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average value of three repeated tests
**Figure 9**

- Control group
- Low oxygen stimulation
- UV light stimulation
- Vitamin A
- Vitamin B
- Vitamin C
- Vitamin D
- Mixture of vitamins A, B, C, and D
- Optimized physical and chemical stimulation

*average value of three repeated tests*
average value of three repeated tests
[Figure 11]

average value of three repeated tests
[Figure 12]

After 30 days of application

After 60 days of application
[Figure 15]

after application

[Figure 16]

control fibroblast adipose-derived stem cell culture medium
culture medium 50% 100%
← collagen type I
1 With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, the international search was carried out on the basis of

   a type of material
      [X] a sequence listing
      [ ] table(s) related to the sequence listing

   b format of material
      [X] on paper
      [ ] in electronic form

   c time of filing/furnishing
      [X] contained in the international application as filed
      [X] filed together with the international application in electronic form
      [ ] furnished subsequently to this Authority for the purposes of search

2 In addition, in the case that more than one version or copy of a sequence listing and/or table relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished

3 Additional comments
A. CLASSIFICATION OF SUBJECT MATTER

C12N 5/08(2006.01)I, C07K 4/12(2006.01)I, C12N 5/02(2006.01)I, A61K 35/12(2006.01)I, A61K 38/00(2006.01)I

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 8 C12N 5/08, C07K 4/12, C12N 5/02, A61K 35/12, A61K 38/00

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Korean Patents and applications for inventions since 1975

Korean Utility models and applications for Utility models since 1975

Japanese Utility models and application for Utility models since 1975

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

eKIPASS (KIPO internal), NCBI PubMed, Delphi (growth factor, adipose tissue, isolation, stem cell, serum free, and similar terms)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>McPHERRON, A., and LEE, S. J. &quot;GDF-3 and GDF-9 Two New Members of the Transforming Growth Factor-β Superfamily Containing a Novel Pattern of Cysteines&quot; In J Biol Chem (February 1993), Vol 268(5) 3444-3449. See abstract, page 3444 right column lines 11-14, and figures</td>
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<td>WO 2003/059272 A2 (HENRY FORD HEALTH SYSTEM) 24 July 2003 See the whole document</td>
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Further documents are listed in the continuation of Box C

See patent family annex

* Special categories of cited documents
   "A" document defining the general state of the art which is not considered to be of particular relevance
   "E" earlier application or patent but published on or after the international filing date
   "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of citation or other special reason (as specified)
   "O" document referring to an oral disclosure, use, exhibition or other means
   "P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance, the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance, the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search
11 JANUARY 2007 (11.01.2007)

Date of mailing of the international search report
11 JANUARY 2007 (11.01.2007)

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Facsimile No 82-42-472-7140

Authorized officer
AHN, Kyu Jeong
Telephone No 82-42-481-8158

Form PCT/ISA/210 (second sheet) (April 2005)
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