Efficacy of the Controlled Release of Concentrated Platelet Lysate from a Collagen/Gelatin Scaffold for Dermis-Like Tissue Regeneration

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Introduction: A collagen/gelatin scaffold (CGS) can provide a sustained release of basic fibroblast growth factor (bFGF), which promotes wound healing. However, bFGF is approved for clinical use in Japan and China only. One potential alternative to bFGF is platelet lysate (PL), a safe and easily attainable source of a wide range of growth factors necessary for tissue repair. In the present study, we investigated the use of PL with CGS to repair wounds and identified the optimal concentration of PL for wound healing.

Materials and Methods: We generated PL from concentrated platelets harvested from individual healthy donors. We measured growth factors in PL. Transforming growth factor (TGF)-β1, platelet-derived growth factor (PDGF)-BB, vascular endothelial growth factor (VEGF), and bFGF were selected because they were the major growth factors contained in platelets and showed the greatest ability to enhance the maturation of newly formed blood vessels. Pieces of CGS impregnated with PL solution (×1, ×2, ×3, or ×4 concentrated) or normal saline solution (NSS) were implanted into full-thickness skin defects on the backs of mice. We evaluated the wound area, neoepithelium length, and total area of newly formed capillaries in the implanted CGS.

Results: Our release experiments revealed that PDGF-BB and TGF-β1 were released from CGS incubated with collagenase in a sustained manner. CGS impregnated with concentrated PL was more effective than CGS impregnated with NSS in all evaluated items. The ×2 concentrated PL accelerated wound healing and enhanced cell proliferation and vessel growth in granulation tissue.

Conclusion: Our findings indicate that CGS can bind to TGF-β1 and PDGF-BB and release these growth factors in a sustained manner. ×2 concentrated PL-impregnated CGS accelerates the formation of dermis-like tissue.
and the variability of growth factor profiles among different donors. Because PL and the growth factors dissolve into blood stream within days after their injection, it is difficult to maintain their presence in the tissue long enough to promote effective tissue regeneration.

The purpose of the present study was to investigate the efficacy of PL-impregnated CGS in promoting dermis-like tissue formation. To conduct this investigation, we used a freeze-dry procedure to generate concentrated PL that would offset the variability in growth factor profiles as well as a controlled-release system utilizing biodegradable CGS that facilitates concentrated PL’s sustained release of major growth factors to maintain their effect.

Materials and Methods

Animals

Eight-week-old C57BL/6Jcl mice (CLEA Japan, Inc., Osaka Japan) were maintained in the Institute of Laboratory Animals, Graduate School of Medicine, Kyoto University, in compliance with protocols established by the university’s Animal Research Committee.

Preparation of CGS

We used gelatin isolated from pig dermis with an isoelectric point (IEP) of 5.0 and molecular weight of 99,000 (Nipp, Osaka, Japan) and atelocollagen isolated from pig tendons with an IEP of 8.5 and molecular weight of 300,000 (Nitta Gelatin, Osaka, Japan) to create the CGS. We prepared CGS as described previously. Briefly, CGS with gelatin concentrations of a mass percentage of 10% of the total solute was prepared by mixing a mass percentage of 3% gelatin solution into a mass percentage of 0.3% collagen solution. The foaming solution was poured into a mold, frozen rapidly, and freeze-dried. The resulting collagen-gelatin sheet was stabilized by immersing it in a crosslinking agent (0.2% glutaraldehyde) for 24 h at 4°C and then freeze-drying it. We used silicone solution (Shin-Etsu Chemical, Tokyo, Japan) to make 200-μm-thick silicone sheets that would be used to protect the implanted CGS.

Generation of human supplements

PLs were generated from concentrated platelets that had been harvested from 6 healthy humans. The concentrated platelets were frozen at −80°C and thawed at 37°C twice to lyse the platelets for release of growth factors. To remove the membrane fragments, we centrifuged the lysates at 3000 g for 30 min and filtered the supernatant through a 0.2-μm Steradisc (Kurabo, Osaka, Japan). To prevent gelatinization, we added 2 U/mL heparin to the media, which was then stored at −80°C until use.

Concentrated PLs were prepared using a freeze-dryer (VD-250R; TAITEC Co., Saitama, Japan) according to the manufacturer’s instructions and stored at 4°C until use. To create a ×4 concentration of PLs, we reconstituted freeze-dried PLs with an amount of normal saline solution (NSS; Otsuka Pharmaceutical, Tokyo, Japan) that was one-fourth the original volume of the PL media. To create a ×3 concentration of PLs, we added NSS to the ×4 concentration to produce a 4/3-fold dilution of the ×4 concentration. To create a ×2 concentration of PLs, we added NSS to the ×4 concentration to produce a two-fold dilution of ×4 concentration.

Measurement of growth factors in PL

The platelets contained in α-granules release the following factors: angiopeitotin-2, calcium, coagulation factors, epithelial cell growth factor, epidermal growth factor, fibronectin, fibrinogen, bFGF of FGF, histamine, insulin-like growth factor-1, interleukin-1, platelet-derived angiogenesis factor, platelet-derived endothelial cell growth factor, PDGF (AA, BB, and AB isomers), platelet factor 4, serotonin, TGF-β1, PDGF-BB, VEGF, and bFGF as representative platelet growth factors because they were produced in the largest amounts and showed greatest ability to enhance the maturation of newly formed blood vessels.

To quantify the amounts of the growth factors in the PLs, we used enzyme-linked immunosorbent assay (ELISA) and a transforming growth factor (TGF)-β1, platelet-derived growth factor (PDGF)-BB, vascular endothelial growth factor (VEGF), and bFGF kits (R&D Systems, Minneapolis, MN) according to the manufacturer’s instructions. Samples of PL from six humans were added to 96-well plates coated with anti–growth factor antibodies. After incubating the plates and removing the unbound substances, we added an enzyme-coupled secondary antibody to the well plates. After the color reactions occurred, we used a spectrophotometer to measure the optical densities of each product at the appropriate wavelength.

Release test of PL from CGS

To evaluate the release of PL from CGS, pieces of CGS (1.0 × 1.5 × 0.3 cm) were added to 250 μL of phosphate-buffered saline (PBS; Invitrogen, Tokyo, Japan) and stored in a CO2 incubator at 37°C. For 24 h, we removed 200 μL of the PBS release media for sampling and replaced it with 200 μL of the PBS release media for sampling at different times and replaced the media with 200 μL of fresh PBS. At 24 h, we removed 200 μL of the PBS release media for sampling and replaced it with 200 μL of 2 U/mL collagenase type A1 (Sigma Chemical, St. Louis, MO). We continued to remove 200 μL of the PBS release media and replace it with 200 μL of 2 U/mL collagenase type A1 at different times for the next 24 h. The experiment was performed independently for three samples and three controls (NSS) at every sampling time.

To quantify the amounts of TGF-β1 and PDGF-BB in the samples, we used ELISA and TGF-β1 and PDGF-BB ELISA kits (R&D Systems, Minneapolis, MN) according to the manufacturer’s instructions. The samples (n = 3) and controls (NSS, n = 3) were added to 96-well plates coated with anti–growth factor antibodies. After incubating the plates and removing the unbound substances, we added an enzyme-coupled secondary antibody to the well plates. After the color reactions occurred, we used a spectrophotometer to measure the optical densities of each product at the appropriate wavelength.

Impregnation of CGS with PL

Pieces of CGS (8-mm diameter, 1.5 mm thick) were placed in 10-cm tissue culture dishes (Falcon; Falcon, NY). We
applied 50 μL of NSS (control), PL, or ×2, ×3, or ×4 concentrated PL to the CGS and then incubated the impregnated CGS overnight at 4°C.

Implantation of CGS

To determine the concentration of PL in CGS that is most effective in promoting wound healing, we first implanted pieces of CGS impregnated with different concentrations of PL into full-thickness skin defects on the backs of mice. Eight-week-old C57BL/6JcJ mice were anesthetized by intraperitoneal injection of 26.3 mg pentobarbital (Abbott Laboratories, North Chicago, IL). After depilating the mice, we used an 8-mm-diameter skin punch biopsy tool (Kai Industries, Gifu, Japan) to create a full-thickness skin defect (including the panniculus carnosus) on the back of each mouse. After dividing the mice into five groups of six mice each, we implanted pieces of CGS impregnated with NSS (control), PL, or ×2, ×3, or ×4 concentrated PL into the defects and sutured the CGS to the margins of the skin wound with 5-0 nylon sutures (Johnson & Johnson, Tokyo, Japan). The silicone sheets were then placed over the defects to keep the inner sponge layer wet and prevent bacterial infection.

Assessment of the wound area and histologic assessment of neoeplithelization

Mice were killed by carbon dioxide inhalation 1, 2, or 3 weeks after CGS implantation. The silicone sheets were removed, and each treated wound area, which was expressed as a percentage of the area of the original 8-mm wound, was photographed and measured using ImageJ software (version 1.38; Wayne Rasband, NIH, Maryland). We used a scalpel and scissors to harvest the implanted CGS. The CGS specimens were fixed with 20% formalin, embedded in paraffin, cut axially into 4-μm-thick sections, and then stained with hematoxylin and eosin. We then used a light microscope to measure the length of each specimen’s neoeplithelium from the outermost hair root of the marginal skin on each side of each cross section.

Immunohistological staining of newly formed capillaries and evaluation of capillary area

To detect newly formed capillaries in the implanted CGS, we performed immunohistological staining of von Willebrand factor. After the harvested CGS sections had been deparaffinized and rehydrated, they were incubated in PBS and then incubated with primary antibody (1:250 dilution), and En Vision +Rabbit/HRP (Dako Japan, Tokyo, Japan) was used as the secondary antibody. The sections were exposed to 3-3'-diaminobenzidine-4HCl (Dako Japan) for 2 min at room temperature and then counterstained with hematoxylin.

Digital light micrographs of sections beyond the muscle layers were taken at 100× magnification. The cross-sectional area of stained capillaries beyond the muscle layer in each wound was measured using the NIS-Elements D imaging software program (version 2. 20; Nikon, Tokyo, Japan). Measurements were performed in the central region of a 500-μm diameter in each section.

Statistical analysis

We used an analysis of variance and a Fisher protected least significant difference test to compare differences between groups. All data are expressed as means±standard errors. p values <0.05 were considered statistically significant. The Microsoft Excel software program with the Statcel software add-in (Oms publishing, Inc., Tokyo, Japan) was used for all statistical analyses.

Results

Growth factors in PL

The concentrations of the principal growth factors released from PL are shown in Table 1. Various growth factors were released from PL after the freeze–thaw activation. The amounts of growth factors in concentrated PL are shown in Figure 1.

PL release from CGS

Our growth factor release experiments revealed that CGS could bind to and release TGF-β1 and PDGF-BB in vivo and that this release depended on the biodegradation of collagenase (Fig. 2). During the initial 24-h incubation period, CGS incubated with PBS without collagenase did not release TGF-β1 or PDGF-BB in a sustained manner but did release initial bursts of small amounts of the growth factors. In contrast, during the second 24-h incubation period, CGS incubated with collagenase-containing PBS released TGF-β1 and PDGF-BB in a sustained manner. During the first 24-h period, the CGS incubated with PBS without collagenase released up to 50% of the growth factors; during the second 24-h period, collagenase degradation caused the CGS to release the remaining 50% of the growth factors. A similar addition effect of collagenase occurred on the time profile of CGS degradation. The time profiles of TGF-β1, PDGF-BB, and CGS degradations had good correlation.

Wound area

The gross appearances of wounds 2 weeks after the implantation of CGS are shown in Figure 3A. Three weeks after CGS implantation, the wounds treated with PL-impregnated CGS had almost completely epithelized. The percentage of original wound area remaining over time is shown in Figure 3B. One, two, and three weeks after CGS implantation, the areas of the wounds in the mice treated with PL-impregnated

| Table 1. Concentration of Growth Factors in Platelet Lysate |
|-----------------|-----------------|
| **Growth factor** | **Concentration (pg/mL)** |
| TGF-β1          | 103449±7137     |
| PDGF-BB         | 5112±1200       |
| VEGF            | 1568±521        |
| bFGF            | 75.1±11.1       |

The concentrations of TGF-β1, PDGF-BB, VEGF, and bFGF released from PL. Samples of PL (n=6) were measured using enzyme-linked immunosorbent assay. TGF, transforming growth factor; PDGF, platelet-derived growth factor; VEGF, vascular endothelial growth factor; bFGF, basic fibroblast growth factor; PL, platelet lysate.
CGS were significantly smaller than those in the control group. Two weeks after CGS implantation, the areas of the wounds treated with CGS impregnated with \( \times 2 \) concentrated PL were significantly smaller than those of the wounds treated with CGS impregnated with \( \times 3 \) or \( \times 4 \) concentrated PL.

**Histologic assessment of neoeptithelium length**

We observed newly formed epithelia in wounds treated with PL-impregnated CGS 2 weeks after CGS implantation (Fig. 4A). One and two weeks after CGS implantation, the lengths of neoeptithelium in wounds treated with CGS impregnated with \( \times 2 \) concentrated PL were significantly longer than those in the control group (Fig. 4B).

**Evaluation of newly formed capillaries in wounds**

CGS that had been immunohistologically stained for von Willebrand factor is shown in Figure 5A. Matrices of CGS were present in wounds treated with NSS-impregnated CGS but not those treated with PL-impregnated CGS. The gross area of the capillaries in the group treated with CGS impregnated with \( \times 2 \) concentrated PL was significantly larger than those in the control group and groups treated with CGS impregnated with \( \times 3 \) or \( \times 4 \) concentrated PL (Fig. 5B).

**Discussion**

In the present study, we found that CGS can bind to TGF-\( \beta 1 \) and PDGF-BB and release these growth factors in a...
sustained manner. Both growth factors were released according to the degradation of CGS, even when concentrated PL was applied. These results suggest that the release of multiple growth factors in concentrated PL from CGS helps heal wounds.

Although freeze-dried platelets have been reported to be effective for tissue regeneration, no study has investigated the efficacy of concentrated PL for wound healing. In living tissue, various growth factors interact with components of the extracellular matrix, such as acidic polysaccharides and collagen, through various intermolecular forces. On the basis of this natural physicochemical interaction, we developed biodegradable CGS impregnated with bFGF to be used for full-thickness skin or mucosal defects.

CGS can release concentrated PL in a sustained manner and facilitate the release of growth factors, which results in better angiogenesis and dermis-like tissue formation. In the present study, we found that ×2 concentrated PL effectively accelerated wound healing, enhancing the proliferation of cell and vessel growth in granulation tissue without obvious inflammatory reactions. Moreover, the use of ×2 concentrated PL

FIG. 4. (A) Hematoxylin and eosin staining micrographs of wounds treated with NSS (a), ×1 concentrated PL (b), ×2 concentrated PL (c), ×3 concentrated PL (d), or ×4 concentrated PL (e) 2 weeks after CGS implantation. Scale bar: 100 μm. The black arrows indicate the neoeipithelium. (B) The time course of the change in the mean neoeipithelium lengths (μm) ± standard deviations of groups of six mice treated with NSS (a), ×1 concentrated PL (b), ×2 concentrated PL (c), ×3 concentrated PL (d), or ×4 concentrated PL (e) 2 weeks after CGS implantation. *p < 0.05 versus NSS; **p < 0.01 versus NSS, ×1 concentrated PL. Color images available online at www.liebertpub.com/tea

FIG. 5. (A) Immunohistological staining of the newly formed capillaries beyond the muscle layer in wounds treated with NSS (a), ×1 concentrated PL (b), ×2 concentrated PL (c), ×3 concentrated PL (d), or ×4 concentrated PL (e). The black arrows indicate newly formed capillaries. Scale bar: 100 μm. (B) The mean gross areas of capillaries ± standard deviations of groups of six mice treated with CGS impregnated with NSS (a), ×1 concentrated PL (b), ×2 concentrated PL (c), ×3 concentrated PL (d), or ×4 concentrated PL (e). Scale bar: 100 μm. *p < 0.01 versus NSS; **p < 0.01 versus ×1PL, ×3PL, ×4PL. Color images available online at www.liebertpub.com/tea
elicit results that were not significantly different from those of our previous study, in which we used bFGF. In the present study, our finding that high concentrations of PL have not accelerated wound healing is consistent with those of previous studies, which showed that very high concentrations of PL have an inhibitory effect on the proliferation and viability of different cell types. For example, one study found that PL significantly enhanced the wound closure of keratinocyte-based tissue in a dose-dependent manner; however, this healing effect was retarded at concentrations 2.5 times the physiological level and higher. Furthermore, in wounded cell monolayers, optimal concentrations of PL have been found to induce a rapid actin cytoskeletal reorganization in agreement with the cells assuming an active cell migratory phenotype.

Our findings regarding PL were not dissimilar from those of earlier studies of bFGF. For example, bFGF treatment has been reported to elicit a bell-shaped dose-response; an excess amount of bFGF over the optimal range decreased the proliferation of keratinocytes and prolonged wound closure in vivo. Similarly, the results of the present study indicated that PL elicited a bell-shaped dose-response pattern with a peak at ×2 concentrated PL, and the excess amount of platelet growth factors such as TGF-β and PDGF-BB over the optimal range reduced cell proliferation and prolonged wound closure in a manner similar to that of bFGF. The reduction in cell proliferation may have been due to the PL’s release of thrombospondin (TSP-1), which is significant in quantities in PL and has been reported to be an important angiogenesis inhibitor. High concentrations of TSP-1 have been found to markedly inhibit cell proliferation. Therefore, platelets’ abundant secretion of TSP-1 might contribute to the antiproliferative effect via cell apoptosis. Moreover, several growth factors become intrinsically involved in angiogenesis. PDGF-BB has a potent arteriogenic effect that promotes differentiation of endothelial cells, and VEGF is known to stimulate angiogenesis after ischemia. TGF-β promotes cell mitosis. However, other studies indicated that several growth factors, such as PDGF-BB and TGF-β, inhibit the angiogenic effect of bFGF. More studies are needed to determine what is responsible for the antiproliferative effect of high concentrations of PL.

On the other hand, different growth factors act in synergy to enhance tissue regeneration. For example, Hao et al. reported that the release of both VEGF and PDGF enhanced the maturation of newly formed blood vessels more than the release of only VEGF did. PL can release any drug capable of promoting the regeneration of various tissues. Thus, the findings of the present study could point the way to using platelets instead of bFGF in CGS for wound treatment. In addition, because CGS can simultaneously release agents in a sustained manner, the combination of concentrated PL- and bFGF-impregnated CGS could 1 day be used to further accelerate dermis-like tissue formation in wounds.

Conclusion

Our findings indicate that CGS can release these growth factors in a sustained manner. The ×2 concentrated PL-impregnated CGS accelerates the formation of dermis-like tissue.

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Disclosure Statement

No competing financial interests exist.

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