INTRODUCTION

Chronic wounds can develop in response to arterial and venous insufficiency, diabetes, pressure, and radiation [1]. Radiation injury disrupts all phases of the wound-healing process. In the early stages of radiation injury, erythema, dry desquamation, and hyperpigmentation occur, and in the later stages, tissue fibrosis and blood vessel damage delay healing, causing wounds to become chronic [2]. Chronic wounds are generally difficult to manage using traditional wound management. Cell therapy is an effective method for managing intractable wounds, and several studies have investigated the effects of adipose-derived stem cell (ASC)-mediated wound healing [3,4]. However, the use of ASCs for cell therapy...
requires cell expansion in vitro, which is time-consuming and increases the risk of contamination and differentiation [5]. In contrast, the stromal vascular fraction (SVF), a collection of non-expanded heterogeneous cells (mesenchymal stem cells, hematopoietic stem cells, immune cells, fibroblasts, endothelial cells, etc.) derived from enzymatically digested adipose tissue, is easy to isolate and free of ethical concerns. Several studies have demonstrated the potential of SVF as a therapy to improve wounds [6,7].

When SVFs are simply delivered to a wound in solution, rapid diffusion into extracellular fluids results in low cell retention. Ji-ang et al. [8] loaded SVFs into a three-dimensional collagen scaffold, applied it to a diabetic wound, and reported drastically increased retention and wound healing. In addition, human acellular dermal matrix, hydrogel, chitosan, and porcine small intestinal submucosa matrix have been used as carriers to deliver ASCs [9-12].

Amniotic membranes have long been used in the treatment of burns and have been recently used for tissue engineering due to their structural and biological properties [13]. Amnisite BA (Bi-oland, Cheonan, Korea) is an acellular bovine amniotic membrane that is stored at room temperature in a freeze-dried state. Storing Amnisite in a freeze-dried state confers stability, longevity, and ease of use while being equivalent in action to that of fresh amniotic membrane [14].

The purpose of this study was to develop a process to retain SVF on Amnisite as a scaffold and to confirm the delivery of SVF to wounds. We also investigated the effects of SVF with Amnisite on radiation-induced chronic wounds in a porcine model.

METHODS

One 7-month-old female pig weighing 30 kg, in good health and without skin disease, was obtained from Minipig; Medi Kinetics Pyeongtaek, Korea and used for this study. This pig was given standardized feed once daily. It was housed in an isolated room maintained at 24°C with 65% humidity. The Institutional Animal Care and Use Committee approved all experimental procedures (IACUC No. KM-2015-46).

Cellular experiments

Isolation of porcine SVF

At 5 days after the operation, which is described below, adipose tissue was harvested from a 6-cm-long × 4-cm-wide paraspinous cutaneous flap that was elevated in areas that were not irradiated. Adipose tissue samples were trimmed and transferred to sterile 50 mL conical tubes that contained 25 mL of phosphate-buffered saline (PBS). These adipose tissue samples were washed twice with PBS and minced using scissors. The minced adipose tissue had a final volume of roughly 20 mL. The minced adipose tissue samples were subsequently digested with 0.075% collagenase type I (Worthington Biochemical Corp, Lakewood, NJ, USA) in PBS at 37°C for 1 hour under steady moderate agitation. Then, culture medium containing low-glucose Dulbecco’s modified Eagle’s medium (DMEM) and 10% fetal bovine serum (FBS) was added to stop enzymatic activity. Centrifugation was performed, after which the supernatant was discarded and the pellet was resuspended and filtered through a 100-μm cell strainer to remove tissue debris. The suspension was re-centrifuged at 1,500 rpm for 5 minutes and resuspended in low-glucose DMEM supplemented with 10% FBS, seeded into 100 Ø culture dishes, and incubated at 37°C with 5% CO₂ for 2 days.

Retention of SVFs on Amnisite BA

The authors asked the manufacturer to produce Amnisite without a slit and used it in the experiment.

In order to prevent the curling effect that the Amnisite membrane typically exhibits during hydration, PBS was used. To find an efficient hydration time (the shortest time necessary to prevent curling), SVFs (1 × 10^7/cm²) were seeded on each of four hydrated Amnisite membranes (1 × 1 cm²) for 30 minutes, 1 hour, 12 hours, and 24 hours in a dish filled with PBS and then incubated at 37°C with 5% CO₂ for 24 hours. Each membrane was then stained with crystal violet and observed using an optical microscope to confirm cell attachment.

To determine the ideal incubation time, SVFs were seeded on Amnisite using the method described above. The membranes were incubated at 37°C with 5% CO₂ for 1, 3, 6, 12, and 24 hours. The success of cell attachment and the bond strength between Amnisite and the SVF were then compared. The ideal incubation time was defined as the time necessary to form a bond strength resilient to just drops of PBS but broken by hydrostatic positive pressure using a pipette. We observed cell attachment using an optical microscope after crystal violet staining.

The success of delivery to the wound and the viability of engrafted cells were evaluated in vivo. Before seeding, SVFs were labeled with PKH67 (PKH67-GL; Sigma-Aldrich, St. Louis, MO, USA). PKH67 labeling was performed in single-cell suspensions containing 0.6–1 × 10^7 SVFs that had been washed once in serum-free DMEM. The staining procedure followed the manufacturers’ instructions. To summarize, PKH was added and the cells were mixed by constant inversion of the tube for 5 minutes. Unbound PKH67 molecules were then blocked with 100% FBS (1:1). The suspension was then centrifuged for 5 minutes at 1,600 × g. The supernatant was carefully aspirated.
and cells were resuspended in the culture medium for seeding onto Amnisite.

**Animal experiment**

**Delivery of irradiation and wound creation**

A radiation-induced chronic wound model was established following protocols described in previous studies [15,16]. The pig received a single dose of 18 Gy with a 6 MeV electron beam per 18 × 8 cm area. Three radiation areas (total radiation area: 432 cm²) were selected on the paraspinal dorsal skin surface of the pig: two were located on the left side of the spine and one was located on the right side of the spine. These areas were selected to ensure sufficient non-radiated tissue around each wound for the areas to stay independent of each other while avoiding the spread of infection and skin necrosis. Over 90% of the prescribed irradiation dose was limited to a maximum depth of 2 cm.

After 5 weeks of irradiation, a skin defect wound model was designed as follows: First, tiletamine/zolazepam (Zoletil; Virbac, Carros, France) was injected intramuscularly for sedation. Then, xylazine hydrochloride (Rompun; Bayer Animal Health, Monheim, Germany) was injected intravenously for induction. After shaving the hair, a 4 cm² square excisional wound site was marked with an oil ink pen on the irradiated zone. The site was 2 cm away from the boundary. A total of six wounds were made. Each wound was positioned 6 cm apart from the other wounds. The excision tissues included the skin and the superficial fat layer. The deep fat layer was not included to prevent the spread of infection via the muscular fascia (Fig. 1).

**Preliminary animal experiment**

A foam dressing (Therasorb, Wonbiogen Co. Ltd., Gumi, Korea) was maintained on the excisional wounds until practical experiments. At 7 days post-operation, the wounds were randomly divided into three groups: group S (SVF with Amnisite, n = 2), group A (Amnisite group, n = 2), and group C (control group, Therasorb, n = 2) (Fig. 2). In group S, 1 × 10⁶/cm² SVF suspended in Amnisite was applied to the wounds, which were then covered with Vaseline gauze and polyurethane foam. In group A, Amnisite was applied to wounds dressed with the same method used for those in group S. In group C, only polyurethane foam was applied to wounds. The body of the pig was then covered with a snug-fitting non-adherent tube gauze dressing (Tubifast; Mölnlycke Health Care, Gothenburg, Sweden), which was tailored into a vest to further protect wounds.

**Measurement of wound contraction**

To compare the rate of wound contraction ([(original wound size (at 7 days post-operation))–wound size measured at that time/original wound size] × 100) among the three wound groups, photographs of the wound region were obtained with a digital camera every time dressings were changed. These photographs were analyzed using ImageJ software (National Institutes of Health, Bethesda, MD, USA). All photographs were taken with a sterilized paper ruler that was used for calibration, standardization, and subsequent quantitative analysis.

**Histological analysis**

Biopsies were taken sequentially from each of three sides of the square at 10, 14, and 21 days post-operation. The specimens were full-thickness incisional biopsy specimens (including skin, superficial and deep subcutaneous tissue, and granulation tissue).
at the margin of the wound. They were 5 mm wide and 10 mm long, and included approximately 4 mm of unwounded skin. All tissue samples were fixed in 10% neutral buffered formalin.

All scoring and interpretations of the immunohistochemical results were performed by one examiner without knowledge of the clinical data. For the histological analysis, hematoxylin-eosin staining was used to determine the grade of inflammation grade and degree of re-epithelialization. The severity of acute and chronic inflammation was graded (mild, 1; moderate, 2; severe, 3). Neutrophils and eosinophils were considered to indicate acute inflammation, whereas the presence of lymphocytes and plasma cells was deemed to be a sign of chronic inflammation. The degree of re-epithelialization was categorized as 0 (none), 1 (incomplete epithelialization, in less than half of the wound), and 2 (more than half to complete epithelialization). The neo-vascularization level of wound granulation tissues was evaluated by immunochemical analysis using CD-31 staining. Sections were scanned at a low magnification (× 40). Areas with the highest concentration of microvessels (vascular hot spots) were selected in each biopsy specimen. In each selected area, five high-magnification fields (× 400) were evaluated and the average of all counted microvessels (mean microvessel density) was calculated [17].

RESULTS

We observed that hydrating Amnisite for 30 minutes enabled the retention of cells and prevented curling (Fig. 3). In addition, we found that incubating Amnisite with the SVF for 1 hour resulted in successful cell attachment with appropriate bond strength (Fig. 4). Successful delivery of the SVF to wounds was observed with a fluorescence microscope. In group S, the PKH67-labeled SVF released specific green fluorescence in every biopsy specimen (Fig. 5).

Compared with the other two groups, group A showed a markedly higher wound contraction rate. Group S demonstrated the lowest wound contraction rate. No group healed completely during the experimental periods (Fig. 6).

On day 10, the re-epithelialization scores were 0 in groups A and C, and 1 in group S. The re-epithelialization scores were 1, 1.5, and 0.5 in groups A, S, and C on day 14, respectively. However, there was no difference in the re-epithelialization score (1.5) between any group on day 21. At 10 days post-operation, group S showed early re-epithelialization, while the other groups did not. Re-epithelialization increased in group S until 14 days post-operation, and then remained constant (Fig. 7).

The acute inflammation scores on day 10 were 1.5, 1.5, and 2 in groups S, A, and C, respectively. On day 14, the acute inflam-

![Fig. 3. Stromal vascular fraction retention over hydration time. (A) Gross findings of the amniotic membrane (AM) (1 × 1 cm²) according to hydrating time. (B) When seeding stromal vascular fractions (1 × 10⁶/cm²) on each membrane, the dried AM (0 minutes) curled due to surface tension, whereas the AM hydrated for at least 30 minutes appeared flat without any curling effect. (C) Microscopic findings after incubation at 37°C with 5% CO₂ for 24 hours (crystal violet stain): although 0 minutes showed aggregated and floating cells (arrow) without attaching to the AM, other groups showed well-attached cells.](image-url)
mation score was the highest in group A, at 2.5. The acute inflammation score on day 21 was the lowest in group C, at 0.5. Acute inflammation was the lowest in group S until 14 days post-operation. Instead of showing acute inflammation, group S had the highest chronic inflammation score at 10 days post-operation. This score decreased until day 14, but fat necrosis increased and foreign body giant cells were still present. On the 21st day, the chronic inflammation score was the same in all groups, at 1.5 (Fig. 8).

On the 10th day, the microvascular density was highest in group S, at 7.7, but by the 14th day, group A was highest, at 11.05. By the 21st day, both groups had stabilized and the densities were well matched (Figs. 9 and 10).

**DISCUSSION**

Fresh human amniotic membrane has been widely used for wound dressings [18]. However, a disadvantage of fresh amniotic membrane is that it is difficult to store. Freeze-dried amniotic membrane is a ready-to-use product with all advantages of fresh human amnion, but without its disadvantages [14]. Some ready-to-use products derived from freeze-dried amniotic membrane are currently available, such as Amnisite BA and Biovance (Alliqua BioMedical, Langhorne, PA, USA) [19]. These products were mainly developed for skin or corneal dressings; however, since amniotic membrane contains growth factors with anti-inflammatory and anti-microbial effects, it also makes an effective scaffold. There are a number of other advantages to using amniotic membrane as a scaffold. First, the membrane has an extracellular matrix, which allows cell adhesion and growth. Additionally, amniotic membrane has plasticity and other biological properties, including several growth factors with anti-inflammatory and anti-microbial effects [13]. For cell therapy, attaching cells to the scaffold is a significant step. Cell seeding is the first step in cell adhesion, and its efficiency and distribution affect the final outcome. Gholiapourmalekabadi et al. [20] seeded human ASCs on a decellularized human amniotic membrane and incubated it at 37°C with 5% CO₂ for 1, 2, 3, and 7 days. Mahmoudi-Rad et al. [21] showed...
excellent growth of allogenic fibroblasts on an acellular amniotic membrane scaffold, which they produced and incubated for 24 hours. Diaz-Prado et al. [22] seeded human chondrocytes on a cryopreserved human amniotic membrane and incubated it for 3–4 weeks. They then used this membrane to support chondrocyte proliferation for transplantation therapy to repair osteoarthritis cartilage in vitro. While the purposes of those experiments were different from the purpose of the present study, the...
processes of preparing the amnion scaffold and incubating the cell-scaffold complex took one day to several days. In addition, the prior studies were all conducted in vitro, without confirming whether actual cells were delivered in vivo. In contrast, we completed amnion scaffold preparation, cell seeding, and incubation in a total of 90 minutes and confirmed cell viability and successful attachment of SVF to Amnisite. Furthermore, we confirmed successful cell delivery to radiation-induced chronic wounds using an in vivo experiment. In clinical practice, the authors’ method can be used to harvest and process cells at the start of surgery. These cells can then be loaded into the scaffold and delivered to the wound at the end of surgery.

Cell delivery was confirmed in preliminary animal experiments, but improved wound healing through the use of cell-loaded Amnisite was not confirmed. Histopathological examinations showed that the SVF had positive effects on re-epithelialization until 14 days post-operation, without significant changes thereafter. These positive effects of SVF might gradually decrease, with SVF losing its effectiveness after 14 days post-operation. This change might be due to the anti-inflammatory effect of SVF, which promotes revascularization [23,24].

The epithelialization at the wound edge was better in group S than in group A, but the wound contraction was better in group A. The acute inflammation score was lower in group S than in group A, but the chronic inflammation score was higher in group S than in groups A and C. These results suggest that SVF-loaded Amnisite temporarily reduces acute inflammation and promotes epithelialization at the beginning of application, but its effect does not last long. Moreover, since radiation-induced chronic wounds already have many chronic inflammatory cells and have limitations in extracellular matrix production and wound contraction, it was difficult to analyze and interpret the results.

As this was only a preliminary in vivo experiment, there are some limitations. The sample size of animals was small (n = 2). In order to better elucidate the effectiveness of cell therapy on radiation-induced chronic wounds, the timing and conditions of cell therapy should be planned in more detail. Even a single

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**Fig. 9.** Microvascular density score. Group S showed the highest level of revascularization at 10 days post-operation. This effect then gradually decreased until 21 days post-operation, when group S showed the least revascularization among the three groups. Group A, Amnisite group; Group S, SVF with Amnisite; Group C, control group.

**Fig. 10.** Histological and immunohistochemical findings of microvascular density at 10 days. In group C, slightly increased vascular structures with retained acute and chronic inflammatory cells. In group A, inflammatory cells slightly increased but the number of vessels did not increase compared to group C. In group S, markedly increased vascular density was noted, with remarkably reduced inflammatory cells in the stroma. Group A, Amnisite group; Group S, SVF with Amnisite; Group C, control group.
cell delivery would have been insufficient to make a difference. In addition, the histological grading of the wound must be based on the meticulous assessment of different parameters according to each wound healing phase to assess the wound status accurately [25].

As far as the authors are aware, this is the first study providing evidence for the effectiveness of freeze-dried bovine amniotic membrane as a scaffold for cell delivery. In this preliminary study, we developed a process to retain SVF on an amniotic membrane as a scaffold and confirmed the delivery of SVF to radiation-induced chronic wounds in a porcine model. We demonstrated that SVFs had acceptable adherence on freeze-dried bovine amniotic membranes, showing that this complex can potentially be an effective scaffold with a high level of convenience for soft tissue engineering applications.

NOTES

Conflict of interest
No potential conflict of interest relevant to this article was reported.

Ethical approval
Animal testing was conducted in accordance with the ethical standards of the Institutional Animal Care and Use Committee of Keimyung University (IACUC No. KM-2015-46).

Author contribution
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