

Evaluation of an Amniotic Membrane-Collagen Dermal Substitute in the Management of Full-Thickness Skin Defects in a Pig

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Background To minimize the inflammatory reaction and improve healing, a new modified dermal substitute composed of an atelocollagen, chondroitin-6-sulfate, and amniotic membrane (AM) was applied to full-thickness skin defects in a pig. Atelocollagen was extracted from bovine skin, and two modified dermal substitutes were generated according to the crosslinking type.

Methods The AM-collagen dermal substitutes were characterized and compared with currently used dermal substitutes in a pig skin defect model. There were five experimental groups: dehydrothermal (DHT) cross-linking atelocollagen with the AM on the top (AM-DHT), DHT and chemical cross-linking atelocollagen with the AM on the top (AM-DHT/chemical), Terudermis, Integra, and AlloDerm. After 3×3 cm full-thickness skin defects on the back of a pig were created, each dermal substitutes dermal substitutes was randomly grafted on the defects. Two weeks after grafting, autologous partial-thickness skin was over-grafted on the neodermis. The take rate of the dermal substitutes, skin, and histological sections were all assessed at 1, 2, and 4 weeks postoperatively.

Results More rapid healing and a higher take rate were evident in the AM-DHT and Terudermis groups. Histological examination revealed fewer inflammatory cells and more fibroblast hyperplasia in these two groups. Four weeks after surgery, the amount of newly formed collagen was significantly more appropriate in the AM-DHT group.

Conclusions These observations provide supporting evidence that a newly developed amnioticcollagen dermal substitute may inhibit inflammatory reactions and promote wound healing.

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INTRODUCTION

Up to the present, autologous skin grafts or flaps have been wide-

ly used for repairing skin and soft tissue defects. However, improved reconstruction poses greater donor site problems. To resolve these problems, skin substitutes have been developed using

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tissue engineering [1]. However, in order for skin substitutes to be applied to patients with full-thickness skin loss, the substitutes should function as an alternative to autologous skin, form an effective barrier against bacterial invasion, minimize inflammation and scar formation, improve fibrovascular tissue ingrowth, and have excellent reproducibility [2,3].

A dermal substitute, which is the most extensively developed of the current skin substitutes, consists of a collagen scaffold that comprises mainly collagen containing either dermis extracellular matrix or cultured fibroblasts. A bilayered skin substitute that contains epidermis-forming keratinocytes attached on a collagen scaffold with cultured fibroblasts is ideal. However, its take rate is problematic.

We have generated a novel dermal substitute containing a collagen scaffold by combining chondroitin-6-sulphate (C6S) with atelocollagen (ATC) extracted from calf dermis. To lessen the antigenicity of ATC, the telopeptides located at the terminal of the collagen molecule were removed using pepsin. C6S was added to increase stability and the scaffold was treated by physical cross-linking in a dehydrothermal (DHT) treatment alone or with an additional chemical cross-linking involving 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC). We used an amniotic membrane (AM) instead of silicone membranes that temporarily served as epidermis, to protect the wound from the external environment, to keep the wound moist, and to aid in the promotion of wound healing through various growth factors.

This study was undertaken to generate a new dermal substitute and to compare it with previously designed dermal substitutes, in order to confirm its usefulness in clinical practice through animal experiments.

METHODS

Preparation of an atelocollagen dermal substitute Purification of ATC solution

The extracted collagen from calf skin was treated with 2 mol/L NaCl and agitated, and the pellets were obtained using centrifugation (6,000 rpm for 20 minutes). The pellets were dissolved in 1 mol/L NaCl/0.05 mol/L Tris at pH 7.4, and the supernatant was obtained using centrifugation (3,500 rpm for 60 minutes). The supernatant was salted out for 3 hours with an addition of 4 mol/L NaCl, and the pellets were retrieved using centrifugation (6,000 rpm for 20 minutes). The pellets were dissolved in a pH 2.0 HCI solution, treated with 2 mol/L NaCl, and agitated. After centrifugation (6,000 rpm for 20 minutes) the pellets were again dissolved in the HCI solution. The solution was dialyzed in a dialysis tube and retrieved. After titration at pH 3, the solution was filtered using a 0.2 µm filter to obtain 0.5% ATC.

Preparation and cross-linking of AM-collagen scaffold

The AM was harvested from a healthy bovine placenta and disinfected with 70% ethanol and 0.05% sodium hypochlorite. During the processing, the epidermis was removed and the remaining stromal layer was incubated in an aqueous solution containing 0.25% (w/v) trypsin, 0.02% (w/v) ethylene diamine tetraacetic acid, and 0.9% (w/v) NaCl at pH 7.4 for 60 minutes to remove the cellular components. After the cells were removed from the tissue, the acellular AMs were dried to preserve their biochemical and structural integrity. The purified 0.5% ATC was mixed with 1% C6S equivalent to 10% of the dried weight of the collagen and was agitated at 1,500 rpm at 4°C for 5 minutes. The cream-colored ATC-C6S solution was poured into a square-shaped disk mold, in which an AM $(12.5 \times 12.5 \text{ cm})$ was attached using a cell scraper. The bubbles were completely removed using an air drier, and the collagen scaffold was frozen at -80°C and then lyophilized for 48 hours.

The lyophilized collagen scaffold was cut into 3×3 cm sections using a surgical scalpel. Each section was placed in a bottle, and DHT cross-linking was performed in a vacuum oven at 110°C. For additional chemical cross-linking, the sections were treated with an EDC solution at pH 5.5, rinsed sufficiently with a 0.1 moL/L NaHPO₄ solution, and hydrated with purified water. The sections were then frozen at -80°C and dried using a lyophilizer. The ATC dermal substitute was sealed using an air-permeable material and sterilized with ethylene oxide gas.

Characterization of the AM-collagen dermal substitute Amino acid analysis

The ATC solution (1.31 mg) was hydrolyzed and labeled with phenylisothiocyanate (PITC) using a Pico-Tag Kit (Waters Associates, Milford, MA, USA). The chromatograms were obtained from 10 μ L aliquots of the 1,600 μ L volume of the PITC-labeled specimens by high performance liquid chromatography (Waters Associates).

Electron microscopy

Both DHT cross-linked and DHT/chemical cross-linked collagen scaffolds were treated with gold using a Polaron E5100 Coating System (JEOL, Tokyo, Japan) and their cut surface was examined by scanning electron microscopy (SEM) using a JSM-6300 microscope (JEOL) at a magnification of × 150.

Biochemical analysis of amniotic membrane

Freeze dried bovine placental amniotic membrane was used as raw material before making the AM-collagen dermal substitutes and processed AM was used for biochemical analysis. Two kinds of tissue samples were frozen in liquid nitrogen, ground with a freezer mill (SamplePrep Model 6750, SPEX, Metuchen, NJ, USA), and homogenized in 1 mL of phosphate-buffered saline. The protein levels were measured using an enzyme linked immunosorbent assay system (fibroblast growth factor [FGF], DFB50; keratinocyte growth factor [KGF], DKG00; R&D Systems, Minneapolis, MN, USA). The total proteins were measured with the Bradford assay.

Animal experiments

The experiments were conducted in 12 2- to 3-month-old Yorkshire female pigs free of skin diseases. The animals were sedated using an intramuscular injection of 2.2 mg/kg azaperone (Stresnil, Korea Yansen, Seoul, Korea), and the skin was prepared. Xylazine HCl (3 mg/kg; Rumpun, Bayer Korea, Seoul, Korea) and ketamine HCl (8 mg/kg; Ketalar, Yuhan Yanghang Pharmaceutical, Seoul, Korea) were administered through the auricular vein.

Creation of full-thickness skin defects and application of the dermal substitutes

The site of the wound was designated with a marking pen on the skin over the paravertebral muscle of the pig, with a square size of 30 mm. The full-thickness skin above the fascia of the muscle was excised with a No. 10 scalpel (Fig. 1). Two kinds of the novel AM-collagen dermal substitute (group 1 and 2) and three other groups of currently available dermal substitutes were prepared: group 1, DHT cross-linking atelocollagen scaffold with the AM on one side (AM-DHT group); group 2, DHT and chemical cross-linked atelocollagen scaffold with AM on one side (AM-DHT/chemical group); group 3, the Terudermis group (Terumo, Tokyo, Japan); group 4, the Interga group (Life-Sciences, Plainsboro, NJ, USA); and group 5, the AlloDerm group (LifeCell, Woodlands, TX, USA) (Table 1). Five different types of dermal substitutes from the five experimental groups were all randomly grafted into the full-thickness of every pig on



its skin defects. The surgical wound was draped with Allevyn and Opsite (Smith & Nephew, London, UK). The dressings were further fixed with cotton balls and a plastic cover and were changed every 3 days. Twelve pigs were operated on sequentially under the same protocol and were categorized into three equal groups of four according to time of examination (n=4). Each group was euthanized for biopsy on postoperative weeks 1, 2, and 4 after inspection and gross examination. For the animals in the 4-week group, autologous partial-thickness skin from the groin area was applied to the neodermis 2 weeks after the dermal substitute grafting in order to ascertain whether the neodermis was vascularized enough to accept a free skin graft.

Gross findings and histological examination

The take rates of the dermal substitute grafts of the five groups were assessed at 1 and 2 weeks after grafting. With the 4-week pig group, the partial-thickness skin grafts that had been applied to the neodermis, which were induced by the dermal substitutes at 2 weeks, were assessed for the take rate. Excluding the principal investigator and co-investigators of this study, five residents of the Department of Plastic and Reconstructive Surgery of our hospital who were blind to this study rated the take rate of the grafts using an ordinary scale method. An 8-point scale was used, with 1 being worst and 8 being best. Each of the scores were totaled and categorized as excellent (score of 31 to 40), good (score of 21 to 30), fair (score of 11 to 20), or poor (score of \leq 10).

After the animals were euthanized, the grafted dermal substitutes were harvested for histological examination including the underlying muscle and fascia and 10 mm of normal skin around the wound edge. Four specimens (n=4) were obtained from

Group	Temporary epider- mal component	Dermal component			
Group 1 (AM-DHT)	Amniotic membrane	Atelocollagen and C6S (bovine skin)			
		DHT cross-linked			
Group 2 (AM-DHT/chemical)	Amniotic membrane	Atelocollagen and C6S (bovine skin)			
		DHT/chemical cross-linked			
Group 3 (Terudermis)	Synthetic polysiloxane polymer	Denatured atelocollagen and fibrillar atelocollagen (bovine skin)			
		DHT cross-linked			
Group 4 (Integra)	Synthetic polysiloxane polymer	Atelocollagen and C6S (bovine tendon) Chemical cross-linked			
Group 5 (AlloDerm)	None	Acellular human dermal tissue			
AM, amniotic membrane; DHT, dehydrothermal; C6S, chondroitin-6-sulphate.					

Table 1. Summary of the experimental groups according tocomposition

each experimental group. The specimens were fixed in 10% neutral buffered formaldehyde and stained with hematoxylineosin, Masson's trichrome, and α-SMA immunohistochemical staining. The specimens obtained 1 and 2 weeks after surgery were assessed for inflammatory cell infiltration, neovascularization, fibroblast hyperplasia, and the replacement extent of the dermal substitutes with the neodermis. The specimens obtained 4 weeks after surgery were assessed for the arrangement of collagen in the neodermis, and the degree of similarity between the grafted substitute and the normal skin was determined.

RESULTS

Characterization of the AM-collagen dermal substitute Antigenicity of the ATC solution

The antigenicity of the purified ATC was evaluated through the analysis of its amino acid composition (Table 2). The percentage of tyrosine relative to the total amino acids was 0.12%, which was lower than that of a criterion for nonantigenic collagen (0.2%) [4].

Electron microscopy findings of the collagen scaffold

The cut surfaces of representative collagen scaffolds were examined using SEM, and the structure of collagen fibers was examined using TEM. The DHT cross-linked atelocollagen scaffolds

Table 2. Analysis of amino acid o collagen	composition in the atelo-
Amino acid	Composition (%)
Glycine	33
Proline	12
OH-proline	10
Tyrosine ^{a)}	0.12
Cysteine	< 0.1
^{a)} Tyrosine (%), antigenicity of atelocollagen.	

Table 3. Clinical analysis of the take rates of the dermal substitute grafts at postoperative week 1 and 2, and the partialthickness skin grafts at postoperative week 4 (ordinary scale method, n = 4)

Group	Ро	/k)				
Group	1 ^{a)}	2 ^{a)}	4 ^{a)}			
AM-DHT	38	36	35			
AM-DHT/chemical	28	31	32			
Terudermis	37	35	34			
Integra	24	28	33			
AlloDerm	8	17	12			
AM ampiatic memory DIT debudyethermol						

AM, amniotic membrane; DHT, dehydrothermal. ^{a)}Excellent, 40–31; good, 30–21; fair, 20–11; poor, ≤10. revealed the cut surface of the collagen scaffolds to be similar to that of normal skin, but the DHT/chemical cross-linked atelocollagen scaffold exhibited a somewhat crushed appearance (Fig. 2).

Biochemical analysis of the amniotic membrane

Ouantities of FGF in a freeze dried bovine placental AM and processed AM were 919.8 pg/mg and 41.7 pg/mg, respectively. The amount of KGF was similar before and after cell exclusion, but the proportional levels of the extracted protein pre- and postcell removal were 31.01 pg/mg and 238.64 pg/mg, respectively. In other words, the protein level was 8 times higher after cell extraction. Therefore, it was confirmed that KGF was preserved in the processed AM.

Animal experiments

Gross findings

One week after surgery, groups 1 and 3 showed better take rates of the dermal grafts than the other groups. Overall, the take rate was better in the DHT cross-linked group than in the DHT/ chemical cross-linked group (Fig. 3). Two weeks after surgery, the take rate of the dermal grafts was slightly better in the DHT cross-linked groups than in the DHT/chemical cross-linked groups, but the difference in the take rate had decreased considerably.

Four weeks after surgery, the take rates of the partial-thickness skin grafted on the neodermis induced by the dermal substitute were excellent in all of the groups with the exception of group 5 (Table 3).

Histological findings

One week after surgery, group 1 had scanty exudate under the AM, good fibroblast hyperplasia, and good neovascularization, and most of the undersurface of the collagen scaffold of the dermal substitute was replaced with newly formed dermis. Group 3

Representative scanning electron micrographs of the DHT crosslinked and DHT/chemical cross-linked collagen scaffolds (×150). The DHT/chemical cross-linked collagen scaffold collapsed. (A) Cross section of group 1 (AM-DHT). (B) Cross section of group 2 (AM-DHT/ chemical). DHT, dehydrothermal; AM, amniotic membrane.



Fig. 3. Macroscopic findings at postoperative week



Table 4. Summary of the microscopic findings at postoperative weeks 1 and 2 (n=4)

Group	AM-DHT	AM-DHT/chemical	Terudermis	Integra	AlloDerm		
Inflammatory cell infiltration (wk)							
1	++	+	++	+	++		
2	+	+	+	+	++++		
Fibroblast proliferation (wk)							
1	+++	++	+++	+	+		
2	++++	+++	++++	++	+		
Angiogenesis (wk)							
1	+++	++	+++	+	+		
2	++++	+++	++++	++	+		
AM, amniotic membrane; DHT, dehydrothermal; ++++, severe; +++, moderate; ++, mild; +, rare.							

also showed good fibroblast hyperplasia and good neovascularization on most of the collagen scaffold, except on a part of the upper surface. However, in group 2, a considerable amount of the grafted dermal substitute remained unchanged and showed fibroblast hyperplasia only on the undersurface of the collagen scaffold. In group 4, it remained unchanged and showed little inflammatory cell infiltration or delayed graft take at the undersurface of the collagen scaffold due to focal penetration of fibroblasts. Group 5 showed severe inflammatory cell infiltration and calcification (Fig. 4, Supplemental Fig. 1).

Two weeks after surgery, groups 1 and 3 showed remarkable fibroblast hyperplasia, good neovascularization, and good maturation of newly formed collagen. Groups 2 and 4 revealed good penetration of fibroblasts into the collagen scaffold and good fibroblast hyperplasia, whereas neovascularization was slightly delayed and the upper surface of the collagen scaffold remained partially unchanged. Group 5 showed poor inflammatory cell infiltration, severe calcification, and lymphadenoid changes (Table 4, Supplemental Fig. 2).

Four weeks after surgery, in groups 1 and 3, the arrangement



and amount of newly formed collagen were appropriate and similar to normal dermis, showing that the number of cells was small compared with the amount of collagen. The remaining groups showed similar findings but revealed a rather unstable arrangement of newly formed collagen fibers (Supplemental Figs. 3, 4).

DISCUSSION

Dermal substitutes should be physically strong and easy to han-

dle, endure contraction through durability, continue to attach viable fibroblasts during the whole wound healing process, enable the reconstruction of the dermal lattice with fibroblasts, and should endure hydrolysis during their presence in the wound until they are finally absorbed after complete wound healing [2,5].

The use of a collagen scaffold as a dermal substitute reduces wound contraction and scar formation, and promotes epithelialization. Collagen has attracted much attention because it stimulates wound healing through its strong affinity to the soft tissue and blood vessels, and it can be replaced with host cells and vessels through acceleration of their penetration into the scaffold [6]. Because chemical and physical changes in collagen may affect the recipient tissue morphologically and physiologically, we attempted to investigate whether changes in collagen would affect its affinity to host cells.

In this study, the novel substitute displayed many advantages over substitutes currently in use. We used the AM instead of a silicone membrane to obtain the physical protection barrier functions as well as the anatomical and physiological functions similar to epidermis. The AM consists of three layers: the epithelial layer, basement membrane, and mesenchymal layer. The epithelial layer consists of a single layer of epithelial cells, and the basement membrane is a thin membrane composed of reticular fibers that attach to the epithelial layer by interdigitation. The mesenchymal layer is an avascular collagen matrix that is divided into compact, fibroblast, and sponge layers [7,8]. Although the AM has some variations in structure, it has histologically similar structures to the skin composed of the multilayered epithelium and basement membrane and, thus, has advantages in wound healing such as promotion of epithelialization, reduction in inflammatory reactions, and inhibition of scar formation and bacterial growth [9,10]. Promotion of epithelialization using AM has been clinically utilized for various skin lesions such as burns and ulcerations, and many growth factors have been identified in the AM by experimental studies. Koizumi et al. [11] documented that the AM expresses mRNAs of 8 growth factors that remarkably influence epithelial regeneration. Hao et al. [12] demonstrated that the AM shows mRNA expression of the interleukin-1 receptor antagonist and interleukin-10. In addition, Tseng et al. [13] suggested that inflammatory reactions are significantly reduced in the cornea, and limbus grown on the AM, probably due to reduced mRNA expression of transforming growth factor (TGF) receptor II and TGF- β -1, 2, and 3 through the inhibition of the TGF- β -signaling system. Inhibition of the TGF- β signaling system can reduce scar formation through the activation of fibroblasts into myofibroblasts. It has been reported that the amnion and chorion have an antibacterial effect against hemolytic Streptococcus group A, Escherichia coli, and Pseudomonas [14].

AM has been used as a temporary substitute because it protects skin wounds from the external environment, it decreases the loss of water, electrolytes, protein, and energy, it reduces pain, it compensates for the disadvantage of dermal substitutes in current use that are prone to infection, and it promotes wound healing through various growth factors [15].

We generated collagen scaffolds by combining C6S with atelo-

collagen extracted from calf dermis. To lessen the antigenicity of the atelocollagen, the telopeptides located at the terminal of the collagen molecule were removed using pepsin. In this atelocollagen, the percentage of tyrosine in relationship to the total amino acids was 0.12%, which was significantly lower than that of the criterion for non-antigenicity of collagen (0.2%), suggesting that inflammatory reactions can be adequately lessened when grafted in human skin. C6S is a high-molecular substance that inhibits proteolytic enzymes, plays a crucial role in the binding and regulation of growth factors, and influences chemotaxis, migration, hyperplasia, and differentiation of cells [16]. Thus, the dermal substitute we generated by the addition of C6S could reduce biodegradation of collagen caused by proteolytic enzymes, and could lead to easy migration of cells from the surrounding tissue into this collagen scaffold due to the increased elasticity, durability, and porosity compared to a collagen scaffold alone [17].

DHT/chemical cross-linking inhibits the rapid degradation of a collagen scaffold, lessens its antigenicity, and enforces its physical properties when it is grafted on human skin. A DHT cross-linked collagen scaffold is more stable, maintains the same porosity as that prior to the treatment without distortion, and does not induce cytotoxic substances. This collagen scaffold is distorted in an EDC solution due to hydrophobicity induced by the initial DHT-treatment. Although this phenomenon can be prevented to some extent when crosslinking is conducted in the presence of ethanol, such chemical substances can induce cytotoxic substances *in vivo*. However, an EDC-treated collagen scaffold can prevent rapid degradation because it becomes resistant to proteolytic enzymes [16].

In this study, the results were significantly different between the experimental groups. Groups 1 and 2 (the DHT-treated groups) showed a more rapid wound healing than the other groups, suggesting that DHT-treated collagen scaffolds have a strong affinity to cellular components of the surrounding soft tissue and blood vessels, which was evident from the electron microscopic findings (Supplemental Fig. 3). The results support the idea that it is conceivable that atelocollagen may inhibit inflammatory reactions by lessening antigenicity, whereas C6S may promote wound healing by reducing biodegradation caused by proteolytic enzymes, which increases physical stability and induces the ready migration of cells from the surrounding tissue into the collagen scaffold through its increased porosity. In addition, AM used as a temporary protection membrane can promote wound healing, probably through the secretion of various growth factors.

In conclusion, our atelocollagen dermal substitute that was generated using atelocollagen, AM, and C6S may be more reliable than collagen dermal substitutes in current use.

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Supplemental data can be found at: http://e-aps.org/src/sm/aps-40-11-s001.pdf