Experimental Study on Ostrich Acellular Dermal Matrix in Repair of Full-Thickness Wounds of Guinea Pig

Fereshteh KHOMEJANI FARAHANI 1 Hamidreza FATAHIAN 1,  Abdol-Mohammad KAJBAFZADE 2

1 Department of Clinical Science, Faculty of Specialized Veterinary Science, Science and Research Branch, Islamic Azad University, Tehran - IRAN
2 Department of Pediatric Urology, Children's Hospital, Tehran University of Medical Sciences, Tehran - IRAN

INTRODUCTION

Skin suffers from different injuries and losses. When tissue loss is extensive, the healing process may not be enough to cure, and treatment with flaps and grafts should be considered. Skin grafts, may be classified as autologous, homologous or xenogenic, such as those of porcine or bovine origins. Different procedures for tissue replacement have been introduced with the purpose of reducing scar formation and speeding up healing time [1,2]. Different...
methods of wound closure are in use, and each has its own advantages and disadvantages. The use of xenogenic skin as a source of ADM (Acellular dermal matrix) might alleviate some problems and make ADM more readily available for use in reconstructive surgery [2]. Due to excellent success in a variety of different biomedical applications, the performance of ADM, derived from full-thickness skin treated and removing of cells and cellular components, recently has drawn the attention of researchers in many fields [3]. Acellular dermal matrices are used as a suitable tissue for covering and also treatment of extensive burns [3]. In the wound-healing process, dermal substitutes play a guiding and supporting role during the cell and blood vessel recovery process [4]. Dermal substitutes serve as a scaffold into which cells can migrate and repair the injury [3]. Acellular dermal tissues are obtained by removing cellular components that are present in the tissue, since the cells are responsible for the activation of the immunological response. The absence of cellular structures result in little inflammation, and prevents rejection of the transplanted tissue [2]. As a dermal regeneration template, they also promote the guided regeneration function of the tissue, reduce scar hyperplasia and improve the quality of wound-healing [4]. The usefulness of ADM, which is derived from full-thickness skin following of removing the cells and cellular components concentrated in skin defect and cosmetic improvements. Because of absence cell particles of skin and diminishing of its antigenicity, it has been one of the most ideal substrate for a microenvironment for the cell movement and proliferation with no rejection. This experimental study deliberate to research of the effect of ostrich ADM in healing of full-thickness wounds of guinea pig.

**MATERIAL and METHODS**

Eighteen (n=18) adult male guinea pigs (230±17 g) were conducted. They were housed in cages with appropriate same food, humidity, light and bedding. Animals were fasted for 6 hours before surgery. All animals were administrated subcutaneously by saline-dextrose solution (15 ml/kg) one hour before surgery for supporting energetic deficit of guinea pigs during surgery. All surgical processes were performed using aseptic technique. Anesthesia consisted of combination of acepromazine (1 mg/kg) and ketamine HCL 10% (40 mg/kg) administrated intramuscularly to maintain deep anesthesia.

Left flanks of the animals were shaved and prepared aseptically. By placing the surgical drape, surgical site was exposed. Skin excision was made as 2×2cm surgical wounds in both groups by 15 no scalpel blade. ADM and normal skin were placed on the wounds and grafts sutured by simple interrupted pattern using a 4-0 nylon suture. The wounds were dressed with cotton gauze and fine cover bandage. Enrofloxacine 0.1 mg/kg, sc. and meloxicam 0.2 mg/kg orally were administrated as antibiotic and NSAID to all animals for 3 days after surgery, respectively.

**Preparation of ADM**

Fresh ostrich skin was obtained from the cadaver. After complete cleaning and excision of subdermal fat tissues, the skin was used for preparation of ADM. The preparation of ADM was consisted of washing the ostrich skin with PBS (Phosphate-buffered saline) then the skin was kept in DW (Distilled Water) for 24 h and then in Triton X-100 at 4°C temperature for 24 h. After that the proceed skins were washed with DW and then skins were kept in SDS (Sodium Dodecyl Sulfate) for 15 h and then the skins were washed with DW and then were kept in SDS for 10 h. Finally the skins were washed with PBS. The cocktail of antibiotic was used after acellular protocol to preclude microbial growth.

**Scanning Electron Microscopy (SEM)**

Normal and ADM samples were treated with routine protocols for preparing SEM examinations and samples were loaded on to aluminum studs and coated with gold. Collagen morphologies were examined under a scanning electron microscope (Philips XL30, Netherlands). Samples were scanned, and the micrographs were recorded. Comparisons were made of morphological changes to collagen fibers before and after ADM protocols.

**Biochemical Assessment of Normal and ADM Samples**

Collagen and sGAG content of the tissues were quantified using the Sircol and Blyscan assay kits, respectively. Total collagen and sGAG of all tissues were quantified and expressed as μg/mg wet skin tissue.

**Gross and Histologic Measurement and Assessment**

Wound characteristics were measured grossly on day 26 after surgery. Direct measurements of wounds area and wound contractions were determined from photographs taken intermittently. All healed wound in each group on day 3, 7 and 26 post surgery were biopsied and then dehydrated with an increasing series of alcohol dehydration and after completing the process, embedded in the paraffin finally. Paraffin-embedded samples were sectioned at a thickness of 6 μm. After H&E staining, samples were examined by light microscopy. A blinded pathologist evaluated all samples histologically for epithelization, fibroblast infiltration and degree of inflammatory cell infiltration, collagen generation and new vessels production. The DNA in samples were assessed by 4',6-diamidino-2-phenylindole (DAPI) staining, and then observed under light microscope.

**Data Analysis and Statistics**

Data were analyzed statistically by ANOVA and Tukey test, the Kruskal-wallis and Mann Whitney U tests. All
analysis was performed using SPSS software (Version 18.0). P value was considered for statistically significance.

The present study has been approved by the Animal Ethics Committee of the Iranian Laboratory animal ethic frameworks under the reference code IAEC 3-12/03.

RESULTS

There is not clinical difference in appearance of ADM and normal skin such as color, texture and size.

**Histologic Observation of Normal and ADM Samples**

Unprepared sample showed presence of epidermal layer in H&E and massive nucleus in ECM in DAPI staining. Decellularized sample showed no epidermal layer in H&E and few positive staining of residual nuclear materials in DAPI staining (Fig. 1).

**Scanning Electron Microscopy (SEM) Evaluation**

The histologic evaluation using SEM examination of normal samples structure showed a dense and integral matrix. After decellularization and underwent a period of processing with Triton, PBS and SDS, the structural pattern of collagen fiber still remained without microscopic changes (Fig. 2).

**Biochemical Assessment of Normal and ADM Samples**

The decrease of collagen content per wet weight of decellularized skin (42.11±2.5 µg/mg) was significant \((P<0.05)\) in compare with native tissue (57.86±4.7 µg/mg), as well as in compare with native (1.8±0.22 µg/mg), sGAG content of decellular skin (1.03±0.05 µg/mg) was significantly decrease (Table 1).

**Gross Evaluation of Wound Contracture**

Dimensions of the implanted graft and contracture show at day 26 after surgery for each group (Table 2). Decreased wound contraction assessed by original wound area. Graft contraction in both groups manifested no statistical difference macroscopically at the end point of study \((P>0.05)\).
Experimental Study on Ostrich ...

Histologic Evaluation of Wound in Days of Sampling

Both wounds in groups had infiltration of with fibroblast and vascularized by 3 days after surgery. Increasing in infiltration of fibroblast and neovascularization were evident in wound bed under collagen scaffold. No significant difference was shown by day 7 after surgery in infiltration of fibroblast between groups \( (P=0.06) \).

Histologically, inflammation was seen in the wound samples of both groups. Comparing ADM and normal implanted skin show no significant difference between the groups \( (P=0.031) \). However extensive inflammation involving the entire biopsy specimen was observed in wound implanted with ADM and normal samples at endpoint of study. Histologic epithelialization in both groups was seen but the quantification of neo-dermis of ADM samples was thicker than normal graft (Fig. 3). There is a significant difference \( (P<0.034) \) between groups in collagen bundles at endpoint of study (Table 3).

**DISCUSSION**

In extensive deep burns and the other full-thickness skin wounds, permanent replacement of lost skin remains a major challenge. The different methods of wound closure are in use, and each has its own advantages and disadvantages. Porcine skin and preserved cadaver skin are
used for temporary wound coverage, but 1 week to 2 weeks after grafting, these tissues undergone immune-mediated rejection. Permanent wound coverage is usually accomplished using meshed split-thickness autograft harvested from undamaged regions of skin. The extensively burned patients have limited donor source, so thin split-thickness autografts are harvested repeatedly from the same sites. This results in substantial donor-site problems resulting from pain, infection, scarring and some keloid formation. Very thin meshed autografts can be used, but the lack of sufficient dermal bed often results in extensive wound contraction at the recipient site. Therefore, deepidermized xenograft porcine or human skin which cell components have been extracted and removed, as a dermal substitute (Acellular Dermal Matrix) has been used with the least immuno-antigenicity and alleviate some of these problems and make ADM available for use in surgical procedure [2].

Human cadaver is expensive, limited and have economic problems, hence, available tissue are restricted as allograft. Also in islamic countries porcine products are forbidden so its skin is not used currently as xenograft. Nowadays ostrich farms spread and its products such as meat and leather are used widely in most countries. Therefore the use of xenogenic skin of ostrich as a new and wide source of ADM by alleviating the problems make it more readily available for use in surgical procedure in skin deficit. Several previous studies have suggested using different materials for producing acellular dermal matrix of porcine or human skin. As Livesey et al.[8] extracted most or all cellular component from porcine, Srivasta et al.[2] used Dispase and Triton to produce acellular porcine skin. Mizuno et al.[27] used human skin for producing acellular matrix by using SDS (Sodium dodecyl sulfate) and hypertonic NaCl. Troung et al.[8] produced acellular human skin by using Dispase and Triton. In recent years Zuo et al.[8] in an experimental study, used EDTA-Na (Ethylenediaminetetraacetic acid), Triton and Trypsin for decellularizing rat skin. In this study, we prepared xenogenic ostrich ADM from fresh ostrich skin using SDS (Sodium dodecyl sulfate), PBS (Phosphate buffered saline) and Triton, used as xenograft in guinea pigs.

As previously mentioned, at the end point of study, wound contracture did not show significant different in two groups, that the findings agreed with Srivaste et al.[2] and Heo et al.[9] studies, which observed no significant contracture in wounds implanted with grafts. Also Walden et al.[10] study showed the dermal grafting was prevented contracture of wound. This data was confirmed our results. Absence of prepared dermal matrix causes the fibroblast initially synthesizes an immature matrix and subsequently remodeled to form a hypertrophic scar or scar contracture [8]. Although decellularizing removes most or all of cellular components, leaves structural and functional molecules such as collagen and sulfated glycosaminoglycan (sGAG), which facilitate the communication of the adjacent cells and with external environment. GAG in ECM have special impact on delaying wound contraction and inducing regeneration [8].

Srivasta et al.[2] and Hoyama et al.[1] were observed infiltration of inflammatory cells around grafts. Srivasta et al.[2] expressed the cause of infiltration of inflammatory cells was the presence of type IV collagen, laminin, some fibronectin, and glycosaminoglycans remains in ADM, albeit in reduced amounts compared with normal skin, it seems unlikely the source of antigenicity. Histological results of presence of inflammatory cells in our study agreed with Srivasta et al.[2] and Hoyama et al.[1] studies.

Troung et al.[5] by using human acellular matrix as xenograft in rat evaluated epithelization in wound bed, also Heo et al.[9] with the aim of evaluating healing potential of different shape of ADM and comparing of the epitelization, defined that although all full-thickness wounds implanted with/without acellular dermal matrix produced epithelialized healed wound by day 28 after surgery that the quantity of wound healing was different between the groups. Our study showed significant difference in epithelization formation between two groups. Early research with acellular dermal tissues was based on theory that if in presence of an appropriate matrix, normal cells of normal adjacent tissue would migrate towards the matrix and performing their functions as if they were in normal tissues. The other studies revealed these expectations, showing the presence of normal fibroblast, collagen and elastin with normal maturity and orientation [1]. Livessey et al. by using porcine acellular dermal matrix found that ADM attached well with the wound bed and promote the ingrowth of fibroblast on the surrounding normal tissue, also after 1 week new blood vessels were formed as a permanent dermal replacement [8]. We found that proliferation of fibroblast and formation new vessels are migrated under ostrich acellular matrix.

In conclusion, epithelization on wound bed of group implanted with acellular ostrich skin demonstrate that ostrich acellular dermal matrix as a temporary substitute may have optimal results in reconstruction of epithelial on wound bed under acellular scaffold.

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Experimental Study on Ostrich ...


