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In vitro assessment of a novel, hypothermically stored amniotic membrane for use in a chronic wound environment

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Amnion; Chronic wound healing; Chronic wounds; Hypothermically stored amniotic membrane; Regenerative healing

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Abstract
Chronic wounds require extensive healing time and place patients at risk of infection and amputation. Recently, a fresh hypothermically stored amniotic membrane (HSAM) was developed and has subsequently shown promise in its ability to effectively heal chronic wounds. The purpose of this study is to investigate the mechanisms of action that contribute to wound-healing responses observed with HSAM. A proteomic analysis was conducted on HSAM, measuring 25 growth factors specific to wound healing within the grafts. The rate of release of these cytokines from HSAMs was also measured. To model the effect of these cytokines and their role in wound healing, proliferation and migration assays with human fibroblasts and keratinocytes were conducted, along with tube formation assays measuring angiogenesis using media conditioned from HSAM. Additionally, the cell–matrix interactions between fibroblasts and HSAM were investigated. Conditioned media from HSAM significantly increased both fibroblast and keratinocyte proliferation and migration and induced more robust tube formation in angiogenesis assays. Fibroblasts cultured on HSAMs were found to migrate into and deposit matrix molecules within the HSAM graft. These collective results suggest that HSAM positively affects various critical pathways in chronic wound healing, lending further support to promising qualitative results seen clinically and providing further validation for ongoing clinical trials.

Introduction
Diabetic foot ulcers (DFUs) are a major comorbidity of diabetes, often resulting from poorly controlled hyperglycaemia. There are 29.1 million type II diabetics in the USA (9.3% of the total population) (1), and with DFUs at a lifetime incidence rate of 15–25% (2), this complication accounts for health care costs in excess of 9–13 billion dollars annually (3,4). In many cases, DFUs lead to amputations, which are associated with mortality rates between 30% and 50% at 1 year (5). DFUs are not only difficult to close, owing to the complex diabetic wound environment, but they are also expensive to treat, with the cost of treating a DFU estimated to run in excess of $18 000 per case (6). Improving healing rates and tissue quality of regenerated tissue in patients with these chronic wounds would not only improve patient quality of life but also reduce the overall burden to the health care system.

Under normal conditions, wound healing proceeds through four specifically timed phases: haemostasis, inflammation, repair and remodelling (7). Chronic wounds develop when the healing process becomes stalled, most often in the inflammatory stage, and fails to progress though the subsequent phases.
of healing (8). Chronic wounds are characterised by a prolonged expression of high levels of pro-inflammatory cytokines, including tissue necrosis factor-alpha (TNF-α), interleukin-6 (IL-6) and interleukin-1 beta (IL-1β). This imbalance in inflammatory cytokines then leads to a prolonged recruitment period for macrophages, neutrophils and mast cells at the wound site. These cells, in turn, produce high levels of reactive oxidative species (ROS) and proteases.

Of the proteases produced within a chronic wound, matrix metalloproteinases (MMPs) are of interest because of their multifunctional role. MMPs are an integral part of the normal wound-healing process, serving to function in several capacities including eliminating damaged proteins, destroying the provisional extracellular matrix (ECM) and facilitating cell migration and tissue remodelling (9). However, in the chronic wound environment, the influx of cells and the increase in pro-inflammatory cytokines leads to elevated and prolonged expression of MMPs and down-regulated expression of MMP inhibitors [tissue inhibitors of matrix metalloproteinases (TIMPs)]. Many studies have shown that levels of MMPs are higher in the exudates of chronic wounds than in those of acute wounds, pointing to a potential mechanism underlying the chronic nature of DFUs (10). In a study by Muller et al. (9) evaluating MMPs longitudinally in DFUs, it was found that despite similar initial levels of MMPs, good healers were characterised by a spike in MMP-1 at 2 weeks and a marked decrease in MMPs by week 4 (compared to no significant changes in MMP levels over time in poor healers). These studies point to the careful regulation of these proteases as an essential component to the successful healing of chronic wounds as increased levels of proteases collectively cause an excessive break down of ECM, growth factors and growth factor-specific receptors (8,11).

A key consequence of the destruction of growth factors and their receptors is a drop in the mitogenic activity of cells and a decrease in the recruitment of additional cells to the wound bed (12,13). Fibroblasts, which are known to be the building blocks of dermal tissue, are one example of cells affected by the chronic wound environment (14). Fibroblasts from diabetic patients have been shown to have a decreased responsiveness to growth factors (15), resulting in a lower proliferative response and deficiency in reorganising the ECM. The chronic wound environment also detrimentally affects keratinocytes. Comparisons of keratinocytes obtained from healthy skin and DFUs have indicated that there are significant changes in many of the signalling pathways associated with apoptosis, migration and proliferation (16). Keratinocytes analysed along the margins of DFUs have been found to be hyper-proliferative and lacking in their ability to migrate and completely differentiate (17–19).

The cellular processes involved in effective wound healing require a significant amount of oxygen, and this oxygen is received through normal blood flow. In many chronic wounds, the oxygen levels needed to proceed with efficient wound healing are unavailable because of underlying ischaemic conditions. Ischaemic conditions are often a consequence of diabetic vascular complications that can affect the blood flow to the wound site (20) and cause deficiencies in blood vessel formation in peripheral tissues (21). Another consequence of the proteolytic environment within chronic wounds is the inhibition of angiogenesis. This occurs through the destruction of ECM proteins and cell adhesion molecules necessary for blood vessel formation and the destruction of angiogenic growth factors (22), consequently affecting the delivery of oxygen to the wound microenvironment, impeding normal healing (23).

Clinically, there are many types of advanced dressings and biological products used to treat difficult-to-heal DFUs. One category that has grown in utilisation recently is cellular tissue products, specifically those derived from placental tissues. These membranes in their native configuration serve to surround and protect the foetus; placental-derived grafts consist of one or more layers of amnion and/or chorion. With consent from the mother, these membranes are collected post-delivery of the full-term, healthy baby. These placental-derived allografts are especially suited for chronic wound healing because they are known to naturally address many of the contributing factors of chronic wound development, including suppressing dysregulated/uncontrolled inflammatory responses, increasing levels of MMP inhibitors in the wound environment, stimulating proliferation and migration of important cell types and promoting angiogenesis (24–26). Current processing technologies focus on dehydration or cryopreservation of placental-derived membranes for clinical use; the goal of this paper is to evaluate wound-healing responses elicited using a novel hypothermically stored amniotic membrane (HSAM).

**Methods**

HSAM (Affinity®, NuTech Medical®, Birmingham, AL) is a hypothermically stored allograft composed of amniotic membrane derived from human placenta. Placentas were donated with informed consent after planned caesarean sections, and all processing was completed in accordance with the Food and Drug Administration’s (FDA) Good Tissue Practices and the American Association of Tissue Banks’ standards. All donors were tested to check if they were free of infectious diseases, including human immunodeficiency virus, human T-lymphotropic virus I/II, hepatitis B and C and syphilis. HSAM is aseptically processed and stored in a proprietary hypothermic storage solution using the AlloFresh™ process.

In order to determine the growth factor content present within HSAMs, multiple 1 cm² samples from nine donors were assessed for a variety of cytokines and growth factors by utilising a quantitative multiplex enzyme-linked immunosorbent assay (ELISA) proteomics microarray (RayBiotech Inc., Norcross, GA). Growth factors evaluated in this study are thought to be relevant to wound healing and have previously been identified within placental-derived tissue (25,26). For the purposes of this evaluation, cytokines have been categorised into general functional areas (detailed in Table 1).

For the proteomics array, HSAM grafts were selected from nine donors unless otherwise specified. Grafts were washed in saline and then homogenised using a Retsch cryomill (Verder Scientific Inc., Newtown, PA). After cryomilling, the tissue was incubated overnight in total protein extraction buffer with a protease inhibitor cocktail (EMD Millipore, Billerica, MA) at 4°C with agitation. Following overnight incubation, the supernatant was removed and loaded into the microarray chambers as per the manufacturer’s instructions. The slides were imaged using a
Assay media conditioned with HSAM. Conditioned media (CM) (Lonza, Walkersville, MD) were cultured in the presence of commercially available adult human fibroblasts or keratinocytes using methods described above. HSAM grafts were then evaluated using a proteomics array.

Table 1 Relevant cytokine categories for wound healing

| Category          | Cytokine                                                                 |
|-------------------|--------------------------------------------------------------------------|
| Angiogenic        | Transforming growth factor beta 1 (TGF-β1)                                |
|                   | Transforming growth factor beta 2 (TGF-β2)                                |
|                   | Insulin-like growth factor-1 (IGF-1)                                     |
|                   | Insulin-like growth factor-2 (IGF-2)                                     |
|                   | Angiopoietin alpha (TGF-α)                                               |
|                   | Tissue Inhibitor of metalloproteinase 1 (TIMP-1)                        |
|                   | Tissue Inhibitor of metalloproteinase 2 (TIMP-2)                        |
|                   | Interleukin 1 receptor antagonist (IL-1ra)                               |
|                   | Interleukin 10 (IL-10)                                                   |
| Regenerative      | Angiopoietin (ANG)                                                       |
|                   | Thrombospondin 1 (TSP-1)                                                 |
|                   | Angiopoietin-2 (ANG-2)                                                   |
|                   | Placental growth factor (PIGF)                                           |
|                   | Angiopoietin-like 4 (APL4)                                               |
| Anti-inflammatory  | Transforming growth factor beta 3 (TGF-β3)                               |
|                   | Insulin-like growth factor-binding protein 1 (IGFBP-1)                  |
|                   | Insulin-like growth factor-binding protein 5 (IGFBP-5)                  |

GenePix 4000B microarray scanner (Molecular Devices, Sunnyvale, CA), and scanned images were imported and analysed using GenePix Pro 7 software (Molecular Devices, Sunnyvale, CA).

To investigate the release of growth factors from HSAM grafts over time, samples were cut into 1 × 1.5 cm² sections and placed in a 48-well plate in a serum-free medium. The HSAM samples were cultured under standard conditions for 7 days with gentle rocking. At 4, 12, 24, 48, 72, 96, 120 and 168 hours, the supernatant was collected, sterile filtered and stored at −80°C. At the end of the experiment, cultured HSAM grafts were collected and also stored at −80°C. Culture media and HSAM grafts were then evaluated using a proteomics array using methods described above.

To evaluate the effect of HSAM on cell proliferation, commercially available adult human fibroblasts or keratinocytes (Lonza, Walkersville, MD) were cultured in the presence of assay media conditioned with HSAM. Conditioned media (CM) was obtained by incubating HSAM in assay media for 5 days at 4°C at a concentration of 1 cm² HSAM membrane per millilitre (ml) of assay media. Following incubation, assay media was filter sterilised prior to use. Media was stored at 4°C for up to 14 days. Cells were then seeded into 48-well plates at a concentration of 3300 and 10,000 cells per well for fibroblasts and keratinocytes, respectively. CM was added to assay media at concentrations of 50%, 25% and 10% (v/v) at 1, 4, 7 and 10 days of culture immediately following evaluation with AlamarBlue. AlamarBlue assays were conducted according to manufacturer’s instructions (Invitrogen, Carlsbad, CA) by incubating cells in 350 μl of a 10% AlamarBlue working solution in Dulbecco’s Modified Eagle Medium with 2.5% fetal bovine serum (FBS) for 4 hours under standard culture conditions. Following incubation, fluorescence was measured in a plate reader from 100 μl samples using a fluorescence excitation wavelength of 540–570 nm and fluorescence emission wavelength of 580–610 nm.

Cell migration assays were conducted with adult human dermal fibroblasts, keratinocytes and human umbilical cord endothelial cells (HUVECs) using a standard Boyden chamber assay (27). Three lots of fibroblasts were used, one from ZenBio (ZenBio Inc., Research Triangle Park, NC) and two from Lonza (Lonza, Walkersville, MD). HUVECs were purchased from Lonza and Life Technologies (Life Technologies, Carlsbad, CA). Keratinocytes were purchased from Lonza. Prior to the start of the assay, cells at less than or equal to 80% confluence were serum-starved overnight. Reservoirs were loaded with assay media alone (negative control), assay media + 10% FBS (positive control) or CM from HSAM at concentrations of 50%, 25% and 10% (v/v). Prior to adding cells to the migration chambers, both the cell inserts and reservoirs were coated with 5 μg/ml Fibronectin (ThermoFisher, Waltham, MA) overnight at room temperature to promote initial cell attachment. Inserts and reservoirs were then washed thrice with PBS. Next, cells were trypsinised and added to the top of the inserts at a concentration of 10,000 (for fibroblasts) or 20,000 (for HUVECs and keratinocytes) cells per insert and then incubated for 24 hours to allow for migration. Non-migrating cells remaining at the top of the cell inserts were removed with a cotton tip applicator, and cells that had migrated to the bottom of the insert were fixed with a 4% solution of neutral buffered formalin in PBS for 10 minutes prior to staining with a crystal violet solution. Images of the inserts were taken with an inverted microscope (Nikon Eclipse Ti, Tokyo, Japan), and representative images were used to count the number of migrated cells.

Tube formation assays were conducted using the angiogenesis starter kit (Life Technologies) as per the manufacturer’s instructions. Two lots of HUVECs (passages 3–5) were used in these experiments (Life Technologies) (28,29). HUVECs were expanded in Medium 200 with large vessel endothelial supplement (Life Technologies) and were passaged at least once after thawing and prior to beginning the tube formation assay. Once cells had reached 80% confluency, HUVECs were seeded into 48-well plates with a reduced growth factor basement membrane matrix (Geltrex, Life Technologies) at a concentration of 25,000 cells/cm². Cells were then cultured overnight with either CM from HSAM or assay media alone (endothelial basal media EBM). CM were obtained as described in detail above. After overnight culture, cells were fixed in methanol for 1 minute, rinsed thrice in distilled water and imaged via phase-contrast with an inverted microscope (Nikon Eclipse Ti). Images were then imported into ImageJ (NIH, Bethesda, MD), and the average tube length was quantified.

For all quantitative mechanistic assays (proliferation, migration and angiogenesis), statistical analysis was conducted using a one-way analysis of variance ANOVA with a post-hoc Tukey test, where P < 0.05 was considered significant. Throughout, * denotes P < 0.05, ** denotes P < 0.01, *** denotes P < 0.001, and **** denotes P < 0.0001. For qualitative imaging, representative images of all groups were taken and presented.
To qualitatively analyse how fibroblasts interacted with HSAM, 50,000 human dermal fibroblasts were seeded onto HSAM grafts (2.5 cm × 2.5 cm). Grafts were then cultured in six-well plates for 2 weeks under standard culture conditions. Post the 2-week culture, grafts were fixed in 4% neutral buffered formalin for 24 hours. Following paraffin embedding, 5-μm thick serial sections were cut from the tissue blocks and floated onto charged glass slides (Super-Frost Plus, Fisher Scientific, Pittsburgh, PA) and dried overnight at 60°C. Sections were stained for haematoxylin and eosin (H&E), Masson’s trichrome, Verhoeff’s stain and Alcian blue. All sections for immunohistochemistry were deparaffinised and hydrated using graded concentrations of ethanol to deionised water. For fibronectin immunohistochemistry, antigen retrieval was performed using a solution of proteinase K (P6556, 1:50 dilution in TE-CaCl₂ buffer pH 8) for 15 minutes at 37°C. For collagen I and III immunohistochemistry, tissues sections were subjected to antigen retrieval using a 0.01 M Tris-1 mM EDTA buffer (pH 9) at 70°C for 20 minutes. Following antigen retrieval, all sections were washed gently in deionised water, then transferred to a 0.05-M Tris-based solution (TBST) in 0.15 M NaCl with 0.1% v/v Triton-X-100, pH 7.6 (TBST). Endogenous peroxidase was blocked with 3% hydrogen peroxide for 20 minutes. To reduce further non-specific background staining, slides were incubated with 3% normal goat serum for 20 minutes. To reduce further non-specific background staining, slides were incubated with 3% normal goat serum for 20 minutes. To reduce further non-specific background staining, slides were incubated with 3% normal goat serum for 20 minutes.

Results

Proteomic microarrays confirmed physiologically relevant levels of numerous growth factors and cytokines important for wound healing (Table 1). These growth factors were broken down using general categories to describe their activity, including angiogenic, regenerative and anti-inflammatory growth factors (Figure 1; average ± standard deviation, n = 9 unless otherwise noted). There is some sample-to-sample variability observed in the levels of growth factors and cytokines present in the tissue. This level of variability is expected and has been observed in previous studies of dehydrated human amnion-chorion membranes (dHAMC) (25,30) and of cryopreserved human amniotic membrane (cHAM) (31–33). Many of the same growth factors are found to be present in cHAM, dHAMC and HSAM. However, differences have been noted that may be caused by several factors, including degradation of growth factors or cytokines occurring as a result of tissue processing (dehydration, lyophilisation and sterilisation) and the layers included in the product (amnion versus amnion and chorion), as well as study-related factors such as protein extraction methods, methods for evaluation of growth factors and sample size. The most abundant growth factors present in the HSAM were thrombospondin-1 (TSP-1), angiopoietin-like 4 (APL4) and insulin-like growth factor-1 (IGF-1) in nearly all of the samples that were tested, suggesting that while specific levels of growth factors in the tissues may vary to some degree, relative concentrations of these most abundant factors are consistent.

Subsequently, we were interested in evaluating the time course for the release of growth factors from HSAMs over a 7-day period. All growth factors analysed in Figure 1 were included in this analysis, and the total growth factor and cytokine release over 7 days was approximately 70.6 ± 19.9% of the total measured cytokines (Figure 2A). Key growth factors, including basic fibroblast growth factor (bFGF), hepatocyte growth factor (HGF), interleukin 1 receptor antagonist (IL-1ra), transforming growth factor beta 3 (TGF-β3) and TIMP-1/2, were evaluated and graphed individually (Figure 2B–F). After 7 days, 91.7 ± 5.6% (n = 3) of the...
Figure 2 Release of growth factors from hypothermically stored amniotic membrane (A) cumulative release curve for all 25 growth factors listed in Table 1, (B) basic fibroblast growth factor (bFGF), (C) hepatocyte growth factor (HGF), (D) interleukin 1 receptor antagonist (IL-1ra), (E) transforming growth factor beta 3 (TGF-β3) and (F) tissue inhibitor of metalloproteinases 1 and 2 (TIMP-1/2). Cytokine concentrations are shown as average ± standard deviation for all time points, and total % average ± standard deviation released at 7 days is inset for all release curves (n=3).
bFGF had been released; bFGF is widely known to accelerate wound healing, primarily through its mitogenic effects on cells (34) (Figure 2B). HGF is particularly important to the wound-healing environment because of its role in stimulating proliferation and cell motility in cells of epithelial origin (35,36); in our study, 89.9 ± 4.5% (n = 3) of HGF was released from HSAM after 7 days (Figure 2C). IL-1ra is a natural inhibitor of pro-inflammatory IL-1β and is thought to also play a role in reepithelialisation. Of interest, delayed healing has been observed in IL-1ra-deficient mice as evidenced by problems with collagen deposition and vascularisation (37). Over the course of 7 days, approximately 97.8 ± 3.7% (n = 3) of TGF-β3 in HSAM had been released (Figure 2E). TGF-β3 is thought to assist in wound healing by reducing fibrosis and scarring at the wound site (38,39).

Another important component of the wound-healing process is appropriate regulation of protease activity. Over-expression...
Assessment of a novel amniotic membrane

Figure 4 Evaluation of hypothermically stored amniotic membrane (HSAM) effects on human epidermal keratinocytes (HEKs).

(A) Quantitative analysis of proliferation of HEKs cultured under various conditions, including growth media, assay media and HSAM conditioned media (50%, 25% and 10%). Migration of HEKs in response to growth media, assay media and 50%, 25% and 10% HSAM CM evaluated quantitatively (B) and qualitatively with crystal violet staining (C). For quantitative assays, significance compared to assay media is denoted by \(* P < 0.05, ** P < 0.01, *** P < 0.001\) and \(**** P < 0.0001\).

of MMPs is thought to be one of the causative factors for chronic wounds; therefore, inhibitors of MMPs (TIMPs) and their availability are of particular interest (40). We found that the majority of TIMP-2 (93.6 ± 4.3%, \(n = 3\)) was released from the membrane by day 7; we also found that TIMP-1 was released from the membrane. However, only 10.7 ± 5.4% \((n = 3)\) of the total quantity of TIMP-1 was released by day 7. The low release rate of TIMP-1 compared to TIMP-2 may be, in part, explained by the interaction of TIMP-1 with the β1 integrin (41). Because of its interaction with the ECM, the release profile of TIMP-1 is particularly likely to be much different in vivo because of interactions of the graft with the chronic wound environment and subsequent break down of the ECM.

In addition to studying the growth factor/cytokine profile and their respective release characteristics, we also investigated the effects released cytokines have on cell types important to wound healing. Specifically, we investigated the effects of cytokines released from HSAM on the behaviour of human dermal fibroblasts, human keratinocytes and human endothelial cells. Fibroblast migration and proliferation is thought to be an indicator of healing in wounds (7,42,43). In Figure 3A and B, it was found that fibroblasts proliferate more quickly in media
conditioned from HSAM than in assay media alone, which represents the baseline control. Quantitatively, it was found that 50%, 25% and 10% CM significantly increased fibroblast proliferation at 7, 10 and 14 days, although there were no measurable differences between the CM groups of different concentrations. Qualitative images were taken at 14 days, and these images confirmed that HSAM CM groups had a higher density of cells present than the assay media group (Figure 3B).

Additionally, the migration of fibroblasts through a micro-porous membrane was evaluated. Fibroblast migration assays showed that the presence of HSAM CM resulted in a significant increase in migration relative to assay media alone. Interestingly, the dose of CM (50%, 25% or 10%) did not affect the increase in fibroblast migration, with all groups showing between a 0.91- and 0.95-fold increase over assay media alone. Crystal violet-stained cells qualitatively confirmed greater migration in groups with HSAM CM than in assay media alone (Figure 3D).

Human epidermal keratinocytes (HEKs) are essential to wound closure; in the normal healing process, HEKs proliferate and migrate into the wound and form an epithelial layer (44–46). To model this in vitro, we assessed the effect HSAM CM would have on HEK proliferation and migration (Figure 4). After 7 days, all the CM groups evaluated (10%, 25% and 50% HSAM CM) resulted in significantly increased cell proliferation relative to assay media alone. Migration of keratinocytes is considered an important step in wound healing as they play an integral role in reepithelialisation of the wound bed (47). To model this effect in vitro, HEK migration was evaluated using a Boyden chamber assay (Figure 4B). It was found that 50% and 25% HSAM CM stimulated significantly increased cell migration compared to assay media, 340% and 232%, respectively. Qualitatively, these changes were confirmed with the crystal violet staining of cells attached to the Boyden chamber membrane (Figure 4C).

Angiogenesis, or the formation of new blood vessels, is essential for wound healing (48). However, this process is often disrupted in chronic wounds. In this study, HSAM CM or assay media was applied to a monolayer of HUVECs, and the formation of tubules was assessed (Figure 5). HSAM CM at concentrations of 50% and 25% significantly increased average tubule length by 104% and 95%, respectively. HSAM CM of 10% and 1% showed no appreciable increases in tubule length. Representative phase-contrast images are shown in Figure 5B for each of the groups analysed, with qualitative increases in tube formation shown in higher concentrations of HSAM CM.

To model the interaction of fibroblasts with HSAM, we cultured fibroblasts on HSAM for up to 2 weeks and subsequently evaluated the tissue using H&E, Masson’s trichrome, Verhoeff’s stain and Alcian blue (Figure 6). Immunohistochemistry was also conducted for collagen I, III and fibronectin (Figure 7). Images show a series of histological results for HSAM with (Figures 6B, D, F, H and 7B, D, F) and without (Figures 6A, C, E, G and 7A, C, E) fibroblasts seeded for up to 2 weeks. Based on the H&E and Masson’s trichrome images, it is clear that fibroblasts readily infiltrate and proliferate within the HSAM grafts. Masson’s trichrome and Verhoeff’s staining showed deposition of collagen and elastin into the HSAM by fibroblasts. While HSAM contains a high quantity of glycosaminoglycans, Alcian blue images do not indicate the deposition of additional glycosaminoglycans by fibroblasts. HSAM was found to contain high amounts of collagen I throughout stromal layer, collagen III specifically localised to the basement membrane layer and the spongy layer and fibronectin within the basement membrane layer and around cells. Fibroblasts cultured on HSAM deposited additional collagen III and Fibronectin throughout the HSAM. While we also expected to see fibroblasts-producing collagen I, the high levels of native collagen I found throughout
Figure 6 Qualitative evaluation of fibroblast interaction with hypothermically stored amniotic membrane (HSAM) was completed using staining. HSAM was cultured with or without fibroblasts for 2 weeks, and samples were processed and stained with haematoxylin and eosin (H&E), Masson's trichrome, Alcian blue, or Verhoeff's stains. Representative images were taken of H&E (A and B), Masson's trichrome (C and D), Alcian blue (E and F) and Verhoeff's stain (G and H).

Discussion

The use of placental-based tissues is proposed to aid in healing through various mechanisms, including reduction in inflammation, enhancement of cell migration into the wound environment, stimulation of cell proliferation, increased production of ECM and improved angiogenesis (25,49–51). A significant part of the function of these tissues is attributed to the growth factor content released from the membranes. Selected growth factors and cytokines of interest are shown in Table 1; while this list is not comprehensive of all molecules found in placental-based tissues, these molecules were selected because of the important role they play in wound healing. Some growth factors and cytokine concentrations are lower than in singular recombinant growth factor therapies used for chronic wounds, but it is important to note that there may be a substantial benefit in the sustained delivery of multiple cytokines with different roles/signalling processes in the healing wound (52,53).
The management of inflammation is essential to resolving chronic wounds. Amniotic membranes and the cells within these membranes have been shown to elicit an anti-inflammatory response. Cells derived from amniotic membranes have been shown to down-regulate the production of certain inflammatory cytokines, including TNF-α, CXCL10, CXCL9 and CCL5 (54,55). The suppression of TNF-α is of particular interest as TNF-α has been found to be elevated in non-healing DFUs (56). Amniotic membranes have been shown to suppress the production of transforming growth factor beta 1 (TGF-β1), Interleukin 8 IL-8 and granulocyte macrophage colony-stimulating factor by fibroblasts, which are known to be stimulated under inflammatory conditions (57). Another relevant cytokine known to trigger inflammatory responses within chronic wounds, IL-1β, was found to be significantly reduced in gingival crevicular fluid following the application of amniotic membranes with bone grafting for periodontal pockets in a 30-patient clinical study (58). In a mouse bleomycin injury model, amnion-derived epithelial cells have been shown to limit macrophage infiltration into the lungs and to also drive a majority of pulmonary macrophages to the anti-inflammatory M2 phenotype (59). The shift to M2 polarisation in response to amnion-derived cells is relevant to DFUs as M2 and M2-like macrophages induce anti-inflammatory, regulatory and reparative functions that lead to wound closure (60–62). While the exact mechanisms responsible for these anti-inflammatory effects remain unclear, potential sources include the release of interleukins from the membrane, including both interleukin 10 (IL-10) and IL-1ra, which were measured in HSAM. IL-10 is known to counteract the effects of several pro-inflammatory cytokines, including IL-6, IL-1, IL-8 and TNF-α (63), and IL-1ra is a potent inhibitor of IL-1, shown to suppress the inflammatory response triggered by IL-1 (64). While not measured in this study, we suspect that the presence of heavy chain hyaluronic acid (65), prostaglandin E2 (66), and macrophage migration-inhibitory factor (67,68) may also play a role in the anti-inflammatory effects observed with HSAM.

An essential part of resolving chronic wounds is the reduction of proteases within the wound environment to prevent the destruction of growth factors and ECM proteins necessary for wound healing. MMP-9 has been found to be elevated in non-healing DFUs (56), and the addition of inhibitors to MMPs (TIMPs) to the chronic wound environment has been shown to be beneficial to wound healing (9,25,69). Amniotic membranes...
have been previously shown to inhibit MMP activity (70,71), and here, we have demonstrated the presence and the release curves of TIMP-1 and TIMP-2 within HSAM. In addition to TIMP-1 and 2, possible mechanisms behind the protease inhibition observed with amniotic membranes include contributions by type-I plasminogen activator inhibitor and TSP-1 (24).

The destruction of essential growth factors and their receptors is another consequence of excessive protease levels within chronic wounds. This consequently leads to cellular dysregulation, especially with fibroblasts and keratinocytes. For fibroblasts, a marked decrease in cellular migration, proliferation and ECM deposition has been observed (72,73). Here, we have demonstrated that CM from HSAM significantly improves fibroblast proliferation and migration. Fibroblasts were also shown to migrate into and deposit ECM proteins within HSAMs. The likely cause of this proliferative and migratory response of fibroblasts to CM is from the release of FGF, HSAMs. The likely cause of this proliferative and migratory response is another consequence of excessive protease levels within chronic wounds. Here, we have demonstrated that HSAM significantly improves fibroblast proliferation and migration. Fibroblasts were also shown to migrate into and deposit ECM proteins within HSAMs. The likely cause of this proliferative and migratory response of fibroblasts to CM is from the release of FGF, HSAMs, which have been shown to lead to the activation and migration of fibroblasts and stimulate their proliferation (42). Keratinocytes along the borders of chronic wounds have also been found to demonstrate abnormal behaviour, including failure to differentiate and migrate (17). Here, we have demonstrated that HSAM contains several growth factors known to stimulate keratinocyte migration and proliferation, including epidermal growth factor (EGF), platelet-derived growth factor BB (PDGF-BB) and TGF-β1. More importantly, we have shown that growth factors/cytokines found within CM from HSAM are active and can promote both proliferation and migration of keratinocytes. In a recent study by Zhao et al., human amniotic epithelial cells were found to improve keratinocyte migration and proliferation. This increase in keratinocyte migration and proliferation was found to be primarily the result of EGF and keratinocyte growth factor KGF produced by these cells, which, in turn, activated the extracellular signal-regulated kinases ERK, jun N-terminal kinase JNK and protein kinase B AKT pathways in keratinocytes (74). Fibroblasts and keratinocytes are known to interact in the wound environment (35,75). Suppression of fibroblast proliferation and migration in chronic wounds have been shown to result in changed cytokine and protease release profiles (76,77), and these changes result in deleterious impacts on keratinocyte proliferation and migration (73). We expect the combined effects of HSAM CM on both fibroblasts and keratinocytes to promote significant improvements clinically in chronic wound healing.

Prolonged hypoxia in the wound environment can prevent wounds from healing normally; therefore, angiogenesis is critical to restore oxygen and nutrients to the injured tissue (7). Because many patients with chronic wounds, especially DFUs, have underlying vascular disease, lack of effective angiogenesis can be particularly problematic in healing these patients. In addition to these underlying issues, there are several anti-angiogenic factors present within chronic wounds, including inflammatory factors, proteases and cell death mediators, which deter pro-angiogenic processes (22). We expect that HSAM may mitigate both the prolonged pro-inflammatory response and high protease content within the wound to create an environment more favourable to angiogenesis. HSAM contains numerous angiogenic growth factors, including FGF, vascular endothelial growth factor (VEGF), PDGF-BB, angiopoietin (ANG), placental growth factor (PLGF), APL4 and TSP1. The results from in vitro tube formation assays reflected the activity of high levels of angiogenic growth factors; these results support the hypothesis that the application of HSAMs to chronic wounds may promote angiogenesis resulting in improved wound healing and closure.

Placental-derived membranes have a long history in wound care (78,79); however, most current commercially available membranes are processed in such a way that they do not maintain native viable cell populations. While these membranes have shown positive effects on healing in chronic wound care (51,80), the absence of a viable cell population prevents these grafts from dynamically responding to the chronic wound environment (81). Numerous studies have shown that the various cell types populating the amniotic membrane, which are present within HSAM, produce factors that act to reduce inflammation, improve cell proliferation and migration and improve angiogenesis (54,55,59,66,67). Recent clinical data suggest that cell containing amniotic grafts may be more effective in treating chronic wounds compared to non-viable placental-derived alternatives (82). Future studies will focus on how the cell populations within HSAM respond to chronic wound environments through in vitro models in the context of inflammation and in vivo studies utilising an ischaemic animal model, which more closely mimics the chronic wound environment (83).

Conclusion

HSAMs offer a new treatment alternative to promote healing in chronic wounds; these membranes deliver a native ECM, growth factors and cytokines and a viable cell content. In this study, we have focused on the overall activity of the grafts by primarily using media conditioned by the grafts to study mechanisms of action. We have reported concentrations of numerous growth factors and cytokines known to contribute to wound healing and detailed their released from HSAM over 7 days. Furthermore, we have shown that HSAM CM significantly increases fibroblast and keratinocyte proliferation and migration, suggesting its potential to improve healing in chronic wounds where the normal activity of these cells is known to be suppressed. Finally, this study shows that HSAM promotes increased tube formation in HUVECs suggesting a potential mechanism for HSAM to promote and improve angiogenesis. In sum, we believe hypothermically stored amniotic grafts are a compelling option for promoting pro-healing activities spanning various functions, including reducing inflammation and protease activity as well as promoting angiogenesis and cellular responses.

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