Identification of Extracellular Matrix Components and Biological Factors in Micronized Dehydrated Human Amnion/Chorion Membrane

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Objective: The use of bioactive extracellular matrix (ECM) grafts such as amniotic membranes is an attractive treatment option for enhancing wound repair. In this study, the concentrations, activity, and distribution of matrix components, growth factors, proteases, and inhibitors were evaluated in PURION® Processed, micronized, dehydrated human amnion/chorion membrane (dHACM; MiMedx Group, Inc.).

Approach: ECM components in dHACM tissue were assessed by using immunohistochemical staining, and growth factors, cytokines, proteases, and inhibitors were quantified by using single and multiplex ELISAs. The activities of proteases that were native to the tissue were determined via gelatin zymography and EnzChek® activity assay.

Results: dHACM tissue contained the ECM components collagens I and IV, hyaluronic acid, heparin sulfate proteoglycans, fibronectin, and laminin. In addition, numerous growth factors, cytokines, chemokines, proteases, and protease inhibitors that are known to play a role in the wound-healing process were quantified in dHACM. Though matrix metalloproteinases (MMPs) were present in dHACM tissues, inhibitors of MMPs overwhelmingly outnumbered the MMP enzymes by an overall molar ratio of 28:1. Protease activity assays revealed that the MMPs in the tissue existed primarily either in their latent form or complexed with inhibitors.

Innovation: This is the first study to characterize components that function in wound healing, including inhibitor and protease content and activity, in micronized dHACM.

Conclusion: A variety of matrix components and growth factors, as well as proteases and their inhibitors, were identified in micronized dHACM, providing a better understanding of how micronized dHACM tissue can be used to effectively promote wound repair.

Keywords: amniotic membrane, wound healing, MMP, protease inhibitors

INTRODUCTION

The treatment of large, chronic wounds (e.g., foot ulcers, venous leg ulcers) is challenging, particularly in diabetic patient populations. Approximately one-quarter of diabetic patients will develop a chronic wound in his or her lifetime, which, if not

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managed successfully, may require amputation. In fact, 81% of amputations in diabetic patients have been attributed to inadequate wound healing.1

Physiological wound healing is a well-coordinated sequence of events in which signaling and remodeling of the extracellular matrix (ECM) are integral components. After injury, the wound progresses through hemostasis, inflammation, proliferation, and repair/remodeling, ultimately resulting in the regeneration of functional tissue. In contrast, a chronic wound may result from delayed resolution of the inflammatory phase. Inflammatory cells recruited to the site of injury produce elevated reactive oxygen species and secrete an abundance of proinflammatory cytokines, which, in turn, recruit additional inflammatory cells, resulting in a perpetual cycle of inflammation.2 The combination of excessive proinflammatory cytokines and reactive oxygen species induce upregulation of proteases and matrix metalloproteinases (MMPs) that not only degrade ECM but also alter the activity of growth factors and proteins that are necessary for progression to tissue repair/regeneration.3 A large clinical study involving a variety of complex wounds in a diverse patient population measured elevated MMP levels in 28% of chronic wounds.4 In a prolonged inflammatory state, tissue inhibitors of metalloproteinases (TIMPs) and other inhibitors may not be sufficient to modulate the effects of the disproportionately high concentration of MMPs.

MMPs are a class of proteinases with diverse roles in tissue remodeling. Aside from the well-known role of ECM proteolysis, the downstream functions of MMPs include release, activation, and inactivation of growth factors and cytokines, all of which influence cell–cell and cell–matrix interactions, and ultimately cell proliferation, migration, and differentiation.5 MMP expression is increased during physiological tissue remodeling, including bone turnover and fracture healing, pregnancy, angiogenesis, and wound healing.5,6 However, unregulated MMP expression can result in excessive tissue breakdown and pathological conditions; therefore, inhibitors of MMPs are necessary to modulate MMP activity.7

Successful closure of chronic wounds often requires advanced therapies to resolve the persistent inflammatory state and to repair the lost and/or damaged tissue. However, standard compression bandaging is often inadequate, as noted when progress toward wound closure is not observed after 3–4 weeks of treatment.8 In these patients, treatment with naturally derived, bioactive ECM allografts such as amniotic membrane is an attractive alternative therapy to accelerate repair. During pregnancy, the amniotic sac forms an adjustable biocapsule that supports and protects the developing fetus.9 The complex role of the amniotic sac is facilitated by the unique structure and functionality of the ECM components that are contained within this specialized tissue. The dynamic and biologically active tissue matrix provides tensile strength and integrity, anchors cells, and influences cell morphology and function via interaction with cell surface receptors, and it is a reservoir for growth factors. The amniotic sac is composed of amnion and chorion layers that are constantly remodeling through synthesis and degradation of the ECM. The amnion layer is a thin (35–60 μm) avascular membrane that is composed of epithelium, basement membrane, compact layer, and fibroblast layer. Each contains many ECM macromolecules (e.g., collagen types I and IV, fibronectin, laminin, heparan sulfate proteoglycans [HSPGs], hyaluronic acid [HA], and glycoproteins) that provide mechanical support for the tissue, provide sites for cell attachment and motility, and sequester growth factors.10 The chorion layer ECM is composed of fibronectin, laminin, and types I and IV collagen that support the cells and sequester growth factors/cytokines of the chorion.11 An intermediate spongy layer separates the amnion and the chorion. Throughout pregnancy, the amniotic membrane ECM undergoes physiological remodeling—a highly coordinated process involving the secretion of growth factors, cytokines, and matrix proteins—to maintain structural integrity while allowing for expansion during growth of the fetus. As in response to injury, the balance of MMPs and their inhibitors is crucial for maintenance of the ECM during pregnancy.

The amniotic membrane has a long history of use in wound-healing applications due to the dynamic nature of the tissue and its growth factor/cytokine-rich ECM. In particular, dehydrated human amnion/chorion membrane (dHACM) has demonstrated clinical efficacy in the enhancement of wound repair compared with standard compression therapy.12 dHACM contains an array of growth factors, cytokines, and chemokines, and it has been shown to promote proliferation and migration of bone marrow-derived mesenchymal stem cells, adipose-derived stem cells (both healthy and diabetic), and hematopoietic stem cells.13–17

The treatment of complex, full-thickness wounds may be augmented by the use of a micronized formulation of dHACM (in addition to membrane dHACM) to fill the wound bed and to provide additional bioactive molecules to the site of injury. Moreover, the larger surface area afforded
by the micronized dHACM may facilitate cell–
tissue interactions and growth factor presentation/ 
availability, ultimately promoting processes such 
as angiogenesis, keratinocyte migration, and stem 
cell recruitment. Micronized dHACM has been 
applied directly in powder form to wounds, and it is 
inhaled after rehydration at wound edges. Micron-
ized dHACM was previously used as an adjuvant 
to membrane dHACM in three patients with 
chronic wounds that had not healed after aggressive 
treatment with other wound care strategies. In 
these challenging clinical scenarios, this combina-
tion therapy resulted in successful closure of the 
wounds.15

CLINICAL PROBLEM ADDRESSED

The tissue-healing effects of the amniotic mem-
brane on chronic wounds are believed to be due, at 
least in part, to the inherent capacity of the mem-
brane to coordinate ECM remodeling, an essential 
function not only during pregnancy but also for 
wound healing. The amniotic membrane presents 
specialized ECM components in their native 
structure, and key regulatory molecules in the 
form of growth factors, cytokines, and proteins 
contained therein. This work evaluated the concen-
trations and distribution of vital ECM compo-
nents and growth factors, along with MMPs and 
their inhibitors, in micronized dehydrated human 
amnion/chorion membrane (dHACM), and it serves 
as the first exploration into the structure and 
composition of the micronized form of the tissue. 
Ultimately, an understanding of how amniotic 
membrane therapies such as dHACM are effica-
cious in the treatment of chronic wounds is crucial 
to advancing the standards of wound care.

MATERIALS AND METHODS

dHACM Tissue Processing

Membrane and micronized formulations of dehy-
drated human amnion/chorion membranes (dHACM, 
EpiFix®; MiMedx Group, Inc.) processed by a pro-
prietary PURION® Process19 were used in this study. 
Briefly, amnion and chorion were isolated from hu-
man placentas that were donated under informed 
consent from scheduled Cesarean section births. The 
tissue layers were gently cleansed, laminated, and 
dehydrated. For micronized dHACM, the tissue was 
subsequently cryomilled and sieved by using filters of 
180 and 25 μm pore size.

Immunohistochemistry of dHACM

For ease of maintaining tissue orientation during 
histological sectioning, membrane dHACM 
tissue samples (n=3) were used for immunohisto-
chemistry (IHC). Tissues were hydrated in 0.9% 
NaCl for 10 min, embedded in TissueTek optimum 
cutting temperature compound (Sakura Finetek), 
and cryosectioned at a thickness of 5 μm. Sections 
were fixed by incubation in chilled acetone for 
10 min and dried at room temperature for at least 
30 min. The sections were washed with phosphate-
buffered saline (PBS), blocked, and incubated with 
primary antibodies (mouse anti-human collagen 
IV, 1:100; Abcam, mouse anti-human collagen I, 
1:1,000; sheep anti-human HA, 1:100; Abcam) 
overnight at 4°C. After washing with PBS, sections 
were stained with a secondary antibody (Alexa Fluor® 660 goat anti-mouse; Alexa Fluor 555 don-
key anti-mouse; Alexa Fluor 647 donkey anti-
sheep, respectively; 1:100; Life Technologies) for 
30 min at room temperature and mounted in Fluorogel (Electron Microscopy Sciences).

Micronized dHACM Extraction and Digestion

Micronized dHACM (n=4–7) samples were ex-
tracted in 4 M guanidine hydrochloride (Sigma) 
and 0.1 M sodium acetate (Sigma) containing Pro-
tease Inhibitor Cocktail Set III (Calbiochem; EMD 
Millipore) at 4°C overnight and pH 6.0. Solid ma-
terial was removed by centrifugation and the su-
pernatant was frozen at –20°C, whereas the pellet 
was resuspended in a second volume of extrac-
tion buffer overnight. Supernatants from both ex-
tractions were combined and dialyzed in 50 kDa 
MWCO dialysis tubing (Spectra/Por® 6 Standard 
RC, Spectrum) against deionized water for 2 days 
and, subsequently, frozen and lyophilized. Sam-
ple were analyzed for fibronectin, laminin, and 
HSPGs by using single-factor enzyme-linked im-
unosorbent assays (ELISAs) (fibronectin, R&D 
Systems; laminin, Abcam; HSPGs, Cusabio).

Collagen was extracted from the remaining mi-
cronized dHACM pellets and membrane dHACM 
pelets that had been processed in the same manner (n=3). Pellets were rinsed and acidified before 
an overnight pepsin (Worthington) digestion. Col-
lagen type I was purified from the pepsin digests 
through a 0.7 M salt precipitation. The precipitated 
collagen was frozen, lyophilized, and prepared for 
separation by sodium dodecyl sulfate-polyacrylamide 
gel electrophoresis (SDS-PAGE) (ThermoFisher) by 
addition of a tris-glycine-dithiothreitol sample buffer 
(ThermoFisher) and heating at 100°C for 5 min. The 
volume loaded was based on the measured bulk mass 
after lyophilization. Separated proteins contained 
within the gel were stained with Coomassie Blue 
(ThermoFisher).
To determine levels of HA in micronized dHACM, tissues \((n=4)\) were digested in a 1 mg/mL papain (Worthington) buffer containing 0.2 M sodium acetate, 1.1 mM EDTA, 5.5 mM cysteine hydrochloride, and 0.067 mM \(\beta\)-mercaptoethanol overnight at 60°C. Solutions were boiled and cooled before dialysis in 50 kDa MWCO tubing against deionized water overnight. Solutions were collected, frozen, and lyophilized. HA content in dry samples was evaluated by using a single-factor ELISA (R&D Systems).

**Analyses of Growth Factors, Cytokines, Proteases, and Inhibitors**

Micronized dHACM \((n=4–5)\) was incubated in lysis buffer (RayBiotech) containing Protease Inhibitor Cocktail Set III at a concentration of 35 mg/mL with gentle agitation overnight at 4°C. Lysates were centrifuged and filtered (0.22 \(\mu\)m pore size; EMD Millipore). Growth factors, cytokines, proteases, and protease inhibitors were evaluated by using multiplex ELISA Quantibody® arrays (Human Cytokine Array Q1000 and Human MMP Array Q1, RayBiotech), and single-factor ELISAs (alpha 2-macroglobulin \([\alpha2M]\) and alpha-1 antitrypsin \([\alpha1AT]\); RayBiotech). Amounts of each molecule were reported as average moles/mg of tissue. For each donor, Ten nanograms of recombinant MMP-2 key’s pair was incubated in Tris 2-macroglobulin \([\alpha2M]\) and alpha-1 antitrypsin \([\alpha1AT]\); RayBiotech). Amounts of each molecule were reported as average moles/mg of tissue. For each donor, Ten nanograms of recombinant MMP-2 (R&D Systems) that had been activated for 1 h at 37°C with 4-aminophenylmercuric acetate (APMA) was used as a control, and a Benchmark Pre-stained protein ladder (Novex) was included with each gel. Each sample was loaded into individual wells of a 10% gelatin zymogram gel (Novex) and run at 125 V for 110 min. Gels were collected and incubated in 1 \(\times\) Renaturing buffer (Novex) for 30 min, followed by 1 \(\times\) Developing buffer (Novex) overnight at 37°C. Gels were stained with SimplyBlue™ SafeStain (ThermoFisher) for 30 min and destained in 20% NaCl for 24 h. Images were taken by using an Amersham Imager 600 (GE Healthcare Life Sciences) and analyzed by using ImageQuant TL software. Band densities were quantified and normalized to the recombinant MMP-2 control.

**Detection of MMPs by Gelatin Zymography**

Micronized dHACM \((n=4)\) was incubated in Tris glycine SDS (ThermoFisher) overnight at 4°C with gentle agitation at a concentration of 40 mg/mL. Samples were centrifuged and filtered before dilution to concentrations of 20, 10, and 1 mg/mL for each donor. Ten nanograms of recombinant MMP-2 (R&D Systems) that had been activated for 1 h at 37°C with 4-aminophenylmercuric acetate (APMA) was used as a control, and a Benchmark Pre-stained protein ladder (Novex) was included with each gel. Each sample was loaded into individual wells of a 10% gelatin zymogram gel (Novex) and run at 125 V for 110 min. Gels were collected and incubated in 1 \(\times\) Renaturing buffer (Novex) for 30 min, followed by 1 \(\times\) Developing buffer (Novex) overnight at 37°C. Gels were stained with SimplyBlue™ SafeStain (ThermoFisher) for 30 min and destained in 20% NaCl for 24 h. Images were taken by using an Amersham Imager 600 (GE Healthcare Life Sciences) and analyzed by using ImageQuant TL software. Band densities were quantified and normalized to the recombinant MMP-2 control.

**Gelatinase/Collagenase Activity Assay**

Micronized dHACM \((n=5)\) was incubated in the EnzChek® Gelatinase/Collagenase Activity Assay kit reaction buffer (ThermoFisher) at 40 mg/mL overnight at 4°C with gentle agitation. Extract samples were collected, centrifuged, and filtered before dilution to concentrations of 20, 10, and 1 mg/mL. Extracts were assayed alone and in combination with Clostridium collagenase (ThermoFisher) at 0.0625 U/mL, or with 0.1 mM of the inhibitor 1, 10-phenathroline (ThermoFisher). All samples were incubated with 10 \(\mu\)g/mL DQ gelatin substrate for 45 min and read with a plate reader (Synergy M3; Biotek) at an excitation wavelength of 485 nm and an emission wavelength of 528 nm. Readings from samples with added collagenase were normalized to a Clostridium collagenase control at 0.0625 U/mL. Readings from samples only and samples with added inhibitor were normalized to a control that contained both inhibitor (0.1 mM) and collagenase (0.0625 U/mL).

**Statistical Analysis**

All values were reported as mean ± standard deviation, and statistical analyses were performed in Minitab (v17.1). Zymography data and the comparison of gelatinase/collagenase activity to collagenase controls were evaluated by one-way analysis of variance (ANOVAs). The comparison of gelatinase/collagenase activity to inhibitor + collagenase controls was performed by using a two-way ANOVA with factors of extract concentration and inhibitor condition. For each ANOVA, pairwise comparisons were made by using a Tukey’s post hoc test. Significant differences were assigned when \(p \leq 0.05\).

**RESULTS**

**Composition of ECM Collagen and HA in dHACM**

To visualize ECM components found in dHACM tissue, immunofluorescent staining was performed for collagen IV, collagen I, and HA. Collagen IV was observed in the amnion basement membrane and throughout the entire chorion layer of the dHACM membrane tissue (Fig. 1A). Collagen I was localized mainly to the amnion layer and the reticular portion of the chorion layer (Fig. 1B), whereas HA was primarily observed in the chorion trophoblast layer (Fig. 1C).

SDS-PAGE was run on digested micronized and membrane dHACM tissues to confirm the presence of collagen and to further identify collagen subunits. All donors of both dHACM formulations exhibited bands at both 139 and 129 kDa (Fig. 1F), molecular weights that correspond to the \(\alpha1\) and \(\alpha2\) collagen subunits of collagen type I. Although some donors exhibited fainter bands of the two
Figure 1. Visualization of collagen and ECM components through immunohistochemistry and SDS-PAGE. (A) Immunofluorescent staining of collagen IV observed in pink. (B) Collagen I staining observed in green. (C) HA staining observed in red. (D) DAPI staining of cell nuclei observed in blue. (E) Overlay of collagen I, HA, and DAPI staining. (F) SDS-PAGE of extracted collagen from micronized and membrane dHACM, n=3, scale bar = 50 µm. dHACM, dehydrated human amnion/chorion membrane; ECM, extracellular matrix.
units, overall, collagen I α1 and α2 subunits were observed.

**Quantification of Matrix Components**

It was determined that $1.84 \pm 1.05 \times 10^3$ ng/mg of fibronectin, $1.82 \pm 1.71 \times 10^3$ ng/mg of HA, $4.54 \pm 4.14$ ng/mg of HSPGs, and $1.46 \pm 0.91$ ng/mg of laminin were measured in the micronized tissue extract (Table 1).

**Analysis of Growth Factors, MMPs, and Inhibitors**

Multiplex and single-factor ELISAs were utilized to identify cytokines and regulatory proteins (nmol/mg of tissue) in micronized dHACM. A table of these growth factors, cytokines, and regulatory proteins is provided in Figure 2B. Of the factors identified, 46.7% are classified as tissue growth factors that can promote wound healing. This includes insulin-like growth factor binding protein-3 (IGFBP-3), placental growth factor (PIGF), platelet-derived growth factor-AA (PDGF-AA), transforming growth factor beta 1 (TGF-β1), and vascular endothelial growth factor (VEGF). Inflammatory cytokines and chemokines were 0.8% and 4.5%, respectively, of the total concentration of factors. Lastly, protease inhibitors such as TIMP-1, TIMP-2, α2M, and A1AT accounted for 46.3% of the identified factors; whereas MMPs such as MMP-2, MMP-8, and MMP-9 comprised 1.7% of the detected molecules (Fig. 2). Further analysis measured plasma protease inhibitors and TIMPs in high quantities: α2M at 92.18 moles/mg, and TIMP-1 and TIMP-2 at ~13 moles/mg each (Table 2). In contrast, MMP-9, MMP-8, and MMP-2 were the most prevalent proteases, at concentrations of $2.67 \pm 1.99$, $1.03 \pm 0.79$, and $0.94 \pm 0.97$ moles/mg, respectively.

**MMP Gelatin Zymography**

To demonstrate the activity of the MMPs found in micronized dHACM, a gelatin zymogram gel was run on dHACM extracts at concentrations ranging from 40 to 1 mg/mL. This assay denatures and dissociates complexed gelatinases within the sample before renaturing after gel electrophoresis; therefore, this technique is able to detect MMPs in both their latent and inhibitor-bound form. The MMP-2 control revealed one band at approximately 62 kDa and one band at 72 kDa, corresponding to the molecular weights of pro MMP-2 and active MMP-2, respectively. For extract concentrations of 40, 20, and 10 mg/mL, bands were observed at 92 and 72 kDa (Fig. 3A), which correspond to the molecular weights of MMP-9 and MMP-2 in their inactive (pro) form. No bands were observed in 1 mg/mL samples. When the band density was quantified and normalized to the optical density of the control MMP-2 sample (band at 72 kDa), no significant effect of concentration was observed for either MMP (Fig. 3B, C).

**MMP Activity**

To characterize gelatinase and collagenase activity in micronized dHACM, degradation of a fluorescently tagged gelatin substrate was measured when treated with the following: dHACM extract, extract with added inhibitor (+I), and extract with added collagenase (+C). After normalization to the control inhibitor + collagenase, in which minimal collagenase activity was expected, extract-only groups at multiple concentrations were not significantly different than the inhibitor + collagenase control (Fig. 4A). In addition, activity levels were not significantly different between extract-only groups and their corresponding extract + inhibitor group. When extract groups were spiked with collagenase, higher extract concentrations exhibited lower collagenase activity: $0.72 \pm 0.14$ U/mL at 1 mg/mL extract concentration, and $0.002 \pm 0.1$ U/mL at 40 mg/mL extract concentration (Fig. 4B). Normalized activity levels for 40, 20, and 10 mg/mL were significantly lower than the collagenase-only control.

**DISCUSSION**

This study identified ECM components, growth factors, cytokines, and MMPs/inhibitors in micronized dHACM that can play a role in the wound-healing process. In addition, the activity of MMPs and protease inhibitors in the tissue was analyzed. Evaluation of ECM components in dHACM tissue revealed, in particular, the localization of collagens I and IV and HA as well as fibronectin, laminin, HA, and HSPGs in high quantities. A broad array of factors (growth factors, cytokines, chemokines, proteases, and inhibitors) were detected in the tissue. The zymogram assay revealed MMPs in their latent form, and the Enzchek gelatinase/collagenase

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**Table 1. Quantified amounts of fibronectin, HA, heparin sulfate proteoglycan, and laminin in guanidine-extracted micronized dHACM**

<table>
<thead>
<tr>
<th>ECM Component</th>
<th>ng/mg Tissue</th>
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<tbody>
<tr>
<td>Fibronectin</td>
<td>$1.84 \pm 1.05 \times 10^3$</td>
</tr>
<tr>
<td>Hyaluronic acid</td>
<td>$1.82 \pm 1.71 \times 10^3$</td>
</tr>
<tr>
<td>Heparan sulfate proteoglycan</td>
<td>$4.54 \pm 4.14$</td>
</tr>
<tr>
<td>Laminin</td>
<td>$1.46 \pm 0.908$</td>
</tr>
</tbody>
</table>

Values reported as moles of that molecule/mg of tissue ± standard deviation, n=4–7.

dHACM, dehydrated human amnion/chorion membrane; ECM, extracellular matrix; HA, hyaluronic acid.
activity assay suggested an excess of active inhibitors in the tissue. Overall, the results from these studies demonstrate that the tissue is composed of an array of structural materials, signaling molecules, and remodeling enzymes that are usually involved in a metabolically active, growing tissue, and provide a better understanding of potential roles of micronized dHACM in wound-healing applications.

In amniotic membranes, the primary ECM components in the amnion layer include a basement membrane that is composed of collagen types IV and VII, fibronectin, and laminin. The adjacent compact and fibroblast layers also contain collagens and provide further mechanical support to the tissue.20 In addition to their roles in situ, these ECM components may provide critical structural and regulatory cues to support and enhance healing in chronic wounds that are treated with dHACM grafts. Immunostaining allowed for visualization and localization of ECM components in the original amnion/chorion membranes before micronization and confirmed that dHACM tissue contains collagens I and IV, and HA, molecules that are present in the native amniotic membrane.20 In addition, collagen IV was observed in the basement membrane of the amnion and throughout the chorion regions of dHACM, similar to its orientation in the native amniotic membrane.20 Gel separation of collagen extracts revealed comparable banding patterns between micronized and membrane dHACM, indicating that both formulations of dHACM tissue contain similar collagen content.

Table 2. Quantified amounts of MMPs, TIMPs, and protease inhibitors found in micronized dHACM

<table>
<thead>
<tr>
<th>Type of Biomolecule</th>
<th>Biomolecule</th>
<th>Moles/mg Tissue</th>
</tr>
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<tbody>
<tr>
<td>Tissue Growth Factors</td>
<td>BDNF, bFGF, EGF, EGF-R, EG-VEGF, FGF-4, GH, HGF, IGFBP-1, IGFBP-2, IGFBP-3, IGFBP-4,</td>
<td>0.42 ± 0.18</td>
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<td></td>
<td>IGF-1, IGF-2, PDGF-AA, PDGF-BB, VEGF-A</td>
<td>0.94 ± 0.97</td>
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<tr>
<td></td>
<td>TIMP-1, TIMP-2, TIMP-4, α2M, A1AT</td>
<td>0.47 ± 0.29</td>
</tr>
<tr>
<td>Inflammatory Cytokines</td>
<td>G-CSF, IL-1α, IL-1β, IL-6, IL-8, IL-12, IL-18, TNF-α</td>
<td>1.03 ± 0.79</td>
</tr>
<tr>
<td></td>
<td>MCP-1, IL-6, IL-18, TNF-α</td>
<td>2.67 ± 1.99</td>
</tr>
<tr>
<td></td>
<td>MMP-1, MMP-2, MMP-3, MMP-8, MMP-9, MMP-10, MMP-13</td>
<td>0.10 ± 0.07</td>
</tr>
<tr>
<td></td>
<td>TIMP-1, TIMP-2, TIMP-4, α2M, A1AT</td>
<td>0.01 ± 0.0008</td>
</tr>
<tr>
<td>Inhibitors</td>
<td>TIMP-1</td>
<td>12.91 ± 1.22</td>
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<tr>
<td></td>
<td>TIMP-2</td>
<td>12.90 ± 4.22</td>
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<tr>
<td></td>
<td>TIMP-4</td>
<td>0.11 ± 0.06</td>
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<tr>
<td></td>
<td>α2M</td>
<td>92.18 ± 53.33</td>
</tr>
<tr>
<td></td>
<td>A1AT</td>
<td>41.30 ± 23.91</td>
</tr>
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Values reported as moles of that molecule/mg of tissue ± standard deviation, n=4.

MMP, matrix metalloproteinase; TIMP, tissue inhibitor of metalloproteinase.
Other ECM components present in micronized dHACM include fibronectin, laminin, HA, and HSPGs, all of which have been shown to play specific roles in wound healing. Fibronectin is a key component of the ECM and functions as an adhesive molecule that provides binding sites for cells, promoting growth and migration. Laminin, found in the basement membrane, mediates binding of bioactive molecules and cells, which, in turn, regulates re-epithelialization, cell shape and movement, and angiogenesis during the wound-healing process. Immunostaining of HA demonstrated that the majority of the HA in dHACM was localized to the chorion layer, with relatively weak staining observed in the amnion. HA, a nonsulfated glycosaminoglycan, is a primary component of the ECM and is known to activate fibroblast proliferation and angiogenesis during the process of wound closure. HSPGs are cell surface proteoglycans that can bind growth factors such as fibroblast growth factor-2 (FGF-2), resulting in protection from degradation and enhancing cell response and activation. The roles of these ECM components are pertinent in wound-healing processes, where it has been shown that heparin sulfate-deficient mice displayed impaired angiogenesis in a corneal micropocket assay, as well as decreased granulation tissue and delayed closure of wounds.

Previous studies characterizing membrane dHACM tissue have identified growth factors and cytokines that were also observed in micronized dHACM tissue in this study (Fig. 2). The tissue growth factors detected in micronized dHACM are known to regulate cell migration, proliferation, and synthesis of ECM components such as collagen, fibronectin, and HA. A variety of these growth factors, including epidermal growth factor, FGF, VEGF, and PDGF, are capable of initiating cell signaling and transcriptional activation of MMPs. In addition, many of the cytokines and chemokines that are identified participate in the inflammatory phase to activate monocytes and regulate leukocyte migration and proliferation, all of which are vital processes in wound healing.

During pregnancy, the amniotic membrane ECM must continue to expand and remodel to maintain structural integrity while accommodating the developing fetus and amniotic fluid through term. Furthermore, tissue remodeling requires protease activity to break down existing ECM components, whereas tissue degradation is offset by a balance of TIMPs and ECM production that supports overall tissue growth. Notably, MMP-9-deficient mice exhibited impaired wound neovascularization and collagen deposition, along with increased inflammatory cell infiltration. Due to their important roles naturally, as well as in wound healing, this study evaluated MMPs and TIMPs released from micronized dHACM tissue and quantified overall MMP activity in tissue extracts. Collagenases (MMP-1, MMP-8, MMP-13), gelatinases (MMP-2, MMP-9), and stromelysins (MMP-3, MMP-10) were detected in the tissue via ELISAs, whereas activity assays revealed that MMPs may...
exist either in their inactive form or complexed with inhibitors contained within the tissue. Typically, MMPs are synthesized in a latent form, at which point they are either secreted or bound to cell surfaces. Thus, the zymogram assay was employed to separate latent and active forms of MMPs based on molecular weight, and it revealed the activity of the enzymes after activation in a renaturing buffer. This was observed with pro MMP-9 and pro MMP-2 bands at 92 and 72 kDa, respectively, in micronized dHACM extracts (Fig. 3). Although significant differences were not observed as a function of dose, the number of samples used may have limited the ability to detect differences.

From the Enzcheck activity assay, no differences in MMP activity were observed (at any concentration of tissue extract) between the extract only and extract + inhibitor, or between these groups and the inhibitor + collagenase control, suggesting minimal protease activity in dHACM tissue (Fig. 4A). Active MMPs play critical roles in wound healing, including regulation of epithelial tissue architecture by degradation of the basement membrane and disruption of intercellular junctions, along with support of keratinocyte migration and wound contraction. However, spatiotemporal control of MMP activity is necessary to maintain the balance of degradation and production of ECM, to prevent further breakdown that could contribute to a chronic wound environment.

TIMPs are known to bind different MMPs; for example, TIMP-2 has been shown to bind MMP-2 and MMP-9, resulting in modulation of MMP activity and regulation of cell migration during wound healing. With ELISAs, multiple TIMPs and two large plasma protease inhibitors (α2M and A1AT) were detected, and each (except TIMP-4) was present at molar ratios in excess of all MMPs measured (Table 2). α2M is known to inhibit most proteinases of all classes (metallo-, serine, carboxyl, thiol), and A1AT can act on serine proteases, collagenases, and elastase. The effect of extract concentration on MMP activity, whereby greater collagenase activity was observed at the lowest concentration of extract (1 mg/mL), further implied the presence of active inhibitors in dHACM tissue extract (Fig. 4B). It has previously been shown that immediate intervention of alkali burns of the cornea with amniotic membrane resulted in lower proteinase activity and subsequent epithelial healing compared with groups without amniotic membrane administration. Furthermore, in both normal acute wounds and chronic leg ulcers, MMP activity decreased significantly as wound closure occurred. Therefore, the administration of protease inhibitors may promote epidermal healing. With the delivery of micronized dHACM, inactive MMPs in the tissue could be activated by MMPs or growth factors in the wound bed; however, the complexation of excess TIMPs and plasma protease inhibitors with both the endogenous MMPs and the MMPs present in the wound is expected to promote resolution.

**INNOVATION**

Due to the dynamic nature of the tissue and its biologic factor-rich ECM, amniotic membranes have demonstrated efficacy in wound-healing applications. These results identified ECM components and characterized a variety of growth factors, cytokines, proteases, and inhibitors present in the dHACM tissue. We believe that at an injury site, the administration of micronized dHACM tissue will deliver matrix molecules, a cocktail of growth factors and cytokines, and
protease inhibitors that collectively promote healing. Ultimately, the use of micronized dHACM in the treatment of chronic wounds may advance the current standards of wound care.

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REFERENCES


KEY FINDINGS

- dHACM tissue contained the ECM components collagens I and IV, HA, heparin sulfate proteoglycans, fibronectin, and laminin, as well as numerous growth factors, cytokines, chemokines, proteases, and protease inhibitors that are known to participate in the wound-healing process.
- Though MMPs were present in dHACM tissues, inhibitors of MMPs overwhelmingly outnumbered the MMP enzymes by a ratio of more than 28:1.
- MMPs in the tissue existed in their latent form and may have been complexed with active inhibitors.
- Overall, this work provides a better understanding of how micronized dHACM tissue can be used to effectively promote wound repair.


Abbreviations and Acronyms

ANOVA = analysis of variance
dsHACM = dehydrated human amnion/chorion membrane
ECM = extracellular matrix
ELISA = enzyme-linked immunosorbent assay
FGF = fibroblast growth factor
HA = hyaluronic acid
HSPGs = heparan sulfate proteoglycans
MMP = matrix metalloproteinase
PAGE = polyacrylamide gel electrophoresis
SDS = sodium dodecyl sulfate
TGF = transforming growth factor
TIMP = tissue inhibitor of metalloproteinase
VEGF = vascular endothelial growth factor