Preserved Amniotic Membrane Transplantation for Ocular Surface Reconstruction

a report by

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Maintenance of the ocular surface is dependent on the complex interplay of the lids, lacrimal gland, tear film, conjunctiva, cornea, and neural network. Although multifactorial in etiology, severe ocular surface disorders (OSDs) commonly result in progressive inflammation, vascularization, scarring, and/or loss of visual function, often refractory to conservative medical therapy. Unique properties of human amniotic membrane (AM) confer several advantages in the management and treatment of various OSDs. AM is the innermost layer of the placenta and is composed of an epithelial monolayer, an underlying thick basement membrane, and an avascular stroma.

The first ophthalmic utilization of fresh placental tissue by de Röth was reported in 1940 with limited success. Since then, AM transplantation (AMT) has been used successfully as a temporary patch and/or permanent graft for corneal and/or conjunctival cells in the management of numerous conditions including persistent epithelial defects, corneal ulcers, descemetocele formation, corneal perforation, limbal stem cell deficiency, symptomatic bullous keratopathy, band keratopathy, chemical injury, thermal injury, repair of conjunctival defects, pterygium surgery, bleb repair, scleral perforation, and high-risk corneal transplantation.

Although the exact mechanism of action is unclear, AM demonstrates physiological properties that promote epithelial and stromal wound healing while suppressing inflammation, fibrosis, and vascularization. The AM basement membrane contains laminin, fibronectin, and collagen type IV and VII, similar to the basement membrane of conjunctival tissue, but more than corneal epithelium. The basement membrane provides a substrate that promotes epithelial cell adhesion, migration, and differentiation and prevents epithelial cell apoptosis. The underlying stromal matrix contains proteins that suppress transforming growth factor (TGF)-β signaling in fibroblasts, and consequently inhibit the proliferation and differentiation into myofibroblasts, extracellular matrix production, and scar formation. The resultant downregulation is responsible for the antifibrosis effect. The anti-inflammatory activity of AM is attributed to numerous mediators, including interleukin (IL)-1 receptor antagonist, IL-10, activin, inhibin, and inter-α-trypsin inhibitor. AM has also been shown to induce apoptosis of inflammatory cells. Vascularization is inhibited by the anti-inflammatory effect of AM and by the antiangiogenic proteins present within AM, including endostatin, thrombospondin-1, and inhibitors of metallo-proteases. Finally, a unique advantage of AM is the low immunogenicity of the tissue and, therefore, the consequent lack of rejection. The immunosuppressive ability of AM to inhibit a mixed lymphocyte reaction appears to be partly mediated by a soluble factor, resulting in decreased alloreactive T-cell proliferation and Th1 and Th2 cytokine production.

Advances in technology have allowed surgeons to move away from the disadvantages and inconveniences of fresh AM to cryopreserved, dehydrated, or freeze-dried AM tissue. In the US, the two different AMs are commercially available as epithelialized, cryopreserved AM (Ammiograft®, Biotissue Inc., Miami, Florida) or de-epithelialized, dehydrated AM (AmbioDry™, IOP®, Inc., Costa Mesa, California). AM is obtained under sterile conditions after elective Cesarean section. The cryopreserved AM contains epithelial cells and is disinfected with antimicrobial agents, attached to nitrocellulose paper with the basement membrane facing upward, stored in a deep freeze at -80°C, and thawed for five to 10 minutes before use. Although the AM is not sterilized during its processing, the tissue undergoes microbiological testing prior to its release. The orientation is distinguishable by the sticky stromal side attached to the nitrocellulose paper. When stored properly at -80°C, the membrane can be used for up to two years from the time of procurement. The dehydrated AM undergoes de-epithelialization, dehydration, sterilization with electron beam irradiation, and storage free-standing at room temperature (50–85°F). As free-standing tissue, the AM does not require careful separation from nitrocellulose paper, and the orientation is clear by the imprinted text ‘IOP.’ In the dry state, the AM can be cut, manipulated, and placed onto the surgical site before being activated with a saline solution or an adhesive agent. The AM can be used for up to two years from the time of processing. Compared with AM epithelial cells, AM denuded of epithelial cells may provide a

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Anterior Segment  Ocular Surface

The key to successful treatment centers on suppression of inflammation. Pro-inflammatory cytokines (IL-1β, tumor necrosis factor [TNF]-α) can promote cultured fibroblast proliferation and overexpression of select matrix metalloproteases (type I, III), which allows for a more invasive cellular phenotype that could increase the likelihood of recurrence. Therefore, the potential anti-inflammatory, antifibrotic, and antiangiogenic benefits of AM may provide several advantages in ocular surface reconstruction after pterygium excision. Using indocyanine green angiography (ICGA), delayed vascularization in the AM graft has been compared with conjunctival autograft transplantation (CAT), which may be attributable to the antiangiogenic effects of AM. These findings may contribute to the delay in recurrence with AMT compared with limbal CAT (LCAT), which has been postulated as a barrier phenomenon of AMT. In addition to the inherent physiological properties of AMT, other benefits include convenience, less technical skill needed, shorter operative time, and fewer complications compared with harvesting a conjunctival (and/or limbal) graft. Another distinct advantage of AMT is the ability to preserve conjunctival tissue in precluding the use of a CAT, particularly for future glaucoma-filtering procedures. AM grafting is considered the preferred adjunctive treatment in large pterygia where insufficient conjunctiva for a graft is available or in cases that are at high risk for recurrence, as AM transplantation can be repeated.

Another technological innovation that has furthered the use of AMT in pterygium surgery is the development of biological tissue adhesives, such as fibrin glue (FG) (Tisseel VH fibrin sealant, Baxter Corporation, Irvine, California). A tissue bioadhesive can be used to supplement sutures to reduce the number needed or can be used in place of sutures. The initial success of FG in conjunction with CAT led to an increased interest in AM grafts. Implementation of FG to secure the AM graft compared with sutures has been associated with shorter surgical times, technical ease of use, and less post-operative discomfort, particularly foreign body sensation. The decreased operative time facilitates the use of topical anesthesia. Additionally, FG avoids the suture-related complications of globe perforation, hemorrhage, graft button-hole, suture abscess, tissue necrosis, and giant papillary conjunctivitis and bypasses the need to remove sutures post-operatively, as the fibrin naturally biodegrades.

Lower rates of recurrence have been found for FG (5.3–8%) compared with sutures (13.5–20%) to secure conjunctival autografts in pterygium surgery. Kheirkhah et al. found significantly greater conjunctival inflammation after AMT and MMC with sutures compared with FG. More severe inflammation has been found to be related to higher rates of recurrence. AM has been modified to broaden its utility. Minimal amounts of FG are recommended due to its potential pro-inflammatory effect. To address this concern a bio-adhesive-coated AMT (fibrinogen and thrombin) was created, maintaining the biological characteristics of AM and bypassing the need for sutures. In addition to use as a graft, AM has been used as a temporary patch for the surgical treatment of primary treatment without recurrence (n=20). In conjunction with a fixed AMT graft, a temporary AMT patch may be beneficial after pterygium surgery, similar to sutureless AMT for partial limbal stem cell deficiency. AM is commercially available to be used as a temporary patch with
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cropyamed AMT fastened to a sylmbephon ring (ProKera®, Bio-Tissue Inc, Miami). Other modifications to AMT include ex vivo epithelial cell expansion, which is beyond the scope of this article.

The literature to support the utility of AMT in ocular surface reconstruction continues to expand. Prospective clinical studies are needed to evaluate the efficacy of AMT in comparison with currently available alternatives.

Additionally, research to better elucidate the mechanism of action behind the physiological properties of AMT continues to be of great interest. Although limitations of AMT exist, particularly in cases of total stem cell deficiency and severe keratoconjunctivitis sicca, we can expect that future modifications in surgical technique, concomitant medical therapy, and technological advances will develop to optimize the results of procedures using AMT.