Mucosal Substitutes for Periodontal Soft Tissue Regeneration

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Abstract

Periodontal plastic surgery would be defined as the surgical procedures performed to prevent or correct anatomic, developmental, and traumatic or disease induced defects of the gingiva, alveolar mucosa or bone. The introduction of oral mucosal equivalents also called as mucosal substitutes or mucosal fillers, composed of the structured triad of scaffold, cells and signaling molecules could be considered the essence or a culmination of the notion of periodontal regeneration. For the purpose of soft tissue augmentation, various mucogingival surgical procedures that are usually performed are: connective tissue grafts (gold standard) and its modifications, lateral pedicle flap, free gingival grafts, semilunar flaps, coronally positioned flaps. Several mucosal substitutes have been fabricated and tried with varying results. Tissue engineered mucosal constructs owe their origin to the treatment of ulcers, diabetic foot and burns. There is an increasing interest in the field of dentistry too. Mucosal substitutes are made of cell sources that have not caused acute and chronic tissue reaction and have self-renewal properties that can re-grow and differentiate to a new tissue. With all the advantages of reproducible tissue engineering technologies, these mucosal substitutes/ oral mucosal equivalents appear to be the imminent hope for tissue repair, replacement or regeneration in regenerative medicine in the treatment of lost tissues, impaired functions and soft tissue defects that are caused by congenital abnormalities, trauma, diseases or aging processes.

Keywords: Periodontal; Mucogingival surgery; Soft tissue augmentation; Mucosal substitutes; Mucosal constructs

Introduction

Tissue engineering or regenerative medicine has been defined as an interdisciplinary field that applies the principles of engineering and life sciences that contributes towards the development of biological substitutes for the repair or regeneration of tissue or organ function [1,2]. The concept of tissue engineered constructs has led to a paradigm shift in treatment options from using synthetic implants and tissue grafts to a tissue engineering approach that uses degradable porous material scaffolds integrated with biological cells or molecules to regenerate tissues. The introduction of oral mucosal equivalents also called as mucosal substitutes or mucosal fillers, composed of the structured triad of scaffold, cells and signaling molecules could be considered the culmination of the notion of periodontal regeneration.

Background

Soft tissue defects that are of concern and usually require treatment are:

- Gingival augmentation
- Root coverage
- Correction of mucosal defects at implants
- Removal of aberrant frenulum
- Augmentation of edentulous ridge
- Prevention of ridge collapse associated with tooth extraction
- Crown lengthening
- Gingival preservation at ectopic tooth eruption site

The most common and prevalent issues faced today are those of gingival recession and inadequate width of attached gingiva. The presence of an adequate width of attached gingiva is a prerequisite for a healthy gingiva. An adequate width varies for each individual and an inadequate width is a complicating factor in periodontal therapy. Gingival recession is defined as the apical migration of the marginal gingiva, creating esthetic as well as pathological complications.

For the purpose of soft tissue augmentation, the various mucogingival surgical procedures that are usually performed are connective tissue graft and its modifications, lateral pedicle flap, free gingival grafts, semilunar flaps and coronally positioned flaps. However, the disadvantages like second site morbidity and patient discomfort with these procedures have led us towards the use of mucosal substitutes.

Mucosal Substitutes/Scaffolds

Mucosal substitutes or scaffolds have following features:

- They serve as a temporary supporting structure (extracellular matrix), the initial architecture, on which the cells can grow three-dimensionally into the desired tissue.
- They provide the environment needed for cellular growth and differentiation.
- They provide the strength to withstand mechanical stress and guide their growth.
- They are biodegradable and degrade at the same rate as the tissue regenerates to be optimally replaced by the host tissue.

These scaffolds could be categorized as:

- Naturally Derived Scaffolds: Acellular Dermis, amniotic membrane
- Fibroblast-populated Skin Substitutes: Dermagraft™, Apligraf™, Orcel™, Polyactive™, Hylograf 3D™
- Gelatin-based Scaffolds
- Collagen-based Scaffolds

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e) Fibrin-based Scaffolds
f) Hybrid Scaffolds: Skin substituteS based on a combination of synthetic and natural materials.
g) Synthetic Scaffolds: Polyethylene terephthalate membranes (PET membranes), Porous polyglycolic acid

Several mucosal substitutes have been fabricated and tried with varying results, are described below.

Mucograft prototype (Geistlich Mucograft)

It is a 3D collagen matrix specifically designed for soft tissue regenerative purposes. It consists of two structures: the compact macro-structure provides stability and favors open healing whereas the spongy micro-structure supports blood clot stabilization and the in growth of soft tissue cells. This new collagen matrix (CM) aimed to increase keratinized gingiva/mucosa, when compared with the free connective tissue graft (CTG), and has been tested by many randomized longitudinal parallel controlled clinical trials. Only few clinical trials have been conducted comparing CM with CTG. All the trials have suggested the unique properties of mucograft for soft tissue augmentation with added advantages of lower patient morbidity and reduced surgery time [3-5]. However one study has indicated that CM results in more tissue contraction (67%) as compared to CTG (60%) [4] (Table 1).

Collagen matrix 10826 (bi-layered collagen nano-structured membrane prototype)

This is a collagen matrix of porcine origin fabricated by Geistlich Pharma AG (Switzerland) To evaluate fundamental cell functions, such as adhesion, IL-6 production and proliferation of human gingival keratinocytes cultured, a study was performed on this newly engineered collagen. Functional tests revealed that keratinocytes adhered to CM-10826 and up-regulated their basal IL-6 production. The type of keratinocytes used expressed cytokeratin 14. Proliferation experiments demonstrated that the best cellular response was observed in the presence of Collagen I, the main component of CM-10826. No undesired effects were observed as regards for keratinocyte viability, morphology or differentiation.

The results demonstrated that CM-10826 has a favorable biological effect on the in vitro response of gingival keratinocytes in terms of IL-6 production, cell growth and adhesion, thus encouraging a possible use of this collagen membrane as a tissue which, alone, may substitute for autologous gingival grafts thereby overcoming the limitations of autologous tissue [5]. No in vivo studies have been performed with this substitute.

Matriderm

Matriderm by Medskin solutions is of bovine origin, consists of 1 mm-thick structurally intact native collagen matrix coated with α-elastin hydrolysate from the ligament, freeze-dried and non-cross-linked. This scaffold was introduced to treat deep and full-thickness EVPOME suitable for intraoral grafting. This whole procedure took a month. The EVPOME clinically has showed changes indicating vascular ingrowth and cytological evidence of the persistence of grafted cultured keratinocytes on the surface. An in vivo study conducted to assess the efficacy and safety of the grafted EVPOME in producing a keratinized mucosal surface epithelium concluded that it has the capability to augment keratinized tissue around the teeth [16].

Fibroblast derived dermal substitute (HF-DDS)

Cultured epithelium fabricated with living mucosal cells and epithelial sheets prepared by cultivating fibroblasts onto scaffold to have been used successfully for gingival augmentation. Human fibroblast-derived dermal substitute (HF-DDS) is an example of a tissue engineered construct designed to increase the amount of

Table 1

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<th>Scaffolds Type</th>
<th>Properties</th>
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<tr>
<td>Mucograft</td>
<td>Biologically stable, favors open healing</td>
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<tr>
<td>Collagen matrix 10826</td>
<td>Stability with added advantages of lower patient morbidity and reduced surgery time</td>
</tr>
<tr>
<td>Matriderm</td>
<td>Biocompatible, color match, no scar formation</td>
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Table 2

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<tr>
<td>Mucograft</td>
<td>Root coverage and soft tissue enhancement</td>
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<tr>
<td>Collagen matrix 10826</td>
<td>Soft tissue augmentation</td>
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<tr>
<td>Matriderm</td>
<td>Regrow following tissue destruction</td>
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Table 3

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<th>Mucosal Substitutes</th>
<th>Applications</th>
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<td>Human amniotic membrane</td>
<td>Root coverage and soft tissue enhancement</td>
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<tr>
<td>Emdogain</td>
<td>Induction of proliferation, migration, adhesion, mineralization and differentiation of cells in periodontal tissue</td>
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<tr>
<td>Ex vivo produced oral mucosa equivalent (EVPOME)</td>
<td>Root coverage and soft tissue regeneration</td>
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Ex vivo produced oral mucosa equivalent (EVPOME)

In vitro cultured equivalents have become very promising in the field of oral and periodontal surgery. EVPOME was made from primary human oral keratinocytes, harvested from palatal keratinized mucosa. It was developed by University of Michigan. This was expanded in vitro in an environment free of serum, transformed irradiated xenogenic feeder cells, and pituitary extract in a defined culture medium [15]. After sufficient oral keratinocytes were produced, they were seeded onto AlloDerm (an acellular dermal matrix; LifeCell) to produce a full-thickness EVPOME suitable for intraoral grafting. This whole procedure took a month. The EVPOME clinically has showed changes indicating vascular ingrowth and cytological evidence of the persistence of grafted cultured keratinocytes on the surface. An in vivo study conducted to assess the efficacy and safety of the grafted EVPOME in producing a keratinized mucosal surface epithelium concluded that it has the capability to augment keratinized tissue around the teeth [16].

Human amniotic membrane

The amniotic membrane (AM) is considered an important potential source for scaffolding material. The AM represents the innermost layer of the placenta and is composed of a single epithelial layer, a thick basement membrane and an avascular stroma. The special structure and biological viability of the AM allows it to be an ideal candidate for creating scaffolds used in Tissue Engineering (TE). Epithelial cells derived from the AM have the advantages of stem cells, thus are a more suitable source of cells for TE than stem cells. The extracellular matrix components of the basement membrane of the AM create an almost native scaffold for cell seeding in TE. Various case reports have observed that amnion membrane has the properties of antibacterial, biocompatibility, color match, no scar formation and many more favoring new epithelium formation [7-9]. Thus, it has been suggested that amnion membrane is a suitable alternative to free gingival grafts for root coverage and soft tissue enhancement (Table 2).

Emdogain

Emdogain or Enamel matrix derivative (EMD) developed by Straumann is an extract of porcine fetal tooth material used to bio mimetically stimulate the soft and hard tissues surrounding teeth to regrow (in a process known as regeneration) following tissue destruction. A commercially prepared and purified extract of enamel matrix proteins, EMD is composed primarily of amelogenin and has been shown to promote PDL fibroblast proliferation and growth [10]. The effects of Emdogain are thought to be the induction of proliferation, migration, adhesion, mineralization and differentiation of cells in periodontal tissue. EMD mimics normal root development by stimulating release of autocrine growth factors from periodontal ligament undifferentiated mesenchymal cells [11]. It has been shown to promote periodontal wound healing and/or regeneration when applied to tooth root surfaces in soft tissue dehiscence models. EMD application has been found to be an effective alternative to achieve root coverage together with a gain in height of keratinized gingiva, in interdental papilla reconstruction and new cementum formation [12-14] (Table 3).
keratinized tissue around teeth that do not require root coverage. The tissue engineered HF-DDS graft has been found to be safe and capable of generating keratinized tissue without the morbidity and potential clinical difficulties associated with donor site surgery [17-19] (Table 4).

**Celtx™ (Organogenesis)**

Celtx™ is a living cellular construct comprised of human fibroblasts, keratinocytes and extracellular matrix proteins. The living cells found in Celtx produce a wide array of growth factors and cytokines that in turn stimulate the patient's own cells to regenerate new tissue that is clinically significant and aesthetically appealing.

The temporal expression of angiogenic biomarkers during wound healing of soft tissue reconstructive procedures was investigated by comparing living cellular constructs (LCC) with autogenous free gingival grafts. 44 human participants bilaterally lacking sufficient zones of attached keratinized gingiva were randomly assigned to soft tissue surgery plus either LCC or autograft. Wound fluid samples were collected at baseline and weeks 1, 2, 3, and 4 post-operatively and analyzed for a panel of angiogenic biomarkers: angiogenin (ANG), angiostatin (ANT), PDGF-BB, VEGF, FGF-2, IL-8, TIMP-1, TIMP-2, GM-CSF, and IP-10. Results demonstrated a significant increase in expression of ANG, PDGF-BB, VEGF, FGF-2, and IL-8 for the LCC group over the autograft group at the early stages of wound repair. Although angiogenic biomarkers were modestly elevated for the LCC group, no clinical correlation with wound healing was found. This investigation demonstrated that, during early wound-healing events, expression of angiogenic-related biomarkers is up-regulated in sites investigated during early wound-healing events, expression of angiogenic-related biomarkers is up-regulated in sites investigated during early wound-healing events, expression of angiogenic-related biomarkers is up-regulated in sites investigated during early wound-healing events, expression of angiogenic-related biomarkers is up-regulated in sites.

**Gintuit™**

GINTUIT is an allogeneic cellularized scaffold product, a thin cellular sheet made of human fibroblasts, keratinocytes, human extracellular matrix proteins and bovine collagen developed by Organogenesis Company. FDA has approved this cell-based product for generating new and aesthetically appealing oral tissues.

Organogenesis completed a multi-center, randomized, pivotal clinical trial in 2012 to determine the efficacy and safety of GINTUIT to regenerate oral soft tissue in patients with gingival recession. The GINTUIT-treated sites generated a clinically significant amount of...
keratinized oral soft tissue. Moreover, GINTUIT generated gingival tissue that better matched the color and texture of the patients' surrounding tissue versus traditional palatal grafting procedures. Importantly, patients overwhelmingly preferred GINTUIT over traditional palatal grafting procedures. Results of the study demonstrated that GINTUIT provided a more natural looking graft with a better color match and better, tissue integration than traditional palatal grafting procedures. The tissue engineered HF-DDS graft was safe and capable of generating keratinized gingiva. The tissue engineered HF-DDS graft was safe and capable of generating keratinized gingiva. The tissue engineered HF-DDS graft was safe and capable of generating keratinized gingiva.

**Table 3: Emdogain.**

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<tr>
<th>Investigators</th>
<th>Study design</th>
<th>Method</th>
<th>Results</th>
<th>Conclusion</th>
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<tbody>
<tr>
<td>Mohammadi et al. [17]</td>
<td>Case report</td>
<td>Tissue-engineered gingival graft (HF-DDS) was used for regenerating facial gingiva around an implant at lower left first premolar area with insufficient attached gingiva.</td>
<td>The histological features demonstrated a fully keratinized tissue supported by dense connective tissue. The width of keratinized gingiva and attached gingiva was more at 3 months compared to baseline.</td>
<td>The tissue engineered gingival graft is safe and capable of generating keratinized gingiva.</td>
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<td>Wilson et al. [18]</td>
<td>Split mouth case control clinical trial</td>
<td>Thirteen patients. Each patient had Miller Class I or II bilateral facial recession defects ≥3 mm on two non-adjacent teeth. The test tooth received an HF-DDS graft Control site: CTG was placed Control group: placebo (propylene glycol alginate) + CPF</td>
<td>Amount of root coverage and Width of keratinized tissues was slightly greater in control group.</td>
<td>Human fibroblast-Dehuman fibroblast-derived dermal substitute may offer potential as a substitute to the connective tissue graft for covering areas of facial Miller Class I or Class II gingival recession in humans derived dermal substitute may offer potential as a substitute to the connective tissue graft for covering areas of facial Miller Class I or Class II gingival recession in humans.</td>
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<tr>
<td>McGuire et al. [19]</td>
<td>Split mouth case control clinical trial</td>
<td>25 patients with insufficient attached gingiva associated with at least two teeth in contra lateral quadrants of the same jaw were treated Control teeth: GA test teeth: HF-DDS graft Control group exhibited an average of 1.0 to 1.2 mm more keratinized tissue over time and about half as much shrinkage as the test group</td>
<td>No difference in gain of width of gingiva in both groups.</td>
<td>The tissue engineered HF-DDS graft was safe and capable of generating keratinized tissue without the morbidity and potential clinical difficulties associated with donor site surgery.</td>
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**Table 4: Fibroblast derived dermal substitute.**

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<tr>
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<th>Study design</th>
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<th>Results</th>
<th>Conclusion</th>
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<tbody>
<tr>
<td>Jaiswal et al. [12]</td>
<td>Randomized case control study</td>
<td>Twenty patients with a total of 46 gingival recession defects. Test group: 10 patients with 22 recession defects treated with Emdogain with CPF Control group: 10 patients with 24 gingival recession defects, was treated with 24% EDTA with CPF</td>
<td>Emdogain gel resulted in a statistically significant increase in root coverage, gain in the clinical attachment level (CAL), and probing pocket depth (PPD). No difference in gain of width of gingiva in both groups.</td>
<td>Emdogain is an effective alternative for root coverage procedures</td>
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<tr>
<td>Berlucchi et al. [13]</td>
<td>Randomized case control study</td>
<td>13 recessions, Emdogain was used in combination with a coronally advanced flap (CAF+EMD group). In the other 13 recessions, Emdogain and the flap were used in combination with a subepithelial connective tissue graft (CAF+CTG+EMD group).</td>
<td>CAF+CTG+EMD group displayed good clinical results in terms of root coverage, increased width of keratinized gingiva, Recovery width (P=0.027) and probing pocket depth (P=0.046) exhibiting a higher reduction in the EMD group.</td>
<td>EMD has appreciable wound healing and regenerative properties.</td>
</tr>
<tr>
<td>Hägewald et al. [14]</td>
<td>Blinded, split-mouth, placebo-controlled and randomized design.</td>
<td>30 patients, aged 22-62 years, with 2 paired buccal recession defects of at least 3 mm. Control group: placebo (propylene glycol alginate) + CPF Control group: placebo (propylene glycol alginate) + CPF</td>
<td>Recession width (P=0.027) and probing pocket depth (P=0.046) exhibiting a higher reduction in the EMD group.</td>
<td>EMD has better long term results.</td>
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**Dermagraft**

It is composed of cryopreserved human-derived fibroblasts and collagen applied to a bioabsorbable mesh. Dermagraft was developed by Advanced Tissue Sciences and has been approved by the FDA for repair of diabetic foot ulcers and for use in the treatment of wounds related to dystrophic epidermolysis bullosa. The use of tissue-engineered dermal replacement in the vestibular extension instead of palatal donor tissue or (split-thickness) skin graft has been described in a recent study. In three patients the living human-derived Dermagraft was implanted on the wound surface after mucogingival junction and suprastructure dissection. Following application of Dermagraft, vestibular depth was increased and no scarring occurred. Tissue engineered dermal replacement consisting of living human fibroblasts has proved to be a useful substitute for autogenous grafts in pre prosthetic surgery. Offering the advantage of unlimited availability, good colour match and no donor site morbidity [22].

**Aongen™**

Aongen™ by Body Organ Biomedical Corporation is a collagen based tissue repair and bone graft augmentation material. It is a bioabsorbable 3D matrix made of Type-I atelocollagen which is designed for successful soft tissue regeneration and stimulation of dental bone proliferation at an accelerated rate. Aongen™ Dental is indicated for application as an aid in the management of extraction sockets, periodontal defects, bridges, dentures, implants, sinus lift osteotomies, soft tissue augmentation and
oral wounds such as oral ulcers, denture sores and trauma or surgical wounds [23,24]. The porous structure of Aongen™ Dental provides an ideal scaffold and interface for space maintenance, allowing the surrounding cells to seed and proliferate [24,25]. The matrix not only enhances the hemostasis process, but also stimulates and modulates the migration and proliferation of cells. Aongen™ Dental is absorbed within 30 days without adverse effects or production of any toxic substances. At the implant site, the course of action may persist for a longer duration depending on the characteristics of the stressed site and the predetermined remodeling mechanisms. It does not require prior preparation to best achieve its singular heal-expansive effects. The bullet and tape shapes allow uncomplicated placement at the insertion site. To prevent dislodgement at the site of placement, it can be trimmed and left in situ, suturing by peripheral gingival-soft tissue borders, or by other means. In terms of regeneration, it acts as a natural biological matrix designed to provide an excellent healing environment and acts as a guide for the regeneration of vital soft and bone tissue [25].

**Few Products that have a Potential to be used as a Dental Constructs**

**Apligraf (Organogenesis)**

It is a bilayered living cell therapy composed of an epidermal layer of living human keratinocytes and a dermal layer of living human dermal fibroblasts. Apligraf is supplied as needed, in one size, with a shelf-life of 10 days. It was FDA-approved in 1998 for use in conjunction with compression therapy for the treatment of non-infected, partial- and full-thickness skin ulcers [26].

**OrCel™ (Forticell Bioscience)**

It is formerly called Composite Cultured Skin. It is an absorbable allogeneic bi-layered cellular matrix, made of bovine collagen, in which human dermal cells have been cultured. It was approved by the FDA premarket approval (PMA) for healing donor site wounds in burn victims and under a humanitarian device exemption (HDE) for use in patients with recessive dystrophic epidermolysis bullosa undergoing hand reconstruction surgery to close and heal wounds created by the surgery, including those at donor sites.

**Epicel (Genzyme Biosurgery)**

It is a cultured epithelial autograft and is FDA-approved under a humanitarian device exemption (HDE) for the treatment of deep dermal or full-thickness burns comprising a total body surface area of greater than or equal to 30%. It may be used in conjunction with split-thickness autografts, or alone in patients for whom split-thickness autografts may not be an option due to the severity and extent of their burns. A study by of the Baltimore Regional Burn Center of the Johns Hopkins University compared the outcomes of therapy in patients with massive burns with or without cultured epidermal autografts (CEAs). There was found to be a significant reduction in mortality in the CEA group compared with controls, from 48% to 14% (p<0.007) [27].

**Integra® Dermal Regeneration Template (Integra LifeSciences)**

It is a bovine, collagen/glycosaminoglycan dermal replacement covered by a silicone temporary epidermal substitute. Integra® Dermal Regeneration Template is an advanced skin replacement system, designed to provide immediate wound closure and permanent regeneration of the dermis. The product is placed in direct contact with the excised wound and consists of a complex three-dimensional porous matrix that acts as a scaffold for cell migration and allows for regeneration of the dermal layer of the patient’s skin. It is FDA-approved for use in post excisional treatment of life-threatening full-thickness or deep partial-thickness thermal injury where sufficient autograft is not available at the time of excision or not desirable because of the physiologic condition of the patient. Integra® Matrix Wound Dressing and Integra™ meshed Bilayer Wound Matrix are substantially equivalent skin substitutes that are FDA-510(k) approved for other indications. No clinical trial by any institute has been conducted on this product yet.

**TransCyte (Advanced Tissue Sciences)**

It consists of human dermal fibroblasts grown on nylon mesh of Biobrane combined with a synthetic epidermal layer and was approved by the FDA in 1997. TransCyte is intended to be used as a temporary covering over burns until autografting is possible. It can also be used as a temporary covering for some burn wounds that heal without autografting. A prospective, randomized, comparison study of silver sulfadiazine and TransCyte was performed with the use of paired wound sites on 14 patients to evaluate the role of TransCyte for the treatment of partial-thickness burns. Wounds treated with TransCyte healed more quickly (mean 11.14 days to 90% epithelialization vs 18.14 days, p=0.002). A non-comparison evaluation was then done for an additional 18 patients, and it confirmed excellent wound healing and an absence of infections. There were no infections in the 32 wound sites treated with TransCyte. In the first study group, late wound evaluations (3, 6, and 12 months postburn) were performed with use of the Vancouver Scar Scale. The results indicated that wound sites treated with TransCyte healed with less hypertrophic scarring than sites treated with silver sulfadiazine (p<0.001 at 3 and 6 months, p=0.006 at 12 months) [28]. Similar results were obtained in another prospective study [29].

**OASIS® Wound Matrix (Cook Biotech)**

It is a xenogeneic collagen scaffold derived from porcine small intestinal mucosa. It was cleared by the FDA’s 510(k) process in 2000 for the management of partial and full-thickness wounds including pressure ulcers, venous ulcers, diabetic ulcers, chronic vascular ulcers, tunneled undermined wounds, surgical wounds, trauma wounds, and draining wounds. A prospective, randomized, controlled multicenter trial was conducted with the objective of comparing the effectiveness of OASIS wound matrix with compression vs. compression alone in healing chronic leg ulcers within 12 weeks. 120 patients with at least 1 chronic leg ulcer were randomly assigned to receive either weekly topical treatment of OASIS plus compression therapy (n=62) or compression therapy alone (n=58). Ulcer size was determined at enrollment and weekly throughout the treatment. Healing was assessed weekly for up to 12 weeks. Recurrence after 6 months was recorded. The primary outcome measure was the proportion of ulcers healed in each group at 12 weeks. After 12 weeks of treatment, 55% of the wounds in the OASIS group were healed, as compared with 34% in the standard-care group (p=0.0196). None of the healed patients treated with OASIS wound matrix and observed for the 6-month follow-up experienced ulcer recurrence. Thus it was suggested that the SIS wound matrix, as an adjunct therapy, significantly improves healing of chronic leg ulcers over compression therapy alone [30].

**Conclusion**

Tissue engineered mucosal constructs owe their origin to the treatment of ulcers, diabetic foot, burns followed by an increasing interest in the field of dentistry too. Mucosal substitutes are made of cell sources that have not caused acute and chronic tissue reaction and...
have self-renewal properties that can re-grow and differentiate to a new tissue. Further, these mucosal substitutes can serve as biodegradable scaffolds and with the appropriate bioreactors, the quality of tissue may also be controlled. With all the advantages of reproducible tissue engineering technologies, these mucosal substitutes/oral mucosal equivalents appear to be the imminent hope for tissue repair, replacement or regeneration in regenerative medicine in the treatment of lost tissues, impaired functions and soft tissue defects that are caused by congenital abnormalities, trauma, diseases or aging processes.

References