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Review Article

Chronic Wound Healing: A Review of Current Management and Treatments

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ABSTRACT

Wound healing is a complex, highly regulated process that is critical in maintaining the barrier function of skin. With numerous disease processes, the cascade of events involved in wound healing can be affected, resulting in chronic, non-healing wounds that subject the patient to significant discomfort and distress while draining the medical system of an enormous amount of resources. The healing of a superficial wound requires many factors to work in concert, and wound dressings and treatments have evolved considerably to address possible barriers to wound healing, ranging from infection to hypoxia. Even optimally, wound tissue never reaches its pre-injured strength and multiple aberrant healing states can result in chronic non-healing wounds. This article will review wound healing physiology and discuss current approaches for treating a wound.

Keywords: - Biofilms, Chronic wounds, Growth factors, Hyperbaric oxygen, Negative pressure wound therapy, Skin infection, Skin substitutes, Wound dressings, Wound healing

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INTRODUCTION

he process of cutaneous wound healing is incredibly complex, dependent on an intricate interplay between a number of highly regulated factors working in concert to restore injured skin towards repaired barrier function. This sequence of events plays out normally in the vast majority of superficial wounds; however, it can go awry at numerous steps along the pathway, especially with underlying disease states such as diabetes. When wound healing does not progress normally, a chronic wound may result and this is at significant burden to both the patient and the medical system. It has been estimated that a single diabetic ulcer carries a cost of nearly US\$50,000⁽¹⁾ and chronic wounds as a whole cost the medical system over US\$25 billion per year, with the number of patients affected growing yearly from 6.5 million, given the increasing prevalence of diabetes and other chronic diseases that may affect wound healing ⁽²⁾. Aside from the

burden of a chronic wound, even simple wounds created after minor procedures such as outpatient surgeries require proper attention and care, and understandably, patients may be concerned about possible resultant scarring. It is no surprise, then, that wound healing has received a great deal of attention, both from a basic science standpoint and a business perspective. The basic science of how a wound heals is fascinating, with new discoveries elucidating mechanisms of physiologic wound repair constantly being reported. Meanwhile, wound healing is also a huge commercial enterprise, with the market for wound care products exceeding US\$15 billion and treating wound scarring another US\$12 billion⁽³⁾. In this review, basic concepts of wound healing will be discussed, with a focus on current practice in treatment of wounds and future directions in wound care. This article is based on previously conducted studies and does not involve any new studies of human or animal subjects performed by any of the authors.

Physiologic Wound Healing

After a superficial wound, a myriad of systems are activated at the site in order to clear foreign material, as the primary barrier function of the skin is lost, and to eventually restore the normal structure of the skin. While this may only be successful to a limited degree—a wound will never reach the maximum tensile strength of unwounded skin, and at best reaches about 70% ⁽⁴⁾-most of the essential functions of the skin will be returned to a wound. This does, however, require the delivery of various inflammatory cells, chemokines, cytokines, matrix molecules, and nutrients to the wound site with a concordant increase in metabolic demand. These processes occur simultaneously and are generally divided into three main phases of wound healing: inflammatory, proliferative, and remodeling.

The inflammatory phase of wound healing starts shortly after hemostasis is achieved, and the primary goal of this phase is to clear pathogens as well as foreign material from the wound and to contain the damage to a localized area. Vascular permeability increases with vasodilation, allowing neutrophils and monocytes to localize to the wound site. A complex interplay of cytokines also helps to regulate this phase, culminating in monocyte conversion to macrophages, often thought of as the master regulator of this inflammatory phase of wound healing ^[5]. The macrophages not only phagocytose and digest tissue debris and remaining neutrophils but also secrete growth factors and cytokines that promote tissue proliferation and cell migration. After about 3 days from the initial wound, the proliferative phase centers around fibroblasts and production of both collagen and ground substance that will form the basis for the tissue scaffold of the previous wound area. Meanwhile, endothelial cells enter a rapid growth phase and angiogenesis occurs within the granulation tissue, creating a rich vascular network supplying this very active area of healing. After about 2-3 weeks, the wound transitions to a remodeling, or maturation, phase where the collagen type is restored to usual (type I, rather than type III seen in a new wound) ⁽⁶⁾ and the wound tissue matures, resulting in full cross-linking and restoration of a somewhat normal structure. The vascular network rapidly regresses as well ⁽⁷⁾. As previously discussed, the wound strength never reaches its normal, preinjury state.

An important consideration In physiologic wound healing is oxygen supply and oxygen tension in the wound bed. Wound healing requires oxygen to interact with numerous cytokines, supply the actively proliferating cells, as well as provide an effector for the neutrophil respiratory burst. It has been estimated that a wound requires at least a tissue oxygen tension of 20 mmHg to heal⁽⁸⁾ and non-healing wounds have been measured to have oxygen tensions as low as 5 mmHg ⁽⁹⁾. These effects seem to compound one another-in situations of low oxygen tension, not only will there be more necrotic debris to facilitate bacterial growth but the primary mechanism of the immune system in combating these microbes is compromised. Thus, special care must be taken with wounds resulting from peripheral vascular disease and also in cases where vascular compromise may play a role, such as a diabetic ulcer. Additionally, the systemic optimization of nutritional status should be evaluated in wound healing ⁽¹⁰⁾. Numerous nutrients have proven to be important in wound healing, especially protein intake . This was illustrated in a study of elderly patients with pressure ulcers where change in ulcer area was significantly correlated with protein intake ⁽¹¹⁾,however, other factors are also important, such as vitamins A and C, and zinc ⁽¹²⁾. The fine balance of these nutrients has to be taken into account, though. For example, with vitamin E, an important lipophilic antioxidant, conflicting studies have both shown reduced tensile strength and collagen content of experimental wounds ^(13 14) and, conversely, increased wound strength and collagen ⁽¹⁵⁾. As with all aspects of wound repair, a fine balance is necessary to ultimately achieve proper healing.

Given the tight regulation of a multitude of factors required for proper wound healing, it is not surprising that chronic wounds are rather common. After an acute wound such as trauma, surgery, or even a bug bite, the above wellcoordinated series of events come into play. The ultimate time course and outcome will depend on the nature of the acute wound-its location, size, depth, and type. However, when other pathologic factors come into play, such as an underlying disease state, a chronic wound can form (see Fig. 1). This refers to a wound that has somehow deviated from the previously described natural physiologic course of events and has stalled at some point. The underlying mechanism varies greatly, but includes factors influencing blood supply (peripheral vascular disease), immune function (such as immunosuppression or acquired immunodeficiency), metabolic diseases (such as diabetes), medications, or previous local tissue injury (such as radiation therapy). External factors, such as sustained pressure, temperature, and moisture, also play an important role in allowing a wound to heal. As the pathophysiology of normal acute wound healing has been well described, this review will mainly

focus on chronic wounds and their treatment.



Figure:-1 Chronic ulcers of peripheral vascular disease

Aberrant wound healing can be seen rarely in normal healthy subjects, but is usually associated with an underlying process, ranging from diabetes to cancer to malnutrition. Of the major concerns for chronic wounds, perhaps none is as menacing or as important as diabetes. In the USA, the number of people with diabetes already reaches 20 million and is expected to double by the year 2030 ⁽¹⁶⁾. Diabetic foot ulcers affect 15% of these patients and precede the vast majority of amputations in this patient population ⁽¹⁷⁾. A wide variety of factors is thought to contribute to this problem, affecting all phases of wound healing and seemingly nearly every molecule involved in this process ⁽¹⁸⁾, and evidence is emerging that proper glycemic control can have a significant

impact on the rate of wound healing in a diabetic patient ^{(19).} This effect was mainly seen in patients with markedly elevated hemoglobin A1c levels, but it underscores emerging evidence that diabetes plays a multifactorial role in wound healing. It has been well described that the neuropathy experienced by diabetics can lead to a loss of protective sensation, producing wounds that eventuate into ulcers; however, there is an emerging role of advanced glycation end products in contributing to not only this neuropathy but also the wound healing cascade and small vessel disease ⁽²⁰⁾. While achieving good glycemic control can certainly be challenging, it also serves to reinforce the need to address lifestyle choices and integration of the primary care provider into treating a chronic wound.

Wound Infection

The most common preventable challenge to wound healing is possible infection, and topical antimicrobials have long been used empirically to attempt to prevent wound infection. While bacteria are a normal part of the skin flora and thus wounds, a critical threshold of 105 bacteria has been proposed as the delineation between colonization and a clinically relevant infection that may impede wound heimportance ⁽²¹⁾. It is also necessary to distinguish between an incidental positive culture and a true pathogen affecting a wound. Repeat surface cultures in a wound are of limited use, neither confirming nor ruling out a continued infection; rather, clinical diagnosis of an infected wound remains of primary importance ⁽²²⁾. Deep tissue cultures are somewhat more controversial. While they have better sensitivity and specificity in isolating a causative organism in an infected wound, it is still not perfect; isolates from different parts of the same wound have even been shown to have different organisms ⁽²³⁾. Additionally, the practitioner is, in essence, exacerbating the initial wound with an even deeper wound, but this may still be a worthwhile trade-off if it guarantees appropriate antimicrobial coverage.

There are many approaches towards both treatment and prevention of wound infections. Silver has been used as adjunct in wound care for over 2000 years ⁽²⁴⁾ and remains a popular wound care ingredient today. It has a broad spectrum of activity and is available in numerous forms. Newer advances in using silver for wound healing have focused on allowing for sustained release of silver in high enough concentrations to allow for retained efficacy. Nanocrystalline silver dressings were developed with this in mind and help to address the shortcoming that silver nitrate has-to work properly, it would have to be administered 12 times a day ⁽²⁵⁾. Furthermore, a recent review found no convincing evidence that silver sulfadiazine has any effect on wound healing overall, despite its common use among practitioners ⁽²⁶⁾. Similarly, iodine-containing compounds have long been used in wound healing but there have been some concerns with toxicity of iodine-containing compounds, especially over large wound areas. For limited wounds, though, cadexomer iodine (iodine within a starch lattice formed into microbeads) has a good deal of data supporting its use as a cost-effective adjuvant for wound healing (27 28).

Numerous topical formulations of antibiotics have also been developed to apply to wound sites. They remain popular, even though emerging evidence has shown that the benefit of

this wholesale application of antibiotic ointments may not be necessary and the only real indication for topical antibiotics is a clinically infected wound, such as purulent drainage, erythema, warmth, pain, tenderness, or induration ⁽²⁹⁾ Numerous recent studies have echoed this sentiment, with routine administration of antibiotic ointment leading to no better outcomes but often resulting in patient discomfort, along with the possibility of antibiotic resistance and contact dermatitis ^(30 31). This is in conflict with a few earlier studies where children with minor scratches and insect bites had reduced rates of infection with topical antibiotic ointment ⁽³²⁾ ³³⁾, although this cannot be clearly generalized to all patients. Even after Mohs micrographic surgery, a prospective study found the rate of infection after clean surgical technique to be less than 1%, with the highest rate of infections in flap closures ⁽³⁴⁾. Overall, the consensus seems to be that in dermatology, use of topical antibiotics should be reserved for conditions such as impetigo or a clearly infected wound and not for general prophylaxis ⁽³⁵⁾.

Wound Care and Dressing

Wound care has become increasingly important given the rise of chronic wounds and the morbidity associated with them. An important concept in wound care is the role of debridement, or the removal of non-viable tissue material. This can be achieved through surgical or autolytic/enzymatic mechanisms in either case, the goal is to expose healthy, well-perfused tissue that is able to proliferate and populate the wound bed via epithelial cell migration, rather than keeping necrotic debris which only serves as fuel for infection and impedes wound healing. The optimal timing and frequency of surgical debridement are still unclear, as they are likely to vary greatly depending on the type of wound being treated, but there is general agreement that surgical debridement is an important component of wound care. Autolytic debridement refers to the self-activation of endogenous enzymes involved in fibrin degradation generated in a moist wound environment and seen with some types of wound dressings ⁽³⁶⁾. While this can have some utility in wound healing, it is certainly not capable of removing devitalized tissue as well as surgical debridement and, as such, cannot serve as adequate replacement for surgical debridement. Recently, there has been some renewed interest in the use of so-called bio surgical debridement, or the application of larvae/maggots to a wound. This is intriguing in that it achieves both a mechanical/surgical debridement and an enzymatic debridement at the same time, whilst having the capability to eliminate pathogenic organisms and stimulate fibroblast proliferation (37 38). Further studies are needed, but it is certainly an intriguing concept.

Many wound dressings have been developed to try to both protect the healing wound from infection and also to help promote the wound healing process itself (Table 1). A moist occlusive dressing helps support the inflammatory phase by creating an environment with low oxygen tension (thereby activating such factors as hypoxia-inducible factor-1) ⁽³⁹⁾ and also increases the rate of re-epithelialization ⁽⁴⁰⁾. Additionally, a limited amount of exudate retained on the wound allows for autolytic debridement, which serves to further promote successful wound healing. However, traditional dry gauze wound dressings may degrade this

[97]

process while also causing further injury when removed. Low adherent dressings and semipermeable films (i.e., Tegaderm) represent the basic types of wound dressings commonly in use, with the goal of restricting liquid and microbial penetration but allowing air and water vapor through. Hydrocolloids and hydrogels take advantage of a hydrophilic material that absorbs a certain amount of exudate but keep a moist environment; hydrocolloids are furthermore impermeable to air and are somewhat more long-lasting, but should not be used on exudative wounds because of its impermeable nature. Hydrogels may additionally be used to help promote moisture in an otherwise dry wound. Another option is alginate dressings, seaweed-derived non-woven fibers that are generally reserved for highly exudative wounds because of their ability to absorb large amounts of fluid. As such, adverse effects can be seen in dry wounds dressed with alginate ⁽⁴¹⁾. Similarly, foams have some absorptive capacity and can be used on moderately exudative wounds, especially helpful because they minimize trauma during dressing changes. Lastly, collagen products have been used on recalcitrant wounds and chronic ulcers. While this collagen is not intended to be a direct replacement for new production of collagen in wounded tissue (as it can be derived from multiple sources, including bovine and porcine collagen), it is thought to help facilitate an environment attracting cell types critical to wound healing while depleting negative effectors such as free radicals and proteases ⁽⁴²⁾.

Table 1: Currently	available	wound	dressings
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pe of dressing	mmercially available products	tes	
uze	rity, Vaseline Gauze, Xeroform	xpensive, drying, may cause further injury on changin	
ms	pclusive, Blisterfilm, Cutifilm, Flexigrid, OpSite, Tegaderm	clusive, retains moisture, only for non-exudative woun	
drocolloids	uacel, Comfeel, DuoDERM, Granuflex, Tegasorb	ng times between changes, fluid-trapping, occlusive, ected wounds	
drogels	rrasyn, Curagel, Nu-Gel, Purilon, Restore, SAF-gel, XCell	hydrates dry wounds, easy removal/changes, may caus tration	
ams	I Adhesive Foam, Allevyn, Lyofoam, Tielle	derately absorbent, insulating	
ginates	gisite, Kaltostat, Sorbsan, Tegagen	ghly absorbent, hemostatic	
drofibers	uacel Hydrofiber	ghly absorbent	
sue engineered skin su	ibstitutioderm, Apligraf, Biobrane, Bioseed, Dermagraft, Epicel, E alograft, Integra Omnigraft, Laserskin, Myskin, Trans Cyte	Z dresses deficient growth factors and cytokines, exp c of infection, antigenicity	

Several more recent developments in wound dressings have focused on integrating antimicrobial compounds into the wound dressing itself. These materials combine traditional wound dressings such as foams or hydrogels with antimicrobial compounds such as silver, betaine, chitin, or polyhexamethylene biguanide (Kendall AMD). These materials, as discussed previously, may not be appropriate for broad-spectrum application to healing wounds, but may be appropriate in chronic lower leg ulcers where infection can be a problem, especially with formation of biofilms. Many pathogens are able to adhere together into biofilms, which represent tightly packed masses surrounding a polymeric matrix, thus helping to evade destruction by antibiotics. This creates not only a physical barrier to wound healing but one in which the normal resolution of the inflammatory phase may be prolonged, and addressing biofilms has become a major challenge in wound healing. Both in vitro tests and patient data show that these antimicrobial wound dressings can be helpful in chronic pressure/venous ulcers that may be affected by these biofilms (43 44)

Skin Substitute

While skin substitutes have long been used in grafts (mostly from autologous sources) either for significant surgical defects or burns necessitating replacement of a large surface area of tissue, advances in this field have allowed for the development of novel bioprosthetic skin substitutes. These materials generally consist of a biologically derived

substance combined with a material to allow for its placement on a wound. Overall, these dressings are quite costly, representing a significant barrier to widespread adoption. However, several studies have been conducted to look at cost-effectiveness and many of these have shown an these expensive substitutes, advantage with using considering that reducing even a single day of a hospital admission represents an enormous cost saving, not to mention being much better for the patient. While multiple options have focused on a mesh material coated with porcine collagen or polypeptides (Biobrane and Transcyte, the latter of which also contains newborn fibroblast cells) or a porcine xenograft (EZ Derm), it is not clear whether they represent a cost-effective option for general application, although one study suggested that it was associated with decreased cost of care and improved outcome in a relatively limited scope of facial burns ⁽⁴⁵⁾. Perhaps more interesting, however, is a material developed using fibroblasts derived from newborn foreskin tissue, extracellular matrix, and a bioabsorbable polyglactin mesh (Dermagraft). The fibroblasts in this material are able to generate growth factors, collagen, cytokines, and glycosaminoglycans to help support the wound healing environment. Numerous studies have borne out its efficacy (46,47), especially for burn wounds and venous/pressure ulcers, but a few downsides are a theoretical risk of rejection (although unreported thus far) and hypersensitivity (as bovine serum may be contained in trace amounts in the preparation). This concept was taken one step further with Apligraf, an allogenic bilayered cultured skin

equivalent, where a dermal layer of cultured fibroblasts and bovine type I collagen is combined with an epidermal layer of cultured keratinocytes, generating a material that can be absorbed much like a graft and generates a similar environment to a normally healing wound. While no dermal appendages or blood vessels are contained, it was shown that this material actually significantly increased blood flow by over 70% to the base of diabetic foot ulcers ⁽⁴⁸⁾. This comes at a significant cost, however, as a single application of Apligraf can cost over US\$1000 for a 7.5-cm-diameter circular disk. Again, as previously noted, this cost has been analyzed, taking into account shortened time to wound closure, and actually resulted in a cost savings over 1 year in wounds that fail to respond to traditional wound care ^[49]. However, this must be taken with the caveat that the study population consisted of recalcitrant, chronic, non-healing wounds, somewhat limiting the applicability in general of these products.

Another option which received a new FDA indication in 2016 for diabetic foot ulcers is the Integra Dermal Regeneration Template (Omnigraft). This material is an bilayer matrix consisting of acellular collagen. glycosaminoglycan, and chondroitin-6-sulfate, with a layer of silicone for barrier function ^[50]. It had been used since 1996 for burn wounds with a later approval for scars from burns, but more recently gained the approval for diabetic foot ulcers on the basis of a pivotal study from 32 sites and over 300 patients. The data showed that treatment with Omnigraft increased wound closure rate by 50% compared to standard of care, with time to wound closure reduced by 5 weeks. A major benefit of Omnigraft is that in the vast majority of cases (72%), only one application of the substrate is needed for complete wound healing, with over 90% of wounds requiring two or less applications ⁽⁵⁰⁾.

Negative Pressure Wound Therapy

Negative pressure wound therapy, or vacuum-assisted closure, has received a good deal of attention since first being used in the USA in 1997. By maintaining a moist environment, optimizing blood flow, removing exudates, and applying pressure to promote wound closure, these devices are able to mitigate numerous factors that may be deficient in a chronic wound. Additionally, numerous studies have shown that these devices are associated with reduced rates of infection in these wounds (51). Meanwhile, dressing changes may be simplified and significant cost savings can be associated with a reduced need for surgical debridement over the long term. As a rule, these devices consist of a foam dressing (that requires changes, up to every other day) and an evacuation tube. Details of the exact material and irrigation fluids that can be run through the system concurrently have been developed, but in general, these devices range from simple portable home units to more complex machines designed for acute care settings such as burn units. One interesting emerging indication for negative pressure ⁽⁵²⁾.

Growth Factors

Growth factors in wound healing have received significant attention in the field of wound healing in the past few decades. However, the only one of these therapies proven to improve healing in a double-blinded randomized controlled trial is platelet-derived growth factor (PDGF) and those results were rather modest (53). Nonetheless, it stands to reason that since a chronic wound contains so many perturbations in growth factors and cytokines, addressing some of these issues may be helpful. For example, a chronic wound tends to show decreased levels of epidermal growth factor (EGF), fibroblast growth factor (FGF), transforming growth factor- β (TGF- β), PDGF, and vascular endothelial growth factor (VEGF) while expressing lower levels of interleukins (IL) 1 and 6, and tumor necrosis factor-a (TNF- α) ⁽⁵⁴⁾. Perhaps most of this is due to the fact that in an environment where so many factors are deficient and dysregulated, simply replacing one is unable to rescue the chronic wound phenotype; alternatively, it is also likely that the "master regulator" of wound healing has yet to be addressed. In any case, the scientific literature is rife with reports of applying a certain growth factor, cytokine, protein, or hormone and achieving remarkable results in various models for wound healing, including all of the aforementioned growth factors and additional targets of interest ^(55,56). Nonetheless, broad acceptance of these factors has remained lukewarm at best, perhaps limited in large by the cost of these factors. The data is also often difficult to interpret, as different models of wound healing are used and practical issues often limit the ability to control the study. It is worth noting that several insurance companies have begun to accept at least some forms of therapy, such as Regranex (becaplermin gel-a PDGF). A review of United Healthcare and Blue Cross Blue Shield policies reveals that at least some of these providers will reimburse for these growth factors given certain conditions (usually diabetic neuropathic ulcers being treated with adjunctive wound care). Work in this field is certainly still progressing, regardless of seemingly promising therapies such as TGF- β supplementation notably failing to produce results such as shown with PDGF⁽⁵⁷⁾.

Hyperbaric oxygen

Hyperbaric oxygen has been used in wound healing on the basis of the principle that it can promote fibroblast proliferation, enhance immune function, and stimulate angiogenesis, among other functions. However, these ideals have not necessarily borne out in practice, leading to some degree of controversy in its use. Importantly, this therapy is applied to the patient in a hyperbaric oxygen chamber as localized delivery of oxygen has not been shown to be effective (Fig. 2), which could lead to significant side effects including myopia, oxygen toxicity in the brain leading to seizures, and pneumothorax ⁽⁵⁸⁾. A Cochrane review was only able to show a modest increase in the likelihood of diabetic foot ulcers healing at 1 year, and even this was based on mostly smaller studies with questionable study design ⁽⁵⁹⁾. At this point in time, it seems that hyperbaric oxygen can only be considered in wounds where an ischemic diabetic ulcer has demonstrated hypoxia (60) and, even then, some of the newer approaches described previously may be of more benefit. This was in some ways underlined in a recent randomized controlled trial where hyperbaric oxygen as an adjunct did help wound healing but even the authors qualified their findings to be applicable only to "selected patients with diabetes"⁽⁶¹⁾. As that study was not powered to look at outcomes such as amputation, a clinical trial is currently underway to answer some of these questions by comparing standard wound care with or without hyperbaric oxygen therapy to prevent amputations in non-healing ulcers in diabetic patients (62)



Figure : 2 Hyperbaric oxygen wound therapy chamber (Sigma 34 Hyperbaric Chamber, Perry Biomedical, Riviera Beach, FL, USA)

Future Direction

With the economic and patient care impacts of wound healing, it comes as no surprise that the field of wound healing research is incredibly active. New research in wound healing comes from seemingly disparate fields ranging from materials science to molecular biology. Some of these have focused on novel molecules of interest in wound healing, such as nitric oxide, with numerous approaches to this end involving innovative material design ^(63,64). While these approaches may be seen as helpful in wound healing and furthermore may, in essence, bypass some of the limitations of single growth factors or cytokines (especially as more broadly acting, gregarious molecules involved in the wound healing cascade are identified), it still will not approach the "holy grail" of wound healing.

It has long been appreciated that fetal wounds do not scar but understanding how this occurs has been much more difficult ⁽⁶⁵⁾. While this is something of a separate issue in terms of wound healing with much of the previous discussion focusing on chronic, non-healing wounds, the idea that a simple wound can heal without scarring or minimal scarring has been enticing nonetheless. In the fetus, a wound heals by regeneration rather than orderly repair ⁽⁶⁶⁾. Perhaps this correlates with the relatively pathogen-free environment seen in utero, and explains why the wound healing "switch" turns on sometime during late pregnancy, in the last trimester ⁽⁶⁷⁾. This has a corollary in the animal kingdom with reptilian limb regeneration and the more recently reported African spiny mouse (Acomys) ⁽⁶⁸⁾. Some researchers have sought to isolate individual factors that are up- or downregulated in the fetal wound, such as TGF, matrix metalloproteinases ⁽⁶⁹⁾, and the such, but perhaps the greatest promise lies in stem cell therapy. Parallel to numerous successful research efforts in chronic wounds focusing on injecting stem cells from bone marrow into non-healing wounds ^(70,71), more recent studies have also focused on using stem cells as scar-attenuating adjuncts to wound healing, leading to tissue regeneration rather than repair with subsequent formation of fibrotic scar tissue ⁽⁷²⁾. Until these therapies bear fruit, however, it may be best to practice the old adage, "protect your skin," and avoid the burden of a wound in the first place.

CONCLUSION

The data indicate a wound healing and antioxidant activities of E. Pruniformis. This study is the first report of flavonoids and wound healing activity of E. Pruniformis.

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