Soft Tissue Wound Healing Review

Introduction

The inflammatory and repair processes are no longer simple events to describe in the light of the increased knowledge in this field. The review that follows is only a brief resume of the salient events associated with tissue repair, particularly concerning the soft tissues. For further information, the reader is referred to recent reviews listed at the end of the paper.

Wound healing refers to the body's replacement of destroyed tissue by living tissue (Walter and Israel 1987) and comprises two essential components - Regeneration and Repair. The differentiation between the two is based on the resultant tissue. In regeneration, specialised tissues is replaced by the proliferation of surrounding undamaged specialised cells. In repair, lost tissue is replaced by granulation tissue which matures to form scar tissue. This review concentrates on the events and processes associated with the repair process.

Probably the most straightforward way to describe the healing process is to divide it up into broad stages which are not mutually exclusive and overlap considerably. There are several different ways to 'divide up' the entire process, but the allocation of 4 phases is common and will be adopted here – these being BLEEDING, INFLAMMATION, PROLIFERATION and REMODELLING.

Overview

A brief overview of each phase is presented here before considering them in any detail. Figure 1 refers to a general arrangement of the phases.

Bleeding Phase

This is a relatively short lived phase, and will occur following injury, trauma or other similar insult. Clearly if there has been no overt injury, this will be of little or no importance, but following soft tissue injury, there will have been some bleeding. The normal time for bleeding to stop will vary with the nature of the injury and the nature of the tissue in question. The more vascular tissues (e.g. muscle) will bleed for longer and there will be a greater escape of blood into the tissues. Other tissues (e.g. ligament) will bleed less (both in terms of duration and volume). It is normally cited that the interval between injury and end of bleeding is a matter of a few hours (6-8 hours is often quoted) though this of course is the average duration after the average injury in the average patient. Some tissues will continue to bleed for a significantly longer period, albeit at a significantly reduced rate. A crush type injury to a more vascular tissue – like muscle - could still be bleeding (minimally) 24 hours or more post trauma.

Figure 1 : Phases of tissue repair

Figure 1 is a gross representation of the key phases of the tissue repair process. The phases identified are shown as separate entities, though in reality, they are interlinked in a very deliberate way. There are events associated with one phase that act as stimulants for the following phase. Some examples of these have been included in the main text.

Inflammatory Phase

The inflammatory phase is an essential component of the tissue repair process and is best regarded in this way rather than as an 'inappropriate reaction' to injury. There are, of course, numerous other initiators of the inflammatory process, though for the purpose of this paper, the injury model will be adopted. The inflammatory phase has a rapid onset
(few hours) and swiftly increases in magnitude to its maximal reaction (2-3 days) before gradually resolving (over the next couple of weeks). It can result in several outcomes (see below) but in terms of tissue repair, it is normal and essential.

**Proliferation Phase**

The proliferative phase essentially involves the generation of the repair material, which for the majority of musculoskeletal injuries, involves the production of scar (collagen) material. The proliferative phase has a rapid onset (24-48 hours) but takes considerably longer to reach its peak reactivity, which is usually between 2-3 weeks post injury (the more vascular the tissue, the shorter the time taken to reach peak proliferative production). This peak in activity does not represent the time at which scar production is complete, but the time phase during which the bulk of the scar material is formed. The production of a final product (a high quality and functional scar) is not achieved until later in the overall repair process. In general terms it is usually considered that proliferation runs from the first day or two post injury through to its peak at 2-3 weeks and decreases thereafter through to a matter of several months post trauma.

**Remodelling Phase**

The remodelling phase is an essential component of tissue repair and is often overlooked in terms of its importance. It is neither swift nor highly reactive, but does result in an organised and functional scar which is capable of behaving in a similar way to the parent tissue (that which it is repairing). The remodelling phase has be widely quoted as starting at around the same time as the peak of the proliferative phase (2-3 weeks post injury), but more recent evidence would support the proposal that the remodelling phase actually starts rather earlier than this, and it would be reasonable to consider the start point at around 1-2 weeks.

The final outcome of these combines events is that the damaged tissue will be repaired with a scar which is not a ‘like for like’ replacement of the original, but does provide a functional, long term ‘mend’ which is capable of enabling quality recovery from injury. For most patients, this is a process that will occur without the need for drugs, therapy or other intervention. It is designed to happen, and for those patients in whom problems are realised, or in whom that magnitude of the damage is sufficient, some ‘help’ may be required in order to facilitate the process. It would be difficult to argue that therapy is ‘essential’ in some sense. The body has an intricately complex and balanced mechanism through which these events are controlled. It is possible however, that in cases of inhibited response, delayed reactions or repeated trauma, therapeutic intervention is of value.

It would also be difficult to argue that there was any need to change the process of tissue repair. If there is an efficient (usually) system through which tissue repair is initiated and controlled, why would there be any reason to change it? The more logical approach would be to facilitate or promote the normality of tissue repair, and thereby enhance the sequence of events that take the tissues from their injured to their ‘normal’ state. This is the argument that will be followed in this paper – the promotion of normality, rather than trying to achieve a better normality.

**Inflammatory Reaction**

Inflammation is a normal and necessary prerequisite to healing (Hardy 1989). Following the tissue bleeding which clearly will vary in extent depending on the nature of the wound, a number of substances will remain in the tissues which make a contribution to the later phases. Fibrin and fibronectin form a substratum which is hospitable to the adhesion of various cells.

The complex chemically mediated amplification cascade that is responsible for both the initiation and control of the inflammatory response can be started by numerous events, one of which is trauma. Mechanical irritation, thermal or chemical insult, and a wide variety of immune responses are some of the alternative initiators, and for a wide range of patients experiencing an inflammatory response in the musculoskeletal tissues, these are more readily identified causes. For the purposes of this review, only the traumatic route will be pursued.

There are two essential elements to the inflammatory events, namely the vascular and cellular cascades. Importantly, these occur in
parallel and are significantly interlinked. Figure 2 summarises the essential elements of the inflammatory cascade. The chemical mediators that make an active contribution to this process are myriad. In recent years, the identification of numerous 'growth factors' had led to several important discoveries and potential new treatment lines (e.g. Wagner et al 2003).

In addition to the vasodilation response, there is an increase in the vasopermeability of the local vessels (also mediated by numerous of the chemical mediators), and thus the combination of the vasodilation and vasopermeability response is that there is an increased flow through vessels which are more 'leaky', resulting in an increased exudate production.

The flow and pressure changes in the vessels allows fluid and the smaller solutes to pass into the tissue spaces. This can occur both at the arterial and venous ends of the capillary network as the increased hydrostatic pressure is sufficient to overcome the osmotic pressure of the plasma proteins. The vessels show a marked increase in permeability to plasma proteins. There are several phases to the permeability changes but essentially, there is a separation of the endothelial cells, particularly of the venules, and an increased escape of protein rich plasma to the interstitial tissue spaces. The chemical mediators responsible for the permeability changes include histamine, serotonin (5-HT), bradykinin and leukotrienes together with a potentiating effect from the prostaglandins.

The effect of the exudate is to dilute any irritant substances in the damaged area and due to the high fibrinogen content of the fluid, a fibrin clot can also form, providing an initial union between the surrounding intact tissues and a meshwork which can trap foreign particles and debris. The meshwork also serves as an aid to phagocytic activity (see below). Mast cells in the damaged region release hyaluronic acid and other proteoglycans which bind with the exudate fluid and create a gel which limits local fluid flow, and further traps various particles and debris (Hardy 1989).

**Figure 2: Inflammatory elements**

**Vascular events**

In addition to the vascular changes associated with bleeding, there are also marked changes in the state of the intact vessels. There are changes in the calibre of the blood vessels, changes in the vessel wall and in the flow of blood through the vessels. Vasodilation follows an initial but brief vasoconstriction and persists for the duration of the inflammatory response. Flow increases through the main channels and additionally, previously dormant capillaries are opened to increase the volume through the capillary bed. The cause of this dilation is primarily by chemical means (histamine, prostaglandins and complement cascade components C3 and C5) whilst the axon reflex and autonomic system exert additional influences. There is an initial increase in velocity of the blood followed by a prolonged slowing of the stream. The white cells marginate, platelets adhere to the vessel walls and the endothelial cells swell.

**Cellular events**

The cellular components of the inflammatory response include the early emigration (within minutes) of the neutrophils (polymorphonucleocytes or PMN's) from the vessels. This is followed by several other species leaving the main flow, including monocytes, lymphocytes, eosinophils, basophils (Lorena et al 2002) and smaller numbers of red cells (though these leave the vessel passively rather than the active
emigration of the white cells). Monocytes once in the tissue spaces become macrophages (Forrest 1983). The main groups of chemical mediators responsible for chemotaxis are some components of the complement cascade, lymphokines, factors released for the PMN’s and peptides released from the mast cells in the damaged tissue (Egozi et al 2003, Vernon-Roberts 1988).

The PMN escapees act as early debriders of the wound. Numerous chemical mediators have been identified as having a chemotactic role, for example, PDGF (platelet derived growth factor) released from damaged platelets in the area. Components of the complement cascade (C3a and C5a), leukotrienes (released from a variety of white cells, macrophages and mast cells) and lymphokines (released from polymorphs) have been identified (see Walter and Israel 1987, Vernon-Roberts 1988, Dierich et al 1987)

These cells exhibit a strong phagocytic activity and are responsible for the essential tissue debridement role. Dead and dying cells, fibrin mesh and clot reside all need to be removed. As a ‘bonus’, one of the chemicals released as an end product of phagocytosis is lactic acid which is one of the stimulants of proliferation – the next sequence of events in the repair process.

The inflammatory response therefore results in a vascular response, a cellular and fluid exudate, with resulting oedema and phagocytic activation. The complex interaction of the chemical mediators not only stimulates various components of the inflammatory phase, but also stimulates the proliferative phase. The course of the inflammatory response will depend upon the number of cells destroyed, the original causation of the process and the tissue condition at the time of insult.

Inflammatory outcomes

**Resolution** is a possible outcome at this stage on condition that less than a critical number of cells have been destroyed. For most patients that come to our attention, this is an unlikely scenario.

**Suppuration**, in the presence of infective microorganisms will result in pus formation. Pus consists of dead cell debris, living, dead and dying polymorphs suspended in the inflammatory exudate. Clearly the presence of an infection will delay the healing of a wound (Zederfelt 1979).

**Chronic inflammation** does not necessarily imply inflammation of long duration, and may follow a transient or prolonged acute inflammatory stage (Vernon-Roberts 1988). Essentially there are two forms of chronic inflammation: either the chronic reaction supervenes on the acute reaction or may in fact develop slowly with no initial acute phase (ab initio) (Hurley 1985). Chronic supervening on acute almost always involves some suppuration whilst chronic ab initio can have many causes including local irritants, poor circulation, some micro-organisms or immune disturbances. Chronic inflammation is usually more productive than exudative - it produces more fibrous material than inflammatory exudate. Frequently there is some tissue destruction, inflammation and attempted healing occurring simultaneously (Hurley 1985, Walters and Israel 1987).

**Healing by fibrosis** will most likely be taking place in the tissue repair scenario considered here. The fibrin deposits from the inflammatory stage will be partly removed by the fibrinolytic enzymes (from the plasma and PMN's) and will be gradually replaced by granulation tissue which becomes organised to form the scar tissue. Macrophages are largely responsible for the removal of the fibrin, allowing capillary budding and fibroblastic activity to proceed (proliferation). The greater the volume of damaged tissue, the greater the extent of, and the greater the density of the resulting scar tissue. Chronic inflammation is usually accompanied by some fibrosis even in the absence of significant tissue destruction (Hurley 1985)

The effects of acute inflammation are largely beneficial. The fluid exudate dilutes the toxins and escaped blood products include antibodies (and systemic drugs). The fibrinogen forms fibrin clots providing a mechanical barrier to the spread of microorganisms (if present) and additionally assists phagocytosis. The gel like consistency of the inflammatory exudate also makes a positive contribution by preventing the spread of the inflammatory mediators to surrounding, intact tissues. Transportation of invading bacteria to the lymphatic system stimulates an immune response whilst the increased blood flow contributes to the increased cell metabolism necessary for the proliferative stage by increasing local oxygen content, supply of necessary nutrients and removal of waste.
products. The leucocytes provide a mechanism for the phagocytosis of foreign material, bacteria, dead cells, with the neutrophils (PMN's) and monocytes (becoming macrophages) making the greatest contribution.

There are several detrimental aspects of inflammation which deserve mention. Firstly the increased local hydrostatic pressure from the oedema can restrict blood flow if the injured tissue space is limited, produce pain and therefore limit function. There have been recent suggestions that free radicals produced as a result of acute inflammatory responses may have detrimental effects on cell membrane processes as may overproduction of lysosomal enzymes from PMN activity.

**Proliferative Phase**

The repair process restores tissue continuity by the deposition of 'repair tissue'. This is initially granulation tissue which matures to form scar tissue. Repair tissue is a connective tissue distinct right from the onset in several ways from the connective tissue native to the site (Forrest 1983). Interesting recent developments have identified that in muscle there is a degree of regenerative activity post trauma, linked to the activation of a mechanosensitive growth factor and subsequent activation of muscle satellite (stem) cells (Hill et al 2003). A range of growth factors have been identified as being active in the processes of proliferation, leading again to some new potential treatments (e.g. Hildebrand et al 1998).

Two fundamental processes involved in the repair are fibroplasia and angiogenesis (Figure 3). The function of the fibroblast is to repair the connective tissue (Vanable 1989). Fibroblasts appear to migrate to the area from surrounding tissue. Fibroblastic activation appears to be chemically mediated, particularly by chemicals released from the macrophages during the inflammatory stage. Fibroblasts migrate into the wounded area and proliferate within the first few days after the tissue damage. Macrophage Derived Growth Factors (MGDF's) are a complex group of mediators responsible, at least in part for the activation of fibroblasts.

Alongside the fibroblastic activation, capillaries in the region of the tissue damage bud and grow towards the repair zone. Loops and arcades are formed together with anastomoses which re-establish a blood flow through the region, providing oxygen and nutrients whilst removing metabolic and repair waste products. Oxygen is critical for many of the reparative processes, but especially for collagen production (Vanables 1989, Niinikoski 1980). A wide range of growth factors and chemical mediators have been identified which exert influences on the developing capillaries. These include macrophage derived factors, PDGF, lactic acid and fibroblast growth factor (Vernon-Roberts 1988). Some of these mediators are produced during the inflammatory phase, thus migrating into damaged tissue from adjacent areas to initiate early wound contraction and increase activity.
making an essential link between the inflammatory and proliferative phases.

Granulation tissue invasion follows the 'demolition' phase (when autolytic enzymes are released from PMN's and dead cells) (Walter and Israel 1987). The activation of fibroblasts and capillary budding would normally occur by about the third day after the tissue insult. The combination of capillary budding and collagen production results in a more vascular than usual repair site. The fibroblasts initially produce predominantly type III collagen which will become type I collagen as the repair matures – during remodelling (Walter and Israel 1987). Fibroblasts also produce fibronectins and proteoglycans which are essential components of the ground substance (Figure 4) (Walter and Israel 1987, Forrest 1983, Hardy 1989).

Myofibroblasts are derived from fibroblasts activated by a variety of chemical mediators, and are responsible for wound contraction and the early strength of the repair. They draw the edges of the wound together, particularly in skin lesions, thus reducing the size of the final scar (Lorena et al 2002, Peacock 1984, Hardy 1989).

Granulation tissue matures with lymphatic development (in much the same way as capillary development), nerve fibre ingrowth and mast cell invasion. Collagen fibres are oriented in response to local stress thus providing tensile strength in the required directions (see Forrest 1983 and Hardy 1989 for useful collagen reviews). As the granulation tissue matures, there is a process of devascularisation with obliteration of the lumen of the vessels.

**Remodelling Phase**

The remodelling phase primarily involves the collagen and its associated extracellular matrix. The initial deposition of collagen produces relatively weak fibrils with random orientation. With maturity, the collagen becomes more obviously oriented in line with local stresses (Culav et al 1999, Gomez et al 1991). A proportion of the original fine (Type III) collagen is reabsorbed (due to the action of collagenases) and is replaced with Type I collagen with more cross links and greater tensile strength (Vanables 1989, Forrest 1983). Collagen synthesis and lysis both occur at a greater rate in a normal wound compared with non wounded tissue as old fibrous tissue is removed and new scar tissue is laid down. The maturing scar is therefore a dynamic system rather than a static one. There are several influential factors during this long phase, including physical stress. This remodelling process is initiated whilst the proliferative stage proceeds, therefore providing a considerable overlap between the phases. Final remodelling may continue for months, and possibly over a year beyond the obvious healing of the damage. See Hardy (1989) for a comprehensive consideration of collagen behaviour in remodelling and Culav et al 1999 for an excellent review of collagen and its roles).

Factors known to delay healing are divided into general and local:

**General:** Age, Protein deficiency, Low Vitamin C levels, Steroids & NSAID's (inhibitory effect), Temperature (lower rate when colder)

**Local:** Poor blood supply / ischaemia, Adhesion to bone or other underlying tissue, Continued inflammation, Drying of the wound, Excessive movement (restarts inflammation)

**Conclusion:** Tissue healing is a complex and dynamic system which enables effective repair of damaged tissue. There is little doubt that appropriate therapy has the capacity to influence the process in a positive way.
References:


Egozi E et al 2003; Mast cells modulate the inflammatory but not the proliferative response in healing wounds. Wound Repair & Regeneration 11(1):46-54


