Management of leg ulcers

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Abstract
Leg ulcer is a leading cause of morbidity among older subjects, especially women in the Western world. About 400 years BC, Hippocrates wrote, “In case of an ulcer, it is not expedient to stand, especially if the ulcer be situated on the leg”. Hippocrates himself had a leg ulcer. The best treatment of any leg ulcer depends upon the accurate diagnosis and the underlying aetiology. The majority of leg ulcers are due to venous disease and/or arterial disease, but the treatment of the underlying cause is far more important than the choice of dressing. The aetiology, pathogenesis, treatment, and the future trends in the management of the leg ulcers are discussed in this review.

Keywords: leg ulcers; compression bandages; skin grafting

There are about 400 000 patients with leg ulcer disease in the UK and at any one time 100 000 have open leg ulcers requiring treatment. This costs about £600 million/year. Venous ulceration of the lower leg is the result of increased venous pressure and its secondary effects on the microvascular system. In patients with venous leg ulcers, nearly half have evidence of a past venous thrombosis while the remainder results from incompetence of valves of the superficial or communicating veins. The venous leg ulcer is an age related disease in the elderly population, especially women. It may be associated with lipodermatosclerosis and eczema. Whereas the venous leg ulcer is usually originated by external trauma, the course is often chronic and/or relapsing.

Epidemiology

INCIDENCE
Community surveys in Lothian, north west London, and Southampton all suggest an overall incidence of about 0.2%. There are about 400 000 patients with leg ulcer disease in the UK and at any one time 100 000 have open leg ulcers requiring treatment. This costs about £600 million/year. Venous ulceration of the lower leg is the result of increased venous pressure and its secondary effects on the microvascular system. In patients with venous leg ulcers, nearly half have evidence of a past venous thrombosis while the remainder results from incompetence of valves of the superficial or communicating veins. The venous leg ulcer is an age related disease in the elderly population, especially women. It may be associated with lipodermatosclerosis and eczema. Whereas the venous leg ulcer is usually originated by external trauma, the course is often chronic and/or relapsing.

PREVALENCE
The prevalence of leg ulcers in Europe is well documented, varying between 0.18% and 1% in different countries. In the UK two major studies demonstrated the prevalence of active ulcers at 0.15% to 0.18% in contrast to 1% in the Scandinavian countries. Leg ulcers occur more commonly in elderly people and their prevalence is likely to increase as the average age of the population increases.

Aetiology

In the Western world, leg ulcers are mainly caused by venous insufficiency, arterial insufficiency, neuropathy, diabetes, or a combination of these factors (table 1). Venous ulcers are the most common type of leg ulcers, accounting for approximately 70% of cases (fig 1). Arterial disease accounts for another 5% to 10% of leg ulcers; most of the others are due to either neuropathy (usually diabetic) or a combination of those diseases.

Pathogenesis of venous leg ulcers

The association between ulceration at the ankle and venous disorders of the lower limbs has been known for more than 2000 years. This connection between deep vein damage and ulceration was noted by Gay and later by...
Homans, who also noticed that venous ulcers often had few visible varicose veins.

The concept of venous stasis suggested that stagnant blood lying within tortuous and dilated veins close to the skin can cause tissue anoxia and cell death. Recently, it has been proposed that the preceding stage of venous ulceration represents a scleroderma-like skin called lipodermatosclerosis. There is a growing recognition that an excessive proteolytic activity by proteases, especially that of matrix metalloproteinases (MMP1, MMP2, MMP8, and MMP9) and fibrinolytic factors of the plasminogen activation system may be a key feature in the pathogenesis of venous leg ulceration. Elevated expression on mRNA and protein level of MMPs and fibrinolytic factors have been detected in liposclerotic skin lesions. Furthermore, MMP2 was imbalanced by locally reduced expression of tissue inhibitors of metalloproteinases (TIMP2) in the basement membrane zone of skin lesions. MMP8 is the predominant collagenase in healing wounds and non-healing ulcers. But in diabetic foot ulcers, increased nitric oxide synthase activity may be responsible for the impaired healing. Furthermore, the increased activity of arginase could account for the characteristic callus formation around the ulcers. In addition, the lower concentrations of transforming growth factor-β1 in diabetic ulcers may explain the raised concentrations of nitric oxide in this condition.

Loots and colleagues demonstrated the important differences in cellular infiltrates and extracellular matrix of chronic diabetic and venous ulcers compared with acute wounds. The CD4/CD8 ratio in chronic ulcers was significantly lower (p<0.0027) due to a relatively lower number of CD4+ T-cells. There was a significantly higher number of macrophages in the edges of both types of ulcers. Hahn et al found that the upregulation of endothelial adhesion molecules (ICAM-1) and dermal infiltration by T-lymphocytes and macrophages in chronic venous insufficiency patients is limited to the region of the ulcer, or at least to skin areas with a severe microangiopathy, and is part of a secondary elimination of necrotic tissue (an “injury and repair” process).

In patients with venous disease, there is incompetence of the valves within the perforating veins connecting the superficial to the deep venous systems in the leg. In normal subjects, venous pressure decreases during exercise. In patients with venous incompetence, pressure remains high during exertion. High venous pressure is associated with capillary proliferation and increased permeability of large molecules into the skin. It is not clear how venous hypertension is transmitted into skin ulceration, but it is apparent that venous ulcers represent a disease of the cutaneous microcirculation as there is rarefaction and dilatation of the skin capillaries (atrophie blanche) together with the changes in the microlymphatics. In 1982, Browse and Burnand proposed that the high ambulatory venous pressure within the calf muscle is transmitted to the capillary circulation in the skin and subcutaneous tissues of the calf. They hypothesised that the distended local capillary bed widened the endothelial pores, allowing fibrinogen molecules to escape into the extracellular fluid, where they form fibrin complexes and plug the capillaries. Pericapillary fibrin has been shown to impede oxygen diffusion in the patients with venous disease.

Various theories have been put forward—for example, venous stagnation, fibrin cuff, white cell trapping, and growth factor trapping theories. In the “venous stagnation” theory, blood stagnates within the veins leading to local ischaemia and ulceration. The “fibrin cuff” theory states that increased venous pressure causes leakage of macromolecules from plasma into perivascular space when fibrinogen changed to fibrin and this acts as a barrier to diffusion of oxygen and nutrients. It is now recognised that the fibrin cuff is seen as an effect but not a cause of the disease, as these lesions are seen in many other chronic wound situations. Pericapillary fibrin cuffs remain a prominent feature, whether they act as a barrier, a marker for endothelial cell damage, or as part of an overall mechanism of macromolecular leakage and trapping.

A recent hypothesis proposed that in venous ulceration, trapped leucocytes occlude capillaries, thereby resulting in cutaneous ischaemia. White cells are normally trapped within the cutaneous microcirculation when legs are dependent. In venous hypertension, these white cells are activated leading to damage and destruction of the capillaries. Falanga and Eaglistein have proposed that macromolecules leaking into the dermis “trap” growth factors and matrix material leading to inadequate tissue repair, which eventually causes ulceration.

The skin may break down, due to three mechanisms: (a) damage to blood vessels leading to ischaemia, (b) inflammation, and/or (c) infiltration. In many instances, it is the inability of the damaged skin to repair itself normally that results in ulceration.

Similar to fibrin cuff theory, growth factors and cytokines are held in the fibrin cuff preventing the maintenance of the normal homoeostatic environment.

More recently, Birgnette Maessen-Visch and colleagues claimed that factor V Leiden mutation is more frequent in patients with venous leg ulceration than in the control group and the general population. Patients with factor V Leiden mutation have an increased risk of developing deep vein thrombosis and recurrent leg ulceration.

Danielsen et al demonstrated that the exotoxin A serum antibodies in patients with chronic leg ulcers, infected with Pseudomonas aeruginosa, may provide a marker for ulcer deterioration, and may support the decision for skin grafting.

There are many predisposing factors, which may prevent healing of leg ulcers, shown in table 2.
Pathogenesis of arterial (ischaemic) ulceration
Arterial or arteriolar occlusion due to any cause can result in ischaemia of the skin and subcutaneous tissues which might lead to ulceration. Peripheral vascular disease due to atherosclerosis, diabetes with microvascular or macrovascular disease and/or vasculitis could lead to ischaemic leg resulting in ulceration.25

There are three mechanisms involved in the pathophysiology of ischaemic leg ulcer25:
(1) Extramural strangulation
(2) Mural thickening or accretion
(3) Intramural restriction of blood flow
Extramural strangulation is usually due to scar tissue and radiodermatitis causing fibrotic bands around the arterioles which may give rise to small but persistent ischaemic ulceration. Mural thickening or accretion of intimal plaques as for example in atherosclerosis, may proceed with an impaired blood flow until atherothrombosis, embolism, or superimposed infection may precipitate complete occlusion, resulting in ulceration. Intramural changes could occlude the small vessels by changes in blood viscosity, platelet adhesiveness and/or fibrinogenesis as in vasculitis, which might lead to leg ulceration.

There is often considerable overlap and the exact pathogenesis cannot be always well defined. Most acute forms of vasculitis and some subacute and chronic forms are likely to cause leg ulceration due to tissue hypoxia and exudation of fibrin-like substances.25

Diabetic foot ulcer
There is usually small vessel disease which interferes with peripheral circulation leading to tissue hypoxia, which may result in ulceration after minor or major trauma. Peripheral gangrene may occur as a large single ulcer characteristically situated at the side or at the back of the ankle due to atherosclerosis. Peripheral neuropathy may coexist, with cold, swollen, and dry feet. Trivial trauma or blister due to any cause may lead to ulceration.26

Haematological disorders
Indolent, non-healing ulcers are a feature of sickle cell disease,27 thalassaemia, and other haemolytic anaemias. These are usually due to blockage of microcirculation. Thrombotic and occlusive diseases—for example, antiphospholipid syndrome, protein C and protein S deficiency, cryoglobulinaemia, etc, lead to rapid skin necrosis and gangrene.26

Familial leg ulcers caused by genetic disorders of coagulation, causing recurrent familial venous thrombosis, may occur as a result of mutations in protein C, protein S, antithrombin III, fibrinogen, and factor V genes.28 30

Infections
Infections may lead to ulceration that is often slow in healing because of associated oedema, cellulitis, thrombophlebitis, or underlying vascular disease or diabetes. Bacterial synergistic gangrene (Meloney’s ulcer) extends rapidly and usually has a burrowing, bluish, and undermined edge.29 Primary uncomplicated streptococcal ulcer is rare, but Streptococcus pyogenes or Staphylococcus aureus may complicate existing ulceration. Tuberculous cutaneous ulcer might occur in erythema induratum or Bazin’s disease, situated usually on the back of the calves.30

Hypertensive ulcer Synonymous Martorell’s ulcer
In hypertension, the density of the capillary beds in the middle and deep dermis of the skin particularly of its lateral aspect is much reduced, which may predispose to ischaemic necrosis when the sparse arterioles are subjected to severe hypertension.31 Hypertension produces superficial ulceration,32 when peripheral pulses are always present, distinguishing the condition from peripheral vascular disease.

Rheumatoid arthritis/vasculitis
The leg ulcer usually develops on the shin or ankle after trauma, is related to vasculitis, and often difficult to heal. Delayed healing is also due to impaired mobility which contributes to poor muscle pump.33 The oedema is often difficult to control. Ulceration of rheumatoid nodules is not common except at pressure points.34

Trophic ulcers
These ulcers may occur as a result of pressure or friction on areas which have become anaesthetic due to peripheral neuropathy. Sometimes they may be associated with underlying vascular abnormalities—for example, diabetes causing small vessel disease.35

Ulceration due to dermatitis artefacta
Ulceration secondary to artefact may be difficult to diagnose. Clinical suspicion might be raised by the following: unusual appearance, unusual site, after trauma at work with possible compensation issues, angulate shape or the apparent indifference of the patient to their ulcer and its implication to their life.29 If the ulcer tends to come and go, a skin biopsy may be necessary to show up “out-side-in” damage, that is, disproportionate damage to the epidermis compared with the dermis.26

Pyoderma gangrenosum
Pyoderma gangrenosum is a relatively common destructive non-infective ulceration of the skin with sloughy base and ragged undermined edge. It may be associated with chronic conditions such as inflammatory bowel disease, inflammatory arthropathies, or myeloproliferative disorders.26 A skin biopsy may be helpful
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**Treatment of leg ulcers**

**INITIAL ASSESSMENT**

Assessment of a patient with leg ulcer consists of the following:

- Comprehensive clinical history and physical examination in order to determine the aetiology of the ulcer.
- Doppler measurement of the pressure at the ankle (dorsalis pedis or posterior tibial artery) and brachial artery to determine ankle:brachial pressure index (if \(< 0.8 = \) presence of arterial insufficiency, if \(>1.0 = \) venous insufficiency, and if \(<0.5 = \) need for urgent referral to a vascular surgeon).
- The wound size and shape must be documented either by measuring the length and width of the ulcer by tracing onto an acetate sheet or by photographic record.
- The ulcer should be swabbed for microbiology.
- Full blood count, random blood sugar, chemical profile, and plasma viscosity should be done.
- Ulcers of unusual appearance should be biopsied.
- Deep ulcers should have plain radiography of the affected areas and computed tomography, magnetic resonance imaging, etc to exclude osteomyelitis, especially in chronic non-healing ulcers.
- Patch tests may be indicated if there is an extensive eczema or local allergy.

**SPECIFIC TREATMENT**

Primary dressing depends on the conditions of the ulcer base. Simple non-adherent gauze should be used for shallow, non-exudating ulcer.\(^4\) Absorbent foam or alginate dressings should be used for exuding ulcers, and hydrocolloids reserved for sloughy, necrotic ulcers. Hyaluronic acid dressing is often used for sloughy, necrotic ulcers.

It is often necessary to treat *S aureus* or *S pyogenes* (haemolytic streptococci) if signs of inflammation are present. Larval (maggot) therapy is now available for debriding resistant sloughy ulcers.\(^5\) Indications of biosurgery (larval therapy) include necrotic or sloughy ulcers.\(^6\) Its preferential indications of the VAC device include leg ulcers, pressure sores, burns, wounds with skin defects, complications of surgical wounds, and delayed healing.\(^7\)

**SECONDARY DRESSING**

Secondary dressing with different types of compression bandages\(^8\) (three layers, four layers of bandages, for example, Charing Cross method) is crucial to the healing of venous ulcers (these bandages must not be used if there is arterial insufficiency).\(^9\) Community based leg ulcer clinics with trained nurses using a four layer bandage is more effective than traditional home based treatment and this is cost effective.\(^10\) A survey of the management of leg ulcers in primary care settings in Scotland revealed that compression bandages were routinely used by 64% of nurses.\(^11\)

**GENERAL MEASURES**

These consist of avoidance of prolonged standing or sitting with the feet dependent. Periods of rest with the legs elevated and graded exercises to improve the leg muscles pump should be practised. Once the ulcer is healed, the patient should wear elastic support hosiery.\(^12\)

The severity of the venous insufficiency and the type of lifestyle ultimately determine the type of hosiery. There are three categories of compression hosiery conforming to the British Standard Institution standard\(^13\) (light pressure at ankle (12 mm Hg/1.6 kPa); medium (20 mm Hg/2.7 kPa) and strong (30 mm Hg/4.0 kPa)). Light compression is mostly inadequate whereas strong compression is effective, but uncomfortable and difficult for many elderly people. European standard compression hosiery is increasingly used. There are four classes of compression: light compression at the ankle (18.4–21.1 mm Hg/2.5–2.8 kPa); medium (25.2–32.3 mm Hg/3.3–4.3 kPa); strong (36.5–46.6 mm Hg/4.9–6.2 kPa); and very strong (>50 mm Hg/7.9 kPa). Light compression hosiery is indicated for treating mild varicosis; medium compression for pronounced varicosis with oedema, after healing of minor leg ulcer and after thrombophlebitis; strong compression for late complications of constitutional or post-thrombotic venous insufficiency, atrophic blanche, dermatosclerosis and after healing of severe or recurrent ulcers; very strong for lymphoedema and elephantiasis. The ideal is “made to measure” for the patient and to receive assistance with dressing. There are many aids available—for example,
Medivalet with short handles, Medi-Export valet with long handles, Medivalet with variable handles, Medi-Hospital valet with adjustable diameter, and Medi-Magnum valet, especially for larger stocking sizes. The dressing aid solves the often difficult problem of putting on compression hosiery.

Patients should stop smoking, if there is an associated arterial disease. Factors causing delay in healing leg ulcers include poor control of leg oedema because of inadequate compression, unrecognised arterial disease, continued smoking, secondary infection (S aureus), anaemia, and/or heart failure. Adequate analgesics should be prescribed if the leg ulcers are painful. The Cadexomer Iodine Study Group demonstrated that cadexomer iodine paste is desirable.58

Treatment of arterial ulcers
The most important aim in the treatment of arterial ulcers is to increase the blood supply to the affected area.59 The patient must stop smoking and diabetes or hypertension should be well controlled. Regular graded exercises should be encouraged to promote development of collateral circulation. Pressure areas may be protected by sheepskin or a special cushion with bony prominences, for example heel, malleoli, etc, being the vulnerable areas for leg ulcers. The head of the patient’s bed should be raised by 4–6 inches to encourage gravity dependent arterial flow and the limbs should be kept warm. Regular use of analgesics should be encouraged for the relief of pain, but if rest pain or acute infection is present, the patient should be referred to a vascular surgeon. Arterial reconstruction surgery may be performed to salvage the limb. After angiographic studies, endarterectomy to remove the clot or atheromatous plaques and/or reconstruction to bypass occluded artery may be performed.59 Balloon angioplasty may be the intervention of choice for ischaemic rest pain, ulceration, and gangrene of the leg. If angioplasty fails, femoropopliteal bypass graft may be tried.59 Cellulitis with or without lymphangitis needs to be treated with systemic antibiotics.

Management of diabetic and neuropathic ulcers
The following instructions should be given to the patient:

- Stop smoking
- Regularly inspect the legs/feet for scratch marks, blisters, etc
- Wash the feet with warm water daily followed by careful drying between the toes
- Apply emollients, for example, Vaseline on the dry skin but not between the toes
- Inspect shoes regularly
- Wear properly fitting shoes
- It is useful to see a chiropodist regularly
- Avoid sandals and pointed shoes which may lead to foot trauma
- Never walk bare footed
- See your doctor regularly if there is any injury or blister of the leg/feet

Neuropathic ulcers are most commonly seen in diabetic patients. In diabetics, the feet should be examined regularly for tinea, deformities, and onychogryphosis. It is important to remove crusted areas, and to inspect and probe the wounds to determine the depth and the extent of tissue destruction.63–65 In case of infection with cellulitis the wound should be cultured and debrided.63 Oral broad spectrum antibiotics should be started and changed according to the sensitivity results. If there is no significant improvement within 48 hours, the patient should be hospitalised. Bacteroides, coliforms, and streptococci are frequently cultured. Non-weight bearing is essential until healing is in progress. Insulin is usually required to ensure good diabetic control and broad spectrum intravenous antibiotics are indicated.66

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In 14 patients treated in a preliminary trial. A fibrinolytic enhancing agent, was found to improve liposclerosis but not venous ulceration. Initial uncontrolled trials suggest success with ulceration. Prostaglandins PGE1 and PGI2 are potent vasodilators and also inhibit platelet aggregation. Initial uncontrolled trials suggest success of intravenous infusions of PGE1 and PGI2 in patients with severe peripheral vascular disease with ulceration. A subsequent large multicentre trial demonstrated no difference between the placebo and the treatment group. Therefore, it should not be prescribed as its role in treating leg ulcers is not evidence based.

Calcium channel blockers

Nifedipine is a potent vasodilator with its mode of action, believed to be due to increased blood flow in the leg and foot in peripheral vascular disease. But the results of its use in the trials are conflicting. To date there is no convincing evidence of its value in the healing of leg ulcers due to peripheral arterial insufficiency. Therefore, from a practical point of view, it is usually not recommended.

Serotonin antagonists

Ketanserin, a serotonin antagonist, reduces peripheral vascular resistance and may improve peripheral haemodynamics in patients with intermittent claudication. Initial trials with ketanserin in treating patients with impending gangrene and peripheral ulceration are promising, but further data are necessary. Topical 2% ketanserin ointment appears to improve granulation tissue formation in skin ulcers. In practice, it is hardly used to treat leg ulcers.

Skin grafting

Pinch grafts

The simple type of split thickness skin grafts are safe and easy to perform. These procedures are particularly advisable in patients with multiple medical problems, taking drugs interfering with the healing of leg ulcers. It can be performed as an outpatient basis and in primary care.

Split thickness skin graft

This method is used for large ulcers under general, spinal, or extensive local anaesthesia. These grafts are usually successful, but the grafts may contract after harvesting and need a large donor site, which may be slow to heal and cause a lot of pain. In grafting large leg ulcers, graft failure may occur because of build-up of exudate underneath the graft; this is best avoided by the use of a meshed graft.

Future trends in leg ulcer treatment

Oral zinc treatment

To determine whether oral zinc treatment was useful for venous leg ulcer, the search strategy of the Cochrane Wounds Group was used to identify the relevant randomised controlled trials. There were six such trials, which were small double blind, randomised, median size 33 (range 10–40 patients). At present there is no evidence to support the use of oral zinc in the treatment of patients with leg ulcers due to venous or arterial disease. There is limited evidence that zinc may be beneficial in the treatment of venous leg ulcers when there is a low serum zinc but recommendations for the dose and duration of treatment cannot be made on the available information.
ULTRASOUND FOR CUTANEOUS WOUND HEALING

Five controlled trials from the Cochrane Cutaneous Wound Healing Group demonstrated that low-dose ultrasound is an effective complementary therapy for wound healing. Further experimental and clinical investigations should test its effectiveness more rigorously and elucidate the mechanisms that might be involved.83 84

LASER THERAPY FOR THE TREATMENT OF VENOUS LEG ULCERS

To assess the effectiveness of low level laser therapy in the treatment of venous leg ulcers the Cochrane Wounds Group search strategy identified four eligible randomised controlled trials which demonstrated no evidence of any benefit associated with low level therapy on venous leg ulcer healing. One small study suggests that a combination of HeNe laser therapy and infrared light may promote the healing of venous ulcers, however more research is needed.85

Compression treatment of leg ulcers: a systematic review

To estimate the clinical and cost effectiveness of compression systems for treating venous leg ulcers, 19 electronic databases including Medline, CINAHL, Embase, relevant journals, and conference proceedings included 24 randomised controlled trials which demonstrated that compression systems improve healing of venous leg ulcers and should be routinely used in uncomplicated venous ulcers. High compression is more effective than low compression,86 but should only be used in the absence of significant arterial disease. There are no clear differences in the effectiveness of different types of compression systems (multi-layer and short stretch bandages and Unna’s boot) have been shown. Intermittent pneumatic compression appears to be a useful adjunct to bandaging. Rather than advocate one particular system, the increased use of any correctly applied high compression treatment should be performed.86 Compression bandages and stockings in the treatment of venous leg ulcers as reviewed by Cochrane Wounds Review Group,87 demonstrated that multilayer high compression bandages were significantly more effective than a single layer bandage. The direct comparisons of the healing rates were described in the two observational studies as 40% (of leg ulcers) in 12 weeks88 and 42% (of limbs) of people attending a leg ulcer clinic.89 The limitations for use of four layer bandaging in a community clinic include the fact that some people are unable to leave their home due to immobility and cannot tolerate or do not like the treatment.80

Cultured epidermal grafting

AUTOGRRAFT

The treatment of venous stasis ulcer can be done by the use of cultured epidermal cells (keratinocytes) as a skin graft. Keratinocyte layers cultured from the patient’s own skin (autografts) have been successfully used for several years in the USA and Europe to cover large burn wounds.91 Other investigators have used cultured autografts for chronic leg ulcers.92 One group reported healing of chronic leg ulcers within 35 days of application of cultured epithelial autografts.82

ALLOGRAFTS

Cultured epidermal allografts can be used in enhancing the healing rate of chronic leg ulcers. Previous trials have demonstrated that more than two thirds of chronic ulcers healed completely within eight weeks of grafting with a mean healing time of 3.3 weeks.93 94 However cultured allografts should be reserved for the treatment of non-healing leg ulcers as cultivation procedure is very complex, labour intensive, and expensive.

Apligraf: a biomaterial for wound healing/human skin equivalent

Apligraf (Organogenesis, Canton, MA; Novartis, East Hanover, NJ, USA) is a new tissue engineered skin product cultured from human dermal fibroblasts and keratinocytes. Histologically it resembles a simplified version of skin. When applied, Apligraf seems to react to factors in the wound and may interact with the patient’s own cells to stimulate the production of growth factors. It is being extensively studied in diabetic ulcers, clean excision wounds, and burns.95 Human skin equivalent appeared to promote wound healing in three ways: (1) apparent graft “take”, (2) temporary wound closure, and (3) stimulation of host healing by acting as a biological dressing. The efficacy of human skin equivalent suggests that it will prove useful for promoting the healing of venous ulcers.96

Electrical stimulation

Electrical stimulation has been demonstrated to enhance wound healing.96 97 The mechanism by which healing occurs remains unknown. They may include inhibition of bacterial growth, effects on fibroblasts motility, or increased expression of transforming growth factor-β on fibroblasts.98 99

Growth factors

In animal models epidermal growth factor promotes granulation tissue formation and wound healing. A variety of growth factors—for example, tumour necrosis factors, fibroblast growth factor, transforming growth factors, α and β, epidermal growth factor, and platelet derived growth factors promote wound healing by angiogenic properties.95 97 98 100 Perhaps in the future combination of growth factors and compression dressings may have a role in the quick healing of the ulcers. Therapeutic angiogenesis may be one of the treatment strategies in the future.

Conclusion

Leg ulcers seem to have a cycle of healing followed by reulceration and rehealing. Recurrence rates of venous ulcers after treatment are high.101 Once the leg ulcer is healed careful skin care, continuous vigilance, and strict use of compression hosiery must be emphasised.
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