Review article

Aloe vera leaf gel: a review update

T. Reynolds a,*, A.C. Dweck b

a Jodrell Laboratory, Royal Botanic Gardens, Kew, Richmond, Surrey, UK
b Dweck Data, 8 Merrifield Road, Ford, Salisbury, Wiltshire, UK

Received 20 April 1999; accepted 20 May 1999

Abstract

Research since the 1986 review has largely upheld the therapeutic claims made in the earlier papers and indeed extended them into other areas. Treatment of inflammation is still the key effect for most types of healing but it is now realized that this is a complex process and that many of its constituent processes may be addressed in different ways by different gel components. A common theme running though much recent research is the immunomodulatory properties of the gel polysaccharides, especially the acetylated mananns from Aloe vera, which are now a proprietary substance covered by many patents. There have also been, however, persistent reports of active glycoprotein fractions from both Aloe vera and Aloe arborescens. There are also cautionary investigations warning of possible allergic effects on some patients. Reports also describe antidiabetic, anticancer and antibiotic activities, so we may expect to see a widening use of alo gel. Several reputable suppliers produce a stabilized alo gel for use as itself or in formulations and there may be moves towards isolating and eventually providing verified active ingredients in dosable quantities © 1999 Elsevier Science Ireland Ltd. All rights reserved.

Keywords: Aloe vera gel; Active polysaccharides; Therapeutic properties

1. Introduction

Aloes have been used therapeutically, certainly since Roman times and perhaps long before (Morton, 1961; Crosswhite and Crosswhite, 1984), different properties being ascribed to the inner, colourless, leaf gel and to the exudate from the outer layers. During the 12 years since the last major review of Aloe vera (L.) Burm.f. gel (Grindlay and Reynolds, 1986) popular interest and use of the gel have increased dramatically. In this country it is now a familiar ingredient in a range of healthcare and cosmetic products widely available and advertised in shops. The preserved but otherwise untreated gel is also sold as a therapeutic agent in its own right as are various concentrated, diluted and otherwise modified products. This need has been met by a number of wholesalers who get their supplies from plantations in Texas, Florida and Venezuela while new ones are

* Corresponding author.
being proposed for Israel, Queensland and East Africa (Jamieson, 1984). This commercial activity has been accompanied by an upsurge of both clinical and chemical research which is reaching more closely towards the active ingredients and their biological activity. There is now less said about doubts as to the efficacy of the material, although there are some warnings of allergic side effects (Klein and Penneys, 1988; Briggs, 1995). Harmful reactions to aloe gel treatment are recorded infrequently (Hunter and Frumkin, 1991; Schmidt and Greenspoon, 1993) but need to be taken seriously. There is still confusion between the leaf exudate and the gel, Morsy and Ovanoviski (1983), Natow (1986) and Duke (1985) where a great number of folk medicine uses are described and Ahmad et al. (1993). However many commentators clearly distinguish between the two parts (Watson, 1983; McKeown, 1987; Capasso et al., 1998) and describe in some detail how the gel is prepared (McAnalley, 1988, 1990; Agarwala, 1997). At one time there was much discussion about the relative efficiency of ‘decolorized’ and ‘colorized’, i.e. with exudate components, gels (Danof, 1987; Agarwala, 1997). There is also a feeling that some of the variable results reported in the literature may be due to treatment of the gel subsequent to harvest (Fox, 1990; Marshall, 1990; Briggs, 1995; Agarwala, 1997).

A number of reviews have appeared in recent years covering various aspects of aloe gel use, as well as much commercial literature. Exaggerated claims are still being made and although doubts as to the substance’s efficacy are more muted, there is still room for the caution which has been voiced (Hecht, 1981; Marshall, 1990). The emphasis is changing towards definition of the active constituent or constituents so that they can be used accurately in formulations (Reynolds, 1998).

There has been a greater willingness to investigate reasons for the recorded variability in curative properties.

Reasons presented for aloe gel efficacy are still varied perhaps because there are in fact several different healing activities operating (Capasso et al., 1998). The action of aloe gel as a moisturizing agent is still a popular concept (Meadows, 1980; Watson, 1983; Natow, 1986; Danof, 1987; McKeown, 1987; Fox, 1990; Marshall, 1990; Briggs, 1995) and may account for much of its effect. More speculative is the presence of salicylates, by implication having an aspirin-like effect (Robson et al., 1982; Klein and Penneys, 1988; Marshall, 1990; Shelton, 1991; Canigueral and Vila, 1993), although the differences between natural salicylates and aspirin, a synthetic product, were pointed out (Frumkin, 1989). Another simple substance, magnesium lactate, is said to inhibit the production of histamine by histidine decarboxylase and is claimed as a gel constituent (Rubel, 1983; Natow, 1986; Marshall, 1990; Shelton, 1991; Canigueral and Vila, 1993). Inhibition of pain-producing substances such as bradykinin or thromboxane is often claimed (Rubel, 1983; Natow, 1986; Danof, 1987; Fox, 1990; Marshall, 1990; Shelton, 1991; Canigueral and Vila, 1993). On a more sophisticated level, action on the immune system has been postulated and to some extent tested (Rubel, 1983; Schechter, 1994; Griggs, 1996). A recent, very interesting book (Davis, 1997), dwells at some length on the immunomodulatory properties of the gel polysaccharides and presents a viewpoint complementary to the present review. Polysaccharides are another group of gel constituents to which activity has been ascribed, particularly in immunomodulatory reactions and one, acemannan, has reached proprietary status (Schechter, 1994; McAnalley, 1988, 1990; Agarwala, 1997). There has been much interest recently in the biological activity of polysaccharides, which is greater and more diverse than previously realized. Although the substances are varied and widespread in plants some are well known as entities, albeit often with uncertain structures (Franz, 1989; Tizard et al., 1989; McAuliffe and Hindsgua, 1997). Also often mentioned are the antibacterial, antifungal and even antiviral properties demonstrated by the gel (Klein and Penneys, 1988; Marshall, 1990; Ahmad et al., 1993), while anti-oxidant effects are becoming of interest. Although A. vera gel is the only one being used commercially, there is the possibility of discovering useful properties among the other 300 or more species (Newton, 1987).

In this update the trend to be emphasized is away from the more naive arbitrary use of the leaf
gel (Reynolds 1996) and towards a quest for deeper, more precise understanding of its constituents and the varied biological activities which they may or may not display (Reynolds 1996).

2. Test systems and clinical trials

2.1. Burns and incisions

For testing the efficacy of aloe gel or its various components on inflammation a number of tests have been used, usually in relation to some sort of deliberate wounding. These need to be distinguished from clinical trials where the injuries already exist and are treated more or less systematically by a number of putative therapeutic agents. The earliest experimentation related to skin burns and arose in relation to clinical observations, going back to the 1930s (Grindlay and Reynolds, 1986). It was often inconclusive due to inadequate controls and replication and an imprecise correlation of cause and effect. One of the most detailed and accurate of these clinical trials took place in 1957 with use of aloe gel against controlled thermal and radiation burns on rats and rabbits compared with clinical studies on human patients (Ashley et al., 1957). This failed to demonstrate any healing properties of the gel. In contrast another careful study but with far fewer replicates gave a positive result (Rovatti and Brennan, 1959). Another approach a little later, was to measure the tensile strength of the healing of a precise incision wound, post mortem (Goff and Levenstein, 1964). An undescribed aloe ‘extract’ speeded healing but had no effect on the final result. A similar wound tensile strength test was used later to compare the effects of steroids and aloe gel on inflammation and healing (Davis et al., 1994b), and later of antibacterial agents (Heggers et al., 1995). Then in the early 1980s both precise experimental scald burns as well as previous injuries were again compared by a variety of criteria (Cera et al., 1980, 1982; Robson et al., 1982) and therapeutic benefits were recorded. This type of test system was returned to later (Heggers et al., 1993) with similar positive results.

A study, with good replication, of healing after a precise skin hole punch demonstrated the anti-inflammatory properties of the gel leading to more rapid healing (Davis et al., 1987a) A different approach was taken whereby the subjects (mice) were fed aloe gel for some time before hole punch wounding and compared with those treated topically after wounding (Davis et al., 1989c). Both methods produced healing. A further variation was the treatment of punch wounds on mice or rats made diabetic by streptozotocin and therefore more slow to heal. Again healing by aloe gel was demonstrated (Davis et al., 1988; Davis and Maro, 1989). A return to wounding by precision burns using a hot metal plate with adequate replication again demonstrated positive healing activity (Rodriguez-Bigas et al., 1988). This technique was further elaborated to produce first, second or third degree burns by precisely timed exposures to the hot metal plate (Buayapraphatsara et al., 1996a).

Two special examples of burns are sunburn and frostbite and these have been used experimentally. Thus, precise UVB burns were produced with a light pen but were unaffected by aloe gel (Crowell et al., 1989). In a later trial to test the effect of the gel on UV-induced immune suppression a bank of UV lights were used (Strickland et al., 1994). Frostbite was produced by exposing rabbit ears to ethanol and solid carbon dioxide (Heggers et al., 1993; Miller and Koltai, 1995) and was relieved by application of aloe gel.

2.2. Irritating compounds producing oedema

Experimental production of swelling, caused by fluid accumulation in a tissue (Oedema) initiated by irritating compounds has been used as an inflammatory model with the mouse ear or rat hind paw as subjects. Croton oil, a powerful irritant, was applied to the right ear with the left remaining as control. Inflammation was measured by weighing a tissue punch sample and was shown to decrease after topical application of aloe gel.
(Davis et al., 1987b; 1989a,b). A subsequent trial demonstrated an even greater decrease when the gel was combined with a corticosteroid (Davis et al., 1991, 1994b). This trial was accompanied by a similar one where mustard as the initiating agent was injected into a rat paw, subsequent swelling being measured volumetrically and fluid withdrawn to determine leucocyte infiltration. In this case aloe gel, with or without steroids was injected previously rather than being applied topically. This study had followed some more or less similar ones, where a variety of irritants, gelatine, albumin, dextran, carrageenan and kaolin had been used (Davis et al., 1989a) and the inflammation successfully treated with aloe gel, orally or topically.

2.3. The air pouch

A modification of these models involving oedema was developed in which air was injected under the skin to form a cavity, a simulated synovium, to which irritants and therapeutic agents could be added. This was taken to be analogous to the joint cavity, containing synovial fluids which becomes inflamed during arthritis. Such an air pouch was produced on the backs of anaesthetized mice, irritated with carrageenan and treated with A. vera gel solution (Davis et al., 1992). Healing effect was measured histologically by counting the number of mast cells in the cavity fluid, decreased by aloe gel treatment and by examining pouch wall vascularity, also decreased by the gel.

2.4. Adjuvant arthritis

An important extension of experiments on lesions caused by applied irritants is the deliberate production of a condition resembling arthritis in an animal model, usually rat. This can then be followed by treatment with putative therapeutic agents to suppress either the inflammation or immunologic consequences. The irritating agent used was a suspension of heat-killed Mycobacterium butyricum in mineral oil which produced inflammation directly in the injected paw and also in the other paw by an immunologic pathway (Saito et al., 1982).

2.5. In vitro studies

A number of experiments have been carried out in which the effects of components of Aloe leaf on various biochemical or microbiological systems, relevant to the four inflammation responses and to wound healing, have been studied. Enzymes in aloe gel destroying the nonapeptide bradykinin which causes vasodilatation and pain production, were the subject of one such investigation, although angiotensin-converting-enzyme received less attention. Prostaglandins were another obvious subject and their presence in Aloe was claimed, as well as a complex lipid inhibiting arachidonic acid oxidation. On the other hand production of the vasodilator histamine from histidine was said to be inhibited by the gel. Effects of aloe gel on separate components of wound healing in tissue culture have been made, notably fibroblast proliferation (Brasher et al., 1969; Danof and McAnally, 1983) and growth of new blood capillaries (Lee et al., 1995). The involvement of immunological effects has been recognized by examining stimulus of phagocyte formation and activity (Imanishi and Suzuki, 1984). Alongside these studies of in vitro systems there were also many attempts to analyse the plant material and to identify the active fraction or fractions.

3. Treatment of inflammation

Inflammation is a tissue reaction by the body to injury and typically follows burns or other skin insults. It is classically characterized by swelling (tumor), pain (dolor), redness (rubor) and heat (calor) as well as loss of function (Macpherson, 1992). It is thus a complex process and investigations into the therapeutic properties of the gel should take account of its effects on these various symptoms. In addition, the gel may have more than one active constituent, which may be addressing different parts of the healing process. Failure to take all this into account may be responsible for ambiguities which may have arisen in the past about the efficacy of the gel. Although inflammatory processes are a natural response to
injury and may hinder healing it may also be undesirable to suppress them in an unstructured way before their purpose is accomplished. Leukocytes accompanied by fluid accumulate in the damaged tissues producing the swelling, these movements being the result of increased capillary permeability. Pain is a complex reaction following the release of short peptides and prostaglandins. The redness and heat are caused by vasodilatation which reduces blood pressure and increases circulation, although this gradually slows. Inflammation can be either caused, or intensified by invasion with micro-organisms. As well as in wounds, inflammation is involved in conditions such as arthritis. Continuing research into inflammation has shown that it is a complex process involving many biochemical pathways and a variety of agents and mediators (summarized in Davis et al., 1989a). In particular these authors distinguish three components,

1. Vasoactive substances; agents causing dilation of blood vessels and opening of junctions between cells of the ultimate capillaries, produced by altering contractile elements in endothelial cells. These factors include vasoactive amines, bradykinin and also prostaglandins.

2. Chemotactic factors; these agents cause increased cell motility, especially of white blood cells (leukocytes) into stressed areas. These include several proteins and peptides.

3. Degrative enzymes; these are hydrolytic enzymes breaking down tissue components. Proteases in particular participate in inflammatory states causing chemotactic factors to be released. It was also shown that aloe gel contained both an inhibitory system and a stimulatory system that influenced both inflammatory and immune responses (Davis et al., 1991a,b).

The healing effects of *A. vera* gel are therefore now being seen as more complex than previously realized (Hörmann and Korting, 1994). It now appears that several activities are operating each with its own part to play in the overall therapy. These activities may well reflect the presence of several different active agents in the gel. There seems to be a need for two types of investigation. The first, the academic, analytical approach, seeks to dissect the processes and reveal the individual biochemical and physiological reactions, while the second, the clinical approach, puts the various processes back together and studies their interactions. The second can only be ultimately successful when the first is well known. In the best, recent, precise experimentation, care has been taken to separate the inner parenchyma of the aloe leaf completely from the outer layers rich in phenolics, both experimentally and conceptually. However in some trials this separation has not been complete and two preparations were deliberately made and tested, one decolorized and the other containing anthraquinones from the outer layers (Davis et al., 1989). These substances were to some degree toxic and reduced for instance suppression of polymorphonuclear lymphocyte infiltration (Davis et al., 1986b). They also greatly reduced the healing of croton oil-induced inflammation (Davis et al., 1989b, 1991). A component extracted from whole *Aloe barbadensis* (sic) leaves and probably originating from the exudate rather than the gel was characterized as a cinnamic acid ester of aloesin and shown to reduce croton oil-induced inflammation (Hutter et al., 1996). A low molecular weight component claimed to be extracted from the gel but probably also of exudate origin, was shown to have cytotoxic effects similar to barbaloin (Avila et al., 1997). This raises the possibility of both irritating and healing agents in the exudate as well as the gel. Even this comparison is not strictly accurate as it is not clear how the anthraquinone-free gel is made and if other substances are removed, or not, at the same time. Mannose-6-phosphate was shown to have anti-inflammatory activity and this was said to resemble the known activity of acetylated mannann, a gel component (Davis et al., 1994a).

Two aspects of inflammation reduction following aloe gel treatment by injection in rats were observed (Davis et al., 1986a,b, 1987c). Mustard induced oedema of the paw was reduced by between 445 and 70% while infiltration of polymorphonuclear lymphocytes into a skin blister was reduced by 58%. Two other substances, RNA and vitamin C synergized with the gel in inhibiting oedema. Similarly mouse ear inflammation induced with
croton oil was reduced by up to 67% following topically applied aloe gel (Davis et al., 1987b). In another study these workers tested the action of topical or injected aloe gel against inflammation produced by a variety of agents which were considered to induce different types of inflammation (Davis et al., 1989a). Thus the gel relieved the inflammatory effects of kaolin, carrageenan (an algal polysaccharide), albumin, gelatin, mustard and croton oil which were said to act either by promoting prostaglandin synthesis or by increasing infiltration of leucocytes. Elsewhere, aqueous or chloroform extracts of the gel reduced a carrageenan-induced inflammation and migration of neutrophils (Vazquez et al., 1996). Aloe gel was less effective against inflammatory agents which produced allergic reactions through the action of bioactive amines such as histamine, even if their synthesis might be inhibited by magnesium lactate. In other ways aloe gel was found to show immunomodulatory properties (t’Hart et al., 1988) so the full picture is still not clear. Another aspect is the use of the gel as a vehicle for application of other active substances to which it may additionally impart its own activity (Davis et al., 1989c).

3.1. Wound healing

If inflammation is a complex process, then wound healing is much more so and the intervention of aloe gel is likely to be multifaceted. A wound to the skin may pierce two layers, the epidermis and dermis as well as damaging appendages. A temporary repair is effected by fibrin clot which is then invaded by a variety of cells, some of which produce the inflammatory response and which eventually carry out a permanent repair (Martin, 1997). It may well be that the repair is not perfect in that scar tissue is produced and appendages do not regenerate. The epidermis is repaired in three phases, migration of cells, proliferation and maturation, while new connective tissue is found in the dermis (Davis et al., 1987a). As well as repair of structure there is an urgency to avoid microbiological entry which can retard wound contraction (Hayward et al., 1992). Increased speed of repair was seen in an early detailed study of healing of a surgical cut which showed that 'A. vera extract in an ointment base' speeded repair but did not alter the ultimate result (Goff and Levenstein, 1964). Perhaps aloe gel removes delaying effects rather than accelerating healing as such. The use of aloe gel to heal wounds is the classic use of the material and one of the first explanations of its efficacy was its high water content which kept the wound moist and increased epithelial cell migration (Morton, 1961; Erazo et al., 1985), although even this has been questioned (Roberts and Travis, 1995). Indeed the beneficial effects on oral wounds where moisture is abundant, indicates that other factors operate (Sudworth, 1997). A report of effective aloe gel healing of pressure sores recorded rapid granulation (Cuzzell, 1986), an effect also noted with various incision wounds in rats and attributed to more rapid maturation of collagen (Udupa et al., 1994).

In a more detailed study, a skin punch wound healed more rapidly when treated with 'decolorized' gel than with 'colorized' gel (Davis et al., 1986). These observations were made on the 7th day after wounding a mouse or rat skin which was said to be optimal for recording healing. Treatment by daily injection of the gel reduced wound diameter, increased skin circulation and seemed to reduce scarring (Davis et al., 1987a, 1989a). Acute inflammation was also inhibited. The colour mentioned is due to anthraquinones from the aloe leaf exudate but details of their removal ('decolorized' gel) were not given. Elsewhere, trials using cultures of human endothelial cells or fibroblasts demonstrated cytotoxicity in gel samples contaminated with leaf exudate (Danof and McAnalley, 1983). In contrast, cytotoxicity was shown to be reduced in neutrophils treated with a low molecular weight fraction of gel, probably exudate-derived, although this was not stated, following inhibition of release of reactive oxygen species (t’Hart et al., 1990). In a further study, 'A. vera' (sic), presumably the gel, was administered orally over two months or applied to the wounded skin in a cream. Both treatments improved wound healing. It was suggested that one of the factors, out of several, enhanced by aloe gel was increased oxygen access.
as a result of increased blood supply. (Davis et al., 1989b). In another trial using topical application, stimulation of fibroblast activity and collagen proliferation was demonstrated (Thompson 1991). Angiogenesis, the growth of new blood capillaries, is a necessary part of tissue regeneration and vascularity of burn tissue of a guinea pig was shown to be reestablished by topical application of aloe gel (Heggars et al., 1992). A low molecular weight component of freeze-dried aloe gel was shown to stimulate blood vessel formation in a chick chorioallantoic membrane, while a methanol-soluble fraction of the gel was shown to stimulate proliferation of artery endothelial cells in an in vitro assay and to induce them to invade a collagen substrate (Lee et al., 1998). Activation of matrix proteinases, which allow penetration, was thought to be involved. Tissue survival following arterial damage in a rabbit ear, which mimicked drug abuse damage, was maintained by topical aloe gel application (Heggars et al., 1993). Healing of an experimental excision wound was promoted by topically applied aloe preparation and this was enhanced when the gel was combined with a nitric oxide inhibitor (Heggars et al., 1997).

Subsequent work showed that another important impediment to wound healing, microbiological activity, infection, was also addressed by aloe gel treatment (Heggars et al., 1995). Here, cuts to rat skin were more rapidly healed by topical applications of aloe gel compared with an untreated control or by applications of potential anti-microbials. Many antibiotic agents are more toxic to fibroblasts than to bacteria and seem to retard the healing process (Lineaweaver et al., 1985). The gel was thought to contain a growth factor which enhanced the breaking strength of wounds. Only buffered sodium hypochlorite solution (0.025%) had therapeutic effects similar to aloe gel (Heggars et al., 1996). Macrophages play a considerable part in controlling microorganisms and it was shown that young active macrophages accelerated the rate of wound healing in aged rats, compared with rates where senescent cells in these animals were left alone to act. Activation of macrophages by acemannan, an aloe gel polysaccharide, was claimed (Maxwell et al., 1996). This followed previous observations on the healing powers of acemannan on wounds in elderly or obese rats also attributed to macrophage stimulation (Tizard et al., 1994). Total regeneration of the skin also requires that ‘difficult’ cells such as neurons are replaced. Proliferation of neuron-like cells and also perhaps cell adhesion, in a culture of rat adrenal cells was stimulated by gel preparations (Boutheil et al., 1995).

Following the idea that there were factors with different types of activity in the gel an attempt was made to separate anti-inflammatory and wound healing components (Davis et al., 1991c). A precipitate formed by treatment with 50% aqueous ethanol seemed to have most of the wound healing activity observed in the raw gel when used against punch wounds in mouse skin. The supernatant contained anti-inflammatory activity which was attributed to glycoprotein. Elsewhere, significant wound healing was produced by mannose-6-phosphate (Davis et al., 1994a) Healing of an incision wound by aloe gel was found to be accompanied by higher levels of hyaluronic acid and dermatan sulphate produced more rapidly. This was suggested to stimulate collagen synthesis and fibroblast activity (Chithra et al., 1998a). There was also increased activity of β-glucuronidase and N-acetyl glucosaminidase which was said to increase carbohydrate turnover in the wound matrix. Fibroblast proliferation in vitro and in vivo was observed after treatment with the acetylated mannan fraction carrisyn™ (McAnalley, 1988). Increased collagen formation in wounded diabetic rats treated orally and topically with aloe gel was later demonstrated (Chithra et al., 1998b). The collagen formed had a higher degree of crosslinking indicating enhanced levels of type III (Chithra et al., 1998c). There were also higher levels of protein and DNA. In another test on surgical cuts in mice, hydrocortisone given by injection, while reducing inflammation, hindered wound healing but when ‘A. vera’ (presumably the gel) was included then wound suppression was reversed and inflammation further reduced (Davis et al., 1994b). Healing and control of acute inflammation, distinct from chronic inflammation, was observed following gel treatment of excision and incision wounds in rats (Udupa et al., 1994).
Widespread acclamation of the healing powers of aloe gel is not however universal. Some earlier studies were unable to demonstrate any curative properties (Ashley, 1957; Gjerstad, 1969; Ship, 1977; Spoerke and Elkins, 1980; Kaufman et al., 1988) and more recently a trial with surgical wounds in human patients even suggested that healing was delayed (Schmidt and Greenspoon, 1993). No effects of aloe gel on re-epithelization or wound contraction of excision wounds in pigs was observed (Watcher and Wheeland, 1989). No healing properties at all were observed with corneal punch wounds (Green et al., 1996), in contrast to earlier positive observations with corneal flash burns (Lawrence, 1984). Elsewhere it was found that acemannan had an equal, not better, healing and bactericidal effect on shave biopsy wounds as the antibiotic bacitracin® (Phillips et al., 1995). The two lines of conflicting evidence may be explained by the fragility of the active ingredients as it appears from several accounts that the treatment of the gel after harvesting is crucial for activity. Effects may also vary with the type and location of wound. Using a proprietary aloe dressing on pad wounds of dogs it was concluded that healing processes during the first 7 days were speeded, an advantage to a wound exposed to weight bearing, although the end result was the same as that with antibiotic treatment alone (Swaim et al., 1992).

3.2. Burn healing

Burn healing can be regarded as a special type of wound healing and most of the skin reactions are the same. It has been pointed out however that conditions for healing would differ according to the depth of the burn wound and that several factors can interfere with the healing process (Kaufman et al., 1989) Thus three zones have been recognized in a burn, an inner zone (coagulation zone) where cell damage is irreversible, a middle zone (statis zone) where damage is severe and an outer zone (hypervemic zone) where recovery is likely. In addition there are three degrees of burns, the first in which the epidermis only is damaged, the second where some dermal damage also occurs but where epithelial regeneration is possible and the third where both epidermis and dermis are irreversibly damaged (Bunyapraphatsara et al., 1996a). Like wound healing it is one of the classic subjects for aloe gel treatment (Ashley et al., 1957; Rovatti and Brennan, 1959; Cera et al., 1982), although as for wound healing some studies demonstrate little benefit (Heck et al., 1981). In a large, double-blind trial using 194 patients with radiation burns, no difference to a placebo was observed (Williams et al., 1996). However other samples of the preparation used in this trial (Fruit of the Earth) were found elsewhere to have no mucopolysaccharide content (Ross et al., 1997). The existence of diverse components of a burn and the diverse components of aloe gel which might be healing the burn, were soon recognized (Robson et al., 1982). Here the gel was said to possess an anaesthetic effect, a bactericidal action and an anti-thromboxane effect. Recognizing the possible multifarious activities of Aloe constitutents, a series of tests of aloe gel on heat burns, electrical burns and frostbite in guinea pigs, rabbits and in clinical studies with humans demonstrated a therapeutic potential across the wide variety of soft tissue injuries (Heggars et al., 1993). The gel was shown to penetrate tissue, relieve pain, reduce inflammation and increase blood supply by inhibiting the synthesis of thromboxane A₂, a potent vasoconstrictor. Hot-plate burns to guinea pig skin healed more quickly after topical aloe gel application and interestingly, the bacterial count was reduced by 60% (Rodriguez-Bigas et al., 1988; Kivett, 1989). A recent study demonstrated healing activity towards gamma-radiation burns but only if applied quickly, when it produced more rapid healing than controls but only because peak reaction levels were reduced (Roberts and Travis, 1995). Here it was speculated that aloe gel affected the induction of the skin reaction but not the later healing phases. In a similar trial using mice, differences were seen in the effect on first, second and third degree burns. Gel preparations delayed the inflammatory response and speeded the recovery time for first and second degree burns and epithelialization was rapid. Third degree burns proved more intractable (Bunyapraphatsara et al., 1996a). A synergism was noted between the gel
and the cream base used. Elsewhere, partial thickness burns were observed to heal more rapidly when treated with aloe gel, compared with vaseline, both growth of epithelial cells and organization of fibro-vascular and collagen tissue being stimulated (Visuthikosol et al., 1995).

3.3. Frostbite

Direct and indirect cellular injury arising from frostbite can be regarded as a type of burn (Heggers et al., 1990), although the stages described differ. One classification distinguished four degrees, the first with numbness and erythema, the second where oedema and blisters occur and thromboxane is released, the third where damage extends to the subdermis and the fourth with full tissue thickness damage. (McCauley et al., 1990). Another classification recognized four phases, the first (pre-freeze phase) with chill but no ice crystal formation, the second (freeze–thaw phase) with ice formation, The third (vascular stasis phase) with plasma leakage and the fourth (ischemic phase) with thrombosis, blood loss and even gangrene (Miller and Koltai, 1995). Thromboxane is a powerful vasoconstrictor and pain producer (McCauley et al., 1983; Miller and Koltai, 1995) and has been implicated in frostbite injuries (Heggers et al., 1987) It was suggested that the main function of aloe gel in healing frostbite is the reduction of thromboxane levels (Raine et al., 1980) and has been used clinically on this assumption to treat the more severe blisters where there was structural damage (McCauley et al., 1983). Topical application of A. vera cream (sic) enhanced tissue survival of frost-bitten rabbit ear. In an accompanying clinical trial with humans, 68% of the aloe-treated patients achieved full healing, while only 33% of those receiving other treatments were fully healed. In the first group 7% required amputation, compared with 33% in the second group (Heggers et al., 1990). In another trial with rabbit ears, 24% survived from those treated with A. vera cream while only 6% of the untreated ears survived (Miller and Koltai, 1995).

3.4. Adjuvant arthritis

One very troublesome instance of inflammation is rheumatoid arthritis where the joints become inflamed and a complex syndrome of pathological effects appears. An experimental model set up to probe this disorder is the so called adjuvant arthritis produced by injection of a substance which unspecifically intensifies the immune response without itself being antigenic. In one experimental design the adjuvant is injected into the right hand paw of a rat where it soon produces inflammation, whereas later inflammation in the left hand paw is held to be an immunologic phenomenon. A whole leaf extract from A. africana (sic), strictly A. ferox Mill. or a hybrid, but probably in fact A. vera, was injected and decreased inflammation (48%) in the right paw also inhibiting the immunological response (72%) in the left paw (Hanley et al., 1982). It was speculated without experimental evidence that these effects resulted from inhibition of prostaglandin synthesis. In another test A. vera extract, described as a 5% leaf homogenate, (also called A. africana in parts of the text) together with ascorbic acid and RNA was applied topically in a hydrophilic cream base, again produced reduction of both immediate inflammation (39%) and subsequent arthritis (45%) (Davis et al., 1985). The gel itself, included in this mixture produced 45% regression (Davis, 1988). A further study attempting to pinpoint active ingredients found that injected aqueous suspensions of anthraquinone, anthracene, cinnamic acid or anthranilic acid inhibited inflammation to various extents (Davis et al., 1986), while anthraquinone and cinnamic acid had some effect on the immune response.

Another experimental model attempting to simulate the synovial cavity in a joint, where inflammatory reactions occur and produce arthritis is the ‘synovial pouch’ where air is injected under the skin to form a cavity. The walls of the cavity are said to resemble the synovial membrane and the action of carrageenan on this is said to resemble arthritic inflammation (Davis et al., 1992). Subsequent injection of aloe gel reduces this inflammation rapidly and then induces fibroblast
growth. The number of mast cells migrating from surrounding connective tissue was also reduced.

3.5. Bradykinin

In studying the effect of aloe gel on skin lesions, one line of research that has been pursued follows the part played by the nonapeptide bradykinin in the inflammatory process. A peptidase, bradykininase, active in breaking down bradykinin to inactive units was isolated from *A. arborescens* Mill. leaves (Fujita et al., 1976) and shown subsequently to be a carboxypeptidase (Fujita et al., 1979) and then a serine carboxypeptidase (Ito et al., 1993). In a separate study with the same species, a glycoprotein with carboxypeptidase activity was isolated (Yagi et al., 1987a). *Aloe arborescens* is much used in Japan in a way similar to *A. vera*, although in these studies it appears that the whole leaf was used in the preparation rather than the isolated gel. A high molecular weight, water-soluble fraction was separated so it could be assumed that this probably came from the gel. However, in yet another study a carboxypeptidase was prepared from the ‘leaf skin’ and partially purified (Obata et al., 1993). This latter preparation alleviated pain after a burn and inhibited the acceleration of vascular permeability. These effects were attributed to the hydrolysis of bradykinin and angiotensin I. It was also shown that intravenous dosing before the burn was more effective than dosing after the burn. Some of these latter authors had previously isolated a bradykinase from yet another species, *A. saponaria* (Aiton) Haw., this time from the gel alone (Yagi et al., 1982).

4. Steroids and prostaglandins

Because of the effect of various prostaglandins in either stimulating or inhibiting aggregation of platelets in relation to wound healing it would clearly be of interest to seek interactions between these compounds and aloe gel components or even presence of the compounds themselves. Steroids are another obvious group of active compounds of interest. The triterpenoid lupeol and the steroids cholesterol, campesterol and β-sitosterol were all found in whole leaf extracts of *A. vera* (Waller et al., 1978, Ando and Yamaguchi, 1990) while β-sitosterol was isolated from *A. arborescens* leaves (Yamamoto et al., 1986) and found to have anti-inflammatory properties in common with some of the exudate compounds (Yamamoto et al., 1991). Lupeol and β-sitosterol were again isolated from *A. vera* leaves, accompanied by β-sitosterol-3-glucoside and its 6-palmitate (Kinoshita et al., 1996). Lupeol, campesterol and β-sitosterol were found to be significantly anti-inflammatory in wounded mice (Davis et al., 1994b). Other, unknown factors in the gel promoted healing which would have been hindered by the sterols alone. Another analysis of the lipid fraction of *A. vera* leaves revealed various common substances, including cholesterol and also a range of more complex polar lipids (Afzal et al., 1991). Among the fatty acids, of which the chief was γ-linolenic acid (42% of total fatty acids), they determined arachidonic acid (3% of total fatty acids), a significant precursor of prostaglandins. This compound in turn is produced in reactions involving phosphatidyl choline and phosphatidyl ethanolamine and these were each found in *A. vera* at levels around 12% of the polar lipids. Although they did not detect prostaglandins as such, they demonstrated the presence of cyclo-oxygenase by the production of prostanooids from radioactive arachidonic acid added to homogenized leaf tissue.

The possible presence of prostaglandins and their effects on platelet activity in wounded tissue is complex. It depends on the molecular species present and other biochemical factors in the tissue (Venton et al., 1991). Some prostaglandins are essential for normal processes in the skin such as cell function and integrity, while others, notably thromboxane A₂ and B₂ can have devastating effects on the cells (Heggers and Robson 1983, 1985). The level of thromboxane B₂ in guinea pig burns was reduced by topical application of aloe gel (Robson et al., 1982). Other studies have suggested that unspecified substances in aloe gel inhibited arachidonic acid oxidation (Penneys, 1982) and thereby reduced inflammation. Against this, another study claimed that aloin, a com-
pound from Aloe leaf exudate, and a common gel contaminant, stimulated prostaglandin synthesis (Capasso et al., 1983). The presence or absence of this compound perhaps explains the apparent contradiction between the results of Afzal et al., (1991) and Penneys, (1982). From a different aspect it was suggested that aloe exudate anthraquinones might act as false substrate inhibitors to enzymes involved in the synthesis of thromboxane A₂, a potent vasoconstrictor (Heggers and Robson, 1985). An aqueous extract from aloe gel inhibited the production of prostaglandin E₂ from arachidonic acid in vitro and steroids were detected in the extract as well as ‘anthraglycosides’ (sic) (Vazquez et al., 1996). Inhibition of thromboxane production and consequent vasoconstriction, was said to be useful in frostbite treatment where restriction of circulation is a problem (Raine et al., 1980; McCauley et al., 1990). A glycoprotein component of the gel, Alocotin A, was shown to inhibit prostaglandin E₂ production but over a relatively long incubation time (Ohuchi et al., 1984), in contrast to drugs such as aspirin, so perhaps the anti-inflammatory factor needs to be sought elsewhere. It is evident that there is much complexity both in the damaged tissues and the plant extracts and that precise mechanisms and pathways have yet to be determined in the field of prostaglandins and their interaction with platelets.

5. Interaction with macromolecules: the immune system

The interactions of large molecules in biological systems play an important part in many life processes. Both polysaccharides and glycoproteins are involved in such activities, especially in connection with the immune system. Some glyco-proteins of non-immune origin, termed lectins, specifically bind to cells causing agglutination, or precipitate macromolecules with specific sugar structures. The immune system itself is very much more complex, having at its centre the reaction of a host’s antibodies with invasive antigens. The many reactions surrounding this and contributing to it are susceptible to interference by certain outside agents, both harmful and beneficial and it is among the latter that aloe constituents may play a part.

High molecular weight substances prepared from A. arborescens extracts were shown to precipitate serum proteins from a range of animals (Fujita et al., 1978a) and described as lectins. Two fractions were purified and both characterized as glycoproteins but with differing biological activities. One, P₂, with a molecular weight of approximately 18 000 Da, had some haemagglutinating activity, precipitated serum proteins and also showed mitogenic activity against human lymphocytes. The other, S₁, with a molecular weight of approximately 24 000 Da showed much stronger haemagglutinating activity against erythrocytes but no other biological properties (Suzuki et al., 1979a). The S₁ fraction was named Alocotin B but its properties remain relatively unknown. The P₂ fraction, named Alocotin A, was shown to agglutinate tumour cells but to show no cytotoxicity (Suzuki et al., 1979b) and to inhibit the growth of fibrosarcoma in vivo but not in vitro (Imanishi et al., 1981). It was also shown to inhibit chemically induced arthritis (Saito et al., 1982) and to inhibit uptake of foreign erythrocytes by activated rat macrophages (Ohuchi et al., 1984).

It was then shown that alocotin A injected intravenously into mice, stimulated the cytotoxicity of harvested spleen cells and peritoneal exudate cells towards tumour cells in vitro, under somewhat limited conditions (Imanishi and Suzuki, 1984). Evidence points to an apparent stimulation of host activity against tumour cells and also elevated levels of plasma proteins (Imanishi and Suzuki, 1986). Lymphokine production as a result of T cell stimulation by alocotin A was increased (Imanishi, 1993). It was then confirmed that the alocotin A-induced killer cells were indeed lymphokine activated and that there was the promise that these cells would be sensitized to tumour antigens (Imanishi et al., 1986). Alocotin A had no direct cytotoxic effect on tumour cells themselves. Later, a range of pharmacological activities was described for alocotin A (Saito, 1993). These included suppression of tumour growth, not apparently directly but by stimulating
host response, increase in macrophage levels and activity and also reduction of gastric lesions and ulcers. A third glycoprotein from *A. arborescens*, molecular weight 40 000 Da, was shown to agglutinate sheep erythrocytes and to stimulate DNA synthesis in cell cultures (Yagi et al., 1985). Yet another lectin fraction from *A. arborescens*, designated ATF1011, distinct from the Alloctins, was shown to activate helper T cells by binding to the cell surface (Yoshimoto et al., 1987). On the other hand, phagocytosis of yeast cells by neutrophils from human asthmatics was stimulated by both glycoprotein and polysaccharide fractions of whole leaves of *A. arborescens* (Shida et al., 1985), while activity was also shown by a mixture of proline and cysteine from the low molecular weight fraction (Yagi et al., 1987c). This in turn contrasts with yet other findings which strongly ascribe enhancement of phagocytosis to a polysaccharide gel component, acemannan, described below (McDaniel et al., 1987).

With another species, *A. vahombe*(sic), it was shown that mice inoculated with a leaf preparation were protected from infection by *Klebsiella pneumoniae* apparently by stimulation of the immune system (Solar et al., 1979). However, it appears that they have obtained their active fraction from the leaf juice which with other workers occurs as a contaminant to the gel as 'colorized' gel. A fraction separated on Sephadex G50 was shown to protect mice against infection by a range of bacteria and fungi (Brossat et al., 1981). The same fraction, now characterized as containing a glucomannan of molecular weight above 30 000 Da suppressed growth, in vivo, of one type of tumour in mice, but not of others (Ralambranto et al., 1982). Later, cell division in lymphocytes in culture was shown to be stimulated (Ralambranto et al., 1987). A commercial aloe product (ALVA) was presented as an immunostimulant, augmenting the production of tumour necrosis factor (Michel et al., 1989).

The therapeutic properties of *A. vera* are of course of prime interest and studies similar to those on *A. arborescens* have been made on that plant. Using a membrane filter a high molecular weight compound of aloe gel above 10 000 Da was separated from a low molecular weight component. The high molecular weight fraction, perhaps polysaccharide, was shown to deplete complement components while the low molecular weight fraction interfered with processes in activated polymorphonuclear leukocytes which led to the production of oxygen free radicals (t'Hart et al., 1988) and subsequent cytotoxicity (t'Hart et al., 1990). The high molecular weight fraction was further separated by gel filtration into two polysaccharide components, B1 (320 000 Da) and B2 (200 000 Da), largely composed of mannose (t'Hart et al., 1989). Both substances show anti-complement activity at the C3 activation step. Elsewhere a proprietary substance isolated from the gel and called acemannan (Carrisyn®) by Carrington Laboratories, Texas was described as an acetylated polymannan at the 1987 meeting of the American Society of Clinical Pathologists and reported to stimulate the immune system (McDaniel and Mcanalley, 1987). It may be that the whole range of activities described for this substance (Table 3) relate to immunological properties. Acemannan was shown to stimulate antigenic responses of human lymphocytes as well as the mitogenic response (Womble and Helderman, 1988) but not to be mitogenic itself. The reaction seemed to be specific for acemannan compared with other polysaccharides and the effect specific for the stimulated generation of T cells (Womble and Helderman 1992). Acemannan injected subcutaneously into irradiated mice (myelosuppressed) stimulated the formation of all types of leucocytes (Egger et al., 1996b), from both spleen and bone marrow, although responses in the two locations were different and depended on the dose rate (Egger et al., 1996a). Mouse macrophages in monolayer culture were stimulated by acemannan to show an enhanced respiratory burst and increased phagocytosis (Stuart et al., 1997). This was reflected clinically by an observed increase in leucocyte count in horses suffering from lethargy syndrome, associated with persistent leucopaenia (Green, 1996). Acemannan injected into mice or added to murine macrophage cultures stimulated the synthesis of a variety of immunologically active interleukins (Merrim et al., 1996). Involvement with interferons was suggested by observations of the induction of programmed cell
death (apoptosis) of macrophages in culture in the presence of IFNγ (Ramamoorthy and Tizard, 1998).

One feature of acemannan-induced macrophages in a chicken bone marrow cell culture was an increase in nitric oxide production said to contribute to cytotoxicity (Karaca et al., 1995). In mouse macrophage cultures, nitric oxide synthase levels were increased by transcriptional activation of the appropriate gene caused by acemannan (Ramamoorthy et al., 1996). In contrast it was shown in vivo that nitric oxide inhibitors enhanced wound healing by limiting generation of oxygen radicals, while addition of aloe gel also enhanced the process (Heggers et al., 1997).

The active gel constituents just described were found to be polysaccharides free of any polypeptide chains. Other work with A. vera went back to the idea of active glycoproteins, lectins. Partially purified fractions prepared by differential step-wise centrifugation from whole leaves of A. vera and A. saponaria were shown to agglutinate human or canine erythrocytes and to stimulate immune reactions against human, canine and baboon sera (Winters et al., 1981) These fractions were also shown to stimulate cell division of lymphocytes (blastogenesis) and that lectin-like haemagglutination was associated with terminal D-mannose (Winters, 1991). Subsequently other gel fractions were found to suppress cell growth (Winters, 1992). Separation by gel filtration enabled these activities to be measured more accurately and suggested that activity resided at the glucose and mannose sites of the glycoprotein (Winters, 1993). They resembled the A. arborescens lectins in many of their properties. Further separation by polyacrylamide gel electrophoresis revealed the presence of 23 distinct polypeptides in whole leaf preparations, of which 13 occurred in the gel (Winters and Bouthet, 1995). Of these, 19 showed lectin activity towards specific antibodies against five known lectins in an immuno-blot assay. Most showed the presence of mannose and glucose in the polysaccharide moiety and a few the presence of galactose and N-acetylgalactosamine. In the same study five major polypeptides were found in whole leaf preparations from A. saponaria. A commercial sample of lyophilized A. vera gel was found to contain 12 polypeptides by the same method (Bouthet et al., 1996). Lectin activity was determined only for the unseparated material and shown as haemagglutination which was glucosamine specific.

6. Effects on gastrointestinal function and ulcers

Aloe gel is offered commercially for oral consumption and many claims are made for benefits in various internal inflammatory conditions. A series of trials on human patients indicated a tonic effect on the intestinal tract with a reduced transit time. Also the bacterial flora appeared to benefit, with a reduction in the presence of yeasts and a reduction in pH. Bowel putrefaction was reduced and protein digestion/absorption improved (Bland, 1985). Preadministration of a water extract of whole A. vera leaves to rats, reversed the inhibition by blood ethanol of alcohol dehydrogenase and aldehyde dehydrogenase activities. It also reversed the increase of lactate/pyruvate ratio which could decrease NAD supply. Thus ethanol levels in the blood were decreased but not uptake (Sakai et al., 1989).

An early trial with human patients found oral administration of aloe gel effective in the treatment of peptic ulcers (Blitz et al., 1963) although the mode of action could not be determined. However, these observations were contradicted later using experimentally induced gastric and duodenal ulcers in rats where both the exudate and the gel were found to be ineffective (Parmar et al., 1986). However, other workers claimed that a component from Cape Aloe exudate named aloe ulcin, suppressed ulcer growth and L-histidine decarboxylase in rats (Yamamoto, 1970, 1973), while another, cruel, experiment where ulcers were induced in rats by severe stress, showed that aloe gel, administered in advance, had a prophylactic effect and was also curative if given as a treatment (Galal et al., 1975) A lectin fraction (glycoprotein) from A. arborescens, Alocitin A, was active against gastric lesions in rats (Saito et al., 1989), while another high molecular weight fraction, not containing glycoprotein, was very effective in healing mechanically and chemically induced ulcers but
not those induced by stress (Teradaia et al., 1993). This fraction contained substances with molecular weights between 5000 and 50000 Da, which were considered to both suppress peptic ulcers and heal chronic gastric ulcers.

Oral ulcers (aphthous stomatitis) are troublesome because of the difficulty of applying and retaining a therapeutic agent. A clinical trial with the polysaccharide Acemannan accelerated healing time and reduced pain without the side effects attributed to other agents (Plemmons et al., 1994).

7. Anti-diabetic activity

Diabetes mellitus is a disorder of carbohydrate metabolism characterized by lowered insulin secretion. It is a syndrome with both hereditary and environmental factors and has been classified into a number of types or groups, among which are the insulin-dependent and non-insulin dependent types. It is evident that causes, symptoms and treatments are varied and need to be carefully distinguished. An early clinical trial in India where over 3000 'mildly' diabetic patients were fed with bread incorporating aloe gel, demonstrated a reduction in blood sugar levels in over 90% of the cases (Agarwal, 1985). A survey of patients in Texas showed that 17% of those of Mexican origin used A. vera in an unspecified way, presumably with satisfaction (Noel et al., 1997). Dried aloe exudate has been used in Arabia in diabetes treatment. Administration to non-insulin dependent human patients in a small trial resulted in a sustained lowering of blood sugar levels (Ghannam et al., 1986). A similar effect was achieved on mice, made diabetic with alloxan treatment (Ajabnoor, 1990). Again, a number of diabetic patients in Thailand were treated orally with 'A. vera juice', to their benefit. Blood sugar and triglyceride levels fell during the treatment period (Yongchaiyudha et al., 1996). In parallel trials, patients that failed to respond to other anti-diabetic medication responded to the aloe treatment in a similar way (Bunyapraphatsara et al., 1996b). On the other hand an A. vera gel preparation was found to be ineffective in lowering blood glucose levels of alloxan-treated rats (Koo, 1994) and in fact seemed to cause an increase. Elsewhere, no effect was found using normal rats (Herlihy et al., 1998b). The question with all these studies is what Aloe leaf constituents are being tested. It is sometimes not clear how rigorous is the separation of the mucilaginous gel and the exudate anthraquinones. Polysaccharide fractions from water extracts of whole leaves of A. vera, A ferox Mill., A. perryi Baker, A. africana Mill. and A. arborescens were found to lower blood glucose levels in normal mice (Hikino et al., 1986). Two polysaccharides were separated from A. arborescens extract and described as Arboran A (molecular weight 12000 Da, 17% O-acetyl groups, 2.5% peptides) and Arboran B (molecular weight 57000 Da, 5% O-acetyl groups, 10% peptides). Both lowered blood glucose levels in alloxan-induced diabetic mice. On the other hand a 'bitter principle' separated from crystalline (sic) aloe, presumed to be from A. vera, produced significant lowering of fasting blood glucose levels when injected into alloxan-treated mice, both in a few hours and after several days (Ajabnoor, 1990). A more detailed study using leaf skin and pulp preparations as well as those from the whole leaves of A. arborescens, showed that the effects were more complex than previously thought (Beppu et al., 1993). By using acetone precipitation to prepare the active fraction they aimed to eliminate anthraquinone material which previous workers might have included. They found that preparations from both the outer regions of the leaf and the inner gel caused a decrease in blood glucose level in mice. With gel components, perhaps glycoproteins, this rapid fall in glucose was followed by a rise when treatment was discontinued. There was also a significant rise in insulin level. The leaf skin preparation also lowered blood glucose and in artificially induced diabetic animals normal insulin production was resumed. High levels of carboxypeptidases were found in this fraction.

Decreased wound healing associated with diabetes is a likely subject for aloe gel treatment. It was demonstrated that in rats an A. vera gel preparation injected subcutaneously promoted diabetic wound healing, reduced abnormal sensitivity to pain and reduced oedema induced by
mustard (Davis et al., 1988). In a following study, both A. vera gel and surprisingly, gibberelic acid, were reported as having almost equal inflammation-reducing properties in chemically induced diabetic mice (Davis and Maro, 1989). In a later trial both excision and incision wounds in chemically induced diabetic rats healed more rapidly after both oral or topical applications of aloe gel. Collagen and hexosamine levels were higher during the early part of healing (Chithra et al., 1998b).

It would seem that at least two processes are being described in these reports. The first results in lowering of blood glucose levels and involves either a leaf exudate component or a glycoprotein and the second results in wound healing, recalling the classic effects of gel polysaccharides.

8. Anti-cancer activity

Agents active against neoplasms are much sought after and aloe preparations are of course obvious candidates. An early report claimed anti-tumour activity in an ethanol-precipitated fraction, aloemicin, from ‘Cape Aloe’ here described as A. ferox, A. vera and A. africana (sic) (Soeda, 1969). A large epidemiologic survey of lung cancer and smoking in Japan suggested that ingestion of aloe ‘juice’, presumably the gel, prevented (sic) pulmonary carcinogenesis and was said to prevent (sic) stomach and colon cancer. Whether it was claimed to also suppress cancers already established was not clear (Sakai, 1989).

Whole freeze-dried leaves of A. arborescens were fed to rats subsequently challenged with either of two carcinogens, an undefined pyrolysis product (the initiation stage) or diethyl nitrosamine (the promotion stage), acting on the liver. The initiation stage was somewhat depressed, while there was a significant reduction in tumour promotion (Tsuda et al., 1993). In 1995, a Japanese patent was filed claiming mutagenesis inhibition by aloe emodin from A. arborescens (Inahata and Nakasugi, 1995). A much earlier paper had described antileukemic activity by aloe emodin from Rhamnus frangula L. (Kupchan and Karim, 1976) and later cytotoxicity against human leukemia cells in culture was observed (Grimaudo et al., 1997).

Activity has been claimed for two fractions from aloes, glycoproteins (lectins) and polysaccharides. Lectin-like substances (sic) from leaves of A. vera and A. sapomaria and a commercial aloe gel were shown to have haemagglutinating properties and fresh preparations also promoted growth of normal human cells in culture but inhibited tumour cell growth (Winters et al., 1981). The commercial aloe gel showed an unspecific cytotoxicity. Interestingly another commercial aloe gel had previously showed cytotoxicity (Brasher et al., 1969). Two glycoproteins, aloctin A and aloctin B were separated from A. arborescens. The first one which is smaller (molecular weight 7500) is water soluble. Thus growth of an induced fibrosarcoma in mice was inhibited by aloctin A perhaps by an immunologic route, as cytotoxicity was not observed (Imanishi et al., 1981). The level of a mouse serum protein named hemopexin was shown to increase during development of some tumours, implying a defensive response. Injection of aloctin A also produced a serum protein increase for a short period and this was correlated with anti-tumour activity (Ishiguro et al., 1984). Another glycoprotein, ATF1011, from A. arborescens, distinct from the aloctins was shown to augment anti-tumour immunity in mice by T-cell activation but not by direct cytotoxicity (Yoshimoto et al., 1987).

A polysaccharide, ‘aloe mannan’, molecular weight 15 000 Da, isolated from A. arborescens inhibited growth of an implanted sarcoma in mice (Yagi et al., 1977). Later another polysaccharide fraction, molecular weight above 30 000 Da from A. vahombe (sic) was shown to reduce the growth of an induced fibrosarcoma in mice, perhaps by stimulation of phagocyte activity (Ralamboranto et al., 1982). Similar effects of the commercial polysaccharide fraction Acemannan™, an acetylated mannan from A. vera, against tumour growth were later noted. Growth of a murine sarcoma implanted in mice, showed regression after acemannan treatment (Peng et al., 1991), probably through an immune attack. Injection of mice with acemannan inhibited the growth of murine sarcoma cells implanted subsequently and
decreased mortality by about 40% (Merriam et al., 1996). Elsewhere, activation of macrophages was again reported (Zhang and Tizard, 1996). Clinical observations on acemannan-treated animals suggested that soft tissue sarcomas initially increased in size but that this was followed by fibrous encapsulation, invasion by lymphocytes and necrosis (Harris et al., 1991). In another clinical survey initial tumour (fibrosarcoma) growth was again observed, followed by necrosis (King et al., 1995).

In a very recent study, carcinogenesis by DNA adduct formation was shown to be inhibited by a polysaccharide-rich aloe gel fraction in an in vitro rat hepatocyte model (Kim and Lee, 1997).

### Table 1

A selection of microorganisms inhibited by aloe gel preparations

<table>
<thead>
<tr>
<th>Organism</th>
<th>Material</th>
<th>Procedure</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streptococcus pyogenes</td>
<td>Gel</td>
<td>Tube dilution</td>
<td>H.</td>
</tr>
<tr>
<td></td>
<td>Exudate</td>
<td>Agar diffusion</td>
<td>L.</td>
</tr>
<tr>
<td>S. agalactiae</td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td>Citrobacter sp.</td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td>Serratia marcescens</td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td>Enterobacter aerogenes</td>
<td></td>
<td></td>
<td>Hk.</td>
</tr>
<tr>
<td>Enterobacter sp.</td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td>Bacillus subtilis</td>
<td>Ethanol extract</td>
<td>Agar diffusion</td>
<td>L.</td>
</tr>
<tr>
<td>Klebsiella pneumoniae</td>
<td>Exudate</td>
<td>Agar diffusion</td>
<td>Hk.</td>
</tr>
<tr>
<td>Klebsiella sp.</td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td>Whole leaf</td>
<td>Liquid culture</td>
<td>G.</td>
</tr>
<tr>
<td></td>
<td>Whole leaf (A., littoralis)</td>
<td>Agar diffusion</td>
<td>GP.</td>
</tr>
<tr>
<td>Escherichia coli</td>
<td>Exudate</td>
<td>Agar diffusion</td>
<td>L.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>H.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>G.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>GP.</td>
</tr>
<tr>
<td>Candida albicans</td>
<td>acemannan</td>
<td>phagocyte culture</td>
<td>H., S.</td>
</tr>
<tr>
<td>Mycobacterium tuberculosis</td>
<td>Exudate (4 species)</td>
<td>Tube dilution</td>
<td>B.</td>
</tr>
<tr>
<td>Corynebacterium xerose</td>
<td>Exudate</td>
<td>Agar diffusion</td>
<td>L.</td>
</tr>
<tr>
<td>Salmonella paratyphi</td>
<td>Exudate</td>
<td>Agar diffusion</td>
<td>L.</td>
</tr>
<tr>
<td>Pseudomonas aeruginosa</td>
<td>Exudate</td>
<td></td>
<td>S.</td>
</tr>
<tr>
<td>Proteus vulgaris</td>
<td>Exudate</td>
<td></td>
<td>S.</td>
</tr>
<tr>
<td>Streptococcus faecalis</td>
<td>Gel</td>
<td>Agar diffusion</td>
<td>R.</td>
</tr>
</tbody>
</table>

several *Aloe* species where barbaloin, however, was said to be the most active component (Dopp, 1953). A later, very careful study, with *Staphylococcus aureus* and *Escherichia coli* in both agar plate and liquid broth cultures failed to show any activity in either the gel or the outer leaf layers (Fly and Kiem, 1963). A vehement rebuttal of this was made the following year, reporting inhibition of bacterial growth by very fresh or freeze-dried *Aloe* 'juice', presumably the exudate (Lorenzetti et al., 1964). Various anthraquinones were found to be inactive but the main such compound in the exudate, barbaloin, an anthrone-C-glucoside, was not tested. Shortly afterwards other workers reported antibacterial activity in various *Aloe* preparations (Soeda et al., 1966; Bruce, 1967; Heggers et al., 1979; Robson et al., 1982). In a clinical trial, aloe gel used to treat burns, controlled bacterial growth which was otherwise present in the untreated controls (Heck et al., 1981) and similar results were achieved in experimental trials (Rodriguez-Bigas et al., 1988; Kivett, 1989), although the relevance of casual microorganisms was challenged (Kaufman et al., 1989). There remained a divergence of opinion as to the active agent, on the one hand said to be in the 'anthraquinonic' fraction (Bruce, 1967, 1975; Anton and Haag-Berrurier, 1980), on the other to reside in the gel (Heggers et al., 1979). The former view was supported by the demonstration of activity against *Bacillus subtilis* in ethanolic extracts of the leaf (Levin et al., 1988). It was suggested here that controversial data could be due to beneficial nutrients present in some aloes preparations.

There are two useful related outcomes if antibacterial activity of *Aloe* can be confirmed. Firstly there is the obvious general antibiotic activity against pathogens exemplified in the very first paper quoted (Gottshall et al., 1949) and secondly there is activity against bacteria which may be hindering the wound healing process and contributing to inflammation (Heggers et al., 1995). A report of clinical cases suggested that the gel was bactericidal towards *Pseudomonas aeruginosa* (Cera et al., 1980). In a much later study, acemannan prevented adhesion of *P. aeruginosa* to human lung epithelial cells in monolayer culture (Azghani et al., 1995). However, some studies failed to demonstrate antibacterial activity, especially in deep wounds which became so heavily infected that death eventually ensued (Bunyaphrathatsara et al., 1996b). In a trial with incision wounds in rats, aloe gel was found to be compared with standard antimicrobials and was found to speed wound healing, while the antimiicrobials had an initial retardant effect (Heggers et al., 1995). It may be that antibiotic factors are released by the healing tissues in response to aloe treatment.

Antifungal activity has received less attention. Unspecified inhibitory activity was reported against *Trichophyton* spp. (Soeda et al., 1966) by *A. ferox* 'juice'. More detailed work demonstrated weak inhibitory activity against spore germination and hyphae growth of *T. mentagrophytes* by high molecular weight components of *A. arborescens* leaves (Fujita et al., 1978b). In subsequent work an antifungal low molecular weight fraction was described, which almost certainly contained barbaloin (Kawai et al., 1998). At the same time inflammation of guinea pig paws infected with *T. mentagrophytes* was reduced by treatment with a whole leaf homogenate. Growth of the yeast *Candida albicans* was also somewhat inhibited by *A. ferox* 'juice' (Soeda et al., 1966) or by a processed *A. vera* gel preparation (Heggers et al., 1979). Later, extracellular killing of *Candida* by acemannan-stimulated macrophages was demonstrated (Stuart et al., 1997).

Anti-viral activity would be more remarkable and especially activity against human immunodeficient virus type 1 (HIV-1), which would arouse topical interest. Indeed, aloe gel was included in nutritional supplements used in a clinical trial with acquired immunodeficient syndrome (AIDS) patients, where it was said to be beneficial, without specific effects being recorded, rather, nutritional supplementation was emphasized as being very important (Pulse and Uhlig, 1990). A polysaccharide fraction from aloe gel, the acetylated mannan acemannan, had previously been used to treat AIDS patients. A 71% reduction in symptoms was recorded, perhaps due to stimulation of the immune system (McDaniel et al., 1987), although some patients seemed to show no response (McDaniel et al., 1988). A further clinical study using cats infected with fe-
line leukemia, a normally fatal disease showed, again, a 71% survival rate over 12 weeks (Sheets et al., 1991) probably through immunostimulation. Acemannan used as an adjuvant to Newcastle disease virus and infectious bursal disease virus antigens in chick, increased virus titre with no toxic side effects (Chinnah et al., 1992). A similar effect was shown with acemannan as an adjuvant for turkey herpes virus vaccine to control Marek's disease in both laboratory and field trials with chickens (Nordgren et al., 1992). Injection with acemannan reduced immunosuppression following revirus challenge, perhaps by macrophage stimulation (Sharma et al., 1994). In a similar trial acemannan was successfully used as an antigen adjuvant against polyomavirus in a variety of birds again with the mildest of side effects (Ritchie et al., 1994). Acemannan also showed antiviral activity in vitro against measles, herpes, feline rhinotracheitis and HIV in monolayer culture (McAnally et al., 1988). In a trial using cats suffering from feline immunodeficiency virus, sepsis was decreased and lymphocyte count increased together with an extension of survival rates, following injection with acemannan (Yates et al., 1992). Infectivity of HIV-1, herpes simplex and Newcastle disease viruses and virus-induced cell fusion in two cultured target cell lines was reduced in the presence of acemannan (Kemp et al., 1990). Again, in human lymphocyte cultures infected with HIV-1, acemannan increased cell viability and reduced viral load, perhaps by inhibiting glycosylation of viral glycoproteins (Kahlon et al., 1991a). Other effects included inhibition of virus-induced cell fusion and suppression of virus release. Acemannan acted synergistically with azidothymidine, enabling lower doses of this agent to be used effectively (Kahlon et al., 1991b). It may well be that many of the antibiotic and also indeed antitumour effects are brought about by stimulation of natural killer cell activity (Marshall and Druck, 1993), an effect also observed with the lectin, Alectin A (Imanishi and Suzuki, 1984). In another trial with advanced HIV patients treated with acemannan, no increase in CD4 cells or viral burden could be demonstrated (Montaner et al., 1996).

Another aloe component, aloe emodin, was shown to disrupt the coating of enveloped viruses such as herpes and influenza virus A, while showing no cytotoxicity to the host cells (Sydiskis et al., 1991). In a clinical trial genital herpes was treated by either an ethanolic extract of freeze dried leaves made up as a cream or the raw gel, both applied topically. Significant healing was achieved but it was not clear if this was a direct action on the virus or some host mediated response (Syed et al., 1996a). Fractions from aloe gel containing lectins were also shown to directly inhibit proliferation of cytomegalovirus in cell culture, perhaps by interfering with protein synthesis (Sahoo et al., 1996).

10. Various activities
10.1. Radiation effects on skin

The modern awakening of interest in aloe gel resulted from treatment of X-ray burns, so it was natural to extend these observations to other forms of radiation. Topical applications of gel were found not to alter the development of either erythema or increased blood flow in human skin exposed to UVB radiation (Crowell et al., 1989). A detailed study of the interactions of UVB and aloe gel on mouse skin demonstrated that the gel prevents immune suppression by UV. This was shown where UV suppressed the immune reaction to either fluorescein or Candida infection but the effect was reversed by gel application. No sunscreen activity was found but the effects of exposure were less deleterious following gel application up to 48 h after exposure. The gel restored the activity of various epidermal cells reduced by UV exposure (Strickland et al., 1994). A much briefer study on photo-ageing of skin indicated that treatment of skin with aloe extracts increased the soluble collagen level (Danof, 1993 quoting Stachow et al., 1984). A later study demonstrated that acetylated mannan from Aloe increased collagen biosynthesis perhaps through macrophage stimulation (Lindblad and Thu, 1994). Elsewhere, a gel component of between 500 and 1000 Da recovered the supression of Langerhans cell acces-
sory cell function induced by UVB radiation (Lee et al., 1997). Damage by free radicals has often been invoked to explain radiation effects and it is interesting that both glutathione peroxidase (Sabeh et al., 1993) and superoxide dismutase (Sabeh et al., 1996) activities have been reported from A. vera gel.

10.2. Cholesterol levels

In a small trial with monkeys it was found that orally administered aloe gel lowered total cholesterol by 61% and also that proportion in the high density lipoprotein (HDL) increased (Dixit and Joshi, 1983).

10.3. Hormone levels

In a trial with rats, ingestion of aloe gel lowered plasma levels of calcitonin and parathyroid hormone (Herlihy et al., 1998b).

10.4. Psoriasis

In a large clinical trial, an A. vera extract, compared with a placebo, significantly cured a large number of patients (Syed et al., 1996b).

11. Deleterious effects

In contrast to clinical reports of no useful activity with aloe gel, there were also a few cautionary accounts of harmful effects. An early report of a single case of an eczema appearing after topical and internal application of A. vera gel (Morrow et al., 1980) was followed by another on A. arborescens gel with a hypersensitive patient (Shoji, 1982) and then with a young child (Nakamura and Kotajima, 1984). On the other hand a patch test trial on 20 human subjects exposed to UV radiation showed only a persistent skin pigmentation (Dominguez-Soto, 1992). The effect of an allergic dermatitis arising in regions remote from the area of application was again described, in some detail (Hogan, 1988), where it hindered the treatment of chronic leg ulcers. In another study of this intransigent lesion, healing with the gel was successful, although local pain was experienced at first, attributed to improved circulation (El-Zawahry et al., 1973). In a study of burns it was suggested that the nature of the healing process depended on the type of damage, which in turn depended on the depth of the wound and that aloe gel could impair some wound healing by not fulfilling all the healing requirements (Kaufman et al., 1988). This multiplicity of factors in the healing of wounds was emphasized in clinical trials where facial skin was deliberately abraded (Fulton, 1990). Here aloe gel-treated zones healed more rapidly and completely than untreated zones although again burning sensations were sometimes noted. In a quite separate case application of aloe gel resulted in a severe burning sensation, followed by long term erythema (Hunter and Frumkin, 1991). With a different type of wound, those following Caesarean delivery, treatment with a proprietary gel fraction delayed healing and was discontinued (Schmidt and Greenspoon, 1993).

A controlled toxicological evaluation of acemannan administered by injection into mice, rats and dogs failed to identify any adverse effects but there was an increase in circulating leucocyte count probably as a result of stimulation of the immune system. There was also a concentration of macrophages in lungs, liver and spleen (Fogleman et al., 1992b). Elsewhere macrophages in culture were shown to be stimulated by acemannan (Zhang and Tizard, 1996). A study of ingestion by rats of a diet containing up to 1% aloe gel showed no adverse effects on growth or pathological effects. (Herlihy et al., 1998a).

A factor which may or may not be relevant to the preceding remarks is the presence of exudate phenolic substances, notably antherone C-glycosides, in the gel as contaminants. It has been shown that the yellow leaf exudate killed fibroblasts in cell culture, whereas the clear gel stimulated cell growth (Danof, 1987). Elsewhere we have the concept of ‘colorized’ and ‘decolorized’gels (Davis et al., 1986a), where the former had much less healing capacity. The decolorized gel reduced wound swelling caused by infiltration of polymorphonuclear leucocytes to a greater extent than colorized gel (Davis et al., 1986b), as
well as reducing wound diameter more quickly (Davis et al., 1987a). Similar studies had compared a gel fresh from the plant and dialysed to remove low molecular weight components with a 'commercial stabilized' gel. Cytotoxic effects of the 'commercial sample' were observed but ascribed to substances introduced during processing (Winters et al., 1981). Some commercial samples were found to contain 'yellow sap' and were cytotoxic in fibroblast cell cultures (Danof and McAnalley, 1983). Later studies on a low molecular weight fraction (< 10,000 Da) from whole A. vera leaves showed that this had a disruptive effect on monolayer cell cultures and inhibited neutrophils from releasing bactericidal reactive oxygen species (Avila et al., 1997). Aloe emodin and aloin (barbaloin) had a similar effect.

12. Aloe gel constituents

During all these discussions on the pharmaceutical properties of aloe's a clear distinction should be made between substances in the colourless, tasteless parenchyma cells, the aloe gel and substances in the bitter exudate from cells associated with vascular bundles in the outer green rind of the leaf (Agarwala, 1997). As mentioned above, this distinction has sometimes been clouded by using extracts of the whole leaf or allowing, during preparation of the gel, exudate compounds to infiltrate. The concept of colored and decolorized gels described above, leads to confusion in ascribing activities to individual components. There may indeed be synergies which would not appear if the fractions were kept separate. In view of the complexities inherent in aloe pharmacology it might be better to be as rigorous as possible in separation, at least initially, and only combine factors at later stages of the investigation.

12.1. Exudate compounds

These are largely phenolic in nature and were reviewed some time ago (Reynolds, 1985). Many of the exudates from around 300 species have been examined chromatographically and about 80 main constituents distinguished. Of these many remain unidentified.

12.2. Gel compounds

Few Aloe species have been examined for gel constituents. Up to 1986 polysaccharides had been extracted and described from A. arborescens, A. vahombe (sic), A. plicatilis (L.)Mill. and A. vera (Grindlay and Reynolds, 1986) and A. saponaria and A. vanballentii Pillans (Gowda, 1980). Later, A. ferox was added to the list (Mabusela et al., 1990). In this species, arabinogalactans and rhamnogalacturonans were conspicuous whereas glucomannans, common in other aloe's, were less so. Work on the components of A. vera gel has of course continued. A study of the rheology of the gel suggested that glucomannans in aloe's were rarely found in most other plants and had plastic properties akin to those of human body fluids (Yaron 1991). In order to maintain the viscosity on storage, addition of other natural polysaccharides was beneficial (Yaron et al., 1992; Yaron, 1993). Table 2 summarises the types of polysaccharide so far reported from the seven species examined. They are largely glucomannans of various compositions, some acetylated and some not, although polymers of other hexoses occur, notably those in A. ferox. Galactose and galacturonic acid polymers are frequently found. It should be noted that observations by different investigators reveal differing polysaccharide structures, especially with A. vera on which most work was done. An acetylated mannann became available commercially and was known as acemannan or Carrisin™ (McDaniel et al., 1987). Its existence was announced at a conference in 1987 but minimal details of extraction, purification and characterization given, although much more information was released in subsequent patents (McAnalley, 1988, 1990). It is an acetylated mannann prepared from A. vera gel with a range of interesting biological activities (Table 3) and appears to consist of three chemical entities. Recently NMR studies have been reported and used as a means of quality control of the gel (Diehl and Teichmüller, 1998). It may be chemically related to the 'aloemannan' isolated somewhat earlier from A. arborescens (Yagi et al., 1977). It was followed in 1988 by another only partially described mannann species (OS2) also only described at a conference
<table>
<thead>
<tr>
<th>Species</th>
<th>Fraction</th>
<th>Type</th>
<th>Molecular weight (Da)</th>
<th>Hexose composition</th>
<th>Linkage</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Aloe vera</em></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1)</td>
<td>A1a</td>
<td>Glucomannan</td>
<td>450 000</td>
<td>Glc:Man:GlcA = 19:19:1</td>
<td>1→4</td>
<td>Farkas (1967)</td>
</tr>
<tr>
<td>2)</td>
<td>A1b</td>
<td>Glucomannan</td>
<td>&gt;2 \times 10^5</td>
<td>Glc:Man = 1:13.5</td>
<td>1→4</td>
<td>Gowda et al. (1979)</td>
</tr>
<tr>
<td></td>
<td>A2</td>
<td>Acetylated glucomannan</td>
<td>&gt;2 \times 10^5</td>
<td>Glc:Man = 1:19</td>
<td>1→4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>Acetylated glucomannan</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3)</td>
<td>A(C1)</td>
<td>Galactogalacturan</td>
<td></td>
<td>Gal:GalA:Rha = 1:20:1</td>
<td></td>
<td>Mandal and Das (1980a)</td>
</tr>
<tr>
<td></td>
<td>A(C2)</td>
<td>Galactogalacturan</td>
<td></td>
<td>Gal:GalA = 1:1</td>
<td>1→6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>A(C5)</td>
<td>Galactogalacturan</td>
<td></td>
<td>Gal:GalA = 25:1</td>
<td>1→4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>A3</td>
<td>Glucomannan</td>
<td></td>
<td>Glc:Man = 1:22</td>
<td>1→6</td>
<td>Mandal and Das (1980b)</td>
</tr>
<tr>
<td></td>
<td>B2(2)</td>
<td>Galactogalacturan</td>
<td></td>
<td>Gal:GalA = 1:5</td>
<td>1→4,1</td>
<td></td>
</tr>
<tr>
<td>4)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acemannan</td>
<td>Frn 1</td>
<td>Acetylated mannan</td>
<td>80 000</td>
<td>Man:Ac = 16:5 (approx.)</td>
<td>1→4</td>
<td>McAnalley (1988)</td>
</tr>
<tr>
<td>(Carrisyn™)</td>
<td>Frn 2</td>
<td></td>
<td>10 000</td>
<td></td>
<td></td>
<td>Manna and McAnalley (1993)</td>
</tr>
<tr>
<td></td>
<td>Frn 3</td>
<td></td>
<td>1 000</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Aloe arborescens*

<p>| 1)             | A        | Mannan                 | 15 000                | 1→4                     | Yagi et al. (1977) |
|                | B        | Acetylated mannan      |                       | 1→4                     |                     |
| 2)             | A        | Glucan                 | 15 000                | 1→6                     | Yagi et al. (1986) |
|                | B        | Arabinogalactan        | 30 000                | 1→6                     |                     |
|                | C        | Acetylated mannan      | 40 000                | 1→4                     |                     |</p>
<table>
<thead>
<tr>
<th>Species</th>
<th>Fraction</th>
<th>Type</th>
<th>Molecular weight (Da)</th>
<th>Hexose composition</th>
<th>Linkage</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>3) Arboran A</td>
<td>Acetylated glucorhamnoglactan Mannogluca</td>
<td>1.2 × 10⁴</td>
<td>Glc:Rha:Gal = 0.3:0.3:1</td>
<td>Hikino et al. (1986)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arboran B</td>
<td>4)</td>
<td>5.7 × 10⁴</td>
<td>Man:Glc = 0.3:1</td>
<td>Wozniewski et al. (1990)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Acetylated glucomannan</td>
<td>1 × 10⁶</td>
<td></td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Acetylated glucomannan</td>
<td>12 000</td>
<td></td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Arabinogalactan</td>
<td>5 000 000</td>
<td>Arab:Gal:Rha:Glc = 43:43:7:7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aloe plicatilia</td>
<td></td>
<td>Acetylated glucomannan</td>
<td>1.2 × 10⁶</td>
<td>Glc:Man = 1:2:8</td>
<td>1 → 4</td>
<td>Paulsen et al. (1978)</td>
</tr>
<tr>
<td>Aloe rahonbe(sic)</td>
<td>1)</td>
<td>Acetylated glucomannan</td>
<td></td>
<td></td>
<td>1 → 4</td>
<td>Radjibi et al. (1983)</td>
</tr>
<tr>
<td></td>
<td>2)</td>
<td>Glucomannan</td>
<td>&gt; 10⁴</td>
<td>Glc:Man = 1:2</td>
<td>1 → 4</td>
<td>Vilkas and Radjibi Nassab (1986)</td>
</tr>
<tr>
<td></td>
<td>II</td>
<td>Glucomannan</td>
<td>100 000</td>
<td>Glc:Man = 7:3</td>
<td>1 → 4</td>
<td>Radjibi-Nassab et al. (1984)</td>
</tr>
<tr>
<td></td>
<td>III</td>
<td>Acetylated glucosylmannan</td>
<td>20 000</td>
<td>Glc:Man = 7:3</td>
<td>1 → 4</td>
<td>Vilkas and Radjibi Nassab (1986)</td>
</tr>
<tr>
<td></td>
<td>IV</td>
<td>Glucomannan</td>
<td>2 500</td>
<td>Glc:Man = 1:4</td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td>Aloe ferox</td>
<td>B1</td>
<td>Arabinorhamnoglactan</td>
<td></td>
<td>Arab:Rha:Gal = 1:2:0:6:1</td>
<td>1 → 4, 1 → 5</td>
<td>Mabuse et al. (1990)</td>
</tr>
<tr>
<td></td>
<td>B2</td>
<td>Arabinorhamnoglactan</td>
<td></td>
<td>Arab:Rha:Gala = 0:7:0:6:1</td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>B3</td>
<td>Xylorhamnoglactan</td>
<td></td>
<td>Xyl:Rha:Gal = 1:6:1</td>
<td>1 → 4, 1 → 5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>B4</td>
<td>Xyloglucan</td>
<td></td>
<td>Xyl:Glc = 2:1</td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>B5</td>
<td>Xyloglucan</td>
<td></td>
<td>Xyl:Glc = 1:3:1</td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AS1b</td>
<td>Glucagalactomannan</td>
<td></td>
<td></td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AS2</td>
<td>Acetylated mannan</td>
<td></td>
<td></td>
<td>1 → 4</td>
<td>Yagi et al. (1984)</td>
</tr>
<tr>
<td></td>
<td>2)</td>
<td>AS1</td>
<td>Acetylated mannan</td>
<td>15 000</td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AS2</td>
<td>Acetylated glucosylmannan</td>
<td>66 000</td>
<td>Glc:Man = 5:95</td>
<td>1 → 4, 1 → 2</td>
<td></td>
</tr>
<tr>
<td>Aloe canbalenii</td>
<td>AV1a</td>
<td>Glucan</td>
<td></td>
<td>Glu:Gal:Mann = 1:0:5:1</td>
<td></td>
<td>Gowda (1980)</td>
</tr>
<tr>
<td></td>
<td>AV1b</td>
<td>Glucagalactomannan</td>
<td></td>
<td></td>
<td>1 → 4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AV2</td>
<td>Acetylated mannan</td>
<td></td>
<td></td>
<td>1 → 4</td>
<td></td>
</tr>
</tbody>
</table>
Table 3
A selection of references to biological activity of Acemannan (Carrisyn™)

<table>
<thead>
<tr>
<th>Activity</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inhibition of bacterial adhesion to human lung cells</td>
<td>Aghaian et al. (1995)</td>
</tr>
<tr>
<td>Adjuvant to virus</td>
<td>Chinnah et al. (1992)</td>
</tr>
<tr>
<td>Stimulation of macrophage formation</td>
<td>Egger et al. (1996a)</td>
</tr>
<tr>
<td>Lack of toxic reactions</td>
<td>Fogelman et al. (1992a)</td>
</tr>
<tr>
<td>Lack of oral toxicity</td>
<td>Fogelman et al. (1992b)</td>
</tr>
<tr>
<td>Stimulation of leucocyte production</td>
<td>Green (1996)</td>
</tr>
<tr>
<td>Necrosis of canine and feline tumours</td>
<td>Harris et al. (1991)</td>
</tr>
<tr>
<td>Suppression of virus replication in vitro</td>
<td>Kahlon et al. (1991a)</td>
</tr>
<tr>
<td>AIDS therapy</td>
<td>Kahlon et al. (1991b)</td>
</tr>
<tr>
<td>Modification of glycosylation of viral glycoproteins</td>
<td>Kemp et al. (1990)</td>
</tr>
<tr>
<td>Regression of fibrosarcomas</td>
<td>King et al. (1995)</td>
</tr>
<tr>
<td>Stimulation of collagen synthesis</td>
<td>Lindblad and Thul (1994)</td>
</tr>
<tr>
<td>Anti-viral activity in cell cultures</td>
<td>McAulay et al. (1988)</td>
</tr>
<tr>
<td>AIDS therapy</td>
<td>McDaniel et al. (1988)</td>
</tr>
<tr>
<td>AIDS therapy</td>
<td>McDaniell (1987)</td>
</tr>
<tr>
<td>Adjuvant to herpes vaccine</td>
<td>Nordgren et al. (1992)</td>
</tr>
<tr>
<td>Regression of murine sarcoma</td>
<td>Peng et al. (1991)</td>
</tr>
<tr>
<td>Healing of oral ulcers</td>
<td>Plemmons et al. (1994)</td>
</tr>
<tr>
<td>Adjuvant to virus antigen</td>
<td>Ritchie et al. (1994)</td>
</tr>
<tr>
<td>Healing of radiation burns</td>
<td>Roberts and Travis (1995)</td>
</tr>
<tr>
<td>Clinical stabilization of feline leukemia</td>
<td>Sheets et al. (1991)</td>
</tr>
<tr>
<td>Stimulation of phagocytosis</td>
<td>Stuart et al. (1997)</td>
</tr>
<tr>
<td>Wound healing</td>
<td>Tizard et al. (1994)</td>
</tr>
<tr>
<td>Induction of cytokines</td>
<td>Tizard et al. (1991)</td>
</tr>
<tr>
<td>Stimulation of lymphocyte response to alloantigen</td>
<td>Womble and Helderman (1988)</td>
</tr>
<tr>
<td>Relief of feline AIDS</td>
<td>Yates et al. (1992)</td>
</tr>
<tr>
<td>Stimulation of macrophages</td>
<td>Zhang and Tizard (1996)</td>
</tr>
</tbody>
</table>

(Eberendu et al., 1988). Apart from technical inconsistencies it appears that the range of carbohydrate types may relate to plants of different geographical origin or possible varieties or even subspecies.

Most of the polysaccharide preparations mentioned above contain very little or no nitrogen. However a fraction from A. arborescens gel was shown to be a glycoprotein, appearing as a single electrophoretic band (Yagi et al., 1986), while two glycoprotein fractions were separated by differential precipitation (Kodym, 1991). Haemagglutinating activity typical of lectins was found in fractions from the gels of A. vera, A. saponaria and A. chinensis (sic) (Winters, 1993).

More recently the polypeptide composition of gel proteins from Aloe species has been determined by sodium dodecyl sulphate polyacrylamide gel electrophoresis (SDS-PAGE) which disrupts oligomeric proteins and sorts the resultant polypeptides according to molecular size (Winters and Yang, 1996). It was shown that A. saponaria, A. vera and A. arborescens had five major polypeptides in common with molecular weights 15,000, 46,000, 65–66,000, 71,000 and 76–77,000 Da. The species had totals of 11, 12 and nine major polypeptides, respectively.

Various biological activities have been ascribed to Aloe proteins, mentioned elsewhere in this review. Lectin activity is prominent among the glycoproteins and two entities, Alocitin A and Alocitin B were isolated from A. arborescens (Suzuki et al., 1979a; Saito, 1993). Alocitin A had a molecular weight of 18,000 Da and was made up of two subunits of 7500 and 10,500 Da with a carbohydrate content of 18%. Alocitin B was 24,000 Da with two subunits of 12,000 Da each and a carbohydrate content of 50%. A different glycoprotein, designated ATTF191, was subsequently prepared from the same species (Yoshimoto et al., 1987), while later another lectin, molecular weight 35,000 Da, was prepared from the outer layers of the leaf (Koike et al., 1995). An aloe preparation which healed excisional wounds in rats was shown to contain a high molecular weight polypeptide (Heggers et al., 1996). Recently a glycoprotein (Pg21-2b) with cell proliferation-promoting activity has been reported.
from *A. vera* gel (Yagi et al., 1997). It has a molecular weight of 29,000 Da and consisted of two subunits. Other protein fractions showed growth inhibiting activity which seemed however to be associated with phenolic contaminants.

A variety of simple substances have been found from time to time in aloe gel (Grindlay and Reynolds, 1986) although there is always the problem of complete separation from leaf exudate components (Agarwala, 1997). A recent, seemingly careful, study reported the presence of aluminium, boron, barium, calcium, iron, magnesium, manganese, sodium, phosphorus, silicon and strontium (Yamaguchi et al., 1993). Among the organic material was β-sitosterol, reported frequently before and large number of long chain hydrocarbons and esters which are more typical of industrial contaminants.

### 13. Gel preparation

It would seem that many of the inconsistent clinical results obtained for therapeutic efficacy of aloe gel result from the history of the sample after removal from the leaf, or even growing conditions of the plant (Yaron, 1993). In the commercial literature there are claims and counter-claims as to the superiority of one or another process. This was supposedly finalised in a report from the United Aloe Technologists Association (Morsy et al., 1983) and was reviewed more recently (Agarwala, 1997). Heat during pasteurization is one of the stresses imposed on the gel and there are advantages in using high temperatures for short times preferably with the addition of an antioxidant such as ascorbic acid (Ashleye, 1983). Muco polysaccharide integrity during storage was found to be preserved by the addition of other natural polysaccharides which act synergistically (Yaron, 1991, 1993; Yaron et al., 1992). An HPLC analysis using size exclusion chromatography, of a number of commercial ‘aloe’ products revealed widely differing levels of mucopolysaccharides (Ross et al., 1997). These processes are also important when the gel is intended for internal use where organoleptic properties are important (Gorloff, 1983) and additives must be carefully chosen (Yamoto, 1983). It may be that irrigation affects gel composition so that leaves from well irrigated plants have less polysaccharide than drier plants (Yaron, 1993). It was also claimed however that plant grown hydroponically had a higher carbohydrate content (Pierce, 1983). There are still other factors operating because a careful analysis of plants from many origins showed great variation in leaf size, pH, fibre content, calcium and magnesium contents and certain HPLC peaks (Wang and Strong, 1993). Finally a continuing problem is the presence of anthraquinone derivatives (‘aloin’) derived from the mesophyll exudate. Methods for addressing all these problems are set out in detail in two US Patents (McAnalley 1988, 1990).

### 14. Commercial production

Use of aloe gel and preparations containing it has become widespread and consequently a large industry has developed, mostly in Texas and Florida. One of the earliest producers was Carrington Laboratories (http://www.carringtonlabs.com/about.html) which used the expertise of staff from Texas A. and M. University and grows its plants in Costa Rica. Among a range of products the preparation named Aecamann or Carrisyn™ was much studied. An associated firm, Mannatech™ Incorporated (http://www.mannatechinc.com/) produced a similar mannose-based mucopolysaccharide from *A. vera*, marketed as Mannapol® by a Carrington subsidiary, Caroloe (http://www.aloevera.com/) backed by HPLC validation. Dr Madis Laboratories of New Jersey is another firm that was early in the field, supplying both the fresh gel and derived products. In view of the many claims made by aloe producers and the variable results achieved, the International Aloe Science Council (http://www.2.iasc.org/iasc/articles.html) was set up in 1981 by the Trade to try to establish standards. One major supporter of the Council is AloeCorp (http://www.aloeCorp.com/aloeCorp.htm) with estates in Texas and Mexico. They support a wide range of research activities and supply products in the form of the gel either in the raw form, concen-
trated or freeze or spray-dried. Another well established (1973) firm is Terry Laboratories (http://
www.terrylabs.com/index.htm) who are major suppliers of gel to many multinational companies
and are major supporters of aloe research and quality control. Dr Madis Laboratories Inc offers
the gel either as a purified extract or in a number of formulations. AloeVera Company UK (Forever
Living Products) (http://www.aloevera.co.uk/home.htm) are active in selling the gel and derived
products by franchise, using aloe grown in Texas. Many firms concentrate the gel by either mild air
drying or freeze drying. Examples are Concentrated Aloe Corporation (http://www.geocities.com/Heartland/Ridge/1396/concentrated_aloe/lisa.html),
CRH International Inc (http://www.aloealoe.com/raw.html) and Valley Aloe Vera Inc (http://
www.quikpage.com/valleyaloe). This is by no means a complete list, there are many other producers,
large and small, some of which have pages on World Wide Web.

An information site 'The Aloe vera studies organization' (http://www.aloe-vera.org/) gives
some interesting hints, although its botany is a little quaint. A similar site has been set up by
'Miracle of Aloe' (http://www.miracleofaloe.com/internal.htm) and another by 'Tripurtic
Laboratories' (http://www.primenet.comp hidden/hayward.html). Although these informative sites
make very positive, often triumphalist statements in favour of the efficacy of aloe gel for a variety of
ills, they do not make the extravagant claims which are a feature of some promotional literature,
even if their scientific descriptions are sometimes a little garbled.

15. Conclusion

The literature covered by the previous review (Grindlay and Reynolds, 1986) contained many
case reports and more or less anecdotal accounts of the healing powers of A. vera gel, especially for
skin lesions but extended by some to a host of other complaints (Bloomfield, 1985). Laboratory
studies indicated that there was indeed in vitro activity present but the relevance to in vivo activ-
ity was not always clear. Since then much more experimental work has been carried out and a
picture of biological activity properties is emerging. One feature that is becoming clear is that the
systems undergoing healing contain several interacting factors, each of which may be affected by
more than one component of the raw gel. It may be that some of the inconsistencies reported are
caused by unknown variation in any of these factors.

It certainly seems that one feature, immunostimulation, is frequently appearing as a major
contributory factor. This is associated with the presence in the gel of polysaccharides. These
substances occur in all plants, often as storage carbohydrates such as starch or inulin or structural
carbohydrates such as cellulose while others have a more limited distribution. Many of these spe-
cialized polysaccharides of unknown function in the plant have been found to be physiologically
active in animals and subjects for new therapies (Franz, 1989; Tizard et al., 1989; McAuliffe and
Hindsaul, 1997). Mucopolysaccharides also occur in saliva and it is fascinating to speculate if
the supposed therapeutic powers of dogs, licking wounds, is due to these substances. In Aloe an
acetylated glucosaminan was found to be biologically active, so much so that it was named ace-
mannan (Carrysin™). If the presence of acetyl groups is necessary for activity, one wonders if
this is because they cover a number of hydrophilic hydroxyl groups and thus make the molecule
more able to cross hydrophobic barriers in the cell. It may also be that some of these ester bonds
are particularly labile, accounting for differences of reported efficacy of different preparations. No
investigations appear to have been made into this or into the possibility of using other residues to
cover hydroxyl groups, except for methylation, described in an American patent (Farkas, 1967)
and this was to confer stability to the polymer chain. It should also be noted however that active
glycoproteins have also been demonstrated in aloe gel and may well play some part in therapeutic
activity, either immunologically as lectins or as proteases such as anti-bradykinins.

There seems to be ever-decreasing doubt that aloe gel has genuine therapeutic properties, cer-
tainly for healing of skin lesions and perhaps for many other conditions. It is also clear that the subject is by no means closed and much needs to be discovered, both as to the active ingredients and their biological effects. These ingredients, acting alone or in concert, include at least polysaccharides, glycoproteins, perhaps prostaglandins, small molecules such as magnesium lactate, infiltrating exudate phenolics and even, simplest of all, water.

Acknowledgements

The authors would like to thank Professors M.D. Bennett and Monique Simmonds for their encouragement and support and Dr N.C. Veitch for critically reviewing the manuscript.

References


Ashley, A.D., 1983. Applying heat during processing the commercial Aloe vera gel. Erde International 1, 40–44.


Azghani, A.O., Williams, I., Holiday, D.B., Johnson, A.R., 1995. A beta-linked mannan inhibits adherence of Pse-

domonas aeruginosa to human lung epithelial cells. Glyco-
biology 5, 39–44.


Bouthet, C.F., Schirf, V.R., Winters, W.D., 1996. Semi-purifi-


Bruce, W.G.G., 1975. Medicinal properties in the Aloe. Exces-
sa 57–68.


toherapy 3, 67–75.

Capasso, F., Mascolo, N., Autore, G., Duraccio, M.R., 1983. Effect of indomethacin on alloxin and 1,8 dioxo-


Cera, L.M., Heggers, J.P., Robson, M.C., Hagstrom, W.J., 1980. The therapeutic efficacy of Aloe vera cream (Der-

maide Aloe(TM)) in thermal injuries. Two case reports.


