Case Report

Anterior Ankle Full Thickness Skin Necrosis Treated with Copper Oxide Dressings without Debridement and Skin Grafting – A Case Report

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Abstract

Full thickness skin necrosis can result from number of causes, including trash emboli, external pressure, burns and severe skin infections and necrotizing fasciitis. Treatment is often dependent on the size and location of the necrotic skin, involvement of adjacent structures, the presence of active infection and the general condition of the patient. The anterior aspect of the ankle is unique in the sense that the structures there (skin, subcutaneous fat, fascia and tendons) are tight without muscles or padding structures to allow healing of skin necrosis without involvement of the deep structures. Therefore, in cases of skin and subcutaneous tissue necrosis the tendons have a very scarce blood underlying facia (e.g. the extensor retinaculum) is often involved, jeopardizing the extensor foot tendons. In such conditions, the treatment normally includes debridement of the necrotic and infected tissues, which often involves resecting the superior extensor retinaculum and the underlying extensors tendons, if they are involved. Since the tendons have a very scarce blood supply, their fate is often doomed either in the first debridement surgery or in the following period. Eventually, the goal of treatment is to fill the area with abundant granulation tissue to afford skin grafting in follow-up surgery. We describe a case in which treatment with Copper Oxide Dressings (COD) prevented infection by its
antibacterial activity while at the same time induced debridement of the necrotic fascia and granulation tissue formation. Subsequently rapid epithelization occurred allowing the wound to heal with normal skin and little scar. Thus, the copper dressing enabled to replace debridement and skin graft surgeries and achieve complete healing within 5.5 months with low cost, very little discomfort and full tendon function. We discuss the mechanisms of COD action that enabled this achievement.

**Keywords:** Copper; Wounds; Dressings; Necrosis; Tendons; Ankle

**Abbreviations:** COD: Copper Oxide Dressings; AI: Artificial Intelligence

1. **Introduction**

Full thickness skin necrosis can result from a number of causes, including trash emboli, external pressure, burns and severe skin infections and necrotizing fasciitis [1-3]. Treatment of skin necrosis is often dependent on the size and location of the necrotic skin, involvement of adjacent structures, the presence of active infection and the general condition of the patient [1]. An important aspect of taking care of a wound or any injury is to retain the function of the affected tissue, limb or any other organ and to protect it from becoming infected until full skin coverage is achieved. Treatment of skin necrosis often includes debridement of the necrotic skin and infected tissues [1]. However, skin necrosis of the anterior ankle necessitates special attention, as the anterior aspect of the ankle is unique in the sense that the structures there (skin, subcutaneous fat, fascia and tendons) are tight without muscles or padding structures. Thus, in cases of skin necrosis of the anterior ankle, the underlying facia (extensor retinaculum) is often involved, jeopardizing the extensor foot tendons (tibialis anterior, extensor hallucis longus and extensor digitorum longus). Since tendons have poor blood supply, once they are exposed, they are prone to microbial infection and granulation tissue rarely and scarcely grow over the tendons. Therefore, in the case of exposed tendons, the race competition between granulation tissue growth and infection of the dysvascular tendon often ends with tendon loss. Hence, the common clinical practice in such cases is debridement of the necrotic tissue and often resecting the extensor tendons if they are infected or exposed. Tendon’s function is most likely lost in cases of anterior ankle necrosis.

2. **Case Description**

Sixty-years-old man was admitted to the hospital and operated on due to infected olecranon bursitis in the right elbow (Figure 1). Past medical history included non-insulin-dependent diabetes mellitus, ischemic heart disease and chronic obstructive pulmonary disease secondary to heavy smoking in the past. On examination, the patient was febrile (39.0°C), and redness plus fluctuation were evident on the right elbow. Laboratory tests were compatible with acute bacterial infection with 12,700 leucocytes and CRP of 30 (normal < 0.5). Aspiration of the abscess revealed pus and methicillin sensitive staphylococcus aureus (MSSA) grew on culture. Incision and drainage were performed in the operating room on two occasions and intravenous (IV) antibiotic treatment was instituted (Cefamizine 2.0 gr x 3 times per day). The patient responded favorably to treatment. The surgical wound at the elbow looked clean and delayed primary suture was carried 10 days following the first surgery.

2.1 **Infected foot description**

The patient did not have prior foot infection or episode of diabetic foot. He did not have neuropathy. Pulses in the foot were normal. Due to poor venous access in the arm, an IV
catheter was installed in the left ankle (Figure 2a). The 12th post admission day, while the patient seemed to heal completely from the elbow infection, he developed local erythema, warm and bulla formation in the anterior ankle region, at the area of the venous catheter portal (Figure 2b). Clinical and imaging findings, were consistent with local infection involving the skin, subcutaneous tissue and fascia deep to the extensor tendons. Sagittal view with soft tissue window of computerized tomography demonstrated swelling and opacity of the subdermal tissues down to the extensor tendons and air bubbles, compatible with cellulitis and necrotizing fasciitis spreading deep to the peri-tendon area (Figure 3).

The patient responded to antibiotic treatment (Augmentin and Ciprofloxacin) and the infection remained contained at the anterior aspect of the ankle. The patient was discharged home with oral antibiotic treatment. The patient returned to the clinic for a follow-up visit. At that time there was full thickness necrosis at the anterior aspect of the ankle (Figure 2c). The wound was covered with thick dry eschar which seemed to be contracted below the level of the adjacent skin surface (Figure 2c). There was no redness surrounding the wound and it seemed that there was no active infection. On direct examination, the wound width (from medial to lateral) was 70 mm and on the long axis of the leg (from proximal to distal) it measured 40-45 mm. Total wound and necrotic tissue area was 20.5 cm², as measured by an artificial intelligence (AI) program (Tissue Analytics software [https://www.tissue-analytics.com/]), used for assessment and management of wounds [4, 5]. The patient was admitted to the ward. The treatment plan was to debride all the necrotic tissue, excise the tendons if they would seem to be compromised or infected, and allow granulation tissue formation growth with subsequent split thickness skin grafting (STSG). An alternative plan, with application of Copper Oxide Dressings (COD), was presented to the patient. The patient was aware of the lack of experience with this approach, but elected the non-operative mode of treatment. The patient was discharged home with 10 cm x 12 cm COD dressing to be changed twice weekly without antibiotic treatment. The patient was instructed should the that if infection flare up, surgical debridement will need to be conducted and he should return immediately for re-evaluation. Normal follow-up was carried out in the clinic at first by weekly intervals and later by bi-weekly and tri-weekly visits.

Hereafter, the time count described is from the beginning of COD treatment onward (Figure 4a). At one week the wound was dry with very little discharge on the dressing (Figure 4b). Gradually the eschar was contracted, and white amorphous tissue of fibrous-necrotic fascia was seen in the interval between the healthy skin and the eschar (Figure 4c). This interval area was covered by an additional layer of COD, to assure that all the wound area and the adjacent skin are protected by COD (Figures 4d and 4e). The white fascial-necrotic tissue underwent autolysis and red granulation tissue proliferated beneath the necrotic tissue and began replacing it, as was seen on Day 35 (Figure 4f). The red granulation tissue became dense at the periphery of the wound and the epithelial tissue began crawling into the wound (Figure 4g). During all this period until final healing, no signs of infection were seen, and antibiotic was not prescribed. At 10 weeks, the eschar was contracted and partially detached (Figure 4g). It was trimmed in the clinic (Figure 4h) to reveal the same whitish fibrous-necrotic tissue throughout the wound bed with granulation tissue forming underneath (Figure 4i). No tendons were visible (Figures 4h and 4i). Area of thick white tissue was bed-side debrided but most of it was left for autolysis by the granulation tissue enzymes, as was seen at 14 and 19 weeks of COD treatment (Figures 4j and 4k). The percentage of granulation tissue coverage of the wound constantly increased reaching...
almost 100% of the area at 14 weeks (Figure 5), as determined by the artificial intelligence program. During this stage, epithelization was rapid and accompanied by deposition of thick corneal layer. At the next visit, at 25 weeks from the beginning of COD treatment, there was no discharge on the dressing (Figure 4) and the wound was closed (Figures 4 and 5). Despite having the wound closed, the patient was asked to continue applying the COD on the closed wound area to assist in better skin regeneration and cosmetic appearance[6]. Nine months after the beginning of COD treatment, there was full function of the ankle including the extensor tendons (Figure 6). The patient’s full tendon activity at the anterior aspect of the ankle was noted (Figure 6). The scar was minimal and hardly discernible at one year following the COD treatment (Figure 7).

**Figure 1:** The patient was operated due to infected olecranon bursitis in the right elbow.

**Figure 2:** a) IV catheter was installed in the left ankle; b) infection developed with local erythema, warm and bulla formation was formed in the area of the venous catheter portal; c) full thickness necrosis developed at the anterior aspect of the ankle.
**Figure 3:** Computerized tomography of the infected area at the anterior aspect of the ankle. The yellow arrows show swelling, air bubbles, and opacity of the sub-dermal tissues down to the extensor tendons revealing the depth of the infection. The white arrows show depression of the skin which became necrotic, dry and shrinked.

**Figure 4:** Pictures taken of the wound from the beginning of the COD treatment of the wound until the wound was closed. The Days from the beginning of treatment are depicted.
Figure 5: Necrotic tissue area, total wound area and % granulation tissue over time as determined by the AI program.

Figure 6: Wound healed with full tendon activity.
3. Discussion

An important aspect of taking care of a wound or any injury, is to retain the function of the affected tissue, limb or area. The illustrated case report shows a case of full thickness skin necrosis anterior to the ankle joint in a diabetic patient, which solely resolved by the management with COD. Skin necrosis at this site is usually full thickness due to lack of padding and no muscle tissues between the deep layers and the fascia. The extensor tendons are just below the fascia, which at this site is the superior extensor retinaculum. The computerized tomography done during the infectious episode illustrated the depth of the infection down to and surrounding the extensor tendons. Blurring and air in the deep tissues at that area (Figure 3) were compatible with severe infection that led to the necrosis of the integuments and facia. The patient responded to antibiotic treatment and the infection did not spread further but the whole epicenter of the infection became necrotic. Although the infection was resolved with eschar formation, the adjacent tendons were prone to infection and necrosis due to poor blood supply. Granulation tissue was not expected to grow over the bare tendons. Regular treatment of debridement, most probably, would result in loss of the tendon function and loss of active extension of the foot and ankle. Nevertheless, conducting tissue surgical debridement of the dead eschar is usually the safest clinical intervention to avoid colonization of the dead tissue with bacteria and infection. The eschar itself, composed of dry coagulated proteins [7], is relatively resistant to infection, and may rightly be referred to as a temporary biological dressing. The infection though usually begins or penetrates through the interface between the eschar tissue (the dry gangrene) and the intact skin. It makes sense therefore to dress this interface with an antibacterial dressing such as dressings containing copper or silver. While dressings containing silver may have good antibacterial properties, there are several reports indicating that silver may impede the wound healing processes (e.g.; [8-10]).

In the current case we used antimicrobial dressings containing copper oxide microparticles to cover the wound [5, 6, 11, 12]. No development of infection occurred even when the interval between the intact skin and the “biological dressing” e.g. the eschar, was interrupted due to contracture and desiccation of
the eschar (Figures 4c-e). Throughout the 5 months to wound closure there was no episode of redness of any other sign of infection and antibiotic was never prescribed. Remarkably, in parallel, intense granulation promoting activity took place at the wound, both at the interface between the eschar and the skin (Figures 4d & 4e), but also underneath the eschar. This was clearly seen when the eschar was trimmed at 10 weeks of treatment (Figure 4f). The necrotic fascial-fibrous tissue was already seeded with large nests of granulation tissue, which emerged at first as islands or pockets or red stains (Figure 4f) to become a uniform layer 4 weeks later (Figure 4g). The induction of granulation tissue is in accordance with experimental studies conducted in full thickness non-infected wounds in diabetic mice, in which induction of angiogenesis was induced by the COD but not by silver dressings or control dressings without an active ingredient [13]. This observation was substantiated in a clinical study in which diabetic foot ulcers that were not healing despite standard of care were enrolled in the trial and treated with COD. Already after one week of using the COD, there was a statistically significant increase in granulation tissue formed [5].

One of the most striking observations of the current case was the apparent induction of autolytic debridement by the COD. This can be attributed to autolytic enzymes, such as matrix metalloproteinases, released by the granulation tissue being formed. It has been found that copper stimulates the expression and secretion of matrix metalloproteinase-1 by dermal fibroblasts, a collagenase that degrades interstitial collagen [14]. The direct effect of copper in promoting collagenase release from leukocytes in the cornea was described [15]. We found that copper induced the upregulation of fibroblast growth factors secretion in full thickness wounds [13]. Thus, the increased number of dermal fibroblasts and blood cells in the newly formed granulation tissue, together with the increased secretion of collagenases by the fibroblasts and possibly by the white blood cells, as was seen in the cornea, following their exposure to copper that reached them in situ beneath the necrotic tissue, may explain the autolytic debridement process we saw in the current case. In any case, the debriding effect of COD on the wound was unequivocally presented.

The effect of copper as a potent regenerative agent was also demonstrated in the increased epithelization. While during the first stages of wound healing, epithelization was not observed and the total wound surface did not significantly change (Figure 5), once granulation tissue prevailed and the necrotic tissue was debrided, the epithelization was very rapid, as can be seen in the transition from 10 to 19th weeks. The positive effect of copper on the epidermal layer expressed itself also in abundant formation of corneal layer (Figure 4g-j). The effect of copper in inducing epithelization was reported [13, 16]. We thus can see that the copper and the COD acted in accordance with the known orchestrated wound healing mechanism. This case thus demonstrates the complimentary concentrated effect of copper oxide, which, on one hand, prevented infection during the healing period due to its antibacterial activity; and on the other hand, at the same time, it stimulated autolytic debridement, saved the surgical procedure, and promoted granulation tissue formation. In addition, once granulation tissue covered the wound surface, rapid epithelization took over, sparing the need for skin graft surgery. The positive effect of COD continued after wound closure allowing elastin formation and more natural skin to be formed with better quality and cosmetic appearance. This is in accordance with previous observations [6], in which the scar formed was minimal and hardly discernible at one year following COD treatment (Figure 7). This is in line with the role that copper plays at the maturation stages of wound healing, with the capacity of copper to penetrate the epidermal layers of the skin
[17, 18], with its stimulation of collagen and elastin secretion by dermal fibroblast [18, 19] and with copper’s involvement with the efficient production and stabilization of the dermal extracellular matrix [20, 21]. Taken together, the combined effects of COD enabled healing without damage to the tendons, allowing complete recovery of the tendons function, and return of the foot and leg to normal function. This not only saved the patient from pain, but was comfortable and easy to use both by the patient and care givers, as well as saving significant cost to the healthcare system and to the patient.

**Conflict of Interests**

G.B. and T.R. are employees of MedCu; E.M. is a medical consultant of MedCu; MedCu developed the COD. A.R. has no conflict of interests.

**References**

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