Combined use of super-oxidised solution with negative pressure for the treatment of pressure ulcers: case report

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Key words
Dermacyn; VAC therapy; Pressure ulcer

Abstract
A 61-year-old patient was affected by flaccid paraplegia for 20 years because of post-traumatic medullar injury, level D10–D11, caused by an accidental fall, with stage IV sacral pressure ulcer for 3 years. The patient later developed stage IV sacral pressure ulcer. After 6 months, a new granulation tissue formation appeared in the wound and a reduction of its diameter was observed (length 20 cm, width 15 cm, depth 5 cm). We therefore treated the wound with PRP (platelet rich plasma) intra-lesion and peri-lesional injections. The wounds were covered with three-dimensional polymerised hyaluronic acid medicated biologic dressing. After the surgery, a moderate reduction in diameter and the depth was observed. Super-oxidised solution (SOS-Dermacyn) was applied to control infection locally together with negative pressure to control the exudate and the local bacteremia, to avoid infectious complications without application of systematic antibiotic therapy.

Key Messages
• a 61-year-old patient was affected by flaccid paraplegia for 20 years because of post traumatic medullar injury level D10-D11, caused by an accidental fall, with stage IV sacral pressure ulcer for 3 years
• the goals of treatment of pressure ulcers should be resolution of infection and promotion of wound healing
• a combination of surgical debridement and medical interventions may be required
• to evaluate the efficacy of the combination Dermacyn® and NPWT, we assessed the following parameters: reduction of exudates with a change in colour and reduction of odour, infection and the size of pressure ulcer
• to conclude, we noticed an immediate reduction of exudate with decreased maceration of surrounding skin
• the association of the two has ensured a significant reduction in healing time by controlling the local bacterial infection until the complete elimination of
regional debridement of the necrosis with formation of granulation tissue

5 cm) (Figures 1–3). We then decided to treat the wound with intra-lesion and peri-lesional PRP (platelet rich plasma) injections. The wounds were covered with a three-dimensional polymerised hyaluronic acid medicated biologic dressing (1, 2). After the operation, a moderate reduction of diameter and depth was observed. However, continuing with the medication and during the antibiotic treatment, no signs of infection were observed. When antibiotics treatment was not used, there was a recurrence of the infection signs in the wound bed, such as an increase of exudate, change of colour, hyperaemia, periwound maceration and fever. The goals of treatment of pressure ulcers should be resolution of infection and promotion of wound healing. A combination of surgical debridement and medical interventions may be required (3).

Figure 1 Preoperative view: at time 0 days.

Figure 2 Dermacyn® application and Polyurethane Sterile Foam Dressing with negative pressure.

Figure 3 Preoperative view: depth 5 cm.

Discussion

The sacral pressure ulcer is a locally damaged area of the skin and subcutaneous tissue caused by pressure, traction, friction and/or a combination of these (working definition EPUAP). Contributions to the onset of pressure ulcer risk factors both local (thermal changes, moisture, maceration, dehydration), and systemic (age, immobility, malnutrition, chronic systemic diseases, acute diseases, vascular diseases, medical–surgical, anaemia, were toxic, hypoalbuminosis). A pressure ulcer is an area of skin that breaks down when constant pressure is placed against the skin and reduces blood supply to that area. After an extended amount of time with decreased tissue perfusion, ischaemia occurs and can lead to tissue necrosis and in the worst case it can reach muscles and bones. Factors that can contribute to the formation of pressure ulcers could be intrinsic (systemic disease, medication, malnutrition, age, dehydration, lack of mobility, incontinence, skin condition, weight) and extrinsic (external influences which cause skin distortion, pressure, shearing forces, friction, moisture) (4,5).

Pressure ulcers can form anywhere on the body. The most frequent formations are in sacral area (30–40%),
heel (15–20%), ischiatic region (5–15%), great trochanter (10–15%), malleolar region (6–7%).

A pressure ulcer is localised injury to the skin and/or underlying tissue usually over a bony prominence, as a result of pressure, or pressure in combination with shear and/or friction. A number of contributing or confounding factors are also associated with pressure ulcers; the significance of these factors is yet to be elucidated (NPUAP, 2007). The ulcers are classified in four stages, starting from a normal erythaema which is characterised by an intact skin with non-bleachable redness of a localised area usually over a bony prominence (stage I), to the worst case with a full thickness tissue loss with exposed bone, tendon or muscle. Eschar may be present on some parts of the wound bed (stage IV).

The European guidelines for pressure ulcer wounds treatment are based on the ‘classes of risk’ include: control of incontinency and moisture, education and training of the patient and family, evaluation of the risk, anti-decubitus system, adequate diet and food tips, personal hygiene, distribution of weight, periwound skin care and control of pain (6–8).

**Device with negative pressure**

The VAC technique is a safe, easy and effective means in chronic wound care management (9), it is a sub-atmospheric pressure device consists of a polyurethane foam dressing that can be either open-cell honeycomb drainage (green) or a honeycomb structure with cells mixed open and closed (white) draining and thermal insulation (10,11). The polyurethane foam, shaped and inserted into the cavity, subjected to vacuum, allows for debriding tissue and stimulates the granulation tissue. Subsequently, the wound is covered with a hydrocolloid adhesive provided with central opening to allow the attachment of the device, which generates the negative pressure, and the subsequent drainage of the fluids of secretion (12,13). The secondary hydrocolloid dressing protects the surrounding skin and is permeable by gas and water vapour, while maintaining a moist environment ideal for the movement of fibroblasts and macrophages from the periphery to the centre of the lesion, with the formation of points where the cells that participate in neo-angiogenesis are deposited to stimulate the new granulation tissue. The dressing is connected to a suction cup that allows you to adjust the negative pressure and allows using it in a continuous or intermittent (14).

**Dermacyn wound care**

Dermacyn® Wound Care is a super-oxidised solution used for debridement, irrigation and moistening of acute and chronic wounds, ulcers, cuts, abrasions and burns. Through reducing the microbial load and assisting in creating a moist environment, it enables the body to perform its own healing process.

This solution is the result of superoxide electrolysis of pure water and sodium chloride (15).

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**Table 1 Results**

<table>
<thead>
<tr>
<th>Date</th>
<th>NPWT + Dermacyn</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exudate</td>
<td>&lt; post 48 hours, disappearance in five applications</td>
</tr>
<tr>
<td>Odour</td>
<td>&lt; post 1 applications</td>
</tr>
<tr>
<td>Infection</td>
<td>&lt; post 3 applications</td>
</tr>
<tr>
<td>Ulcer diameter</td>
<td>&lt;1 cm² for every applications</td>
</tr>
<tr>
<td>Ulcer depth</td>
<td>&lt;post 3 applications</td>
</tr>
<tr>
<td>Granulation tissue growth</td>
<td>Good</td>
</tr>
<tr>
<td>Healing time</td>
<td>&lt;30% compared to others protocols</td>
</tr>
<tr>
<td>Colour</td>
<td>Colour change from green to yellow</td>
</tr>
</tbody>
</table>

In wounds treated with, Dermacyn® starts a process of phagocytosis, which destroys pathogenic microorganisms. Dermacyn® surrounds the bacterium and attacks the cell wall, proteins contained in it are denatured by reactive oxygen species that causes cellular lysis. Come result of osmosis the cell wall breaks and are disintegrated microorganisms are destroyed.

Dermacyn® is immediately ready for use without any dilution, the product must be applied directly on the area to be treated by dipping, spraying spray irrigation pressure or through sterile gauze (16).

The treatment consists in the dip up to 15 minutes daily during the initial phase, in the opinion of the physician responsible, wetting and/or spray at each dressing change according to the specific needs of the individual case. This solution and safe because it does not damage the tissues, not toxic, does not irritate the skin, ready for use without dilution or mixing, is stable and valid for a period of 1 year, neutral pH between 6-2 and 7-8, colourless so it masks the necrotic tissue (17).

**Conclusion**

To evaluate the efficacy of the combination Dermacyn® and NPWT, we assessed the following parameters: reduction of exudates with a change in colour and reduction of odour, infection and the size of pressure ulcer (Table 1). The evaluation of improvements of the wound, in terms of size, was evaluated at a time of 50 days (Table 2). The dressings were performed every 5 days.

To conclude, we noticed an immediate reduction of exudate with decreased maceration of surrounding skin (Figures 4–5). The bacteriostatic action of NPWT has been enhanced by the addition of Dermacyn®. It enabled an immediate (as early as the first application) dispersal of the odour and exudates, change in colour, from green with yellow, also the dissipation of periwound maceration and hyperaemia to allow the growth of peri-lesional and bottom edges of the ulcer and the appearance of granulation tissue.

The association of the two has ensured a significant reduction in healing time by controlling the local bacterial infection until the complete elimination of regional debridement of the necrosis with formation of granulation tissue.
Table 2  Decrease in ulcer depth in relation to dressing changes performed every 5 days

<table>
<thead>
<tr>
<th>Wound depth</th>
<th>5 days</th>
<th>10 days</th>
<th>15 days</th>
<th>20 days</th>
<th>25 days</th>
<th>30 days</th>
<th>35 days</th>
<th>40 days</th>
<th>45 days</th>
<th>50 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 cm</td>
<td>4.8 cm</td>
<td>4.5 cm</td>
<td>4.2 cm</td>
<td>3.8 cm</td>
<td>3.3 cm</td>
<td>2.7 cm</td>
<td>1.9 cm</td>
<td>1.4 cm</td>
<td>0.9 cm</td>
<td>0.5 cm</td>
</tr>
</tbody>
</table>

Figure 4  Postoperative view: depth 0.5 cm.

Figure 5  Postoperative view: at time 50 days.

References


