Successful use of a topical mixture with ozolipoile in the treatment of actinic ulcers

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Purpose: Post-radiation skin damage is the result of alterations produced in the irradiated zone. There are different studies aimed at verifying the effectiveness of several kinds of molecules in the treatment of radiation-induced skin damage. The purpose of this study is to test a mixture with a formulation containing several natural active ingredients on actinic ulcers in patients receiving radiation therapies.

Patients and methods: The authors recruited 13 patients, and randomly divided them into a test group (T₁) and a control group (T₂). The patients in both groups were undergoing radiotherapy. The T₁ group was treated with a mixture called ozolipoile, whereas the T₂ group was administered hyaluronic acid gel followed by non-ablative laser therapy. We evaluated the obtained results, the time of clinical healing, the reduction of pain, and each side effect, comparing all data between the T₁ and T₂ groups.

Results: The average Visual Analog Scale results showed decrease in pain in both groups; however, while the T₁ group showed a significant decrease in the values, the T₂ group reported a more gradual reduction in the values, without ever reaching the minimum values obtained with the treatment with ozolipoile.

Conclusion: Treatment of actinic ulcers with ozolipoile mixture leads to faster control of pain and to better healing of small-size ulcers.

Keywords: actinic ulcers, radiation-induced ulcers, skin wound, ozonide, wound healing

Introduction

Chronic skin lesions show a low tendency to spontaneous healing.¹ Chronic injuries are particularly painful and debilitating for the patient, and they are often the manifestation of an underlying condition with a vascular etiopathogenesis that obstructs healing. These factors cause a slowing of the reconstruction processes of the matrix and of the reepithelialization, favoring, consequently, a process of necrosis.

Chronic injuries are represented by ischemic ulcers, diabetic ulcers, venous ulcers, and pressure ulcers that do not reepithelialize. Post-radiation skin damage is the result of alterations produced in the irradiated zone, although the major site of damage is the germinal layer of the epidermis and the early response is clearly evident in the capillary network of the dermis: the dilation of these capillaries and the release of histamine causes a characteristic skin erythema.² After this transient erythema, a latency period follows. Even this second phase is due to radiation-induced capillary dilation: the capillaries dilate to compensate the reduced oxygen supply due to the peripheral ischemia of the arterioles. The second phase of erythema is accompanied by the loss of the superficial layers of the epidermis: the skin appears similar to that observed in
first-degree thermal burns. The radiation-induced death of the germinative layer of the epidermis causes an extended desquamation.\textsuperscript{2,3}

There are different studies aimed at verifying the effectiveness of several kinds of molecules in the treatment of radiation-induced skin damages, and some of them have, in fact, produced good results by using biomolecules rich in oxygen,\textsuperscript{4,5} thanks to the hyperoxygenation of the tissues. These biomolecules promote the healing of skin wounds and of mucous membranes, regenerating them and forming a new epithelial layer.\textsuperscript{6}

In addition, they also act as antibacterial and antifungal agents due to their oxidizing action.\textsuperscript{7}

Other studies tested, instead, alpha-tocopheryl acetate, to prove its radical scavenging activity:\textsuperscript{8} this liposoluble vitamin blocks the lipid peroxidation of membranes; therefore, it contrasts the free radical species responsible for cell aging and for the damages from excessive exposure to solar radiation. A well-known category of molecules used in the treatment of skin damages is ozonides.\textsuperscript{9} They act on biochemical mediators and enzymes involved in the mechanism of inflammation by activating defense systems.

These different pharmacologically active molecules have been widely tested individually, but there is not much evidence of their synergistic action.

The purpose of this study is to test a mixture with a formulation containing all the active ingredients described above, together with other natural ingredients, to determine whether they, in a single solution, are able to add up their beneficial effects on ulcerated skin tissues in patients receiving radiation therapies.

**Material and methods**

The authors recruited 13 patients, all female, aged from 62 years to 65 years, from private studies mainly located in south Italy. These patients were recruited randomly, were clearly informed regarding the study, and signed an informed consent. The study was conducted in compliance with the “ethical principles for medical research involving human subjects” of the Declaration of Helsinki. The study was conducted in accordance with Italian laws and regulations.

The enrolled patients suffered from second- or third-degree actinic ulcers following a radiotherapy cycle, according to the EORTC classification.\textsuperscript{10}

All the patients were suffering from head and neck oncological pathology.

These patients were randomly divided into a test group (T\textsubscript{1}) and a control group (T\textsubscript{2}). The T\textsubscript{1} group was treated with a mixture called ozolipoile: the treatment tested was a medicament mainly composed of ozone, oleic acid, \(\alpha\)-lipoic acid, vitamin E acetate, panthenol, tocopheryl acetate, and tocopherol. Ozolipoile is patented and already marketed; thus, no experimental studies were carried out on the patients, but we have assessed the efficacy of this treatment.

Instead, the T\textsubscript{2} group was administered hyaluronic acid gel followed by non-ablative laser therapy, as previously described.\textsuperscript{11}

At the enrollment phase, each patient received exhaustive information about personal hygiene and the mode of application of the mixture, along with a card to express numerically the level of pain the patient was suffering according to a Visual Analog Scale (VAS), ranging from 0 (no pain) to 10 (unremitting pain) (Figure 1). Both the protocols should be administered about three times a day, at least 2 hours before and after the radiotherapy, until complete healing. A clinical measurement was performed and archived before the start of the protocols (Table 1). Sterile protective dressings of ulcers were admitted. Clinical observations of the patients were per-

**Figure 1** VAS (Visual Analog Scale) for the monitoring of pain.

**Table 1** Baseline data related to the ulcerate in T\textsubscript{1} and T\textsubscript{2} groups

<table>
<thead>
<tr>
<th>Baseline data</th>
<th>T\textsubscript{1} group</th>
<th>T\textsubscript{2} group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean Area of the ulcer (cm)</td>
<td>3.8</td>
<td>4.1</td>
</tr>
<tr>
<td>Duration in days</td>
<td>5</td>
<td>7</td>
</tr>
</tbody>
</table>

**Notes:** T\textsubscript{1}, test group treated with a mixture called ozolipoile; T\textsubscript{2}, control group treated with a combined administration of hyaluronic acid gel followed by a non-ablative laser/radiofrequency/intense pulsed light therapy.
formed daily during radiotherapy and the data were collected at days 1, 3, 5, and 7 from the beginning. The follow-up was aimed at registering any changes in the perception of pain related to the eventual healing process of the ulcers. In order to not alter the collected data, each patient was monitored by the same physician, who scored the data without being aware of the difference between the treatments of the T₁ and T₂ groups. We evaluated the obtained results, the time of clinical healing, the reduction in pain, and each side effect, comparing all data between the T₁ and T₂ groups.

**Results**

The results carried out from VAS showed a significant difference in the results of the treatments of the T₁ and T₂ groups (Table 2). The first day of treatment achieved best scores in the T₂ group: despite the pain increasing immediately after the application of the laser beam, because of the vascular effect depending on the local heat, at the end of treatment, the patients of this group referred reduced pain in the treated area. However, by the third day of treatment, the VAS scores gave an average of 5.3 in T₁, instead of 6.5 in T₂.

The average VAS results showed a decrease in pain in both groups; however, while the T₁ group showed a significant decrease in the values between day 1 and day 7, the T₂ group reported a more gradual reduction in the values, without ever reaching the minimum values obtained with the treatment with ozolipoile.

In all reported cases, scars of different degrees appeared at the end of radiotherapy, even if only 7 days from the beginning of T₁ treatment, the ulcers appeared less deep and reduced in size (Figure 2). The small-size ulcers were fully healed within 3–5 days; in those cases esthetically more compromised, plastic surgery has been performed with minimally invasive techniques.12

**Discussion**

In second-degree radiodermatitis, erythema usually occurs very early and is accompanied by intense edema and by the formation of blisters and bubbles. On the fifteenth day after the irradiation, it appeared as a richly vascularized area, and this form was associated with intense pain; repairing of the injuries occurred in a few months, but usually functional alterations of the skin, telangiectasia, and skin xeroderma remained. Ulcerative radiodermatitis usually shows no signs of repair, and in the most favorable cases, the healing is slow, with frequent recurrences.1–3

The treatment of radiation-induced ulcers is initially aimed at the removal of necrotic tissue, the elimination of bacteria, and the drainage of infected lesions.13,14 Reconstructive surgery could be indicated in areas deeply atrophic, following the radiotherapy.15,16

Local treatment with detergent and cleaning compounds is based on the use of alkaline solutions: it is advisable to perform all those measures addressed to obtain a gradual cleansing and the development of granulation tissue. Our treatment has shown very good performance not only on small-size ulcers but also on those wider and on the surrounding tissues: the combination of active substances against inflammation, the oxidative killing by the ozonides, the massive action against free radicals carried out by antioxidants, and the barrier effect

**Table 2** Synoptical table comparing the T₁ and T₂ VAS scores

<table>
<thead>
<tr>
<th>VAS scores</th>
<th>Baseline</th>
<th>Day 1</th>
<th>Day 3</th>
<th>Day 5</th>
<th>Day 7</th>
<th>D1–D7 variation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Group T₁</strong></td>
<td>Median</td>
<td>9.0±0.5</td>
<td>7±1.5</td>
<td>5±0.5</td>
<td>2.5±0.5</td>
<td>1±1</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>8.3</td>
<td>5.3</td>
<td>2.5</td>
<td>0.5</td>
<td>7.7</td>
</tr>
<tr>
<td><strong>Group T₂</strong></td>
<td>Median</td>
<td>9.5±0.5</td>
<td>7±1.1</td>
<td>6.5±1.5</td>
<td>3.5±1.5</td>
<td>2±1</td>
</tr>
<tr>
<td></td>
<td>Mean</td>
<td>7.7</td>
<td>6.5</td>
<td>3.5</td>
<td>2.5</td>
<td>5.2</td>
</tr>
</tbody>
</table>

Notes: T₁, test group treated with a mixture called ozolipoile; T₂, control group treated with a combined administration of hyaluronic acid gel followed by a non-ablative laser/rF/IPL therapy.

Abbreviations: VAS, Visual Analog Scale; rF, radiofrequency; IPL, intense pulsed light.

![Figure 2 Actinic lesion with a wide erythematous area: a case–control study. The same case–control study 7 days after beginning of T₁ treatment.](image-url)
by alpha-lipoic acid have created the best conditions for this impressive outcome in our first pilot study.

Conclusion
In conclusion, the treatments of actinic ulcers based on ozonide mixture show important advantages in controlling pain and healing time. The same procedure could be extended to chronic skin ulcers of different etiologies, given that the skin damage is similar, and in the most favorable cases, this treatment may even result more effective than in actinic ulcers.

Disclosure
The authors report no conflicts of interest in this work.

References