**Name of Journal:** *World Journal of Clinical Cases*

**Manuscript NO:** 88785

**Manuscript Type:** ORIGINAL ARTICLE

***Retrospective Study***

**Efficacy of surgical resection and ultra-reduced tension suture combined with superficial radiation in keloid treatment**

Hu XY *et al.* Efficacy studies of keloid lesions

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**Supported by** Qingdao Medical Research Guidance Program Project, No. 2020-WJZD110.

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**Received:** November 10, 2023

**Revised:** December 1, 2023

**Accepted:** December 6, 2023

**Published online:** December 16, 2023

**Abstract**

BACKGROUND

There are many available treatment options for keloid; however, single treatments are usually less effective. Therefore, more scientifically rational and effective combined treatment methods should be sought to solve the pain associated with keloids.

AIM

To explore the efficacy and safety of surgical resection and ultra-reduced tension suture combined with superficial radiation as keloid treatment.

METHODS

Fifteen keloid patients admitted to Qingdao Eighth People's Hospital from June 2020 to January 2022 were enrolled in this retrospective analysis. All patients underwent a comprehensive treatment approach comprising surgical resection, ultra-reduced tension suture incision, and superficial radiation therapy within 24 h postoperatively. The modified Vancouver Scar Scale (mVSS) and Patient and Observer Scar Assessment Scale (POSAS) were used to evaluate the treatment effect, whereas the efficacy, adverse effects, and recurrence rate were observed according to the 12-mo follow-up after treatment.

RESULTS

The mVSS and POSAS scores at 1 and 6 mo after combination treatment decreased compared to before treatment (*P* < 0.001), and the overall response rate was 93.3%. Only one case recurred, yielding a 6.7% recurrence rate. The incidence of local chromour sedimentation rate in 1–3 mo after radiotherapy was 33.3% (5 patients), all subsiding after 6–9 mo, without complications, such as delayed wound healing or dermatitis.

CONCLUSION

Surgical resection, super subtraction sutures, and superficial radiotherapy are treatment methods with short courses, low recurrence rates, and good safety profiles.

**Key Words:** Keloid; Ultrasound suture; Radiotherapeutics; Superficial radiation; Ultra-reduced tension suture; Radiation

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**Citation:** Hu XY, Yang Q, Guan XY, Li JY, Wang LL, Li K, Zhang XT. Efficacy of surgical resection and ultra-reduced tension suture combined with superficial radiation in keloid treatment. *World J Clin Cases* 2023; 11(35): 8310-8319

**URL:** <https://www.wjgnet.com/2307-8960/full/v11/i35/8310.htm>

**DOI:** https://dx.doi.org/10.12998/wjcc.v11.i35.8310

**Core Tip:** This study utilizes the most advanced ultra-reduced tension suture technology combined with superficial radiation therapy, known as "superficial radiation therapy," to reduce the recurrence rate of keloids. Only a few reports on this combination therapy method have been reported in China. Based on the pre-and post-treatment follow-up observations, comparison of photographs, and analysis of quantitative indicators in the scar evaluation table, this combination therapy can achieve a short treatment course, low recurrence rate, and good safety profile.

**INTRODUCTION**

Keloid is a special type of pathological scarring that regularly occurs in some body parts of normal people, such as the earlobe and auricle, double mandible, anterior chest, deltoid area of the upper arm, shoulder, upper back, pubic hair, and around the limb joints, in addition to other areas of patients predisposed to scarring. Although the etiology of this condition is complex, with various available treatments, including freezing, laser, injection, surgery, and radiotherapy, use of only a single treatment is associated with a higher recurrence rate[1]. Therefore, more scientific and effective combined treatment methods should be sought to treat pain in patients with keloid. This study investigated plastic professional technology and radiotherapy-related equipment, adopting cutting-edge super tension suture technology to organically combine surgical treatment and radiotherapy, and develop a feasible comprehensive treatment plan based on the severity and evaluation of common keloids. Additionally, the efficacy of the comprehensive treatment method on keloid was analyzed according to the pre-and post-treatment follow-up observation. Therefore, the optimal treatment plan suitable for most clinical keloid cases was explored to improve clinical efficacy and patient satisfaction, and it is expected to be replicated and promoted in other hospitals. In this study, the most advanced tension suture technique combined with superficial radiotherapy (hereinafter referred to as "superficial release therapy") was applied to reduce the recurrence rate of keloids. However, this combination treatment method has rarely been reported in China. Therefore, this study aimed to compare pre- and post-treatment follow-up observations, compare the photographs, analyze the scar evaluation table, and assess the duration, recurrence rate, and safety of the combination therapy.

**MATERIALS AND METHODS**

***General information***

In total, 15 patients with keloid who were admitted to Qingdao Eighth People's Hospital from June 2020 to January 2022 were included in this retrospective analysis. The inclusion criteria were as follows: (1) Patients fulfilling the diagnostic criteria for keloids[2]; (2) Less than 3 scars; (3) Scar width (vertical to the skin tension line) < 3 cm; (4) Disease course of < 5 years; (5) Secondary lesions; (6) Normal mental state; and (7) Examination and treatment and follow-up. The exclusion criteria were: (1) Age < 16 years; (2) Pregnant or lactating women; (3) History of severe heart, brain, and kidney systemic diseases; (4) Abnormal coagulation function; (5) History of treatment within 6 mo; (6) History of cutaneous malignancy; and (7) History of malnutrition and villainous disease.

Of the 15 patients aged 16–80 years, 3 were male and 12 female, respectively, with a mean age of 38.3 ± 21.7 years. The disease duration ranged from 1 to 40 years. Among seven cases of ear keloids (six and one cases were caused by ear perforation and perineural surgery, respectively), two were bilateral lesions. In two cases of facial keloids, one was acne-induced bilateral lesions, and the other was nevus-cutting surgery-induced. Additionally, in two cases of anterior thoracic keloid, one was acne-induced (frequent lesions), and the other was surgically induced (single lesion). Keloid scarring of limbs caused by local vaccination or surgery occurred in 4 cases. Furthermore, the keloid sizes were 0.6 cm × 0.6 cm × 0.4–9 cm × 4 cm × 3 cm, and scars had a tough texture, obvious bulge, and clear boundary, accompanied by different degrees of pain, itching, and other discomfort symptoms.

***Surgical resection with ultra-reduced tension suture incision***

Surgery was performed under local infiltration anesthesia. In brief, the surgical area was disinfected three times with 0.5% iodophor, and a 1:200000 adrenaline lidocaine solution was locally injected around the keloid. After satisfactory anesthesia, the incision design and suture method were decided based on the keloid’s location and base condition.

Regarding extensive keloid base or a small amount of surrounding tissue, which cannot be directly stitched or affect symmetry after resection, the core, and most scar bumps were stripped from the skin and the edge was maintained at a width of 3–10 mm and the scar tissue flap at a thickness of 2 mm. Next, the most suitable type of slow absorption suture (5-0 to 6-0) was selected according to the scar’s location to perform subcutaneous tension suture 2 mm away from the skin margin, and the incision margin was slightly raised to ensure no tension. The skin was intermittently closed using a 6-0 monofiloral nylon thread, and the drainage strips were placed, as appropriate. Finally, erythromycin eye ointment is externally applied, followed by bandaging.

In contrast, for cases with a narrow keloid base or a large amount of surrounding tissue, direct stitching could be performed after evaluation, and the keloid edge could be completely removed. After the edge of the wound is fully detached, the ultra-reduced tension suture can be performed. Due to the large amount of surrounding tissue, the bulge of the incision margin becomes more obvious than that of the scar flap. The suture details included multiple symmetrical super-tension suture skin points designed at 1 cm from both sides of the incision, with a 1 cm ipsilateral spacing, and the outer skin points 0.5 cm above the horizontal edge of the wound. The adjacent puncture points on both sides of the incision were in one group, while the two adjacent groups were pairs. Subsequently, the appropriate type of slow absorption sutures (4-0 to 5-0) was selected according to the scar’s location, penetrating from one end to point A, passing through the epidermis, dermis, and subcutaneous tissue to the adjacent point B; the suture entered from point B, advanced through point C from the subcutaneous tissue to adjacent point D, and finally enters from point D to the incision site (Figure 1). When tying with the tail line, the skin tissue on both sides should be appropriately pushed to the middle, and the skin surface should be depressed at the puncture point. Additionally, the skin around the depression bulged, enabling the incision to be closely aligned, while the line knot becomes buried under the skin. The suture proceeded successively from one end of the incision to the other end. Subsequently, an intermittent subcutaneous suture was continually applied on the skin margin to smooth the wound. Finally, the skin was closed intermittently with a 6-0 single femoral nylon thread, and the drainage strips were placed as appropriate. External erythromycin eye ointment sterile dressings were applied on both sides of the wound to protect the raised incision, followed by bandaging.

***Radiotherapeutics***

The patient received the first superficial placement treatment within 24 h of surgery. The dressing was opened during treatment, and the wound was disinfected. Subsequent treatment depended on the drainage situation; those with drainage strips were pulled out or retained until the second postoperative day. Mold shielding protection was required outside the radiation area, and postoperative continuous radiotherapy was performed five times. The opening times were 5–7, 8–10, and 12–14 d for the face and ear, trunk, and limbs, respectively.

***Wound care***

Daily care measures for scars in the surgical area, such as sun protection, avoiding spicy food, and no drinking, were implemented after the wound removal for 3–6 mo. Other anti-scar treatments, including silicone preparation and pressure, were not used.

***Observational indicators***

**The observational indicators included the following:** Scar score: Scars were evaluated preoperatively, 1 mo postoperatively, and 6 mo postoperatively using the following: (1) Modified Vancouver Assessment Scale (mVSS): This was achieved by palpating patients’ scar and assessing the severity of the scar using six color aspects, scar thickness, vascular distribution, softness, pain, and itching. The highest and lowest scores were 18 and 0, respectively. Higher and lower scores indicate heavier and lighter scars, respectively; and (2) The Patient and Observer Scar Assessment Scale was measured through self-assessment by the patient or their family members and observation by the attending physician. The Patient Scale (PSAS) (including pain level, pruritus, color, hardness, thickness, and irregularity) and Observer Scale (OSAS) (such as vascular distribution, color, thickness, roughness, softness, and surface area) were assigned separately, and an overall evaluation was determined. The maximum and lowest scores were 10 and 1 point, respectively; higher and lower scores indicated heavier and lighter scars, respectively.

Efficacy assessment: Based on contemporary scarring and following Liu Wenge’s standard for efficacy, this assessment was categorized according to the degree of improvement of symptoms and signs into cure, effective, and ineffective. (1) Cure: This assessment comprises disappeared itching pain and other symptoms; scar softening, not higher than the skin; continuous observation; and no recurrence after 6–12 mo; (2) Effective: This is characterized by significant improvement in itching pain and other symptoms, smaller size, or the severity changed to moderate, moderate to mild, and no reversal achieved 6–12 mo after treatment; (3) Ineffective: Slight or no improvement in itching pain and other symptoms, increased scar texture and volume, or once cured and effective post-treatment, but relapse within 6–12 mo after treatment completion. Effective refers to the sum of cured and effective blocks. The total response rate was calculated using the following formula: (cure + significant)/100%.

Adverse reactions and recurrence rate: Data were collected at follow-up visits in 1, 3, 6, and 12 mo after the combination treatment, and the adverse reactions and recurrence rates were counted.

***Statistical analysis***

All data analyses were performed using the IBM SPSS Statistics for Windows, version 26 (IBM Corp., Armonk, NY, United States). Measuring data with mean ± SD, *t*-test; statistical significance was considered at *P* < 0.05.

**RESULTS**

***Comparison of the mVSS scores at 1 and 6 mo after 19 keloids were observed in 15 patients***

Comparison of before combination therapy and 1 and 6 mo after the combination treatment (*P* < 0.001); Comparison of 1 and 6 mo after combination therapy (*P* < 0.001) (Table 1).

***Comparison of the OSAS scores in the 15 patients***

All 15 patients had significantly lower OSAS scores after the combination therapy than pre-treatment (*P* < 0.001) (Tables 2 and 3). The difference in scores was statistically significant (*P* < 0.01) (Table 4).

***The degree of pain, pruritus, color, hardness, thickness, irregularities, and the overall PSAS scores of the scar compared before and in 1 and 6 mo after the combination treatment***

The PSAS scores of the 15 patients were significantly lower after the combination therapy than before combination treatment (*P* < 0.05 and *P* < 0.01, Tables 5 and 6). The hardness and irregularity degree score decreased in 1 and 6 mo after the combination treatment, which were statistically significant (*P* < 0.05),the other scores were not significant (*P* > 0.05), but the overall evaluation was significant (*P* < 0.05), indicating that the effect 6 mo after the combination treatment was superior to that 1 mo post-treatment (Table 7).

***Curative effect***

The surgical incision of 19 keloids in 15 patients healed initially without complications, such as hemorrhage, hematoma, infection, or delayed healing. The uplift area was generally completely flat 25–40 d after surgery. At 1 and 6 mo after treatment, scar vascular distribution, color, thickness, roughness, softness, and surface area all significantly improved according to the clinical observation evaluation; furthermore, the subjective feeling of itching or pain also decreased and the scar area texture became more soft. Further, there was no swelling hyperplasia or linear mature scars, and patients were satisfied with the appearance. The total effective rate was 94.74%.

***Adverse effects and the recurrence rate***

Overall, 5 of 15 patients experienced hyperpigmentation in the wound area during radiotherapy; however, this hyperpigmentation was resolved 6–9 mo after the end of the radiotherapy. In one case, sagging and pulling caused by the large breast volume surrounding the wound resulted in the recurrence of a new scar postoperatively, which the injection treatment improved. The remaining case had no recurrence.

***Typical case 1***

Case 1 was a female patient aged > 30 years who presented with a keloid on her left earlobe 3 years prior. This left eardrop keloid caused the formation of an ear hole, which gradually increased, with obvious itching that affected the appearance and function. The patient had a keloid before the left earlobe, dark red ellipsoid, 2 cm × 1.5 cm × 1 cm, hard keloid texture, obvious itching, and no local rupture (Figure 2A). The treatment plan included surgical resection, hypercompression suture, and superficial radiation therapy. The patient underwent keloid resection under local anesthesia with the scar valve. Subsequently, 6 Mev-β electron line was administered 16 h postoperatively, and the target area was expanded 1.5 cm outside the surgical incision (Figure 2B), with a single dose of 3 Gy administered over 5 consecutive days, for a cumulative dose of 15 Gy. The incision had healed well by day 7 postoperatively (Figure 2C). Furthermore, the postoperative follow-up at 12 mo revealed that the earlobe had a satisfactory appearance without recurrence or other complications (Figure 2D).

***Typical case 2***

Case 2 was a 70-year-old female patient who underwent total right breast resection and systemic chemotherapy for right breast cancer 7 years prior. Six months postoperatively, scar hyperplasia appeared at the medial half of the incision and gradually widened with pain. The patient sought treatment due to the concern about cancer recurrence in the scar area. During the physical examination, an oblique surgical scar was observed on the right side of the chest wall, approximately 21 cm long; the medial half was approximately 7 cm bulge and dark red; the widest part was approximately 1.4 cm; irregular edge, hard texture; a certain range of motion; obvious pain and itching; and no local rupture was observed (Figure 3A). The established treatment regimen comprising surgical resection plus radiotherapy was planned. The patient underwent scar resection under local anesthesia, and the bilateral tissue extension was legal (Figure 3B and C). Both intraoperative freezing and postoperative pathology suggested scar tissue. β-ray at 6 Mev was initiated 15 h postoperatively, and the target area was expanded 1.5 cm outside the surgical incision (Figure 3D) at a fractionated regimen of 3 Gy for 5 consecutive for a cumulative dose of 15 Gy was administered. The incision healed well 9 d postoperatively, but the color was obvious (Figure 3E). After 40 d, the area was flat, the local color subsidence subsided (Figure 3F), and the color subsidence had completely subsided by 6 mo. The scar was soft and flat without recurrence, with a satisfactory appearance at 12 mo postoperatively (Figure 3G).

**DISCUSSION**

Keloid scarring manifests as a colored mass raised above the normal skin surface. Keloid scars extend beyond the original damage range and exhibit persistent growth, with a hard texture and poor elasticity, frequently accompanied by itching or pain. Keloids are characterized by their resistance to therapy and high recurrence rate after treatment. The keloid formation mechanism has been shown to involve capillary angiogenesis[3], fibroblasts, myofibroblasts, various cytokines, and tissue collagen[4]; as such, a single treatment method cannot achieve significant and lasting outcomes. Previous clinical practice has shown that skin regions with higher tension, such as the chest and back of the shoulder are more prone to developing multiple keloids. When these skin regions experience insults, such as skin disease or wound, the greater tension stimulates collagen hyperplasia, and the consequences of collagen decomposition and synthesis eventually cause the formation of a keloid[5]. Keloids are conventionally treated with surgical resection; however, this procedure can impact the skin and cause tissue defects, thereby increasing the incision tension and leading to postoperative recurrence. Years of clinical observation and professional research have shown that the choice of operation method and process is one of the important factors in preventing keloid recurrence, and a 1:3 reduction of tension in the operation is key[6] to surgical success[6].

In addition to meeting the requirements of wound malocclusion and valgus, ensuring a tension reduction effect is crucial. When the incision tension is large, the inner suture can easily cause local poor blood transportation and pigmentation, and the shallowly buried line head becomes exposed and at risk, sometimes even resulting in scar widening or hyperplasia in the later stage. The improved vertical mattress seam buries the suture knot to a deeper level, resulting in better incision alignment based on providing better tension reduction ability[7,8]. The built-in horizontal mattress suture technique, commonly used in cesarean section, greatly extends the indwelling time of the non-absorbable line to enable a more effective and persistent reduction of the incision tension and improve the incision’s long-term healing[9]. The deep-buried annular mattress suture technique (LBD suture technique) and Zhang's overextension-reducing suture technique can withstand greater tension, resulting in a better healing effect[10,11]. Regardless of the method adopted, the objective is to fully mobilize the tissue surrounding the incision, which seals the deep dermis and superficial fascia layer and maintains the incision without tension to the maximum extent.

In this study, the 15 patients had a medical history of > 1 year. When selecting surgical methods, decisions are based on the location and size of the keloid. This study combines the methods of LBD suture, which uses more non-absorbable thread in the internal suture, and Zhang's super-tension suture, which employs more absorbable barbed thread. Additionally, we investigated the influence of a non-absorbable line on keloid recurrence over time; considering the keloid location, the cost of the barbed line, and other factors, the inner suture was replaced with a slow absorption suture. Simultaneously, the personalized adjustment was made according to the characteristics of the keloid area to achieve the tension state of the incision of scar repair surgery. The authors’ experience involves preserving the aesthetics of the original lesion based on maximizing the resection region.

Using a scar valve simplifies the operation, making surgery easier, resulting in minimal alterations to the appearance and easy acceptance of this approach by the patients[12]. However, the keloid recurrence rate can reach as high as 45%–100%[12]. The retention of the scar flap is also one of the reasons for recurrence; therefore, under the condition of maintaining the ear appearance, the tension of the cutting margin should be significantly reduced to avoid the recurrence of keloids caused by large tension. Regarding the trunk, limbs, and other areas with a large amount of surrounding tissue, complete removal of the scar is achievable with direct suture, provided the edge of the wound remains fully free under the skin, enabling the tensioned suture to be performed.

Radiotherapy is an independent treatment modality for keloids and is commonly used as an adjunct to surgical treatment. Radiotherapy is used to eliminate fibroblasts and vascular endothelial cells in keloids, inhibit the production of collagen and the growth of naive fibroblasts, and block vascular regeneration, thereby directly and indirectly inhibiting keloid growth[13]. However, damage to the skin is a common side effect of radiotherapy which produces skin redness, itching, and even short-term pain. In the long term, radiotherapy can also cause excessive keratosis, skin tissue atrophy, abnormal skin pigment metabolism (pigmentation or depigmentation), and telangiectasia. Generally, low-energy electron rays (β-ray, 6–7 MeV) or X-rays (soft X-ray) are used as the radiotherapy modality, and two prophylactic radiotherapies can be performed within 24 h (and no more than 48 h) after keloid surgery, at a dose of 500–700 cGy/fraction.

In this study, radiotherapy was indicated for scars where the thickness of the lesion area was < 5 mm. All areas that underwent surgical treatment were considered eligible for radiotherapy. Additionally, the β-rays used can penetrate the skin surface and reach the deeper layer of the scar to inhibit its growth. Because the highest dose point of the electronic line is 15 mm below the skin, 80% of the depth dose exhibits rapid decay, and only 8.9% remains below the subcutaneous 30 mm, thereby providing good protection for the deep tissues and organs[14]. A special mold is available to protect the surrounding tissues outside the operation area during radiation to prevent damage to tissues outside the operation area. In our study, some of the patients experienced transient pigmentation in the irradiation area, which subsided from 6 to 12 mo postoperatively.

However, this study had some limitations which should be mentioned. First, the sample size was small due to the intention and choice of treatment and the long duration required for observing the treatment efficacy. Second, it was a single-center study. Therefore, in future studies, as this treatment plan evolves, more cases should be included in the observation, and the patients who prefer surgery and other methods, including injection treatment, should be used as control groups to seek a more scientific and effective comprehensive treatment plan for patients with keloid.

**CONCLUSION**

This study shows that the effect of surgical resection and ultra-reduced tension sutures combined with superficial radiotherapy is obvious; it has a short course, low recurrence rate, and good safety profile, and is easily accepted and preferred by patients, which is worthy of clinical promotion.

**ARTICLE HIGHLIGHTS**

***Research background***

Keloids are a type of pathological scar whose appearance increases the psychological burden to patients; however, in more severe cases, keloids can affect the patient’s body function, disrupting their ability to lead a normal life and even isolating them from society. When treating keloids, a unimodal treatment strategy cannot achieve significant and lasting effects; therefore, the identification of more effective treatment methods is of great significance.

***Research motivation***

Surgical treatment is the modality that achieves the most visible improvement; however, because of the high postoperative recurrence rate, ensuring the effect of the operation and limiting the recurrence rate is crucial. Therefore, this study is based on the well-established single treatment in the department, which can provide safe and effective comprehensive treatment, using cutting-edge super reduction suture technology and radiotherapy equipment, organic combination of surgical treatment, radiotherapy, surgical removal of keloid, super reduction suture combined with shallow radiotherapy, comprehensive treatment, according to the observation of comprehensive treatment of keloid, to elucidate.

***Research objectives***

To investigate the optimal scheme for adjuvant surgical treatment of keloids and improve clinical efficacy and patient satisfaction.

***Research methods***

The suture techniques employed in this investigation are the relatively new LBD suture and Zhang's suture. The ultra-reduced tension suture was placed once the subcutaneous surface of the wound edge was completely free. This study selected several radiotherapy strategies approved in the field to guarantee optimal results; however, all were performed within 24 h of surgery. The Scar Observation Scale of the patient and observer was completed concurrently during the same observation period, considering both patients' subjective experiences and professionals' objective indicators.

***Research results***

This study’s findings demonstrate that maximum resection, super tension suture, and superficial radiotherapy administered within 24 h of surgery can achieve a beneficial effect and ensure a low recurrence rate for keloids congruent with surgical justifications.

***Research conclusions***

Surgical treatment of keloid is the traditional therapeutic modality. This study uses a more cutting-edge tension suture technique. Surgical resection, super subtraction sutures, and superficial radiotherapy are treatment methods with short courses, low recurrence rates, and good safety profiles.

***Research perspectives***

Future studies should observe more cases as treatment options mature. Simultaneously, patients who select surgery and other methods, such as botulinum toxin, platelet-rich plasma, and nano-fat injections, could be selected as other observation groups to seek more scientific, effective, and diversified comprehensive treatment plans for patients with keloids.

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**Footnotes**

**Institutional review board statement:** This study was reviewed and approved by the Ethics Committee of Qingdao Eighth People’s Hospital (Approval no.: QBYLL-KY-2023-013).

**Informed consent statement:** This retrospective study used only anonymous patient data. The patient signed an informed consent form for treatment before surgery, and according to institutional policies, informed consent for this study was waived.

**Conflict-of-interest statement:** All the authors declare no conflicts of interest related to this article.

**Data sharing statement:** The datasets used and/or analyzed during the current study are available from the corresponding author upon reasonable request.

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**Provenance and peer review:** Unsolicited article; Externally peer reviewed.

**Peer-review model:** Single blind

**Peer-review started:** November 10, 2023

**First decision:** November 22, 2023

**Article in press:** December 6, 2023

**Specialty type:** Dermatology

**Country/Territory of origin:** China

**Peer-review report’s scientific quality classification**

Grade A (Excellent): 0

Grade B (Very good): 0

Grade C (Good): C

Grade D (Fair): 0

Grade E (Poor): 0

**P-Reviewer:** Romandini M, Italy **S-Editor:** Liu JH **L-Editor:** A **P-Editor:** Xu ZH

**Figure Legends**



**Figure 1 Schematic diagram of the overtension suture.** A-D: Ultra-reduced tension suture points.



**Figure 2 Example case 1.** A: Before treatment; B: First superficial radiotherapy after surgery; C: 7 d after surgery and superficial radiotherapy; D: 12 mo after treatment.



**Figure 3 Example case 2.** A: Preoperative; B and C: After scar resection and ultra-reduced tension suture; D: First postoperative radiotherapy; E: 9 d after removal; F: 40 d postoperative; G: 12 mo postoperative.

**Table 1 Comparison of the modified Vancouver Assessment Scale scores before, 1 and 6 mo after the combination treatment**

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Score** | **Example number** | **Before combination therapy** | **1 mo after the combination treatment**  | **6 mo after the combination treatment**  | ***t* value** | ***P* value** |
| mVSS | 19 | 13.21 ± 2.20 |  4.68±2.41 | / | 15.12 | < 0.001 |
| mVSS | 19 | 13.21 ± 2.20 | / | 0.89 ± 2.11 | 24.27 | < 0.001 |
| mVSS | 19 | / | 4.68 ± 2.41 |  0.89 ± 2.11 | 7.69 | < 0.001 |

mVSS: Modified Vancouver Assessment Scale.

**Table 2 Comparison of the Observer Scale scores before and 1 mo after the combination treatment**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **Before combination therapy** | **1 mo after the combination treatment** | ***t* value** | ***P* value** |
| Vascularity | 8.20 ± 1.15 | 2.67 ± 1.05  | 16.46 | < 0.001 |
| Color and luster | 7.87 ± 1.13 | 2.87 ± 1.60  | 10.92 | < 0.001 |
| Thickness | 7.73 ± 1.28 | 1.93 ± 0.88  | 19.59 | < 0.001 |
| Roughness | 7.13 ± 1.60 | 1.73 ± 0.80 | 13.92 | < 0.001 |
| Compliance | 7.87 ± 1.30 | 2.80 ± 1.08  | 13.65 | < 0.001 |
| Superficial area | 7.33 ± 1.50 | 2.20 ± 1.01  | 14.66 | < 0.001 |
| Overall evaluation | 7.53 ± 1.36 | 2.47 ± 0.83 | 17.84 | < 0.001 |

**Table 3 Comparison of the Observer Scale scores before and 6 mo after the combination treatment**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **Before combination therapy** | **6 mo after the combination treatment** | ***t* value** | ***P* value** |
| Vascularity | 8.20 ± 1.15 | 1.27 ± 1.03 | 17 | < 0.001 |
| Color and luster | 7.87 ± 1.13 | 1.40 ± 1.06 | 18.47 | < 0.001 |
| Thickness | 7.73 ± 1.28 | 1.47 ± 1.30 | 20.87 | < 0.001 |
| Roughness | 7.13 ± 1.60 | 1.40 ± 1.06 | 13.67 | < 0.001 |
| Compliance | 7.87 ± 1.30 | 1.47 ± 1.30 | 19.09 | < 0.001 |
| Superficial area | 7.33 ± 1.50 | 1.27 ± 1.03 | 17.61 | < 0.001 |
| Overall evaluation | 7.53 ± 1.36 | 1.40 ± 1.06 | 22.41 | < 0.001 |

**Table 4 Comparison of the Observer Scale scores**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **1 mo after the combination treatment** | **6 mo after the combination treatment** | ***t* value** | ***P* value** |
| Vascularity | 2.67 ± 1.05 | 1.27 ± 1.03 | 10.69 | < 0.001 |
| Color and luster | 2.87 ± 1.60 | 1.40 ± 1.06 | 5.05 | < 0.001  |
| Thickness | 1.93 ± 0.88 | 1.47 ± 1.30 | 1.97 | 0.068 |
| Roughness | 1.73 ± 0.80 | 1.40 ± 1.06 | 1.58 | 0.136 |
| Compliance | 2.80 ± 1.08 | 1.47 ± 1.30 | 4.39 | 0.001 |
| Superficial area | 2.20 ± 1.01 | 1.27 ± 1.03 | 3.5 | 0.004 |
| Overall evaluation | 2.47 ± 0.83 | 1.40 ± 1.06 | 4.68 | < 0.001 |

Data are expressed as the mean ± SD.

**Table 5 Comparison of the Patient Scale scores before and 1 mo after the combination treatment**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **Before combination therapy** | **1 mo after the combination treatment** | ***t* value** | ***P* value** |
| Degree of pain | 2.67 ± 2.09 | 1.27 ± 0.59  | 2.82 | 0.014 |
| Degree of itching | 6.27 ± 2.02 | 1.60 ± 0.74  | 9.26 | < 0.001 |
| Pigment | 7.87 ± 1.55 | 2.80 ± 1.66  | 10.95 | < 0.001 |
| Hardness | 8.20 ± 1.08 | 2.40 ± 0.83  | 17.02 | < 0.001 |
| Thickness | 7.73 ± 1.58 | 1.93 ± 1.16  | 12.08 | < 0.001 |
| Irregular degree | 7.33 ± 1.68 | 1.80 ± 1.15  | 14.23 | < 0.001 |
| Overall evaluation | 7.80 ± 1.21 | 2.27 ± 1.10  | 17.2 | < 0.001 |

Data are expressed as the mean ± SD.

**Table 6 Comparison of the Patient Scale score before and in 6 mo after the combination treatment**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **Before combination therapy** | **6 mo after the combination treatment** | ***t* value** | ***P* value** |
| Degree of pain | 2.67 ± 2.09 | 1.13 ± 0.52  | 2.98 | 0.01 |
| Degree of itching | 6.27 ± 2.02 | 1.33 ± 1.05  | 10.02 | < 0.001 |
| Pigment | 7.87 ± 1.55 | 1.73 ± 1.39  | 15.3 | < 0.001 |
| Hardness | 8.20 ± 1.08 | 1.47 ± 1.06  | 20.38 | < 0.001 |
| Thickness | 7.73 ± 1.58 | 1.47 ± 1.30  | 14.19 | < 0.001 |
| Irregular degree | 7.33 ± 1.68 | 1.20 ± 0.78  | 15.78 | < 0.001 |
| Overall evaluation | 7.80 ± 1.21 | 1.53 ± 1.06  | 19.85 | < 0.001 |

**Table 7 Comparison of the Patient Scale scores in 1 and 6** **mo after the combination therapy**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Item** | **1 mo after the combination treatment** | **6 mo after the combination treatment** | ***t* value** | ***P* value** |
| Degree of pain | 1.27 ± 0.59 | 1.13 ± 0.52 | 1.47 | 0.164 |
| Degree of itching | 1.60 ± 0.74 | 1.33 ± 1.05 | 1.17 | 0.262 |
| Pigment | 2.80 ± 1.66 | 1.73 ± 1.39 | 2.98 | 0.1 |
| Hardness | 2.40 ± 0.83 | 1.47 ± 1.06 | 2.71 | 0.017 |
| Thickness | 1.93 ± 1.16 | 1.47 ± 1.30 | 1.45 | 0.169 |
| Irregular degree | 1.80 ± 1.15 | 1.20 ± 0.78 | 2.36 | 0.033 |
| Overall evaluation | 2.27 ± 1.10 | 1.53 ± 1.06 | 2.96 | 0.01 |

Data are expressed as the mean ± SD.



Published by **Baishideng Publishing Group Inc**

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