Prevention of Postsurgical Scars: Comparison of Efficacy and Convenience between Silicone Gel Sheet and Topical Silicone Gel

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INTRODUCTION

Silicone-based products are widely used to limit pathologic scars. Using these products is cost- and time-effective, as well as more convenient and comfortable for patients. To date, a substantial number of studies have assessed the efficacy of these products in preventing scars, but have reached no definitive conclusions. However, in certain randomized, controlled studies, these products were significantly effective in reducing postoperative incision wound scarring (1-4). Currently, several types of silicone-based products are available for clinical use (5). Of these, silicone gel sheets and topical silicone gels are the most popular forms (5-7). However, both have known limitations and can be inconvenient for patients to use, thus posing a risk of misuse or treatment interruption. Specifically, silicone gel sheets are disadvantageous because they cannot be applied to mobile or visible areas of the body, and require additional taping or bandaging (3, 4). Moreover, the sheets cannot achieve and maintain adequate contact with scars when applied to the skin with an irregular contour (3). On the other hand, topical silicone gels take time to completely dry (4). Furthermore, patients have to take extra sunscreen precautions to prevent hyperpigmentation, and must also apply topical silicone gels to scars multiple times per day (4, 8). Few studies have compared the effectiveness of topical silicone gels to that of silicone gel sheets in surgical scar prevention. Consequently, we conducted this prospective study to examine both the efficacy and the convenience of each product.

MATERIALS AND METHODS

Patients and study design

This study enrolled 30 patients between January and November 2012. Participant inclusion criteria were as follows:

1) History of surgery 2 weeks to 3 months before enrollment
2) Scarring of a surface area < 10 × 10 cm²
3) Age 18 yr or older

Exclusion criteria were as follows:

1) Any infection
2) Any wound producing a significant amount of discharge, e.g., still requiring a dressing 2 weeks after surgery or lacking visible signs of normal epithelialization
3) Any systemic disease (e.g., diabetes mellitus, hematologic disorder, or dermatologic disorder)
4) Use of anticancer, psychiatric, or steroid medication
5) Psychiatric disorder
6) Deemed unable to complete the study according to the judgment of the authors.

To date, few studies have compared the effectiveness of topical silicone gels versus that of silicone gel sheets in preventing scars. In this prospective study, we compared the efficacy and the convenience of use of the 2 products. We enrolled 30 patients who had undergone a surgical procedure 2 weeks to 3 months before joining the study. These participants were randomly assigned to 2 treatment arms: one for treatment with a silicone gel sheet, and the other for treatment with a topical silicone gel. Vancouver Scar Scale (VSS) scores were obtained for all patients; in addition, participants completed scoring patient questionnaires 1 and 3 months after treatment onset. Our results reveal not only that no significant difference in efficacy exists between the 2 products but also that topical silicone gels are more convenient to use. While previous studies have advocated for silicone gel sheets as first-line therapies in postoperative scar management, we maintain that similar effects can be expected with topical silicone gel. The authors recommend that, when clinicians have a choice of silicone-based products for scar prevention, they should focus on each patient’s scar location, lifestyle, and willingness to undergo scar prevention treatment.

Keywords: Cicatriz; Postoperative Period; Wounds and Injuries; Prevention and Control
Evaluation criteria and outcome measures
All patients were randomly assigned to and equally distributed between 2 treatment arms (treatment with either a silicone gel sheet [Scarclinic™-Thin, Hans Biomed, Seoul, Korea] or a topical silicone gel [Kelo-Cote™, SejongMedix, Seoul, Korea]). The arms were categorized as Group I (n = 15) and II (n = 15), respectively. First, patients’ scars were photographed and evaluated using the Vancouver Scar Scale (VSS), for which their vascularity, pigmentation, pliability, height, pain, and itchiness were measured as outcomes (Table 1) (6, 9). The aforementioned evaluation was performed by 2 independent observers. We also recorded patients’ questionnaire responses about any scar-related pain, pruritus, color change, hardness, thickness, overall size, irregularity, and inconvenience of use they experienced, and scored them using a 10-point scale (Table 2). Then, we evaluated the VSS scores and patient questionnaire scores at 1 and 3 months and compared them with the baseline measurements.

Table 1. Vancouver Scar Scale

| Feature                  | Score |
|--------------------------|-------|
| Vascularity              | Normal| 0     |
|                          | Pink  | 1     |
|                          | Red   | 2     |
|                          | Purple| 3     |
| Pigmentation             | Normal| 0     |
|                          | Hypo-pigmentation| 1     |
|                          | Mixed-pigmentation| 2     |
|                          | Hyper-pigmentation| 3     |
| Pliability (Elasticity)  | Normal| 0     |
|                          | Supple (flexible with minimal resistance)| 1     |
|                          | Yielding (giving way to pressure) | 2     |
|                          | Firm (inflexible, not easily moved, resistant to manual pressure) | 3     |
|                          | Banding (rope-like tissue that blanches with extension of the scar) | 4     |
|                          | Contracture (permanent shortening of scar, producing deformity or distortion) | 5     |
| Height                   | Flat  | 0     |
|                          | < 2 mm| 1     |
|                          | 2-5 mm| 2     |
|                          | > 5 mm| 3     |
| Pain                     | None  | 0     |
|                          | Occasional | 1     |
|                          | Requires medication | 2     |
| Itchiness                | None  | 0     |
|                          | Occasional | 1     |
|                          | Requires medication | 2     |

Table 2. Patient questionnaire

| Question                                      | Yes | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | No |
|------------------------------------------------|-----|---|---|---|---|---|---|---|---|---|----|----|
| Do you have any pain on your scar?            |     |   |   | 1 | 2 | 3 |   |   |   |   |    |    |
| Is there any pururitus on your scar?          |     |   |   |   |   |   | 1 | 2 | 3 |   |    |    |
| Do you see any change of color on your scar?  |     |   |   |   |   |   | 1 | 2 | 3 |   |    |    |
| Do you feel your scar getting hardened?       |     |   |   |   |   |   |   |   |   |   |    |    |
| Does your scar get thicker or increase in size?|     |   |   |   |   |   |   | 1 | 2 | 3 |    |    |
| Does your scar change into irregular shape?    |     |   |   |   |   |   |   |   |   |   |    |    |
| Was it uncomfortable to use?                  |     |   |   |   |   |   |   |   |   |   |    |    |

Fig. 1. (A) Before and after views of silicone gel sheet use in scar management. The scar has improved in its vascularity, irregularity, and height after 3 months of treatment. (B) Before and after views of topical silicone gel use in scar management. The scar has improved in its pigmentation, irregularity, and height after 3 months of treatment.
Statistical analysis
Statistical analysis was performed using SPSS version 16.0 for Windows (SPSS, Inc., Chicago, IL, USA). All data were expressed as mean ± standard deviation (SD). We used the Mann-Whitney test to compare outcome measures between the two groups. A P value of < 0.05 was considered statistically significant.

Ethics statement
This study was approved by the institutional review board (IRB) of our medical institution (IRB approval number: HC11DSSI0091). All patients were informed of the study details (e.g., objectives, methods, predicted outcomes, adverse effects, and study devices), and submitted a signed informed consent document.

RESULTS

Baseline and clinical characteristics of the patients
We enrolled 30 patients (n = 30) in our study. Of these, 5 dropped out for personal reasons; therefore, 11 patients in Group I and 14 in Group II completed the study. These participants included 9 male and 16 female patients with a mean age of 37.52 yr (total range, 21-75 yr). When recording scar location, we observed 17, 7, and 1 patients with scars in the head and neck region, in the extremities, and in the trunk, respectively. All patients demonstrated tolerability for the treatment; none experienced specific clinical problems. By the conclusion of the study period, all patients’ scars had improved in terms of pigmentation, height, irregularity, and overall size (Fig. 1).

Changes from baseline in VSS scores at 1 and 3 months after treatment onset
At 1 month after treatment onset, we observed a degree of pain score change of -0.455 points in Group I and 0.143 points in Group II. This difference reached statistical significance (P = 0.033). However, with the exception of the pain score, as shown in Fig. 2, no significant differences were observed between either group in either their VSS scores by outcome measure or their total scores at 1 and 3 months from baseline.

Fig. 2. (A) Changes from the baseline in VSS scores at 1 month after treatment onset. (B) Changes from the baseline in VSS scores at 3 months after treatment onset. VSS, Vancouver Scar Scale.

Fig. 3. (A) Changes from the baseline in patient questionnaire response scores at 1 month after treatment onset. (B) Changes from the baseline in patient questionnaire response scores at 3 months after treatment onset.
Changes from baseline in patient questionnaire response scoring at 1 and 3 months after treatment onset
At 3 months after treatment onset, questionnaire response scores for inconvenience of use were 3.818 points in Group I and 1.571 points in Group II. This difference also reached statistical significance ($P = 0.015$). However, no significant differences between the groups were seen in any other outcome measures at 1 or 3 months from baseline (Fig. 3).

DISCUSSION
Several treatment and prevention methods for surgical scarring are available today. These include intraliesional corticosteroids; 5-fluorouracil; bleomycin; cryotherapy; silicone gel; pressure therapy; pulsed dye laser treatment; radiation; and surgical correction (10-12). Emerging scar-reducing therapies include the TGF-β superfamily; COX-2 inhibitors and nonsteroidal anti-inflammatory drugs; collagen synthesis inhibitors; angiotensin-converting enzyme inhibitors; minocycline; and gene therapy (11). Among these methods, silicone has typically been considered the standard noninvasive approach (13).

Since the early 1980s, silicone has been described as having potential effectiveness in treating pathological scars (14). Today, it is considered a conventional scar treatment approach (15-17). Indeed, a number of studies have assessed silicone’s efficacy in preventing scar formation during the postoperative period (1-4); it is frequently used after surgery because it is both non-invasive and causes few adverse effects (8, 16). However, a matter of controversy is whether silicone is truly effective in scar prevention (1, 18).

Various mechanisms have been proposed as possible modes of action for silicone materials. These hypotheses include increased temperature or oxygen tension, direct action of silicone oil, wound hydration, polarization of scar tissue caused by negative static charge and modulation of growth factors (7, 9, 10, 13). Silicone has been produced in several forms, including silicone cream compounds; silicone oil or gel, with additives such as Vitamin E; in combination with other dressing materials, and as custom-made silicone applications (18). Among these formats, silicone gel sheets are the most widely used; however, patients’ compliance with silicone gel sheet use is not always satisfactory. The silicone gel sheets are inconvenient for patients to apply to large mobile areas, such as the area near the joints, and are generally not appropriate to use on visible areas like the face (3). Taping or bandaging is required to secure the silicone gel sheet to the scar, which may cause irritation in patients with pliable skin, particularly the elderly and the young (4). Moreover, silicone gel sheets may also cause excessive sweating in hot and humid climates (4). They must be washed carefully to prevent infections and other complications (3). All of the above factors often lead to interruption of gel sheet treatment. Topical silicone gels, by contrast, are available in a tube, and can be applied in a thin layer on the skin. They form a flexible, gas-permeable and invisible film on the scar (4); they may also be spread on the scar and will dry up within a period of several minutes, allowing patients to then wear clothes or apply makeup. In addition, topical silicone gels are advantageous in that they require no fixation materials. At the same time, these gels do pose disadvantages: they must be used in combination with sunscreen to prevent hyperpigmentation and must also be applied multiple times each day (8). Patients must also be sure that gel applied to parts of the body covered by clothing have dried completely before they get dressed; failure to do so may result in friction that removes the silicone film too early (4). Finally, these gels cannot be applied to the periorcular or perioral area, particularly in younger patients.

Karagoz et al. (8) compared the efficacy of a topical silicone gel, a silicone gel sheet and a topical onion extract in treating postburn hypertrophic scars, and reported that the two silicone-based products were more effective than the onion extract. The authors also found no significant difference in the effectiveness of the two silicone-based products in treating hypertrophic scars. However, they did conclude that silicone gel sheets are the preferable method of treatment. This is not only because the degree of treatment response was relatively higher in the silicone gel sheet group but also because 1 patient in their study showed no response to the topical silicone gel. In addition, Mustoe et al. (15) also deemed silicone gel sheets worthy of consideration as a first-line of therapy for scar prevention.

It was the objective of our study to compare both efficacy and convenience of use in silicone gel sheets and topical silicone gels as surgical scar preventives. Between our two groups, we found no statistically significant differences in either VSS or questionnaire response scores at 1 or 3 months after treatment onset. The only significant difference observed was the degree of change in VSS pain scores at 1 month. While we assume the silicone gel sheet offers a greater degree of stability and protection in the early stages of scar formation, we could find no statistically significant difference in the degree of VSS pain score changes at 3 months after treatment onset. Meanwhile, the patient questionnaire inconvenience of use score was higher in our silicone gel sheet group at 3 months after treatment onset. This means that convenience of use proved greater in the topical silicone gel group. Our results prove that topical silicone gel is as effective as silicone gel sheet for preventing surgical scars and that it is also easier to use.

Our study did face several limitations. Although scar maturation can continue for up to a year after surgery, we followed scar progression for only 3 months. Previous studies that evaluated the efficacy of silicone gel in scar management also typically followed scar progression for a 2- to 4-month period (4, 19-21). We believe this is because many clinicians regard that the early
stage of scar remodeling is the time of greatest change. An additional limitation was that our study population was small and consisted solely of Korean patients. Finally, when using the silicone gel sheet, patients who also used tape or bandaging to secure the sheet in arbitrary fashion may have influenced the pigmentation of their scars.

In conclusion, numerous silicone-based products are used in modern management of postoperative scars. Clinicians should select products for each patient while considering both their efficacy and convenience of use. Scar management is an ongoing process over a course of at least 3 to 6 months. It is therefore essential to ensure patient compliance throughout the treatment process. Consequently, clinicians must also carefully weigh the characteristics, lifestyle, and compliance likelihood of each patient, as well as the location of their scars, in choosing the appropriate management modality. In our study, we have demonstrated that there is no significant difference in the degree of efficacy between either silicone-based product types; at the same time, our results indicate that topical silicone gels are more convenient for patients to use than silicone gel sheets. While previous studies have advocated for silicone gel sheets as first-line therapies in postoperative scar management, we maintain that similar effects can be expected with topical silicone gel. Further long-term, large-scale, prospective controlled studies are likely warranted to confirm our findings.

DISCLOSURE

There are no conflict of interest statements.

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