Effects of a New Wound Dressing Material SG-01 in an Experimental Rat Skin Burn and Decubitus Ulcer Model

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A wound dressing material, SG-01 was developed to treat various skin ulcers. In this study, to clarify the features of SG-01, we investigated the therapeutic effects of this dressing material in rat burn and decubitus ulcer models. In the rat burn model, SG-01 significantly reduced the percent wound area compared to gauze (control group), and decreased the sum of the percent area, suggesting that SG-01 promotes the healing of burns. Subsequently, we investigated the curative effects of SG-01 in the rat decubitus ulcer model using commercially available dressing materials, NUGEL® and DuoACTIVE CGF®. SG-01 significantly reduced the percent wound area compared to gauze (control group), decreased the sum of the percent area, and shortened the interval until healing. There were no significant differences between SG-01 and the above commercially available materials. These results suggest that SG-01 is useful for treating burn and decubitus ulcers.

Key words — wound dressing, hydrogel, rat, burn model, decubitus ulcer model

INTRODUCTION

SG-01 is a wound dressing material consisted by monolithic adhesive including water and ionic cross-linking hydrophilic polymers, cellulose derivative and polycrylate, with aluminum ion, spread onto the backing material.

As this agent has favorable physical properties such as high water absorption, water-maintaining ability, appropriate vapor permeability, adhesiveness, oxygen permeability and flexibility, it is useful for removing exudate, and was developed to treat burns and decubitus ulcer. Furthermore, SG-01 did not cause reactions in a primary skin irritation test, a cumulative skin irritation test, an intracutaneous reactivity test, a skin sensitization test, and a pyrogen test, suggesting its safety.

In this study, we examined the pharmacological effects of SG-01 in rat burn and decubitus ulcer models.

MATERIALS AND METHODS

Test Substance —- SG-01 (2.5 × 2.5 cm) was used as a test substance, and gauze (2.5 × 2.5 cm) containing sterile physiological saline was used as a control substance. In an experimental rat decubitus ulcer model, commercially available materials, NUGEL® (hydrogel A; JOHNSON & JOHNSON MEDICAL Inc., Tokyo, Japan) and DuoACTIVE CGF® (hydrocolloid B; Bristle Myers Squibb Co., Ltd., Tokyo, Japan) (2.5 × 2.5 cm), were also employed as commercially available control substance for comparison.

Animals —- The experimental protocol was approved by the Ethics Review Committee for Animal Experimentation of Nihon University. 7-week-old male Crj : CD(SD)IGS(SPF) rats were obtained from Japan Charles River Co., Ltd. (Kanagawa, Japan). Pelleted chow (CRF-1, Oriental Yeast Industry Co., Ltd., Tokyo, Japan) and tap water were given ad libitum. These animals were acclimated for 1 week under the following conditions: temperature, 22 ± 3°C; humidity, 50 ± 20%; lighting cycle, 12 hr/day; and ventilation cycle, 13 to 17 times/hr. The animals used were without abnormalities in the general condition.

Experimental Methods —- Preparation of Models:

Burn Model: In 8-week-old rats, hair on the back was removed using electric hair clippers and an electric shaver. Under ether anesthesia, a burning iron weighing 52 to 54 g (a modified 100 W soldering iron; 302 to 303°C; diameter, 10 mm; contact pressure, 63.7 to 76.4 g/cm²) was applied on an area on the posterior side 2-cm distant from the right axilla and on the right side 1.5-cm distant from the median line on the back, for 10 sec to induce grade
III burns (heat degeneration involving subcutaneous tissue).7) After 2 days, necrotic tissue (skin) was resected under ether anesthesia, and used as a burn model.8,9

**Decubitus Ulcer Model:** In 8-week-old rats, hair on the skin of the right greater trochanter was extensively removed using electric hair clippers and an electric shaver. For anesthesia, sodium pentobarbital at 50 mg/kg and sodium amobarbital at 30 mg/kg were intraperitoneally administered. On a wooden fixation plate, the animals were fixed in a prone position. As a cushion, a piece of absorbent cotton was placed between the femoral region and the fixation plate, and a stainless steel stick weighing 1.02 to 1.03 kg and measuring 19 mm and 50 cm in diameter and length, respectively, with a gum stopper measuring 12 mm in diameter, was placed on the skin of the right greater trochanter for 24-hr pressure loading (902.3 to 911.2 g/cm²). If necessary, additional anesthesia with ether, sodium pentobarbital, or sodium amobarbital was performed during pressure loading. After 24 hr, pressure loading was removed, and if necessary, 5% glucose was orally administered. After 2 days, the necrotic skin was resected under ether anesthesia in animals with a good general condition and a round necrotic site, and used as a decubitus ulcer (pressure-loaded necrosis) model.8,9

**Grouping:** Resected specimens (wound surface) from the burn and decubitus ulcer models were washed with absorbent cotton containing sterile physiological saline. The maximum and minimum diameters of the wound surface were measured using calipers to calculate the wound area (maximum diameter × minimum diameter). In the burn model, 16 animals were selected, and divided into 2 groups so that the wound area was similar between the two groups. In the decubitus ulcer model, 40 animals were selected, and divided into 4 groups so that the wound area was similar between the four groups.

**Application of the Test Substance:** After grouping, the test substance was applied on the wound surface, and fixed with an elastic bandage (Elastopore®, Nichiban Co., Ltd., Tokyo, Japan). In the control group, 2 sheets of gauze (2.5 × 2.5 cm) containing sterile physiological saline were applied, and similarly fixed. The test and control substance were changed once a day. Before application, the wound surface was wiped with absorbent cotton containing sterile physiological saline. These procedures were performed under ether anesthesia until the macroscopic completion of epithelization of the wound surface.

**Observation and Evaluation Methods:** After the test substance was applied, the maximum and minimum diameters of the wound surface were measured using calipers once a day to calculate the wound area. Measurement was continued until epithelization was completed. Furthermore, we calculated the percent wound area (%) to the area at grouping (when the burn and decubitus ulcer models were prepared), and the sum of the percent area (area under the percent wound area curve) using the following formula. The interval from application of the test substance until the completion of epithelization was regarded as the interval until healing:

\[
\text{Percent wound area} \times \frac{\times 100}{n} + 1) \text{ days}/2
\]

where \( n \) is the interval from the start of application until the day before healing.

**Measurement of Body Weight:** During the study period, body weight was measured at 3- to 4-day intervals.

**Statistical Analysis:** The data are expressed as the mean ± standard deviation. Significance was tested using the F-test. When isovariance was noted, Student’s t-test was employed. When unequal variance was noted, Aspin-Welch’s t-test was used. \( p < 0.05 \) was regarded as significant. In the decubitus ulcer model, the multiple comparison test was performed with consider to the percent wound area, sum of the percent area, and interval until healing, as described below, to compare the efficacy between SG-01 and the two commercially available dressing materials. Isovariance was tested using Bartlett’s method \( p < 0.05 \). When isovariance was noted, one-way analysis was performed. When there was a significant difference, the means were compared using Tukey’s method. When unequal variance was noted, the Kruskal-Wallis H-test was employed. When there was a significant difference, the means were compared using Tukey’s method. \( p < 0.05 \) was regarded as significant.

Effect (a decrease in the sum of the percent area and shortening of the interval until healing) on the decubitus ulcer model was compared based on 90% confidence intervals for differences in the mean logarithm [criteria: Log(0.8) to Log(1.25)], as described for the test methods for biological equivalence.10,11
RESULTS

Effects on Experimental Burns

Figure 1 shows serial changes in the percent wound area.

In the control group, the percent wound area gradually increased to 131.0 ± 18.4% after 2 days of application, but then decreased. In the SG-01 group, the percent wound area similarly increased for 2 days of application, but then decreased. The percent wound area in this group was lower than that in the control group; it was significantly lower on days 4 to 8, 10, and 18 to 20 of application.

Table 1 shows the sum of the percent area (area under the percent wound area curve) and the interval until healing. In the control group, the sum of the percent area was 1142.3 ± 176.7% × days. In the SG-01 group, the value was 979.0 ± 114.6% × days, significantly lower than the control value.

The interval until healing was 22.5 ± 0.9 and 22.0 ± 1.1 days in the control and SG-01 groups, respectively.

Body weight did not change until day 6 in the control or SG-01 groups, and gradually increased after day 9 (data not shown).

Effects on Experimental Decubitus Ulcers

Figure 2 shows serial changes in the percent wound area. In the control group, the percent wound area gradually decreased after one day of application. In the SG-01 group, the percent wound area also gradually decreased after one day of application. It was significantly lower than the control value from day 3 until 21. In the hydrogel A and hydrocolloid B groups, similar to SG-01; the percent wound area was significantly lower than the control value from day 3 until 21.

Table 2 shows the sum of the percent area (area under the percent wound area curve) and the interval until healing.

In the control group, the sum of the percent area was 901.8 ± 42.6% × days. In the SG-01 group, the value was 727.0 ± 76.6% × days, significantly lower than the control value. In the hydrogel A and hydrocolloid B groups, the values were 722.2 ± 56.6 and 710 ± 60.1% × days, respectively, significantly lower than the control value. The interval until healing was 22.2 ± 0.9 and 20.8 ± 1.2 days in the control and
Table 2. Effect of SG-01 in Rat Decubitus Ulcer Model — Sum of the Percent Area, Interval Until Healing —

<table>
<thead>
<tr>
<th></th>
<th>n</th>
<th>Sum of the percent area (% × days)</th>
<th>Interval until healing (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>10</td>
<td>901.8 ± 42.6</td>
<td>22.2 ± 0.9</td>
</tr>
<tr>
<td>SG-01</td>
<td>10</td>
<td>727.0 ± 76.6**</td>
<td>20.8 ± 1.2**</td>
</tr>
<tr>
<td>Hydrogel A</td>
<td>10</td>
<td>722.2 ± 56.6**</td>
<td>21.0 ± 1.1*</td>
</tr>
<tr>
<td>Hydrocolloid B</td>
<td>10</td>
<td>710.0 ± 60.1**</td>
<td>20.8 ± 1.2**</td>
</tr>
</tbody>
</table>

Each value represents mean ± S.D. Significant difference from control (*, **p < 0.05, 0.01).

Table 3. Analysis of Equivalence

<table>
<thead>
<tr>
<th>Parameter</th>
<th>SG-01 vs. hydrogel A</th>
<th>SG-01 vs. hydrocolloid B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sum of the percent area</td>
<td>logarithm</td>
<td>logarithm</td>
</tr>
<tr>
<td>Interval until healing</td>
<td>&gt; 0.99</td>
<td>&gt; 0.99</td>
</tr>
<tr>
<td>Minimum difference in detection (α = 0.1, 1-β = 0.8)</td>
<td>1.89%</td>
<td>2.40%</td>
</tr>
<tr>
<td>Difference in the mean</td>
<td>-0.070%</td>
<td>0.326%</td>
</tr>
<tr>
<td>90% confidence interval</td>
<td>0.926–1.070%</td>
<td>0.968–1.054%</td>
</tr>
<tr>
<td>Evaluation</td>
<td>equivalent</td>
<td>equivalent</td>
</tr>
<tr>
<td></td>
<td>Equivalent</td>
<td>Equivalent</td>
</tr>
</tbody>
</table>

Analysis of equivalence were calculated from results in Table 2.

SG-01 groups, respectively, with a significant difference. In the hydrogel A and hydrocolloid B groups, the values were 21.0 ± 1.1 and 20.8 ± 1.2 days, respectively, significantly lower than the control value.

Effect was compared among the SG-01, hydrogel A, and hydrocolloid B groups based on 90% confidence intervals for differences in the mean logarithm, as shown in Table 3. Effect was considered to be equivalent based on a decrease in the sum of the percent area and shortening of the interval until healing. In the control, SG-01, hydrogel A, and hydrocolloid B groups, body weight did not change until day 3, and gradually increased after day 6 (data not shown).
DISCUSSION

To clarify the pharmacological effects of SG-01, we investigated the effects of this material on burns and decubitus ulcers in experimental rat burn and decubitus ulcer models using the percent wound area, the sum of the percent area (area under the percent wound area curve), and the interval until healing as indices.

As it is important to maintain wet environments on the wound surface for wound healing,\(^\text{12-15}\) we employed rat burn and decubitus ulcer models, which reflect decubitus ulcers and burns in humans, and are commonly used in drug efficacy tests; rat burns and decubitus ulcers require a higher secretion of body fluid for the promotion of healing compared to cutting wounds, and the process of wound healing resembles that in humans.\(^\text{16,17}\) Therefore, we considered that the study using burn and decubitus ulcer models was appropriate for evaluating the features of the wound dressing material.

In the burn model, the percent wound area after 4 to 8 days of application in the SG-01 group was significantly lower than that in the control group. The sum of the percent area in the SG-01 group was significantly lower than that in the control group. The interval until healing was shortened in the SG-01 group. After 4 to 8 days of application, a relatively large volume of exudate containing various factors involved in wound healing, such as cell growth factor and various cytokines, is secreted.\(^\text{18}\) Various covering agents consisting of hydrophilic colloid contribute to wound healing by accumulating exudate on the wound surface and promoting moisturization.\(^\text{19}\) Therefore, SG-01 may have efficiently promoted retention of exudate in this phase, resulting in a significant decrease in the percent wound area. Thus, our results suggest that SG-01, which consists of a hydrophilic polymer, promotes wound healing by accumulating exudate containing various factors involved in wound healing, such as cell growth factor and various cytokines, on the wound surface.

Using the rat decubitus ulcer model, we compared the effects of SG-01 to those of hydrogel A and hydrocolloid B. The percent wound area in the SG-01 group was significantly lower than that in the control group after 3 to 21 days of application. Similarly, the values in the hydrogel A and hydrocolloid B groups were significantly lower than that in the control group after 3 to 21 days of application. The sum of the percent area and the interval until healing in the SG-01, hydrogel A, and hydrocolloid B groups were significantly lower than the control values. Thus, SG-01 promoted healing in the rat decubitus ulcer model. SG-01 decreased the percent wound area when a large volume of exudate was secreted,\(^\text{19}\) as demonstrated for hydrogel A and hydrocolloid B, suggesting that the action mechanism of SG-01 in this model involves retention of exudate on the wound surface,\(^\text{19}\) as demonstrated in the rat burn model.

These results suggest that SG-01 is useful for promoting wound healing in rat burn and decubitus ulcer models. Furthermore, SG-01 has favorable physical features\(^\text{5}\) such as high water absorption, water-maintaining ability, appropriate vapor permeability, adhesiveness, oxygen permeability and flexibility. In addition, after absorption of exudate, the adhesiveness of this agent in the wound site is reduced while maintaining the morphology of cross-linked gel, facilitating the exchange of the covering material and showing paste flexibility-related shock absorbing effects and cooling effects. Therefore, SG-01 may improve compliance, and relieve pain in clinical practice.

REFERENCES

6) Kaneko, T., Hashimoto, A., Hayashi, T., Umehara, N. and Tezuka, M. (2005) Safety studies of a new...


