Wound Healing Effects of Calvacin Gel on Burn Wound in Rats

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Calvacin gel is composed Calvatia gigantea ex Pers and Gentiana macrophylla Pall which usually used for burn and aseptic, lining and cutting wound in traditional Mongolian medicine. The objective of the study to determine technological parameter and total biological active substances and analyze the effect of burn healing Calvacin gel in rats with full-thickness second-degree burn model. Methods: Quantitative determination of the organic acid of the gel of Calvacin was performed using titremeter method respectively. Standard method was used to induce burn wound on in rats and the animals were divided into 4 groups: Control group, Calvacin-1, Calvacin-2, Silver sulfadiazine. After 24 hours of burn model being induced, we have put a thin layer of experimental gel and Silver sulfadiazine once a day. Wound area was measured every three day for 28 days under careful observation, and on the day of 7, 14, 21, 28 blood samples were taken from the animals in order to determine level of VEGF, TGFß1 to evaluate wound healing process. Results: For choosing of appropriate gel former, gels were prepared by 1% and 2% carbomer, 3%, 6% sodium carboxymethyl cellulose. The total content of organic acid 1.3±0.08%, 1.29±0.1%, 1.28±0.09% 1.29±0.09%. The rats treated with Calvacin had significantly increased levels of VEGF (at 14, 21 and 28 day), and TGFß1 (at 28 day). Conclusions: The current study showed that Calvacin has a burn healing activity.

Keywords: Burn Wound; Calvatia Gigantea Batch ex Pers; Gentiana Macrophylla Pall; Traditional Mongolian Medicine.
healing process; as well as preventing from scarring after burn injury. As a result of years of study, our researchers have created preparation named Calvacin from Calvatia gigantea Batsch. ex Pers and Gentiana macrophylla Pall. Calvatia gigantea Batsch. ex Pers has wound healing, antioxidant, cytotoxicity, antibacterial, antifungal, antitumor activity. This study also evaluated the effects of Calvacin on levels of growth factors by observing histopathological features associated with standard method induced second degree burn.

**MATERIAL AND METHODS**

The study was carried out in accordance with the Health Ethics Guidelines issued by the Mongolian Ministry of Health (2018). The study protocol (02/01/2018-06) was approved by members of “The Research Ethics Committee” and by the Institute of Traditional medicine and technology.

**Plant Material**

Spor of Calvatia gigantea Batsch. ex Pers. and herb of Gentiana macrophylla Pall. were collected as a raw material in Arbulag, Khuvsgul province, Mongolia in August 2018 and was identified by prof E. Ganbold, Sc.D in biology (voucher number ?03/2018 and ?04/2018). A voucher specimen of the plant under the number 2018/08/20/105 was deposited in the Herbarium of Pharmacology Department, Center of research, Institute of traditional medicine and technology. The herbs were air-dried and reduced to fine powder suitable for extraction.

**Preparation of Liquid Extracts**

The liquid extract of Calvatia gigantea Batsch. ex Pers. and Gentiana macrophylla Pall. were prepared by dispersing the sodium carboxymethyl cellulose of 3% and 6% w/w in half of total distilled water (20 ± 2°C). The solution of humectant was prepared in the remaining amount of water. The solution of humectants was added at the end of dispersion stage. The pH value of gel bases was measured using pH meter (Hanna HI 98128, Germany) (n = 3). Viscosity of gel bases was measured using BDV-1S digital viscometer (Biobase, Karnataka, India) (n = 3).

**Preparation of Sodium Carboxymethyl Cellulose Gel Bases**

The sodium carboxymethyl cellulose gel bases were prepared by dispersing the sodium carboxymethyl cellulose of 3% and 6% w/w in half of total distilled water (20 ± 2°C). The solution of humectant was prepared in the remaining amount of water. The solution of humectants was added at the end of dispersion stage. The pH value of gel bases was measured using pH meter (Hanna HI 98128, Germany) (n = 3). Viscosity of gel bases was measured using BDV-1S digital viscometer (Biobase, Karnataka, India) (n = 3).

**Preparation of Carbomer Gel Bases**

We prepared various carbomer gel bases with different ratio of gel former, antimicrobial preservative, neutralizer and humectant. At first stage, the solution of antimicrobial preservative and neutralizer were prepared in half of total water. At second stage, carbomer (1% or 2%) was dispersed using mixer (Akira HM-202BSS) in the solution of antimicrobial preservative and neutralizer. At third stage, the humectants were added very slowly at the dispersion stage. Triethanolamine was used as a neutralizer and the pH of the gel systems was adjusted by per cent of 1%, 1.1%, and 1.2% w/w. As a conserver were used 0.18% methylparaben-0.02% propylparaben, 0.1% methylparaben, and 0.3% methylparaben. The pH value of gel bases was measured using pH meter (Hanna HI 98128, Germany) (n = 3). Viscosity of gel bases was measured using BDV-1S digital viscometer (Biobase, Karnataka, India) (n = 3).

**Formulation of Calvacin Gel**

The spor of Calvatia gigantea Batsch. ex Pers and extract of Gentiana macrophylla Pall. was added by 5% (2:1) at the end of the dispersion stage. The gels had specific smell, dark brown to white brown color. The pH value of gels were measured using pH meter (Hanna HI 98128, Germany) (n = 3). Viscosities of gels were measured using BDV-
1S digital viscometer (Biobase, Karnataka, India) (n = 3). The quality of the gel formulation was investigated comparatively by its appearance (color, smell), pH, viscosity, and bacterial contamination. Bacterial and mould contamination was defined according to MNS-5189-2002, MNS-5190-2002, MNS-5193-2002 and MNS-5194-2002.10-14

**Animals**

The wistar rats were purchased from the animal house of research center, Institute of traditional medicine and technology, Mongolia. 90 male wistar rat of weighing between 220-250 gm were purchased from the Experimental Animal Center, Institute of Traditional Medicine and Technology of Mongolia. They were kept under controlled conditions of temperature (20±1°C) and humidity (about 50-60%), with a 12-hour light/dark cycle, and automatic ventilation 8-15 times every hour. Rats could drink ad libitum, and were fed with standard nutrient.

**Burn induction and treatment**

90 male Wistar rats weighing were used in this study. Standard method was used to induce burn wound on in rats,15 and the animals were divided into 4 groups: 1. Control group, 2. Calvacin-1 (Calvatia gigantea Batsch. ex Pers was extracted gel), 3. Calvacin-2 (Calvatia gigantea Batsch.ex Pers not extracted gel), 4. Silver sulfadiazine (1% ointment). After 24 hours of burn model being induced, we have put a thin layer of experimental gel and Silver sulfadiazine once a day. Wound area was measured every three day for 28 days under careful observation, and on the day of 7, 14, 21, 28 blood samples were taken from the animals in order to determine level of VEGF and TGFß1 to evaluate wound healing process. The study results were obtained by evaluating in serum level of VEGF and TGFß1 and histomorphological examination.

**Blood Samples**

After 7, 14, 21, 28 days experimental rats from each group were anesthetized with ketamine hydrochloride (90 mg/kg, intraperitoneally). A 5 ml blood sample was collected from each rat by cardiac puncture. The serum was separated by centrifugation at 3000 rpm for 15 minutes. The level of serum VEGF and TGFß1 SOD was measured by ELISA according following the kit’s instructions (Shanghai MLBIO Biotechnology Co.Ltd).

**Histopathological Examination**

Rats euthanized the end experiment. Burn wounds were excised down to the level of the muscle fascia by sharp dissection and included the surrounding wound margin tissue. Formalin-fixed skin specimens were prepared from four randomly

<p>| Table 1. Quality of Calvacin Gel With Different Concentration and Various Gel Former |
|----------------------------------------|---------------------------------|----------------|----------------|</p>
<table>
<thead>
<tr>
<th>No.</th>
<th>Gel former</th>
<th>Quality criteria for Gel formulation</th>
<th>Appearance (specific smell, dark brown color)</th>
<th>Viscosity (Pa x s)</th>
<th>Ph (Mean± SD)</th>
<th>Total organic acid, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Carbomer, 2%</td>
<td></td>
<td></td>
<td>201.2</td>
<td>6.3±0.03</td>
<td>1.3±0.08%</td>
</tr>
<tr>
<td>2</td>
<td>Sodium carboxymethyl cellulose 3%</td>
<td></td>
<td></td>
<td>15.5</td>
<td>6.1±0.02</td>
<td>1.29±0.1%</td>
</tr>
<tr>
<td>3</td>
<td>Sodium carboxymethyl cellulose 6%</td>
<td></td>
<td></td>
<td>36.7</td>
<td>7.9±0.03</td>
<td>1.28±0.09%</td>
</tr>
<tr>
<td>4</td>
<td>Carbomer 1%</td>
<td></td>
<td></td>
<td>93.6</td>
<td>6.02±0.02</td>
<td>1.29±0.09%</td>
</tr>
</tbody>
</table>

| Table 2. Quality of Calvacin gel with different concentration triethanolamine |
|-----------------------------------------------|----------------|----------------|----------------|
| Neutralizer                                   | Appearance (specific smell, dark brown color) | Quality criteria for Gel formulation | Ph (Mean± SD) | Viscosity (Pa x s) |
| Triethanolamine 1%                           | +              | 6.34±0.03       | 93.6           |
| Triethanolamine 1.1%                         | +              | 6.84±0.05       | 93.7           |
| Triethanolamine 1.2%                         | +              | 7.21±0.06       | 93.7           |
chosen rats per group. Specimens were dehydrated in a series of increasing ethanol concentrations then embedded in paraffin. Tissue sections (5 µm) were stained with haematoxylin & eosin (HE) and Masson-trichrome. At least three slides were prepared from each specimen and blindly examined. Histopathological scoring was achieved via an expert pathologist using Nichon microscope for detection of pathological changes.

### Statistical Analysis

Mean ± standard deviation (SD) were calculated for the observed values in each

<table>
<thead>
<tr>
<th>No.</th>
<th>Antimicrobial preservatives</th>
<th>Microbial contamination</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Total bacteria</td>
</tr>
<tr>
<td>1</td>
<td>Methylparaben-0.1%</td>
<td>3.5x10</td>
</tr>
<tr>
<td>2</td>
<td>Methylparaben-0.3%</td>
<td>3x10</td>
</tr>
<tr>
<td>4</td>
<td>Methylparaben-0.18% and Propylparaben-0.02%</td>
<td>Not detected</td>
</tr>
</tbody>
</table>

Table 3. Microbial contamination of Calvacin gel with various antimicrobials

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**Fig. 1.** Time (post burn wounding day)

**Fig. 2.** Serum level of VEGF in rats
experimental group. Statistical analysis was done by two-way ANOVA followed by tukey post hoc test was performed. Graph Pad Prism-7.0 software was used for statistical analysis with \( p < 0.05 \) considered statistically significant.

RESULTS

For choosing of appropriate gel former, gels were prepared by 1% and 2% carbomer, 3% and 6% sodium carboxymethyl cellulose. The result is shown in [Table 1].

Gels with carbomer gel base were prepared with various concentration triethanolamine and determined quality. The result is shown in [Table 2].

For the determination of the quality of gel dependence on the antimicrobial preservative, we prepared gels using methylparaben-0.18% and propylparaben-0.02% combined methylparaben-0.1%, methylparaben-0.3%. The result is shown in [Table 3].

TLC fingerprints of reference standard gentisic acid and various Calvatsin extracts are showed dark spots in 254 nm. All extracts presented chromatographic band corresponding to that of standard gentisic acid and Rf value was 0.71.

Wound Examination

The burn healing process was observed in 1, 7, 14, 21 and 28 days. A day 7 the Calvacin 1 and Calvacin 2 treated groups exhibited dark-brown, dry, hard scab. After day 14, the Calvacin-1, Calvacin-2 and SSD treated group exhibit small, thick scabs, but the control group showed dry, and dark brown scabs slightly decreased. After day 21, re-epithelialization was observed in all treated groups but the control group. Wounds healed most in the Calvacin 1 and Calvacin 2. But wound of control group a lot scab. By day 28, re-epithelialization was observed in all treatment groups and the control group (Fig. 1).

Table 4. Serum level of TGFβ1 in rats

<table>
<thead>
<tr>
<th>Time (day)</th>
<th>Control (pg/ml)</th>
<th>Silversulfadiazin (pg/ml)</th>
<th>Calvacin-1 (pg/ml)</th>
<th>Calvacin-2 (pg/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7</td>
<td>90.3±3.3</td>
<td>88.5±3.9</td>
<td>79.4±6.0</td>
<td>88.5±7.8</td>
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<tr>
<td>14</td>
<td>82.0±4.4</td>
<td>77.7±4.2</td>
<td>78.6±6.8</td>
<td>77.3±3.7</td>
</tr>
<tr>
<td>21</td>
<td>81.1±3.4</td>
<td>75.0±2.7*</td>
<td>84.4±4.8</td>
<td>80.9±6.6</td>
</tr>
<tr>
<td>28</td>
<td>81.1±5.4</td>
<td>70.4±2.1*</td>
<td>81.0±5.2</td>
<td>87.3±5.8*</td>
</tr>
</tbody>
</table>

*\( p=0.05 \)

Fig. 3. Histological examination of burn wounds stained with hematoxylin and eosin
Level of VEGF

VEGF level in plasma of the rats with full-thickness second-degree burn wound have increased for the following days of 7, 14, 21, 28 from the control group. The growth of the VEGF level in plasma varied within research groups.

For instance, group of Silver sulfadiazine’s plasma VEGF level in 7, 14, 21, 28 days were 343.0±14.4, 370.0±18.5, 391.0±20.1, 472.0±19.4 (pg/ml), Calvacin-1 group was 331.2±13.0, 317.0±19.6, 421.0±13.0, 506.0±17.4 (pg/ml); and Calvacin-2 groups result showed 332.0±18.8, 394.6±19.2, 391.0±12.0, 546.0±19.7 (pg/ml) respectively. Above mentioned group results showed statistically significant improvement (p=0.0032).

Level of TGF-ß1

TGFß1 is a control of cell growth and differentiation, induces fibrosis and scar formation (the process of wound healing).

In the control group, the level TGFß1 decreased burn induction. We found significant differences (p=0.05) between the treatment groups. In the present study, the levels of TGF-ß1 were increased in all groups and decreased after 14 to 21 days in the SSD treated groups.

Histopathological Examination

The study of the effects of Calvacin gel in the full thickness second degree burns in the experimental rats. The first week after the of the microscopic examination showed that all the groups except the Calvacin 2 were exposed to epithelial edema, epithelial and dermis inflammatory cell infiltration rates. (Figure 3 ????., A-D).

After 14 days the burning of the wound, the epidermis layer of the control group and the Calvacin-1 epithelium group was relatively larger than the other groups, and the neutrophils and macrophages observed in the control group were more penetrating than the cells. The presence of over burdishness, the formation of new vessel and collagen fibers in all the groups of experimental and control groups in the dermis layer indicates the wounds in the wound stage 3 or proliferation stage (Fig. 3, E-H). In the 21st days of observation, the epithelial cell regeneration didn’t complete in the control and experimental groups. In the epidermis and the dermis of the skin, neutrophils and macrophages were transverse in the control group of the cells more than in other groups (Fig. 3, I, J, K, L). During the study 28 days, the control group epithelium of the regeneration

Fig. 4. Histological examination of burn wounds stained with masson-trichrome
was not fully completed, and the inflammatory cells were observed in epithelial layer, scab not completely removed. The of the Calvacin-2 group skin epithelium were regenerated, in some parts of the very small size of scab, Calvacin-1 group and SSD group of adult tissue epithelium mature but some parts of the scab were observed. In the skin dermis, accumulation of collagen, accumulation of fibroblast cells, and new vascular formation are indicative of phase 4 regeneration or wound healing (Fig. 3, M-P).

7 days after observed for the group of calvacin-2 groups, was more likely to have swelling and wound in control, SSD and Calvacin-1 groups. The accumulation of new collagen fibers in the dermis layer of all experimental groups was observed (Figure 4 A-D). On the 14th day of study, experimental and control groups did not disappeared the ruptures of the wound and swell, and the formation of new vessels and collagen fibers in the dermis layer was observed, but the SSD, Calvacin-2, and Calvacin-1 groups were slightly more than the control group (Figure 4 E-H). In the 21st day of the study, wound of redness was reduced to all groups, and the epidermal layer of the regeneration Calvacin-2, Calvacin-1 were better compared to other groups, such as control and SSD groups. In the Dermis layer increased the vascular formation of the true skin layer, were significantly larger compared to other groups in the Calvacin-1 and SSD groups (Figure 4 F-I). After the 28th day of the all group, rupture of the wound, which were scarred, were scavenging, epidermis re-mature, and the newly formed collagen fiber in the dermis layer were significantly observed and were more active in the Calvacin-1, Calvacin-2 and SSD group compared to control (Figure 4 M-P).

DISCUSSION

Calvacin gel was prepared with 5% spor of Calvatia gigantea Batsch. ex Pers. and extract of Gentiana macrophylla Pall. The most suitable for the base of gel former was 1% carbomer. Carbomer aqueous gels are more viscous at pH 6 to 11.14 So we used 1.1% triethanolamine as a neutralizer. Calvacin gel containing 1% propylene glycol had good moisturizing effect and did not change viscosity. Calvacin gel base composition is similar to Cacalia gel and Indian researcher’s gel composition.9 They prepared herbal gel containing Clerodendron in for tunatum leaves extract with 1% carbomer, 0.2% methylparaben 0.1% propylparaben, 5% propylene glycol and 1.2% triethanolamine.16 Cutaneous wound healing is an essential physiological process consisting of the collaboration of many cell strains and their products.17 Wound healing and tissue repair in burn injuries are considered as a complex process including inflammation, granulation and remodeling of the tissue. We evaluated for the first time the effect of a gel of Calvacin in the full-thickness second-degree burn wound model. The result of this evaluation revealed that treatment with Calvacin gel significantly increased wound healing. Re-epithelialization and increased migration of myofibroblasts, fibroblasts, and macrophages were more prominent in Calvacin -treated groups showing that natural substances played a prominent role in wound healing process after burn injury. Angiogenesis is an important factor in proliferative phase of wound healing. VEGF is one of the most potent proangiogenic growth factors in the skin. In the last stage of wound healing, VEGF plays a role of promoting scar formation.18 These results suggested that Calvacin-1, 2 could heal scald wounds faster and result in less scarring than other treatments by regulating VEGF in the whole wound healing process. TGF-ß1 is a key growth factor secreted by several cells and is involved in a number of processes in wound healing, i.e., inflammation, angiogenesis, fibroblast proliferation, collagen synthesis, and remodeling of new extracellular matrix.18 In the control group, the level TGFß1 decreased burn induction. In the present study, the levels of TGF-ß1 were increased in all groups and decreased after 14 to 21 days in the SSD treated groups. But Calvacin believes that all stages of the wound healing process are affected. Many natural substances such as flavonoids,18 iridoids,19,20 geniposide21 and tannin22 have shown the effect of healing experimental induced burn via increasing of re-epithelialization cytokine secretion. Because Calvacin is a composed polyphenol, organic acid, iridoid, tannins. These have high wound healing and antioxidant, anti-inflammatory properties.22,23 Therefore, several publications revealed that a ointment and gel had beneficial effects in animal models of burn disorders. The Mebo has been reported to promote
chronic ischemic and neurogenic ulcer healing in patients. MEBO significantly promoted the formation of granulation tissue in cutaneous excisional wounds, shortened the time of wound healing, and increased neovascularization and the number of fibroblasts.26

Arnebia euchroma has exerted a healing effect against standard second degree burn wounds were induced as well as its significant impact on fibroblast proliferation and collagen synthesis.27 These wound healing effects of medicinal plant and ointment related to had significantly stimulatory influence on fibroblast proliferation, collagen bundle synthesis, and revascularization it was similar to our result.

CONCLUSIONS

The current study showed that Calvacin has a burn healing activity.

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