

Original Article

Effects of Topical Application of Royal Jelly on Second-Degree Burn Wound Healing in Male Rats

Zeinab Mahdian¹ Ph.D., Fereshteh Amiri² M.Sc., Zohreh Mohammadi³ M.Sc.,
Parviz Farzadnia⁴ Ph.D., Fahimeh Safizadeh⁵ M.Sc., Zahra Zare⁶ Ph.D.,
Rahimeh Rahimi⁷ M.Sc., Zahra Dehghani⁸ M.D., P.hD.,
Fariba Mohammadi Tahroodi^{9*} M.Sc.

¹ Department of Biochemistry, Faculty of Medicine, Babol University of Medical Sciences, Babol, Iran

² Department of Biology, Science and Reserch branch, Islamic Azad University, Tehran, Iran

³ Department of Microbiology, Islamic Azad University, Kerman branch, Kerman, Iran

⁴ Department of Biology and Anatomical Sciences, Faculty of Medicine, Bushehr University of Medical Sciences, Bushehr, Iran

⁵ Iranian Social Security Organization, Kerman, Iran

⁶ Department of Biology, Farhangian University, Tehran, Iran

⁷ Department of Biochemistry, Faculty of Medicine, Bushehr University of Medical Sciences, Bushehr, Iran

⁸ Department of Pathology, Bushehr University of Medical Science, Bushehr, Iran

⁹ Department of Clinical Biochemistry, Afzalipour School of Medicine, Kerman University of Medical Sciences, Kerman, Iran

ABSTRACT

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Keywords

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Background and Aims: Burn injury remain as a major medical problem throughout the world. This injury is accompanied with inflammatory and wound healing responses. Since royal jelly (RJ) has anti-inflammatory and wound healing activity therefore, the aim of this study was to evaluate the repairing effects of RJ on skin burn- damage.

Materials and Methods: In an experimental study, 40 male Wistar rats (8 weeks old) were engaged. The animals were divided into five equal groups. Group 1 was considered healthy control. Group 2 (positive control) was treated topically with Silver Sulfadiazine Cream, group 3 received Eucerin as negative control, and group 4, 5 treated with RJ (10 and 30%). Sampling was performed after observing the second-degree burns on the first, seventh and fourteenth days. Then after 28 days, rats were sacrificed and their skin tissues were used for morphological and morphometric assessments.

Results: The results of this study showed that the amount and arrangement of collagen type 1 protein was higher in the RJ treatment groups versus control group. Reconstruction and thickening of the epithelium in RJ treated groups confirmed therapeutic effects of RJ. In addition, RJ increased angiogenesis compared to the control group. The wound's surface area was reduced in the RJ treatment groups compared to the control group. In addition, fibroblast cell proliferation was increased in the groups receiving RJ versus control.

Conclusions: It could be concluded that, RJ induce wound healing effects and might be considered as potential treatment option to improve the burn wound healing.

*Corresponding Author: Department of Clinical Biochemistry, Afzalipour School of Medicine, Kerman University of Medical Sciences, Kerman, Iran. Tel: +989131983798, Email: faribamt93@yahoo.com

Introduction

Skin is a one of the body's extensive organ and important physical barrier amongst a creature and its surrounding [1]. This organ protects the human body against various harmful agents. The wound healing phases includes inflammation, hemostasis, differentiation, proliferation, and reformation. Hemostasis initiated before wound inflammation. Macrophages role in proliferation, matrix formation, and angiogenesis through secreting mediators comprises transforming growth factor beta, epithelial growth factor, insulin-like growth factor, and vascular endothelial growth factor has been demonstrated. Proliferation is another phase of wound healing [1-3]. Endothelial cells and fibroblasts are the preceding cell population that infiltrated into the healing wound site. The platelet-derived growth factor as a major chemotactic growth factor imbibes fibroblasts into the wound site. In this phase, proteoglycans and collagens are generating. Indeed, collagen type I and III are the important collagens engaged in the regeneration process [2, 3]. The ultimate stage of wound healing is scar formation that taken place in fibroblastic phase of wound healing [1, 2, 4]. Thermal wounds are continuously considers as one of the hazards that threatens human health [5-7]. In burn, the skin integrity is losing. Indeed, in the second-and third-degree burns, the wound healing procedure is regularly a long-term. Consequently, evaluating new treatment options might be capable to accelerate the wound healing progression of burn wounds and have less side effects that is excessive prominence. Perhaps, it is the purpose why

despite the presence of various therapeutic agents and approaches for the burn wounds treatment, there are numerous novel investigations for finding further effective therapeutic agents for the wound management [8, 9].

Royal jelly (RJ) is a bee product. There are many reports on pharmacological activity of RJ on experimented animals. The previous study indicated that daily application of RJ possesses better wound healing effects than nitrofurazone [10].

To our knowledge, in spite of many investigations that have assessed the effects of administration of RJ on skin problems, there are insufficient experimentations about the topical use of RJ for treatment of burn wounds. The aim of this study was to evaluate the repairing effects of RJ on tissue damage caused by burns and effect of RJ on the induction of wound healing response in Wistar rats. Considering the pathogenesis of second-degree burn wound and healing mechanisms based on that, the existing management has been designed with usage of RJ on these wounds that may be effective and be an appropriate dressing with tissue-healing characteristics.

Materials and Methods

In this study, 40 healthy, intact male weighing 200-250 gr Wistar rats were enrolled. The experimental animals were purchased from Animal Breeding Center, Bushehr University of Medical Sciences, Iran. The animals placed on each group, kept in a standard animal room (Temperature 23°C, 12-12 dark-light cycle). All

the animals had free access to food and water pellets earlier and throughout the experimental tests. The number of each group was obtained as follows:

$$\alpha = 0.05$$

$$\beta = 0.2$$

$$N = (Z\alpha + Z\beta)^2 \times \beta^2 S^2 / d^2$$

$$N = (1.96 + 0.8)^2 \times 2 \times 2 \times 0.932 / 6.72$$

$$N = 8$$

RJ was prepared from Tajdini Natural Honey Company, Kangavar, Kermanshah Province. Topical doses of RJ were selected as the effective dose based on the pilot results. For this purpose, RJ administered as an ointment.

Second-degree burn wounds induction

All the studied protocols were carried out according to the Guide for the Care and Use of Laboratory Animals [11]. For induction of second-degree burn injuries in first, the animals were anesthetized with the intraperitoneal injection of a mixture of xylazine (10 mg/ kg) and ketamine (90 mg/ kg) and the backside hair of the rats, removed. In following the area were antisepticised with the 70% ethanol. Then, a solid aluminum bar with the temperature of 150°C were employed on the area. Two second-degree burn wounds of 1 cm² were made at full thickness of the skin at intervals of 1 cm. The burn plaque plate was placed on rat skin for 15 seconds.

The project consisted of five groups of eight male Wistar rats each weighing 200-250 g, previously weighed in the scales and placed in groups, with the mean weight close to each other. After 1 week of habituation, using a randomized complete block design with weight, they were divided into 5 groups including 8 subjects.

Experimental design

After induction of the second-degree wounds (day 0), the animals randomly were divided to five group. In the experimental groups, rats received an ointment that was a uniform mixture of RJ (1% and 3%) after the wound induction. Rats in negative control group received daily Eucerin ointment for the similar condition as the experimental groups and the rats in positive control group, received silver sulfadiazine treatments. In addition, eight control healthy rats were studied to compare factors in healthy rats with treatment groups. During the treatment, the changes of the wounds appearance were evaluated with imaging by physical examination. To evaluate wound healing process, various parameters such as the dynamic measure of wound healing, the wound healing percent or contraction, were assessed in all groups, in postburn. The percent of contraction in wound surface area was computed.

Microscopic examination of wound area

In the last day of the tests (day 14 for all groups), the wound surface areas, wound healing rate, were assessed. Wound length changes were measured with image G software. The skin samples were applied to measure the amount of inflammatory factors. In addition, the morphological and morphometric examination were studied. In following, the area of wounds fixed in the 10% formalin solution. After skin tissues, processing, the samples embedded in paraffin and the blocks were cut. The sections were stained with Hematoxylin & Eosin and Mason Trichrome methods. The specimens were investigated for the fibroblasts migration,

granulation tissue formation and inflammatory response. The study protocols approved by ethical committee of Bushehr University of medical sciences, Iran by ethic code; Ir.bpums.rec.1396. 193 and Approval code:520.

Results

Macroscopic examination of wound area

After induction of second-degree burn wound in the studied groups, wound surface area pictures were recorded by digital camera in 1, 7, 14 and 21 days after wound induction. As presented in figure 1 and table 1, the wound surface area examined carefully to determine the general appearance of the wound

comprising color, consistency, wound discharge and area of wound closure. Figure 2 showed the wound surface after 7 day of treatment. The skin tissue of negative control group has necrotic pattern and had secretion and much larger wound area in compared to RJ treated groups. In other groups, partial healing was presented and the wound showed clear red appearance that indicate the partial healing progression.

Results showed that the wound surface in 14 days after treatment, in the negative control group had a red inflamed tissue and greater wound area versus RJ treatment groups. In other groups, partial healing and the closed wound pattern was observed (Fig. 3).



Fig.1. A typical second-degree burn wound in the studied groups after the wound induction with digital camera: Negative control group [Eucerin (A)], positive control (Silver sulfadiazine ointment) (B), experimental groups comprising (1% of Royal jelly; C) and 3% of Royal jelly (D) in first day.



Fig. 2. Images of skin burn wounds in different groups: Negative control [Eucerin (A)], Positive control [Silver Sulfadiazine (B)] and experimental groups; 1% of Royal Jelly (C) and 3 % of Royal jelly (D) in seventh day

As presented in Figure 4, wound surface in 21 days after treatment in all experimental groups relatively improved, but the color, extent and appearance of the wounds in treatment groups, particularly those treated with RJ, were more improved and show partially closed pattern. As

shown in Figure 5, wound surface in 28 days after treatment in all experimental groups were improved, but the color, extent and appearance of the wounds in treatment groups, particularly those treated with RJ, were more recovered and show complete closed pattern.

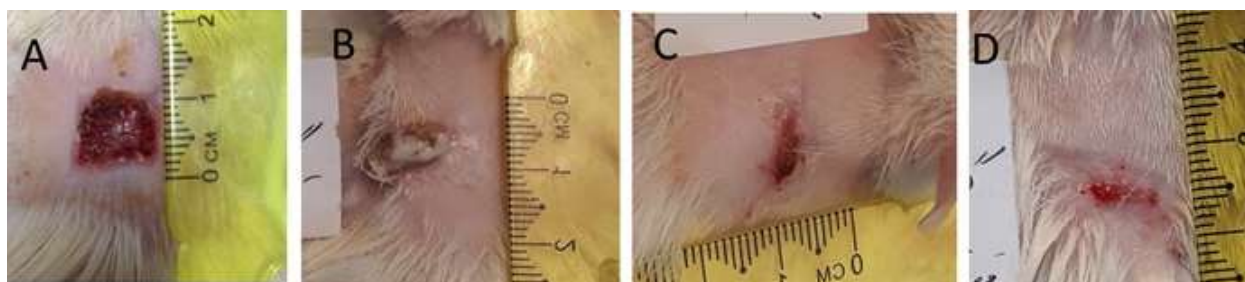


Fig. 3. Images of skin burn wounds in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in Fourteenth day

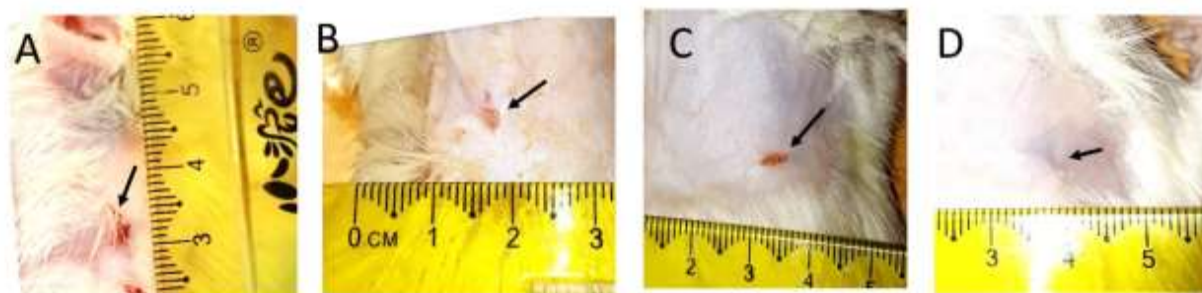


Fig. 4. Images of skin burn wounds in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in 21th day.

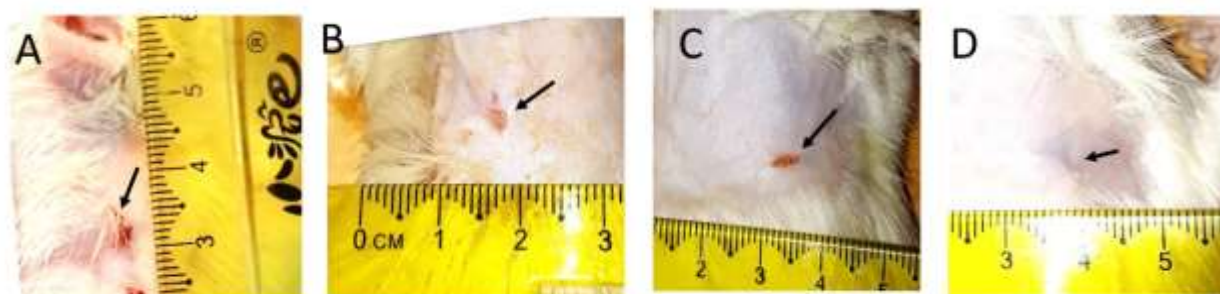


Fig. 5. Images of skin burn wounds in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine(B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in 28th day.

Macroscopic examination of wound area

Microscopic examination in 1, 14 and 28 days after wound induction were performed. Correspondingly, wound structure examined by two conventional staining methods to evaluate the wound healing structure, including the re-epithelialization. Indeed, specific staining for evaluate the regeneration and maturation of the collagen fibers are considered to study the microscopic wound structures at three consecutive time-intervals; beginning, middle and end of treatment period. Skin tissue micrographs in different groups presented in figure 6. The skin tissue in all groups have uniformly normal microscopic patterns and healthy epithelium with the normal tissue structure. Skin tissue micrographs in different groups, after 14 day of treatment were studied. Our data showed

that the skin tissue in treatment groups including Silver Sulfa Diazine ointment and also particularly in RJ treated cases had relatively normal uniform microscopic structure and epithelium have relatively good thickness and growth without any histopathologic changes. In negative control group, collagen fibers repaired but epithelium is not formed (Fig. 7). As presented in figure 8, at the end of treatment (28th day), skin tissue in all groups has relatively normal microscopic structure, good thickness of epitheliums, growth and repair pattern with any specific histopathological changes. Nevertheless, in the positive control groups (Silver Sulfa Diazine ointment) and RJ treatment, there are thicker epithelium structure versus other groups.

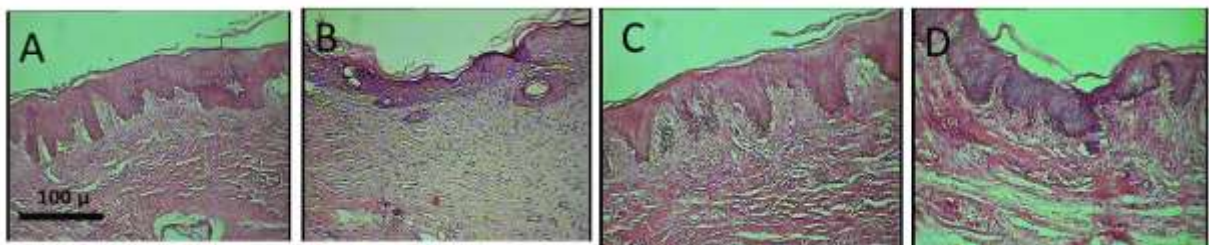


Fig. 6. Skin micrographs from Hematoxylin and Eosin staining in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine(B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in first day. magnification: $\times 100$

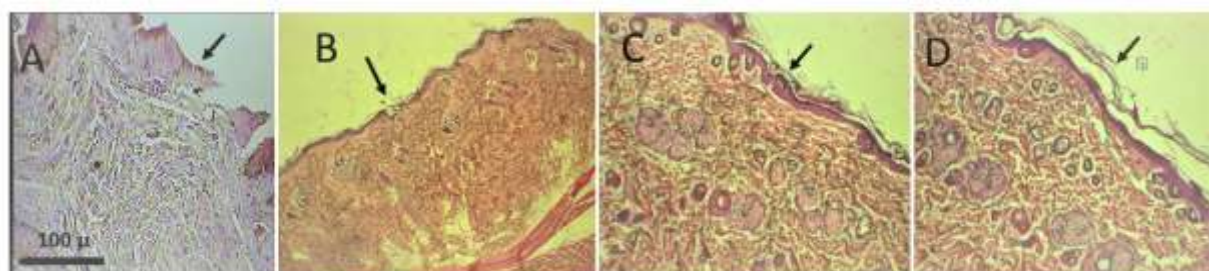


Fig. 7. Skin micrographs from Hematoxylin and Eosin staining in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal Jelly (D) in 14th day. magnification: $\times 100$

According to our results from Mason's Trichrome staining (figure 9), in all experimental groups the skin tissue with strands of graft collagen have uniformly normal microscopic structures. Figure 10 showed the micrograph of skin tissue (dermis area) in different groups on the 14th day. As shown in figure 10, skin tissue with strands of

graft collagen have not regular normal microscopic structure in the negative control group but in the other positive control groups (Silver Sulfa Diazine ointment) and the RJ treated groups (1% and 3% of RJ), skin structure is completely normal with mature collagen fibers.

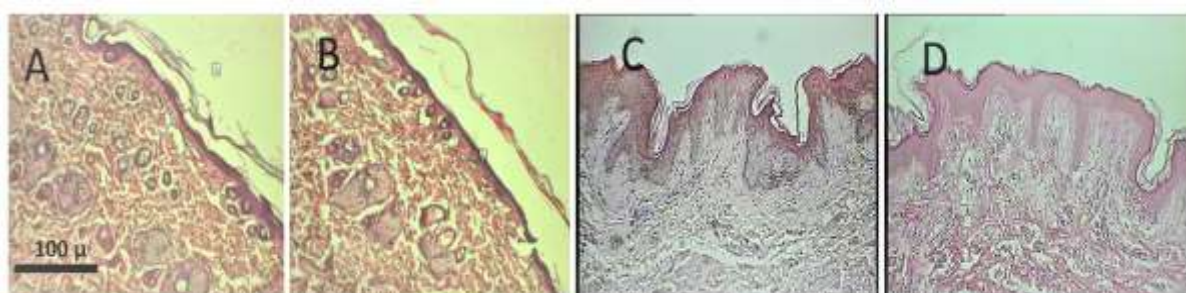


Fig. 8. Skin micrographs in different groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in 28th day. magnification: $\times 100$

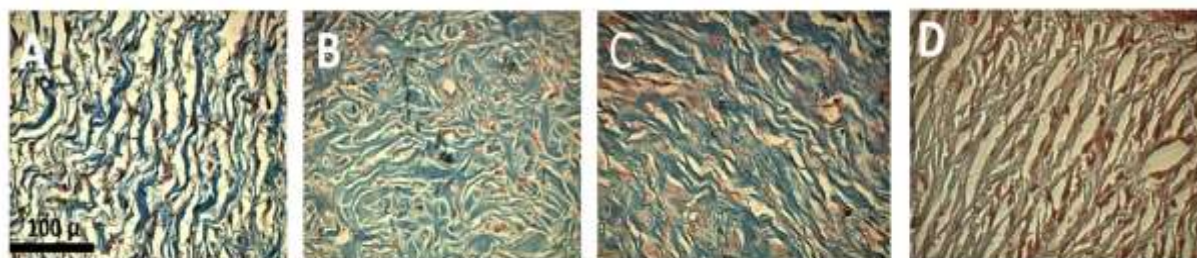


Fig. 9. Skin micrographs of Mason's Trichrome staining in various experimental groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in first day. magnification: $\times 100$

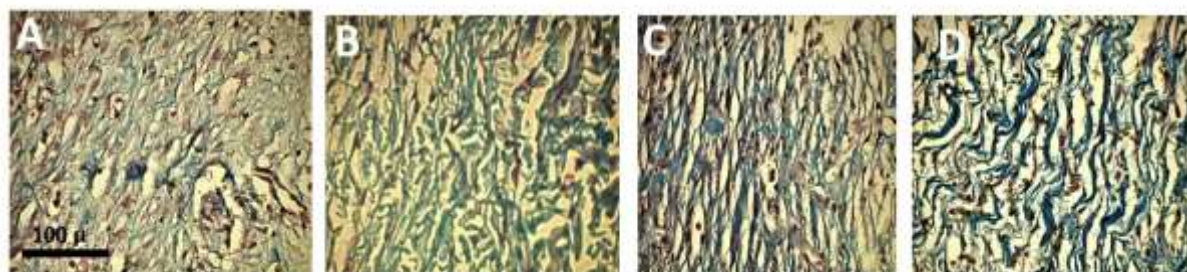


Fig. 10. Skin micrographs of Mason's Trichrome staining in various experimental groups: Negative control [Euserin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in 14th day. magnification: $\times 100$

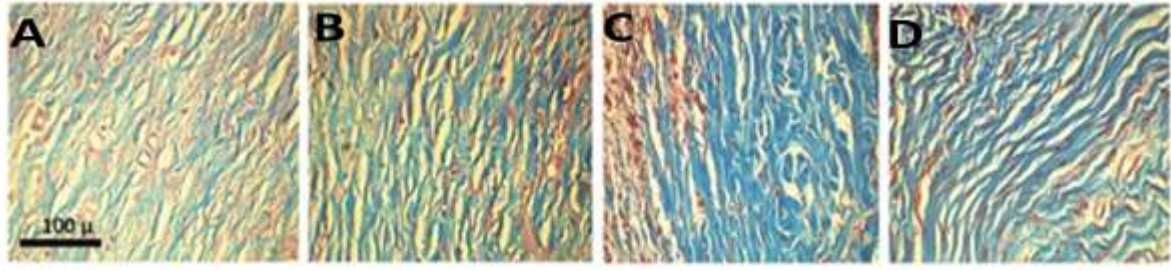


Fig. 11. Skin micrographs of Mason's Trichrome staining in various experimental groups: Negative control [Eucerin (A)], Positive control [Silver Sulfa Diazine (B)] and experimental groups; 1% of Royal jelly (C) and 3 % of Royal jelly (D) in 14th day. magnification: $\times 100$

Table 1. Qualitative changes in the wound's appearance in experimental groups

Parameters	Date(day)	I (Negative control)	II (Positive control)	III (Test)
Epithelium thickness (Micron)*	1	149.11 \pm 43	151.11 \pm 23	152.45 \pm 13
	14	No epithelium	34.11 \pm 3 *	31.21 \pm 5*
	28	160.53 \pm 22	158.36 \pm 21	159.46 \pm 22*
The number of blood vessels**	1	10.21 \pm 2	11.17 \pm (2)	9.31 \pm (0.9)
	14	7.15 \pm (0.8)	12.22 \pm (1)*	15.10 \pm (1)**
	28	11.30 \pm (0.8)	13.17 \pm (2)	14.01 \pm (2)
The number of fibroblast cells**	1	21.16 \pm (110)	22.12 \pm (1.09)	23.45 \pm (2.15)
	14	18.14 \pm (1.1)	33.10 \pm (1.15)*	30.22 \pm (2)*
	28	15 \pm (1.36)	19.26 \pm (1.26)	20.21 \pm (1.12)
Area of wound closure (%)	7	96.32 \pm 11)	98.12 \pm (1.31)	92.20 \pm (4.18)
	14	76.45 \pm (2.50)	66.13 \pm (1.34)*	59.67 \pm (1.46)*
	21	70.14 \pm (.87)	38.01 \pm (4.11)*	31.12 \pm (3.21) *
	28	14.90 \pm (1.21)	8.42 \pm (.96)*	10.14 \pm (2.01)*

Data is presented as mean \pm SD; * Significance with negative control; **Significance with negative control

Further examination in the negative control group (on the 14th day) indicated that the skin tissue (dermis area) with strands of graft collagen still have not any regular normal microscopic structure. Nevertheless, in the other positive control group (Silver Sulfa Diazine ointment) and the experimental groups (1% and 3% of RJ) natural skin structure with mature collagen fibers were observed (figure 11). Morphometric properties comprising the data from wound healing, epidermal thickness, number of small blood vessels, and fibroblasts have been investigated and presented in figure 11.

Discussion

Various natural products apply in wound healing therapeutic strategies [1, 10, 12]. In this regards RJ as a bee product has numerous pharmacological activities on experimental animals [10]. Therefore, in this study we evaluate the effects of RJ on the induction of wound healing response in Wistar rats.

The present study confirmed that in burned-induced wounds of Wistar rats, RJ induce the wound healing and increases the collagen content indicating the triggering the reconstruction and thickening of the epithelium in treated groups.

As an effective common histopathological examination for tissue change after therapeutic intervention in animal models, Hematoxylin and Eosin staining [13, 14] and also Mason's Trichrome staining were conducted. According to our results, RJ improved angiogenesis during wound healing process versus to the control group. The therapeutic effects of RJ confirmed by reduced wound's surface area after RJ exposures.

The present data also revealed that in RJ treated cases, the time required for complete healing of the burn- lesion, was shorter than the control.

Accordingly, RJ may be effective for increasing the healing activity in definite pathway. Commonly, wound healing takes an extended time in some patients.

Generally, anti-inflammatory agents delay wound healing process by decreasing cell proliferation that is an unfavorable properties of these drugs. Nevertheless, RJ acted as an anti-inflammatory drug, although these activities, RJ reduced the time required for wound healing. These activities might be related to one of the numerous components in RJ, such as skin respiratory factor. Nevertheless, another related factor has not yet been recognized [15, 16].

In this regards, Fujii et al., showed some anti-inflammatory activity of RJ. They also indicate that, RJ shortened the period of wound healing and is able to enhance wound healing [16]. In comparable survey, which conducted by Siavash et al. [17], RJ dressing applied as effective technique for treating diabetic foot ulcers. They indicate that this treatment option may promote wound healing as an effective intervention. Also, other study in animal model

shown that RJ could promote wound healing [18] which is in accordance to our data.

In study by Rashidi et al, the impacts of topical usage of mixture of RJ, olive oil, honey, and propolis extract were evaluated on cutaneous wound healing. Overall estimation of wound healing progress of this mixture including RJ was enhanced than those of the control group [19]. Certainly, previous reports indicate that RJ increases wound-healing activity. In both *in vitro* and *in vivo* wound-healing models, in presence of RJ, human fibroblasts were capable to migrate and also RJ increases the levels of sphingolipids. Therefore, RJ shortened the curing time of desquamated skin lesions [20, 21].

Another survey on the RJ administration have also displayed protective action on human skin against ultraviolet B-induced photoaging by inducing collagen production [22]. RJ wound dressing exerted vasodilation impacts nearby the wound, that can help to dilate the blood vessels to increase blood flow [17]. Based on author knowledge the burn-wound healing activity of RJ has not yet been reported. Consequently, based on our findings, RJ induce wound healing effects and might be considered as potential treatment option to improve the quality and quantity of burn wound healing.

Conclusion

The present study focused on the potential wound healing effects of RJ. This product is extremely rich in active ingredients that have biological actions in promoting good health. Accordingly, reconstruction and thickening of the epithelium in RJ treated groups confirmed therapeutic effects. In addition, RJ increased

angiogenesis compared to the control group. In addition, fibroblast cell proliferation increased in the groups receiving RJ compared to the control. Thus, this product could be applied as potent wound healing agent. However, it is crucial to conduct further researches to define the main mechanisms that is associated with pharmacological activities of RJ and the

suitable amounts that could be administered to obtain favorable health incomes.

Conflict of Interests

The authors declared no conflict of interest.

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