The Effects of Topical Carvacrol Application on Wound Healing Process in Male Rats

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ABSTRACT

Carvacrol containing products such as Origanum onites have been used as phytotherapeutic agents in the treatment of serious skin injury. It has been suggested that carvacrol is the active component of these herbs because of its anti-microbial property. With the anti-microbial activity and as an ingredient of these herbs, carvacrol is a promising molecule for the treatment of skin injury. In the present study, we have evaluated the efficacy of carvacrol on healing progress after excisional skin injury. Here, Wistar-Albino rats were divided into two groups and treated with carvacrol and vehicle. Carvacrol was administrated topically at a concentration of %12.5 for the 5 consecutive days after excisional skin injury. Tissue samples were harvested on days 3rd, 8th and 12th after injury. Significant beneficial effect of carvacrol was observed at the end of the experiment. In the acute phase of the injury, carvacrol treatment increased tissue granulation and decreased wound depth moderately. These effect of carvacrol was associated with increased TNF-α. However, at the second half of the experiment the elevated level of TGF-β1 was observed as compared with control animals. The level of IL-1β was increased in carvacrol treated animals only on day 8. Here, we provide evidence that carvacrol improves wound healing by regulating pro-inflammatory molecules TNF-α, IL-1β and TGF-β1.

Keywords: carvacrol; wound healing; TNF-α; IL-1β; TGF-β1; aromatic herbs.

INTRODUCTION

Carvacrol is an essential oil component of aromatic herbs such as Origanum onites.[1–4] In-vitro and in-vivo studies performed with carvacrol showed that it has diverse effects, including antimicrobial,[5,6] antiviral,[7] antifungal,[8–10] antioxidant,[11,12] and antispasmodic.[13–15] In this context, carvacrol is a promising molecule for the treatment of local skin injury.

Wound healing is defined as the completion of closure at the clinically wounded skin area. Wound healing processes develops from some pathophysiological events including haemostasis, inflammation, cellular proliferation, matrix synthesis and remodeling. It has been also shown that IL-1, TNF-α and TGF-β stimulate neutrophile migration, fibroblast activation and reepithelization.[16,17]

In this context, we have investigated the effects of carvacrol on wound healing induced by excisional skin injury model in rats. We have evaluated wound surface area, wound depth, and granulation of the skin in a time dependent manner. In addition, we have studied the effects of carvacrol on the expressions of pro-inflammatory IL-1β, TGF-β and TNF-α.

MATERIAL AND METHOD

Animal experiments

Experiments were performed in accordance with National Institutes of Health Guidelines for the Care and Use of Laboratory Animals, and the study was approved by the local ethics committee (Ankara University, Medical Faculty). In this study, 350±27 gr, male Wistar-albino rats were used in this study. Rats were equally divided into two set of experimental groups and treated with either

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vehicle or carvacrol. Carvacrol was purchased by Sigma (Sigma-Aldrich; Istanbul, Turkey) and dissolved in sunflower oil in a concentration of 12.5%. Two-hundreds microliter carvacrol containing sunflower oil was applied topically on the injured skin just after excision of the skin and repeated for 5 consecutive days. Tissue samples were collected 3 (n=10), 8 (n= 10) and 12 (n= 10) days after induction of wound injury.

The animals were anesthetized with ketamine (60 mg/kg) and xylazine (6 mg/kg). Back, starting from the neck (2 × 3cm), of rats were completely cleaned of hair and wiped with 70% alcohol. Full-thickness skin wounds were formed on the cleaned part, as 1cm away from each of the middle line and each other, through the use of 4 mm punch biopsy tool.

Half of the tissues taken out were fixed with 10% formal solution for histological examination and the other half of the injured tissues collected were immediately frozen within liquid nitrogen for the examination of TNF-α, TGF-β₁, IL-1β, levels with enzyme linked immunosorbent assay (ELISA) method.

Pathological and histological evaluation

Photographs of scar tissues were taken by using a Canon Power shot A410 model digital camera, and surface areas of scar tissues were calculated by using the program AutoCAD (2006). For the histological examination, 5 μm tissue samples were stained with Hematoxilen-Eosin (H&E). Depth of tissue damage and thickness of granulation tissue on ulcer surface were measured by using Olympus BX50 light microscope and microscopic grid. During the evaluations, the depth between healthy epidermis on the side and the deepest area of ulcer surface was measured for determination of the depth of tissue damage. The distance between lower part of ulcer exudates on the surface and the lowest part of fibroblast was measured. In addition, the vascular proliferation and thickness of granulation tissue were evaluated.

Evaluation of TNF-α, TGF-β₁, IL-1β levels by ELISA

Glas-Col homogenizing system with Teflon top (Glas-Col, U.S.A.) was used for homogenization of the tissue samples. 4 ml buffer solution, 200 μl PMSF and 2 μl leupeptin were respectively put into homogenization glass tube. Each two pieces of four each tissues taken out according to days were added into this mixture. Glass tube was put into another container filled with ice. Tissue was ensured to be disintegrated by means of homogenizing system with Teflon top at 3200 rpm for ten minutes. The extract obtained at the end of ten minutes was transferred into a glass centrifuge tube and centrifuged for 10 minutes at 3000xg within a centrifuge device with its ambient temperature adjusted as 4 degrees. The supernatant obtained after the centrifuge operation was transferred into polypropylene tubes and kept at -80 degrees until the time they will be used in ELISA operation.

BIOSOURCE Immunoassay Kit Rat TNF-α KRC3012 (Biosource International, Inc. USA) for TNF-α, BIOSOURCE Immunoassay Kit Rat Multispecies TGF-β₁ KAC1689 (Biosource International, Inc. USA) for TGF-β₁ and BIOSOURCE Immunoassay Kit Rat IL-1β KRC0012 (Biosource International, Inc. USA) for IL-1β were used throughout the study. In the preliminary study conducted for TNF-α and IL-1β, solutions at different dilution rates were obtained from tissue homogenate solutions by using “standard dilution buffer”. For TNF-α and IL-1β, the homogenate was not diluted; for TGF-β₁, the supernatant was 10-times diluted with “standard dilution buffer”.

Statistical evaluations

Variance analysis was applied in repetitive measurements for the purpose of comparing groups with regard to time. Due to the fact that main effects of group and time were found significant in consequence of variance analysis in repetitive measurements, double comparisons between group and time averages were made by examining them with Bonferroni test with confidence interval adjustment.

Parameters on the same day of carvacrol and control groups were analysed by nonparametric Mann Whitney-U test. P values <0.05 were considered statistically significant.

RESULTS

Pathological examination evaluation results

Carvacrol treatment improved wound healing processes. Scar surface measurement were conducted 3, 8 and 12 days after induction of injury. On day 12, carvacrol treatment decreased scar surface significantly (p=0.014) (Fig. 1). Histological examination showed exudate formation on day 3 in the carvacrol and vehicle treated animals (Fig 2). Upon statistical evaluation of the data obtained from microscopic examination of scar tissue samples taken from control and carvacrol groups, wound depth on the 8th day was significantly high in carvacrol group with respect to control group (p=0.002) (Fig. 2), granulation tissue thickness on the same day was found significantly low in carvacrol group compared to control group (p=0.008) (Fig. 2).
Cytokine levels of tissue homogenates obtained from scar tissues in control and carvacrol groups were measured with ELISA. TGF-β₁ levels on the 8th and 12th days were found significantly low in carvacrol group compared to control group (p=0.009) (p=0.03) (Fig. 3A). IL-1β levels on the 8th day were found significantly high in carvacrol group compared to control group (p=0.034) (Fig. 3B).
TNF-α levels of 3rd day were found significantly high in carvacrol group compared to control group (p=0.000) (Fig. 3C).

**DISCUSSION**

Carvacrol is an essential oil which is present in compounds of many herbs. Apart from its known effects, its anti-microbial effects have been studied.\[^{[88]}\] It inhibits prostaglandin synthesis, which plays roles in decreasing postoperative pain.\[^{[18]}\] In this context, we have studied the roles of carvacrol in the pathophysiological events induced by excisional skin injury. First of all, we have defined the effective concentration of carvacrol. Following, we have treated animals with carvacrol at the concentration of 12.5%. Our results revealed that carvacrol treatment improves healing rates of scar tissue surface area on the 12th day after surgery as compared with control. In contrast to increased tissue granulation, the wound depth is consonant with the condition in normal wound healing.\[^{[19]}\] That granulation tissue thickness went up to approximately same levels on the 12th day while it was lower in carvacrol group compared to control group on the 8th day makes one think that carvacrol may affect wound healing between 8th-12th days. As a matter of fact, difference between groups in wound depth measurements on the 8th day changed in favor of carvacrol group on the 12th day, that is to say that wound depth in carvacrol group reached the same values in control group between 8th-12th days. This findings support that carvacrol may affect wound restoration process between 8th-12th days.

Different soluble factors play role in arranging and controlling phases of wound healing.\[^{[20]}\] These factors are cytokines, growth factors, proteases, eicosanoids, quinines and cellular metabolites.\[^{[26]}\] For this reason, it is not possible to affix events in the course of restoration process on effect of a single cytokine or growth factor.

IL-1 is a cytokine that plays a key role in inflammatory phase. It mediates synthesis and secretion of IL-1 and other proinflammatory cytokines (IL-6, G-CSF, GM-CSF) by inducing fibroblast and endothelial cells. It also leads to synthesis and secretion of more interleukin and PDGF by stimulating monocytes on wound area.\[^{[28]}\] Leukocyte chemoattraction affects reepitilization and fibroplasia phases.\[^{[23]}\] IL-1 level reaches to measurable levels in the first 24 hours in experimental wounds, peaks between the 1st and 3rd days and rapidly decreases through the first week.\[^{[22,23]}\] In the study, IL-1β in both of control group and carvacrol group were at measurable levels in the first day and showed a tendency to increase until the 3rd day. IL-1β levels on the 8th day were significantly high in carvacrol group. This study showed that carvacrol may contribute to inflammation during the early period and reepitilization and fibroplasia processes during late period by means of increasing IL-1β levels.

Besides leukocyte chemoattraction, TNF-α increases vascular proliferation and permeability.\[^{[24]}\] TNF-α also contributes to great number of cellular metabolic events such as procurement of nutrients and acute phase protein synthesis which are essential for wound healing.\[^{[28]}\] On the other hand, TNF-α may have negative effects on wound healing. These effects occur as inhibition in collagen synthesis depending upon decrease in productions of collagen hydroxyproline and proalpha-1 chain.\[^{[26,27]}\] Rapala et al. has shown that TNF-α, depending on dose in acute experimental wounds, decreases granulation tissue growth on the 7th day. However, this effect was not observed on the 14th and 21st days.\[^{[28]}\] TNF-α can be locally detected within 12 hours and, peaks after 72 hours in clinically formed wound.\[^{[29]}\] However, our findings showed that TNF-α levels in homogenates obtained from scar tissue have a tendency to decrease until the 3rd day in control group, peaked on the 8th day and gradually decreased later on. On the contrary, TNF-α levels in carvacrol group showed a continuous tendency to decrease.

TGF-β is one of the most significant mediators in wound healing. It is secreted by T lymphocytes, endothelial cells, keratinocytes, thrombocytes in scar tissue, macrophages, smooth muscle cells and fibroblasts.\[^{[30,31]}\] TGF-β1, TGF-β2, and TGF-β3 are 3 isoforms of TGF-β found in mammals. TGF-β1 and TGF-β2 stimulates scar tissue formation by causing extracellular matrix deposition via increasing extracellular matrix production and suppressing proteolysis and catabolism of extracellular matrix. On the other hand, some studies showed that TGF-β1, unlike the other two isoforms, has activities intended to decrease scar tissue formation.\[^{[32–35]}\] However, our results showed that TGF-β1 expression was significantly higher in carvacrol treated animals as compared with control animals.

**CONCLUSION**

Here, we have evaluated the effects of carvacrol on tissue repair and time dependent expression patterns of IL-1β, TGF-β1 and TNF-α after skin injury. Our result indicated beneficial effects of carvacrol on skin injury treatment
and it was associated with a moderately increased tissue granulation in the subacute phase of injury and increased expression of TGF-$\beta_1$.

**DECLARATION OF INTEREST SECTION**

The authors have any financial, consulting, and personal relationships with other people or organizations that could influence (bias) the author’s work.

**REFERENCES**