Effect of ozonated olive oil on experimentally induced skin infection by *Streptococcus pyogenes* and *Staphylococcus aureus* in rats

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Antibiotic resistance problem is one of the most important problems in treating infectious diseases. To overcome such problems, agents apart from antibiotics that can be used in the treatment of infectious diseases and have no resistance problem should be explored. Natural substances, such as vegetable oils and ozone are possible resources. Thus, in the present study, we investigated the effect of ozonated olive oil on experimentally induced skin infection by *Streptococcus pyogenes* and *Staphylococcus aureus* in rats. The study was performed on 3 main groups. Experimental skin infection was induced by *S. pyogenes* (Group 1) or *S. aureus* (Group 2). Animals in Groups 1.1 (n=10) and 2.1 (n=10) received ozonated olive oil, in Groups 1.2 (n=10) and 2.2 (n=10) received ozonated olive oil, in Groups 1.3 (n=10) and 2.3 (n=10) received fusidic acid on the infection side in their skin at 24, 28, 32, 40, 48, 56, 64, 72, 80, 88, 96 hours. Groups 1.4 (n=11) and 2.4 (n=11) did not receive any treatment after infection. Group 3 (n=11) received only serum physiologic at above mentioned periods. After occurrence of the skin infection; ozonated olive oil had anti-microbial, anti-inflammatory effects and observed to have healing activity at 72-80 h after infection. Furthermore, the effects of ozonated olive oil were comparable with the effects of fusidic acid which its activity proved against skin infections with Staphylococcus and Streptococcus.

**Keywords:** Fusidic acid, Olive oil, Ozonated oil, Ozone, Ozone therapy

Staphylococci and Streptococci are gram positive and cocci morphology bacteria1. *Streptococcus pyogenes* and *Staphylococcus aureus* are the most commonly encountered microorganisms of bacterial infections of the skin in humans2,2. Though in animals too *S. aureus* causes bacterial infections of the skin mostly, at times *S. pyogenes* also leads to mono infection of skin3. In the treatment of skin infections caused by bacteria, both systemic and local treatment can be applied4. Overuse, low dose or ineffective duration of antibiotics usage in the treatment of infectious diseases cause resistance5. Ozone therapy has come up as an alternative treatment option that does not cause any such resistance6-11.

Ozone is a natural gaseous molecule consists of three oxygen atoms whereas the oxygen molecule, far more stable and is composed of only two atoms12. Ozone has a strong germicidal activity and destroys bacterial cell membrane integrity by the oxidation of phospholipids and lipoproteins. It can break down the cell wall of many pathogenic microorganisms5,13. Ozone, has been used as the oxygen-ozone mixture, ozonated oil and ozonated water in bacterial skin infections in human12,14-16.

However, effects of ozonated olive oil (OZO) on experimentally induced skin infection by *S. aureus* and *S. pyogenes* in rats have not been reported in literature until now. In the present study, we evaluated the potential effect and antibacterial activity of OZO on experimentally induced skin infection using *Streptococcus pyogenes* and *Staphylococcus aureus* in rats.

**Materials and Methods**

Nintey-three female Wister Albino rats aged between 6 and 8 weeks weighing 80-130 g were grouped into 3 main groups wherein Group 1 was infected with *Streptococcus pyogenes*; Group 2 was infected with *Staphylococcus aureus*; and Group III was negative control group. Infections were made in the skin of the experimental animals as described by Kugelberg et al.17 with some modifications described below. Furthermore, each of Group 1 & 2 were further divided into 4 subgroups.

The animals used in the present study were obtained from the Laboratory Animal Research Centre of the University of Erciyes. Experiment,
hosting and maintenance were made at this centre. Ethical Committee Permission were taken from the Ethical Committee of the Local Animal Experiment Commission, University of Afyon Kocatepe as the 420-15 Reference number and dated on 12.02.2015 with the 4953302/09 number.

Apart from S. pyogenes or S. aureus infection, animals in Groups 1.1 (n=10) and 2.1 (n=10) received 0.1 g, OZO; Groups 1.2 (n=10) and 2.2 (n=10) received olive oil (OL); and Groups 1.3 (n=10) and 2.3 (n=10) received 0.1 g fusidic acid (FA) on the infection side of the skin at 24, 28, 32, 40, 48, 56, 64, 72, 80, 88, 96 h. Groups 1.4 (n=11) and 2.4 (n=11) did not receive any treatment after infection. Group 3 (n=11) received only 0.1 g serum physiologic at above mentioned periods.

Experimental skin infection
Infection, modified from Kugelberg et al.,17 superficial skin infection model. Under general anaesthesia (xylazine 10 mg/kg; basilin 2% bavet, ketamine 50 mg/kg; ketasol 10%, interleas), first hairs were shaved by fixing the epilating strip and then removing it quickly. It was repeated three times. Abraded skin regions were obtained and the area was disinfected with a disinfectant (bacitron) and alcohol. Before inoculation of bacteria, swaps were taken from the abraded skin and were kept at 4°C for microbiological analysis. The Wattman paper (11 millimeter in diameter) was placed on to the abraded skin region. Standard strains of S. aureus ATCC 25923 and S. pyogenes, a clinical isolates, were obtained from the Erciyes University, Medical Faculty, Microbiology Laboratory. About 0.1 mL of S. pyogenes strain and S. aureus ATCC 25923 strain (each 1×10^8 CFU/mL) was prepared and were adsorbed on to the Wattman paper. After the bacterial inoculation, the infection area was covered with an adhesive bandage (Betafix) and closed with adhesive tape to protect animals from removing the adhesive bandage. Bandage and strip were removed after 24 h of infection. To determine the status of infection, swaps were taken from the region before the treatment. Then, treatment was started and continued up to 96 h. Clinical finding scores were made for recovery as 1 (No), 2 (Low), 3 (moderate) and 4 (severe).

Histopathological analysis
Tissue samples under general anaesthesia using 4 mm biopsy punch before treatment and 24, 28, 32, 40, 48, 56, 64, 72, 80, 88, 96 h after the treatment were collected and evaluated in all groups. The biopsy samples fixed in formalin (10%). The formalin-fixed biopsy specimens were embedded in paraffin and stained with haematoxylin (Sigma-HHS32-IL) and eosin (Merck-109844.1000). For identification of the bacteria, the biopsy specimens were gram’s stained (Merck-M111885.0001). Furthermore, swap samples before treatment and only 24 h after treatment were collected to determine the presence of infection in all groups.

Ozonation procedure and Characterization of oils
As required for experimental study, OL and OZO material was obtained from a certified commercial company (Aktifoks, Isik Cosmetics/Turkey). According to the commercial firm Hansler brands (Ozonosan Alpha Plus, Germany), the ozone device (which produce 95% oxygen and 5% ozone mixture) was used to prepare 10 L of OL which this device reported to produce 25 mg/L of ozone per min in the 18-20°C. This procedure was applied for 10 days. Acidity, peroxide, iodine and p-anisidine values of OL and OZO were determined according to AOCS18, AOCS19, AOCS20, AOCS21 respectively. The pH value of OL and OZO were determined at room temperature (24°C). Viscosity of OL and OZO were determined by vibrating viscometer device (AND SV-10 Japan) at 24-40°C. These values are important for obtaining a high quality OZO22,23.

Fusidic acid and stafine cream material
This material obtained from a commercial firm (Kocak Farma AS/Turkey) which is licensed for human use as local application and can be obtained from any chemist shop. In the animal experiment, stafine cream was used.

Determination of antimicrobial activity
The antimicrobial activity was determined by antimicrobial disc diffusion tests with FA, OZO and OL according to the CLSI24 and CLSI25 methods, respectively.

Statistical analysis
The IBM SPPS 21 for Windows software statistical package program was used for statistical analysis. Statistical comparisons with concern to clinical findings in between groups and within group were made using Kruskal-Wallis test. Statistical comparisons of presence of bacteria in between groups were made using Chi-square test. Values were given as mean ± standard deviation. The level of \( P <0.05 \) was used to determine the significance.
Results

Antimicrobial sensitivity and disc diffusion tests are given in Table 1. According to the table both microorganisms used in the present study were resistant against OL but not against FA and OZO (Table 1).

After preparing the animals for infection, swaps were taken before and 24 h after infection. Swaps taken before infection (Groups 1, 2 and 3), inoculated for microbiological analysis. No bacterial growth was observed in any of the groups. However, bacterial growth developed in the swaps taken after 24 h of infection in all the infected samples (Fig. 1 B & C, Fig. 2 B & C, and Fig. 3 B & L).

Presence of bacteria in the dermis in Groups 1, 2 and 3 were given in Table 2 and 3. According to this table, Group 3 which was negative control and it had no bacterial growth. In Groups 1.4 and 2.4, bacterial growth were 100% which were positive controls. The least bacterial growth in dermis were in Groups 1.1, 2.3, 1.3, 2.1, 1.2 and 2.2, respectively (Table 2 and 3), (Fig. 1 B & C, Fig. 2 B & C).

Inflammatory cell ratios in papillary dermis, reticular dermis and subcutis are given in Table 2 and 3. According to these tables, no inflammatory cells were observed in Group 3 histopathologically. Inflammatory cell ratios in Groups 1.4 and 2.4 were

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**Table 1**— *Staphylococcus aureus* ATCC 25923 and *Streptococcus pyogenes* (a clinical isolate) antimicrobial sensitivity tests results

<table>
<thead>
<tr>
<th>Strain</th>
<th>Fucidic acid</th>
<th>Ozonated olive oil</th>
<th>Olive oil</th>
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<tbody>
<tr>
<td></td>
<td>disc (µg/mL)</td>
<td>disc (µg/mL)</td>
<td>disc (µg/mL)</td>
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<tr>
<td>S. aureus</td>
<td>5 µg</td>
<td>10 µg</td>
<td>25</td>
</tr>
<tr>
<td>ATCC 25923</td>
<td>25</td>
<td>26</td>
<td>0.25</td>
</tr>
<tr>
<td>S. pyogenes</td>
<td>30</td>
<td>40</td>
<td>0.03</td>
</tr>
<tr>
<td>clinical isolate</td>
<td>16</td>
<td>0.25</td>
<td></td>
</tr>
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</table>

[(MICs) Minimum inhibitory concentration]

**Table 2**—Presence of bacteria in dermis (%), presence of inflammatory cells [neutrophil and mononuclear cell infiltration (%)] in papillary dermis, reticular dermis and subcutis in Groups 1 and 3

<table>
<thead>
<tr>
<th>Groups 1 and 3</th>
<th>Bacteria</th>
<th>Papillary dermis</th>
<th>Reticular dermis</th>
<th>Subcutis</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. 1.1 (n=10)</td>
<td>18.2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>18.2&lt;sup&gt;b&lt;/sup&gt;</td>
<td>9.1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>27.3&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>G. 1.2 (n=10)</td>
<td>63.6&lt;sup&gt;b&lt;/sup&gt;</td>
<td>63.6&lt;sup&gt;b&lt;/sup&gt;</td>
<td>27.3&lt;sup&gt;b&lt;/sup&gt;</td>
<td>27.3&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>G. 1.3 (n=10)</td>
<td>36.4&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>45.5&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>45.5&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>60.0&lt;sup&gt;bc&lt;/sup&gt;</td>
</tr>
<tr>
<td>G. 1.4 (n=11)</td>
<td>100.0&lt;sup&gt;c&lt;/sup&gt;</td>
<td>100.0&lt;sup&gt;c&lt;/sup&gt;</td>
<td>100.0&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>100.0&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>G. 3 (n=11)</td>
<td>0.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.0&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.0&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>p</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
<td>&lt;0.001</td>
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[Different letters in the same column determine statistical significance (P <0.05)]
Fig. 2.—Histopathological findings Group 2 and Group 3. (A) Negative control group (Group 3), normal skin histology, hematoxylin-eosin (H&E) (24th h); (B-C): infected control group (Group 2.4), (B) Subcorneal pustules (Blue star), intensive mononuclear cell infiltration (Yellow star), (80th h), (C) Gram stain. Subcutaneous tissue in the lumen of the coccoid bacteria clusters (24th h); and (D-F): Treatment group. (D) Olive Oil group (Group 2.2), Epidermal acanthosis, vacuolization in spinosa cells and intensive mononuclear cell infiltration is still severe (Blue star) edema in papillary dermis (Yellow star), (80th h), (E) Fucidic acid treated group (Group 2.3), (80th h), disappeared inflammatory reaction in the dermis but vacuolization in spinosa cells, hyperplasies (blue star) still present, (F) Ozonated olive oil treated group (Group 2.1), (80th h); epithelium and dermis normal view Bar x10 (A,D,E,F) x 40 (B) x 100 (C).

the highest which were infected controls. The lowest inflammatory responses were in the other groups (Fig. 1B and Fig. 2B).

Treatments started 24 h after infections in all groups. Healing started Group 1.1 and Group 2.1 at 40 h (Fig. 3 E & M). Healing completed in Group 1.1 at 72th h (Fig. 1 F & F), in Group 2.1, Group 1.3 and Group 2.3 at 80 h (Fig. 1E, Fig. 2 E & F, Fig. 3 J, P & T). In Groups 1.2, Groups 2.2, Groups 1.4 and Groups 2.4, the healing process did not occur up to 80 h (Fig. 1 B & D, Fig. 2 B & D, Fig. 3 D, H, N & R), (Tables 4 and 5).

Fig. 3.—Clinical findings Group 1, Group 2 and Group 3. Group 1 (A-J), (A) 0th h before bacterial inoculation in Group 1.4 (infected control group), (B-J) After the bacterial inoculation and treatment B: 24th hour in Group 1.4, C: 40th h in Group 1.4, D: 72nd h in Group 1.4, E: 40th h in Group 1.1 (ozonated olive oil treated group), F: 72nd h in Group 1.1, G: 40th h in Group 1.2 (olive Oil group), H: 72nd h in Group 1.2, 1: 40th h in Group 1.3 (fucidic acid treated group), J: 72nd h in Group 1.3, Group 2 (K-T), K: 0th h before bacterial inoculation in Group 2.4 (infected control group), (L-T) After the bacterial inoculation and treatment L: 24th h in Group 2.4, M: 40th h in Group 2.4, N: 80th h in Group 2.4, O: 40th h in Group 2.1 (ozonated olive oil treated group), P: 80th h in Group 2.1, Q: 40th h in Group 2.2 (olive oil group), R: 80th h in Group 2.2, S: 40th h in Group 2.3 (fucidic acid treated group), T: 80th hour in Group 2.3 Group 3 (U-Y), U: 0th h in Group 3 (control group), V: 24th h in Group 3, X: 40th h in Group 3, W: 72nd h in Group 3, Y: 80th h in Group 3.

Discussion

OZO is widely used for the therapeutic efficacy of the antimicrobial activity against bacteria, viruses and fungi. When ozone is used topically on the skin, it has been shown to have an antibacterial effect by some researchers. Lezcano et al. have investigated ozonated sunflower oil (OSO) on to the susceptibility to bacteria, according to the method of microdilution. They found Minimum inhibitory concentration (MIC) 9.5 µg/mL for the strain S. aureus ATCC 25923. Sechi et al. also found similar results.

On the contrary to the earlier studies, here, the OZO on to the susceptibility was determined to be 0.25 µg/mL as MICs value for the strain of S. aureus ATCC 25923 according to the microdilution method (Table 1). Sechi et al., who have also investigated ozonated sunflower oil on to the susceptibility to bacteria according to the method of microdilution found MICs value as 9.5 µg/mL for the S. pyogenes strain which was a clinical isolate.
Table 4—Clinical healing in Group 1 and Group 3 which made according to clinical score

<table>
<thead>
<tr>
<th>Groups</th>
<th>40th hour</th>
<th>48th hour</th>
<th>56th hour</th>
<th>64th hour</th>
<th>72nd hour</th>
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<tr>
<td></td>
<td>( \bar{x} \pm s )</td>
<td>( \bar{x} \pm s )</td>
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<td></td>
<td>(m(min-max))</td>
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<td>(m(min-max))</td>
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<td>(m(min-max))</td>
</tr>
<tr>
<td>G. 1.1 (n=10)</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.28 ( \pm 0.48 )</td>
<td>1.33 ( \pm 0.51 )</td>
<td>1.80 ( \pm 0.44 )</td>
<td>2.67 ( \pm 0.00 )</td>
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<tr>
<td>G. 1.2 (n=10)</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
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<tr>
<td>G. 1.3 (n=10)</td>
<td>0.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.16 ( \pm 0.40 )</td>
<td>1.80 ( \pm 0.44 )</td>
<td>2.00 ( \pm 0.00 )</td>
</tr>
<tr>
<td>G. 1.4 (n=11)</td>
<td>0.00 (0.00-0.00)</td>
<td>0.00 (0.00-0.00)</td>
<td>0.00 (0.00-0.00)</td>
<td>0.00 (0.00-0.00)</td>
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<tr>
<td>G. 3 (n=11)</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
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<td>(&lt;0.001 )</td>
<td>(0.001 )</td>
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</table>

[Different letters in the same column determine statistical significance \( P <0.05 \)]

Table 5—Clinical healing in Group 2 and Group 3 which made according to clinical score

<table>
<thead>
<tr>
<th>Groups</th>
<th>40th hour</th>
<th>48th hour</th>
<th>56th hour</th>
<th>64th hour</th>
<th>72nd hour</th>
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<tbody>
<tr>
<td></td>
<td>( \bar{x} \pm s )</td>
<td>( \bar{x} \pm s )</td>
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<tr>
<td></td>
<td>(m(min-max))</td>
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<td>(m(min-max))</td>
<td>(m(min-max))</td>
<td>(m(min-max))</td>
</tr>
<tr>
<td>G. 2.1 (n=10)</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.28 ( \pm 0.48 )</td>
<td>1.66 ( \pm 0.51 )</td>
<td>2.00 ( \pm 0.00 )</td>
<td>2.75 ( \pm 0.50 )</td>
</tr>
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<td>0.00 ( \pm 0.00 )</td>
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<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
</tr>
<tr>
<td>G. 2.3 (n=10)</td>
<td>0.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.33 ( \pm 0.51 )</td>
<td>1.25 ( \pm 0.50 )</td>
<td>2.00 ( \pm 0.00 )</td>
</tr>
<tr>
<td>G. 2.4 (n=11)</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
<td>0.00 ( \pm 0.00 )</td>
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<tr>
<td>G. 3 (n=11)</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
<td>1.00 ( \pm 0.00 )</td>
</tr>
<tr>
<td></td>
<td>( p &lt;0.001 )</td>
<td>(&lt;0.001 )</td>
<td>(&lt;0.001 )</td>
<td>(&lt;0.001 )</td>
<td>(0.001 )</td>
</tr>
</tbody>
</table>

[Different letters in the same column determine statistical significance \( P <0.05 \)]

Furthermore, in the present study, according to microdilution method, OZO on to the susceptibility was determined 0.25 µg/mL MICs value for the S. pyogenes strain also which was a clinical isolates (Table 1). The difference obtained in the present study may be due to ozonation time, peroxide, acid, iodine, p-anisidine and viscosity values.

Ozone breaks down protein in spite of the amino acid tryptophan residues and molecular weight difference. The implementation of this system, to treat skin infections can be more extensive and comprehensive caused by S. aureus infection. Lezcano et al. reported that OSO does not harm human epithelium and has significant antibacterial property. Diaz et al. have also reported that OSO and OZO, has an effective antibacterial activity against bacteria in human being.

In the present study, histopathological examination has shown the presence of bacteria in Group 1.1 (18.2%) and in Group 1.4, it was 100%. The OZO caused significant reduction \( P <0.001 \) in the presence of bacteria compared to Group 1.4 (Table 2). In Group 2.1, the presence of bacteria was 45.5% and in Group 2.4 it was 100% wherein OZO caused significant reduction \( P <0.001 \) in the presence of bacteria compared to Group 2.4 (Table 3). Our findings for both S. aureus and S. pyogenes were in agreement with literature findings mentioned above.

FA has been used to treat impetigo, infected wounds, abscess, infections of eye, skin and mucosa caused by Streptococcus, Staphylococcus and other gram positive bacteria. Rodrigues et al. made 6 mm wound on the skin of rat and infected the lesion with S. aureus ATCC 6538 strain. They applied OSO (35 mg) on to the wound. They also used neomycin (clostebol) for treatment and for comparison. They found OSO as effective as neomycin in terms of antimicrobial and anti-inflammatory effects. Our findings were in parallel with these findings, in Group 1.1 with concern to presence of bacteria; no statistical difference was observed compared to Group 1.3 (Table 2). Furthermore, Group 2.1 was not different statistically compared to Group 2.3 (Table 3).

Particularly on the wound, ozone causes an increase in the activity of growth factors.
Accordingly, angiogenesis, fibroblast activity and collagen synthesis increased. Platelet derived growth factor (PDGF), transforming growth factor (TGF-β) and vascular endothelial growth factor (VEGF) increase in fibroblast proliferation and collagen synthesis revealed acceleration of wound healing. Kim et al. have compared the local effects of OZO and OL that was used in wound healing in guinea pigs. In their study, they found that OZO to be more effective compared to the control group and OL group. They also reported that OZO caused increase in fibroblast proliferation and collagen synthesis revealing accelerated wound healing, which could be due to PDGF, TGF-β and VEGF at the wound site. Bactericidal effects of ozone, as well as its derivative, are the first phase of wound healing. Also, OZO reduces the presence of phagocytic cells, consequently reduces oxidative burst and increase the oxygen tension and supports dermal wound healing.

Apart from the antibacterial activity of ozone on the skin and wound repair, it also increases wound oxygenation properties. Curro et al. have also determined the effects of an innovative formulation of OZO with vitamin E acetate on male children with Balanitis xerotica obliterans (BXO). OZO with vitamin E showed effective results in children affected by BXO by reducing the inflammatory process and stimulating mechanisms for tissue regeneration of the foreskin. This indicates that in the present study; when groups compared with concern to recovery; between Group 1.1 and Group 1.4 at 40, 48, 56, 64, 72 and 80 h after infection, recovery were also better (P <0.001) in Group 1.1 compared to Group 1.4 likewise, between Group 1.1 and Group 1.2 at 40, 48, 56, 64 and 72 h after infection, recovery were also better (P <0.001) in Group 1.1 compared to Group 1.2 (Table 4).

When groups were compared with concern to recovery; between Group 2.1 and Group 2.4 at 40, 48, 56, 64, 72 and 80 h after infection, the recovery were also better (P <0.001) in Group 2.1 compared to Group 2.4. Likewise, between Group 2.1 and Group 1.4 at 40, 48, 56, 64 and 72 h after infection, recovery were also better (P <0.001) in Group 2.1 compared to Group 2.2 (Table 5).

Furthermore, with concern to dermal lesions, in papillary dermis and subcutis, inflammatory cells was higher (P <0.001) in Group 1.4 compared to Group 1.1. But, statistical importance were not observed in reticular dermis both in Group 1.4 and Group 1.1 and also more number of inflammatory cells were determined (Table 2). Additionally, with concern to dermal lesions, in reticular dermis, inflammatory cells was higher (P <0.001) in Group 2.4 compared to Group 2.1. But, statistical importance was not observed in papillary dermis and subcutis in both Group 2.4 and Group 2.1 and also more number of inflammatory cells were determined (Table 3). This is considered to suggest that OZO acting by both antibacterial and anti-inflammatory effects. These support Valacchi et al. findings.

Estrela et al. reported that ozone is effective on Staphylococcus. Furthermore, Schulz had proven the effect of ozone on burn and wound who was investigated on animals. In addition, Schulz found that ozonated oil had 40% better healing effect applied for 11 days on wound compared to other pomades. Song et al. were determined the microbial effects of topical ozone treatment on S. aureus and MRSA. After their study the inhibition zone diameters of OZO for S. aureus and MRSA were 17 and 13 mm, respectively, which were significantly larger than the control group. And also all cases of skin S. aureus and MRSA infection were completely healed after ozone treatment.

In the present study, according to microdilition method and disc diffusion test, although FA were more effective against S. pyogenes compared to OZO (Table 1), in Group 1.1; healing started at 40th h (Fig. 3E) and complete recovery observed at 72 h after infection (Fig. 1F & 3), but in Group 1.3, healing started at 48th h and complete recovery seen at 80 h after infection. In Group 1.3 at 72 h after infection, absence of inflammatory reactions and presence of hyperkeratosis on the epithelial tissue were observed (Fig. 1E).

Although FA were more effective against S. aureus compared to OZO in disc diffusion test (Table 1), in Group 2.1, healing started at 40th h (Fig. 3O) and complete recovery observed at 80 h after infection (Fig. 2F and 3P), but in Group 2.3, healing started at 48th h and complete recovery seen at 80 h (Fig. 3T) after infection, even though, absence of inflammatory reactions and presence of hyperkeratosis were observed on the epithelial tissue (Fig. 2E). As a result, in ozone applied groups, healing started earlier, and hence complete recovery observed earlier. The reasons could be due to; in addition to antibacterial and anti-inflammatory effect
of ozone, its increasing effect on collagen synthesis, fibroblast proliferation, epithelisation and vascularisation that accelerate healing process.

**Conclusion**

As a result, in the present study, experimental skin infections with *Streptococcus pyogenes* and *Staphylococcus aureus* were made and the effects of OZO in the treatment of the infection were evaluated and results were compared with other treatment methods. After occurrence of the infection, 0.1 g OZO had antimicrobial and anti-inflammatory effect and observed to have healing activity at 72 and 80 h after infection. Furthermore, the effects of OZO were comparable with the effects of FA which is well known and its activity has been proven in the skin infections with Staphylococcus and Streptococcus. Additionally, as far as known microorganisms has no resistance development against ozone and has no important side effects.

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