Antimicrobial efficacy of 2.5% sodium hypochlorite, 2% chlorhexidine, and ozonated water as irrigants in mesiobuccal root canals with severe curvature of mandibular molars

Sergio Luiz Pinheiro¹, Caio Cesar da Silva¹, Lucas Augusto da Silva¹, Marina P. Cicotti¹, Carlos Eduardo da Silveira Bueno¹, Carlos Eduardo Fontana¹, Leticia R. Pagrion¹, Natália P. Dalmora¹, Thaís T. Daque¹, Francisco UF de Campos²

ABSTRACT

Objective: The aim of this study is to evaluate the antimicrobial efficacy of 2.5% sodium hypochlorite, 2% chlorhexidine, and ozonated water on biofilms of Enterococcus faecalis, Streptococcus mutans, and Candida albicans in mesiobuccal root canals with severe curvature of mandibular molars. Materials and Methods: This was an experimental ex vivo study in microbiologic laboratory. Sixty mesiobuccal root canals with severe curvature of mandibular molars were contaminated with standard strains of E. faecalis, S. mutans, and C. albicans. The specimens were randomly divided into four groups (n = 15) according to irrigating solution: SH: 2.5% sodium hypochlorite; CH: 2% chlorhexidine; O₃: ozonated water; and control: double-distilled water. The mesiobuccal root canals of all groups were instrumented with the WaveOne Gold Primary reciprocating system. Three cycles of instrumentation with three short in-and-out brushing motions were performed: (1) in the coronal third, (2) in the middle third, and (3) in the apical third of the canal. A ProGlider file was used before the first cycle. Statistical Analysis: Statistical analysis was performed using one-way analysis of variance followed by Tukey’s multiple comparison test. Samples were collected for viable bacterial counts before and after instrumentation. Results: All groups showed significant biofilm reduction after irrigation (P < 0.01). After instrumentation, sodium hypochlorite (98.07%), chlorhexidine (98.31%), and ozonated water (98.02%) produced a significantly reduction in bacterial counts compared with double-distilled water (control, 72.98%) (P < 0.01). Conclusion: All irrigants tested in this study showed similar antimicrobial activity. Thus, ozonated water may be an option for microbial reduction in the root canal system.

Key words: Endodontic treatment, irrigation, reciprocating system

INTRODUCTION

Disinfection of the root canal system is essential for successful endodontic treatment.[¹] Root canals are usually infected by multiple bacteria.[²⁻⁷]
In general, 2.5% sodium hypochlorite and 2% chlorhexidine are used as irrigating solutions in endodontics. Sodium hypochlorite has been used as an irrigant because of its broad antimicrobial spectrum and ability to dissolve necrotic tissue remnants while 2% chlorhexidine has been used as an irrigant due to its broad antimicrobial activity, substantivity, and low cytotoxicity. An alternative that is currently being considered for disinfection of the root canal system is ozone therapy. Ozone is applied to oral tissues in the forms of ozonated water, ozonated oil, and oxygen/ozone gas either alone or in combination to treat dental disease. Ozone has a high oxidation potential, being 1.5 times more effective than chloride as an antimicrobial agent against several microorganisms, and it can also stimulate blood flow and immune response. An increasing number of irrigants have been proposed in an attempt to achieve optimal irrigation. However, none of the available irrigants showed all the necessary requirements.

No published studies have assessed microbial load reduction in the root canal system with a combination of reciprocating instrumentation and irrigation with ozonated water. Reciprocating instrumentation shapes the root canal system rapidly, thus enabling high-flow, large-volume irrigation, while the biocompatibility and properties of ozonated water make it an interesting alternative irrigant. Within this context, the present study sought to evaluate the antimicrobial efficacy of 2.5% sodium hypochlorite, 2% chlorhexidine, and ozonated water on biofilms of Enterococcus faecalis, Streptococcus mutans, and Candida albicans in mesiobuccal root canals of mandibular molars. The null hypothesis was that equivalent reductions in biofilm formation would be achieved with the use of 2.5% sodium hypochlorite, 2% chlorhexidine, and ozonated water as irrigants.

**MATERIALS AND METHODS**

**Sample selection**

Sixty permanent mandibular molars were selected among teeth donated by patients at dental clinic. Written informed consent was obtained from all patients before donation. The study was approved by Research Ethics Committee (protocol number: 1.841.252).

The sample size was calculated based on a pilot data set using analysis of variance (ANOVA) with statistical power of 0.80 and alpha of 0.05. A sample size of 15 specimens per group was required.

The inclusion criteria were as follows: complete root formation, no previous of endodontic treatment, no pathological external and/or internal root resorption, no root cracks or fractures, root length ≥ 15 mm, distinct root canals for the mesiobuccal and mesiolingual canals, root curvature 25°–40° (severe) (Schneider 1971), and anatomical canal diameter compatible with a size #15 K-file.

**Tooth preparation**

The teeth were radiographed in the buccolingual direction, and the degree of curvature of the mesiobuccal root canal was determined. The crowns were removed with double-sided diamond discs (Microdont, São Paulo, SP, Brazil), and root length was standardized to 15 mm. The mesial root length was determined with a digital caliper (Starrett, Itu, SP, Brazil). The orifice of the mesiolingual and distal canals was sealed with light-curing resin (Z250 XT; 3M, Deutschland, Germany).

The working length was determined visually by inserting a size #15 (Dentsply Maillefer, Ballaigues, Switzerland) into the root canal until its tip was visible at the apical foramen under operating microscope visualization (8X). The file was then withdrawn 1 mm to determine the working length. Before contamination, the canals were manually instrumented with size #10 and #15 K-files (Dentsply Maillefer, Ballaigues, Switzerland) until reaching the working length and irrigated with 5 mL of distilled water (Dinâmica, Campinas, SP, Brazil). The apical foramen and the external surfaces of all roots were sealed. The specimens were sterilized in an autoclave (Sercon, Mogi das Cruzes, SP, Brazil) at 121°C for 15 min.

The root canals were contaminated with standard strains of E. faecalis (ATCC 29212; LabCenter, Campinas, SP, Brazil), S. mutans (25175; LabCenter, Campinas, SP, Brazil), and C. albicans (10231; LabCenter, Campinas, SP, Brazil)). The bacterial suspension was prepared in a tube containing 10 mL of sterile saline (Dinâmica, Campinas, SP, Brazil), matched to a 10 McFarland standard.

A 20-µL aliquot of the final suspension was injected into each root canal using a 0.3-cc insulin syringe (Ultrafine BD, São Paulo, SP, Brazil). Specimens were stored in 24-well cell culture plates (CoStar, New York, NY, USA) in an incubator (Fanem Ltda, São Paulo, SP, Brazil) at 37°C in a 5% CO₂ atmosphere for 21 days.
The viability and purity of the microorganisms within the canals were checked weekly by random sampling of two specimens using sterile paper points.\[14,16,19\] Bacterial samples were collected using sterile paper points (Dentsply Maillefer, Ballaigues, Switzerland) compatible with the anatomical diameter of the root canal before and after instrumentation.\[14,19\] The paper point was inserted into the canal for 30 s and immediately placed in a test tube containing 5 mL of BHI broth (Acumedia Manufacturers, Lansing, MI, USA).

**Canal instrumentation**

Instrumentation was performed using the WaveOne Gold (WOG) single-file reciprocating system (Dentsply Maillefer, Ballaigues, Switzerland), powered by an X-Smart Plus electric motor (Dentsply Maillefer, Ballaigues, Switzerland) set to operate in the WaveOne mode, with a slight pressure in the apical direction of up to 3–4 mm.

Instrumentation was performed according to the manufacturer’s instructions. A size #15 (Dentsply Maillefer, Ballaigues, Switzerland) was used to confirm the path of the canal to the foramen. The glide path was expanded by at least 0.16 mm using a ProGlider file (Dentsply Maillefer, Ballaigues, Switzerland) until the working length was reached. Before the first cycle of WOG file, a ProGlider file (Dentsply Maillefer, Ballaigues, Switzerland) was applied in one in-and-out brushing motion at a speed of 300 rpm with light apical pressure, set at 2 Ncm for torque control. The Primary WOG 25/.07 file was the file that best fitted to the canal and was used to the full working length. The Primary 25/.07 and ProGlider files were single use.

Three cycles of instrumentation were performed. Each cycle consisted of three short in-and-out brushing motions.\[17\]

The canals were irrigated using a 5-mL plastic syringe (Ultradent Products Inc., South Jordan, UT, USA) with a 0.55 × 20 mm hypodermic needle (24G; Ultrafine BD, São Paulo, SP, Brazil), inserted up to the middle third of the canal (10 mm). All canals were instrumented by the same operator.

The specimens were randomly divided (www.random.org.br) into four groups (n = 15) according to irrigating solution: SH: 2.5% sodium hypochlorite (compounded – Farmácia Art Med, Jundiaí, SP, Brazil); CH: 2% chlorhexidine solution (compounded – Farmácia Art Med, Jundiaí, SP, Brazil); O₃: ozonated water (40 µg/mL); and control: double-distilled water.

In all groups, before instrumentation, samples were collected for viable bacterial counts. After use of the ProGlider file (Dentsply Maillefer, Ballaigues, Switzerland) and during instrumentation (each cycle), canals were irrigated with the solutions corresponding to groups SH, CH, O₃, and control, for 3 min, using 5 mL (10) of the irrigating solution, for a total of 20 mL irrigant. After this protocol, a new sample collection was performed for viable bacterial counts.

**Preparation of ozonated water**

An ozone generator was used (Philozo, Camboriu, SC, Brazil). Sterile double-distilled water was cooled at 14°C and kept under refrigeration until use. The ozone gas was adjusted in the generator to the concentration required for this experimental model (40 µg/mL).\[20\]

**Specimen culture**

The specimens were diluted, seeded, and cultured. Total viable bacterial count was determined using a colony counter. Counts were given as colony-forming units (CFU).

**Statistical analysis**

Antimicrobial efficacy was evaluated as bacterial growth (in CFU/mL) and percentage biofilm reduction. Percentage reduction of biofilm was calculated using the following formula:

\[
\text{Percentage reduction} = \frac{\text{Biofilm count before instrumentation} - \text{Biofilm count after instrumentation}}{\text{Biofilm count before instrumentation}} \times 100\%
\]

BCB, biofilm count before instrumentation; BCA, biofilm count after instrumentation.

Results were analyzed using Biostat 4.0 (Sociedade Civil Mamirauá, Belém, Pará, Brasil). All data obtained in CFU/mL were log₁₀ transformed. The log₁₀ transformed and percentage biofilm reduction data were tested for normality using the Shapiro–Wilk test. All data were found to be normally distributed and therefore analyzed using one-way ANOVA followed by Tukey’s multiple comparison test at the 1% significance level.

**RESULTS**

Bacterial counts obtained before and after root canal instrumentation were compared and all groups
Table 1: Mean (standard deviation) values of biofilm counts obtained before and after irrigation with 2.5% sodium hypochlorite, 2% chlorhexidine, ozonated water, and double-distilled water (control) (log_{10} colony-forming units/mL)

|                | Before | After | Before | After | Before | After | Before | After |
|----------------|--------|-------|--------|-------|--------|-------|--------|-------|
| 2.5% SH        | 4.60 (0.09) | 2.88 (0.11) | 4.57 (0.13) | 2.77 (0.14) | 4.56 (0.10) | 2.79 (0.11) | 4.59 (0.07) | 4.02 (0.08) |
| P              |         |       |        |       |        |       |        |       |

ANOVA (with Tukey test): <0.0001

Different uppercase or lowercase letters: P<0.01. SH: 2.5% sodium hypochlorite, CH: 2% chlorhexidine, O₃: ozonated water, ANOVA: Analysis of variance, SD: Standard deviation

Table 2: Percentage biofilm reduction after irrigation with 2.5% sodium hypochlorite, 2% chlorhexidine, ozonated water, and double-distilled water (control) (%)

|                | Before | After | Before | After | Before | After | Before | After |
|----------------|--------|-------|--------|-------|--------|-------|--------|-------|
| 2.5% SH        | 98.07 (0.44) | 98.31 (0.48) | 98.02 (0.60) | 72.98 (3.02) |
| P              |         |       |        |       |        |       |        |       |

ANOVA (with Tukey test): <0.0001

Different uppercase letters: P<0.01. SH: 2.5% sodium hypochlorite, CH: 2% chlorhexidine, O₃: ozonated water, ANOVA: Analysis of variance, SD: Standard deviation

DISCUSSION

The experimental procedure of biofilm analysis of this study was based on the studies of Machado et al., Cord et al., Ghinzelli et al.,[21] and Pinheiro et al.[22] The culture method to evaluate antibacterial activity was based on the study of Alves et al.,[23] who indicated that the culture method can be used effectively in ex vivo studies to test the antibacterial efficacy of treatment protocols and it is equivalent to polymerase chain reaction. Serial dilutions and sowing in BHI agar for counting CFU were performed according to Le Goff et al.[24]

The microbial reduction produced by 2.5% sodium hypochlorite (98.07%) may be explained based on the observations of Rutala and Weber,[25-27] who suggested that, when combined with water, sodium hypochlorite produces hypochlorous acid, which contains active chlorine. Chlorine exerts its bactericidal action through the irreversible oxidation of sulfhydryl groups of essential bacterial enzymes, disrupting the metabolic function of bacterial cells.[8] Sodium hypochlorite may also have a deleterious effect on bacterial DNA, which involves the formation of chlorinated derivatives of nucleotide bases. In addition, there are reports that sodium hypochlorite may induce bacterial membrane disruption.[28] According to Estrela et al.,[29] the tissue dissolution capacity of sodium hypochlorite is based on the reaction with fatty acids and lipids, which are transformed into soap and alcohol. Hypochlorous acid and hypochlorite ions lead to amino acid degradation and hydrolysis.

Chlorhexidine reduced bacterial counts by 98.31% after instrumentation probably due to its ability to adsorb to dentin, acting on bacterial cell walls and cytoplasmic membrane, resulting in the loss of osmotic balance and leakage of intracellular material.[30] Its antimicrobial activity has residual effects ranging from 7 days[31] to 12 weeks. Chlorhexidine substantivity is facilitated by its viscosity, which keeps the solution in contact with the canal walls and dentinal tubules.[32] According to Estrela et al.,[29] the antimicrobial activity of chlorhexidine may be explained by the interaction between its cationic nature and the anionic compound on the bacterial surface (phosphatase groups of teichoic acids in Gram-positive bacteria and lipopolysaccharides in Gram-negative bacteria). According to Gomes,
et al.,\textsuperscript{[32]} low concentrations of chlorhexidine allow low-molecular-weight substances to be released, resulting in bacteriostatic effects. At high concentrations, it has bactericidal effects due to precipitation and/or coagulation of the cytoplasm.

The microbial reduction promoted by ozonated water (98.02\%) is in agreement with the study of Nogales et al.,\textsuperscript{[28]} who stated that ozone was a promising agent for endodontic treatment. Ozone’s oxidizing power is exerted specifically on polyunsaturated fatty acids of the bacterial membrane, increasing oxygen delivery to tissues and modulating the immune system, thereby improving and accelerating tissue repair. According to Goztas et al.,\textsuperscript{[33]} the advantages of ozone in the aqueous phase include lack of mutagenicity, rapid microbial effects, and ease of handling. The authors also reported that ozonated water shows no cytotoxicity and is highly biocompatible compared with other antimicrobial agents. Ozone may have other clinical applications in addition to root canal irrigation, unlike hypochlorite and chlorhexidine, which have no other therapeutic uses in dentistry; this would justify the financial investment in ozonation equipment.

The control group also showed a significant microbial reduction after instrumentation (72.98\%). This result was probably due to instrumentation using the WOG reciprocating system. The design and gold wire alloy in the system make the file more flexible and more efficient in terms of cutting efficiency compared with other nickel–titanium systems. These features increase the capacity to auger debris out of the coronal third of the canal.\textsuperscript{[34]}

**CONCLUSION**

We believe that the properties of ozone, in aqueous phase, render it an important disinfectant in endodontic treatment. In this respect, in addition to its oxidative potential to induce the destruction of bacterial cell walls and cytoplasmic membrane, ozone acts on glycoproteins, glycolipids, and amino acids, inhibiting the enzymatic control system of the cell. This results in increased membrane permeability, which allows ozone molecules to easily penetrate the cell and induce microbial lysis. Ozonated water has been demonstrated in several studies as a solution of good applicability with quite promising results, and the present findings confirm what has been reported in the literature. Moreover, the ozone generator is extremely cost-effective and easy to operate and may be a valuable tool for clinical use by endodontists.

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**Conflicts of interest**

There are no conflicts of interest.

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