Antibacterial Power of Sodium Hypochlorite Combined with Surfactants and Acetic Acid

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The aim of this ex vivo study was to compare the antibacterial power of 1% NaOCl with 1% acetic acid, 5.25% NaOCl and two commercially available NaOCl modified with surfactants in bovine root dentin. A total of 120 dentin tubes prepared from intact bovine incisors were infected for 21 days with Enterococcus faecalis and randomly divided into six groups as follows: 5.25%NaOCl; Hypoclean; Chlor-Xtra; 1% NaOCl with 1% acetic acid; infected dentin tubes (positive control); and sterile dentin tubes (negative control). At experimental times of 0, 7, 14, 21 and 28 days, dentin chips were collected using sequential round burs with increasing diameters in separate test tubes containing 3 mL of freshly prepared BHI. Statistical analysis were performed using parametric methods (one-way ANOVA, and Bonferroni’s multiple comparisons test, α=0.01). After culturing, the number of colony-forming units (CFU) was counted. All the NaOCl solutions showed small number of CFU over 28 days. ChlorXtra and Hypoclean had the smallest number of CFU at all times with greater antimicrobial efficacy than 5.25% NaOCl and 1% NaOCl solution with 1% acetic acid.

Introduction

Bacteria and their byproducts have been long recognized as the primary etiological agents of periapical diseases. The root canal treatment prognosis is influenced by the presence of bacteria in the root canal system (RCS) at the time of filling. Mechanical preparation and disinfection procedures during root canal treatment should completely eradicate all microorganisms in the main root canal and surrounding tooth structure or at least reduce bacterial populations to a level below that necessary to induce or sustain disease. Failure to heal can usually be a result of recontamination of the RCS as well as of residual infection (1).

In fact, microorganisms can penetrate areas that are difficult to clean mechanically and not readily accessible to antimicrobial solutions such as isthmus, ramifications and lateral or accessory canals. Moreover, the dentin wall of an infected root canal can harbor bacteria or bacterial metabolic by-products (1,2). An intense bacterial penetration to about 300 µm deep into the dentinal tubules has been reported (2,3); penetration depths >500 µm (4) and high prevalence in cases of persistent apical periodontitis (5) have been reported for Enterococcus faecalis. This species is also known to extend even more deeply, penetrating 800–1000 µm into the dentinal tubules after 3 weeks of incubation (4). Additionally endotoxins from Gram-negative bacteria can penetrate dentinal tubules approximately to a depth of 300–500 µm (6,7).

Nonetheless, successful microbial elimination from the infected RCS is still challenging. The ability of NaOCl to inactivate bacteria is essential, but it must reach areas of difficult access and deeper penetration into dentinal tubules. The greatest penetration of NaOCl into dentinal tubules has been reported as being 300 µm with a 6% solution for 20 minutes at 45 °C (8), due to its high surface tension (9). Increasing the concentration of NaOCl improved the penetration depth of its antibacterial action, but complete eradication of bacteria from the dentinal tubules cannot be achieved, especially beyond the first 300 µm from the pulp-dentin junction (10). Further, irrigants should ideally have adequate antibacterial action against microorganisms and their products without damaging host tissues (11,12). However, NaOCl does not exert any residual antimicrobial activity (12) so that the recolonization of persistent microorganisms would not be prevented. Thus, infected or contaminated dentin might serve as a potential source of persistent apical periodontitis.

A possible way of improving the bactericidal efficacy of the disinfecting solutions is to incorporate different detergents as surface active agents, enhancing their wetting properties (13,14). High wettability is supposed to enable deeper penetration into the dentinal tubules of the antibacterial solutions. Two commercially available NaOCl-based irrigants modified with surfactants, Hypoclean (Ogna Laboratori Farmaceutici, Muggiò, Italy) and Chlor-Xtra (Vista Dental Products, Racine, WI, USA), showed surface tension values (29.13 mJ/m² and 33.14 mJ/m², respectively) significantly lower than that of regular 5.25% NaOCl (48.90 mJ/m²) (9). In sequence to this line of
reasoning, a recent assay showed the superior wettability of Hypoclean and Chlor-Xtra over dentin in comparison to regular 5.25% NaOCl: NaOCl solutions with lowered surface tension spreaded immediately on dentin surface, yielding a zero-degree contact angle (14). Hypoclean demonstrated a more effective antibacterial action against *E. faecalis* than 5.25% NaOCl at all experimental periods (7, 14, 21, 28 days) in bovine root dentin and displayed no bacterial growth at the two-first cultures after treatment (12). Low surface tension could improve the antimicrobial efficacy of NaOCl by increasing its penetration ability into inacessible areas of the RCS and into dentinal tubules. Nonetheless, for the 6% solutions a modification of the wetting properties appeared to have no effect on their bactericidal power (15). Thus, the available evidence remains inconclusive regarding the effects of reducing the surface tension of NaOCl solutions on antimicrobial efficacy.

The bactericidal action of hypochlorite’s can also potentially be enhanced by lowering their pH, by its association with acids (16) and by stabilizing them in low pH (17-19) (weak acidification of hypochlorite) (17). Acetic acid resulted a preferable choice because of its own antibacterial activity and safety (18). The NaOCl solution at 5.25% solution was shown to be significantly more effective against *E. faecalis* than weaker NaOCl solutions within dentinal tubules (20) also in mixed culture. Nonetheless, lower concentrations of NaOCl tested significantly reduced the microorganisms within dentinal tubules and in the main root canal.

The aim of this study was to assess the in vitro antibacterial power of 1% NaOCl in combination with 1% acetic acid, 5.25% NaOCl, Hypoclean and Chlor-Xtra in bovine root dentin infected with *E. faecalis*. The null hypothesis is that these solutions have a similar antibacterial efficacy.

**Materials and Methods**

The method of this study was based on the procedure previously described by Haapasalo and Orstavik (4). Sixty extracted bovine maxillary anterior teeth were used. All teeth were kept in 0.5% NaOCl solution for up to 7 days. The crown and apical third were removed from each tooth with a rotary diamond saw at 1000 rpm (Isomet Plus precision saw; Buehler, IL, USA) under water cooling. Cementum was removed by using polish paper (Ecomet 3, variable-speed grinder-polisher; Buehler), which resulted in a centre-holed piece of root dentin with a 6-mm outer diameter. The remaining piece of each tooth was then cut into 4-mm thick slices with a diamond saw as above. The root canals of the 4-mm blocks were enlarged (standardized) with an ISO 023 slow speed round bur. In order to prevent dehydration, all teeth and dentin slices were preserved in vials containing tap water during the procedures.

Each dentin block (n=120) was individually treated with 5.25% NaOCl and 17% ethylene diamine tetraacetic acid (EDTA) (pH 7.2) placed in an ultrasonic bath for 10 min, and irrigated under tap water for 1 h to remove the smear layer. The specimens were then placed in brain heart infusion (BHI) broth (Oxoid, Basingstoke, UK) and autoclaved. To monitor the efficacy of the sterilization they were then kept in an incubator at 37 ºC for 24 h. One bottle of Chlor-Xtra (Vista Dental Products) was purchased shortly before the beginning of the project; the solution was immediately transferred into an amber glass bottle, kept at 4C (following the recommendations of the manufacturer) and brought to room temperature before use. One percent and 5.25% NaOCl solutions were freshly prepared by diluting a stock solution of 15% NaOCl (Ogna Laboratori Farmaceutici, Muggiò, Italy) in distilled water.

Hypoclean solution was also freshly prepared by mixing 5.25% NaOCl and surfactants. A total of 120 specimens were randomly divided into 6 groups as follows: Group 1: 5.25% NaOCl (Ogna) (n=25); Group 2: Hypoclean (5.25%NaOCl modified with cetrimide and polypropylene glycol) (Ogna) (n=25); Group 3: Chlor-Xtra (<6% NaOCl modified with Triton X100, proprietary surface modifiers and alkylating agents) (Vista Dental Products) (n=25); Group 4: 1%NaOCl (Ogna) with 1% acetic acid (n=25); Group 5: negative control (sterile dentin tubes) (n=10); Group 6: positive control (infected dentin tubes) (n=10). Mean active chlorine content (3 readings) of all solutions was determinate by titration; pH was determined by using a digital pH meter (Metertom model 827 pH LAB; Metrohm, Varese, Italy) 5.25%NaOCl - pH 12.6 and % chlorine = 5.2 ± 0.01; 1% NaOCl + acetic acid - pH 3.5 and % chlorine = 0.023 ± 0.06; Hypoclean – pH 12.4 and % chlorine 5.1 ± 0.01; Chlor-Xtra – pH 12.7 and % chlorine 5.42 ± 0.02.

Isolated 24-h colonies of pure cultures of *E. faecalis* (ATCC 29212) were suspended in 5 mL of BHI. The bottles containing each specimen in Groups 1, 2, 3, 4 and 6 were opened in a laminar flow chamber. Two milliliters of sterile BHI were removed with sterile pipettes and replaced with 2 mL of bacterial inoculum. The bottles were closed and kept at 37 ºC for 28 days, with the replacement of 1 mL of contaminated BHI for 1 mL of freshly prepared BHI every 2 days, to avoid medium saturation. After the contamination period, the bacterial growth was verified with a histological technique (Brown and Brenn staining method) (Fig. 1).

Then, each specimen was removed from its bottle under aseptic conditions, and the root canal was irrigated with 5 mL of sterile saline and dried with sterile paper points. In order to prevent contact of the medicament with the external surface, the outer surface of the specimens was covered with two layers of nail polish. Thereafter, using
decontaminated sticky wax, specimens were fixed at the bottom of wells of 24-well cell culture plates, which also obliterated the apical surface of the root canal. Finally, the irrigating solutions were delivered into the canal lumen with sterile 3-mL plastic syringes and 27-gauge needles until the dentin tubes were totally filled. Solutions were removed using sterile paper points 10 min after placement into the lumen. The specimens were then incubated at 37 °C for 28 days to maintain humidity. At experimental times of 0, 7, 14, 21 and 28 days, dentin chips were removed from the root canals with sequential sterile low-speed round burs with increasing diameters of ISO sizes: 025, 027, 029, 031 and 033, respectively (Fig. 2). Each bur removed approximately 0.1 mm of dentin around the canal. The powder dentin samples obtained with each bur were immediately collected in separate test tubes containing 3 mL of freshly prepared BHI. Thereafter, 100 μL from each test tube was cultured on blood agar. Growing colonies were counted and recorded as colony-forming units (CFU).

Statistical analysis was performed using parametric methods (one-way analysis of variance). Post hoc pair-wise comparisons were performed using Bonferroni’s multiple comparisons. The level of significance was set at p<0.01. Statistical analysis was carried out using IBM SPSS Statistics 20.0 software (IBM Corp, Armonk, NY, USA).

Results

The number of CFU obtained from five consecutive dentinal layers at each experimental time is presented in Table 1. The number of CFU in all four experimental groups

Table 1. Mean numbers and standard deviations of colony forming units (CFU) for the different groups at each experimental time.

<table>
<thead>
<tr>
<th>Period (days)</th>
<th>Solution</th>
<th>5.25% NaOCl (n=25)</th>
<th>Hypoclean (n=25)</th>
<th>Chlor-Xtra (n=25)</th>
<th>1% NaOCl +AA (n=25)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.43 ± 0.30 a</td>
<td>0.00 ± 0.00 b</td>
<td>0.00 ± 0.00 b</td>
<td>5.52 ± 0.69 c</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>18.38 ± 4.29 d</td>
<td>7.41 ± 1.58 e</td>
<td>10.82 ± 2.00 f</td>
<td>27.59 ± 2.72 g</td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>35.29 ± 5.82 h</td>
<td>18.95 ± 1.12 i</td>
<td>19.00 ± 3.01 j</td>
<td>43.73 ± 3.27 j</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>67.47 ± 7.01 k</td>
<td>31.79 ± 4.36 l</td>
<td>29.34 ± 4.16 l</td>
<td>65.71 ± 3.08 l</td>
<td></td>
</tr>
<tr>
<td>28</td>
<td>95.08 ± 3.71 m</td>
<td>48.38 ± 3.54 n</td>
<td>48.05 ± 4.16 o</td>
<td>96.62 ± 3.84 o</td>
<td></td>
</tr>
</tbody>
</table>

Different superscript letters indicate statistically significant differences among groups (p<0.01). AA: 1% acetic acid.

n: dentin tubes.
was minimum after treatment.

The positive control group showed viable bacteria at all experimental times, which indicated the efficiency of the method. The average number of CFU in each specimen of this group was 120. In contrast, the negative control group showed no viable bacteria at all experimental times. Acidic NaOCl solution showed the highest number of CFU immediately after treatment (p<0.001) and at days 7 (p<0.001) and 14 (p<0.001). However, at days 21 and 28 antibacterial effectiveness against E. faecalis of 1% NaOCl added with 1% acetic acid versus 5.25% NaOCl was not significantly different (p<0.01). Hypoclean and Chlor-Xtra showed the most effective antibacterial action at all five experimental times (p<0.001). The differences for Hypoclean versus Chlor-Xtra were not statistically significant (p>0.01) except for results at day 7 (p<0.001). For all test solutions the intra-group comparisons among findings at the different experimental times showed significant differences (p<0.001). When surfactants were added to 5.25% NaOCl its antibacterial effectiveness increased significantly at all experimental times (p<0.001) with a 38.9 % additional decrease of the CFU recovery at day 28.

Discussion

The use of teeth or dentin blocks in in vitro and ex vivo studies of root canal disinfection is an effort to bring the experimental conditions much closer to the in vivo reality of the root canal than direct contact tests with planktonic bacteria. The experimental model used in the present investigation was proposed by Haapasalo and Orstavik (4). The time of incubation with the bacteria, the frequency of nutrient exchange and the substrate (surface to attach to) have been taken into account. However, the extent of bacterial growth into the dentin tubules was verified (positive control group).

When 1% NaOCl combined with 1% acetic acid, regular 5.25% NaOCl and 5.25% or <6% NaOCl combined with surfactants have been maintained on infected root surface with E. faecalis during 10 min, 40–79% CFU were recovered at days 28. These results are in accordance with those of a previous in vitro study on bovine root dentin (12).

The chemical characteristics of NaOCl have made it the most used and studied endodontic irrigant worldwide, but no consensus exists regarding NaOCl ideal concentration, which ranges from 0.5% to 6%. High concentration hypochlorites have shown significantly stronger antibacterial efficacy against E. faecalis than low concentration hypochlorites into the dentinal tubules (15). In addition, decreased antibacterial properties have been reported for 5.25% NaOCl when it is diluted (17). It is well known that increasing the chlorine concentration of NaOCl solutions increases their cytotoxicity (11). The antimicrobial activity of low concentration NaOCl can potentially be enhanced, with less aggressive action to the periapical tissues, by lowering the pH of the solution (16), because of it depends on the HOCl concentration (21). Surely, larger volumes of irrigant and frequent replenishing would be required with weaker NaOCl solutions to maintain their antibacterial effectiveness. A decrease in pH increases in the NaOCl solution the concentration of chlorine available as HOCl and thus its antimicrobial effectiveness. Kuroiwa et al. (18) indicated acetic acid as a preferable acid for safe preparation of acidic hypochlorite solution. Mixtures with other acids consume available chlorine and reduce capacity for bactericidal activity of the solution; acetic acid had no effect on available chlorine. In the present study a combination of 1%NaOCl with 1% acetic acid was tested with the aim of verifying an improved antibacterial activity of low concentration NaOCl with lowered pH value. Acidic NaOCl solution showed the highest number of CFU immediately after treatment and at days 7 and 14, with a 50,16 % and 75,91% recovery of CFU at days 21 and 28 respectively. Kuga et al. (21) evaluated the penetration of 2.5% NaOCl combined with 17% EDTA, 1% citric acid, and 1% peracetic acid into dentin tubules. The combination of NaOCl with acid solutions did not increase its penetration depth into root dentin. According to this finding, the results of the present study showed a recovery of CFU in all evaluated periods. Moreover, differences among the antibacterial efficacy of 1% NaOCl added with 1% acetic acid versus 5.25% NaOCl were not significant at days 21 and 28. In the present study, % chlorine value of 0.023 ± 0.06 and pH value of 3.5 were determinate by titration for 1% NaOCl added with 1% acetic acid. It is important to point out that the antibacterial effect of NaOCl is directly related to the amount of HOCl available, besides the total chlorine present in the solution. Moreover, the addition of acetic acid to NaOCl in a ratio capable of adjusting the solution pH would have no effect on the rates of available chlorine (18).

In extracted human teeth, hypochlorite solutions with neutral pH have been shown to be more active against E. faecalis than conventional solutions with pH 12 (16,17). Theoretically, with the decrease of 1% NaOCl pH due to the incorporation of acetic acid, the antibacterial power of hypochlorite solutions should increase as a result of the increase of HOCl concentration. Hypochlorous acid dissociation depends on pH, with the clinical equilibrium between HOCl and OCl⁻ being maintained as HOCl is consumed through its germicidal function. Mercade et al. (17) tested in infected ex vivo root canals the antimicrobial efficacy of 4.2% NaOCl at two different pH values by assessing bacterial growth presence or absence following irrigation. A significant increase of the antimicrobial power of 4.2% NaOCl solution by weak
interactions with surface active agents will be greater with contemporary acceleration of the kinetics of the chlorine oxidizing activity (23). Its oxidizing capacity is retained by achieve greater stability, and this translates into chlorine's of the chlorine level of NaOCl solutions (13,25). Chlorine solutions. Nonetheless, surfactants may modify the stability in antibacterial efficacy results among all the examined study had to be rejected as differences could be detected most effective antibacterial action at all five experimental surface tension (Hypoclean and Chlor-Xtra), showed the for 5.25% NaOCl at day 28. The test solutions with lowered 95.08 ± 3.71 CFU (79,23%) versus addition of surface active agents increased significantly for 5.25% NaOCl at day 28. The test solutions with lowered 95.08 ± 3.71 CFU (79,23%) for Hypoclean versus Chlor-Xtra were not statistically significant (p>0.01) at all experimental periods except for results at day 7 (p<0.001). The exact NaOCl and the additional bactericidal effect of cetrimide must be also taken into account to understand these findings. Cetrimide used as single antibacterial agent it was equally effective as 2% NaOCl, 2% chlorhexidine, and iodine potassium iodide against E. faecalis into the dentinal tubules (15).

In conclusion, addition of acetic acid to 1% NaOCl did not enhance significantly the bactericidal activity of hypochlorite solution. Low concentration acidic chlorine-containing solutions seem to have lower antimicrobial effect than the currently used NaOCl concentrations. However, when CFU recovery for 1% NaOCl and 5.25% NaOCl were evaluated in comparison, encouraging results were recorded on the way to improve the antibacterial power of low concentration NaOCl. Thus, prospective studies should be conducted with comparisons between these modified irrigating agents, compared in other models of study, the ideal concentration for effective antibacterial power, adverse effects (possibility of interactions, degradation and formation of byproducts) and tissue tolerance. Based on the methodology employed and considering the limitations of this study, all irrigating solutions showed a small number of CFU during 28 days of evaluation.

Resumo
O objetivo deste estudo foi verificar o poder antibacteriano do hipoclorito de sódio a 1% acrescido do ácido acético 1%, do NaOCl 5,25%, e de duas soluções irrigadoras modificadas com surfactantes, Hypoclean e Chlor-Xtra, em tubos de dentina bovina. Um total de 120 tubos de dentina preparados a partir de incisivos bovinos foram infectados durante 28 dias com o Enterococcus faecalis, e aleatoriamente divididos em seis grupos: NaOCl 5,25%; Hypoclean; Chlor-Xtra; NaOCl 1% com ácido acético 1%; tubos de dentina infectados (controle positivo); e tubos de dentina esterilizados (controle negativo). Em períodos de tempo de 0, 7, 14, 21 e 28 dias, raspas de dentina foram coletadas usando brocas esfericas com diâmetros crescentes, e mantidas em tubos de ensaio contendo 3 mL de infusão cérébro coração (brain heart infusion). A análise dos dados foi feita com testes paramétricos (ANOVA a um fator e teste de comparações múltiplas de Bonferroni, α=1%). Após a cultura, o número de unidades formadoras de colônias (UFC) foi contado. Todas as soluções de hipoclorito de sódio mostraram pequena quantidade de UFC nos períodos por até 28 dias. Chlor Xtra e Hypoclean tiveram menor número de UFC em todos os períodos observados, com maior efetividade antimicrobiana do que o NaOCl 5,25% e do que o NaOCl a 1% com ácido acético a 1%.

References

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