Chemical aspect of sodium hypochlorite activation in obtaining favorable outcomes of endodontic treatment: An in-vitro study

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Abstract

Background. Endodontic treatment is one of the most widely performed procedures in a dental office. New techniques for enhancing the effectiveness of irrigants are being introduced into the dental market. It is crucial to choose a proper method to obtain the highest possible long-term success of performed endodontic treatment. Sodium hypochlorite (NaOCl) appears to be one of the most common and profitable solutions for root canal irrigation. The activation of a solution may be analyzed in 2 fields, physical – turbulence of flow, and chemical – disintegration of irrigant molecules into very active radicals that improve its activity. While the physical alternations of irrigant flow with different techniques are widely studied, there are not many attempts to approach the subject in chemical terms.

Objectives. The aim of the study was to compare the chemical effectiveness of 2 methods of NaOCl activation: ultrasonics vs the Self-Adjusting File system (SAF) as an adjunct to increase the efficacy of the irrigant.

Material and methods. The level of activation was evaluated via a reaction of the activated NaOCl samples, with 9-fluorenol as the starting organic material. The model reaction is based on the oxidation of 9-fluorenol to 9-fluorenon. The evaluation was performed using 1H nuclear magnetic resonance (NMR) spectroscopy, comparing the spectra obtained for the examined mixtures.

Results. Nuclear magnetic resonance studies show that the use of ultrasonics resulted in increased chemical degradation of NaOCl as compared to the SAF system and non-agitated samples. The prevalence of chemical activation in the ultrasonic group over the SAF group was almost 3 times higher, 3.11 to 1.20, respectively. The Kruskal–Wallis rank sum test revealed there is a statistically significant difference in distributions between the groups.

Conclusions. Both SAF and ultrasonics activate NaOCl. Ultrasonic agitation provided higher chemical activation of NaOCl solution than the SAF. The use of ultrasonic agitation of NaOCl in endodontic treatment will allow us to obtain better long-term clinical results.

Key words: NMR, sodium hypochlorite, endodontics, Self-Adjusting File, ultrasonics
Introduction

Sodium hypochlorite (NaOCl) is the most commonly used irrigant in endodontic treatment. Coolidge introduced it as early as in 1919. It is an inorganic compound with the chemical formula NaOCl. Commonly referred to as bleach, it is actually a diluted aqueous solution of NaOCl. It has numerous applications ranging from an oxidizing and chlorinating agent in organic synthesis (used in the chemical industry) to water treatment, bleaching and disinfection.

In dentistry, NaOCl is used in the form of dilute aqueous solutions with concentrations varying from 0.1% to 6.0%. Being a highly oxidizing compound, it shows suitable properties for dissolving remnants of dental pulp as well as acting as an antimicrobial agent. These features prompted the use of aqueous NaOCl in endodontics.

Pécora et al. reported that NaOCl exhibits a dynamic equilibrium in an aqueous solution, according to the equation:

\[ \text{NaOCl} + \text{H}_2\text{O} \Leftrightarrow \text{NaOH} + \text{HOCl} \Leftrightarrow \text{Na}^+ + \text{OH}^- + \text{H}^+ + \text{ClO}^- \]

As an ionic compound, NaOCl dissociates in water to sodium cation (Na⁺) and hypochlorite anion (ClO⁻), which is in equilibrium with its protonated form (HOCI). Both hypochlorous acid and the hypochlorite anion are strong oxidizing agents. HClO and ClO⁻ have been reported to react with proteins, amino acids, peptides, and lipids. Estrela suggests NaOCl acts as a dissolving agent for organic matter, e.g., fats.

An ultrasonic wave is an acoustic wave which transmits energy of a vibrating file to the irrigating agent. Waves in the 20,000–25,000 Hz frequency range, which cause several physical and biological effects, are used in endodontics. The increase of energy in the system treated with ultrasonics results in heating of the surrounding fluid, which subsequently results in a better bactericidal effect. The presence of cavitation and microstreaming effects is equally significant. The emerging waves disrupt bacterial cells and eliminate necrotic remnants. This mixing of the fluid and its turbulent flow has a significant influence on the efficacy of root canal irrigation.

The Self-Adjusting File (SAF) is a rotary system for mechanical instrumentation. It is also the only system that enables instrumentation and irrigation at the same time. This is a hollow cylinder, made of Ni-Ti alloy with fusiform holes along the entire length. The file can compress and decompress according to the topography and morphology of the treated canal. Two sizes, 1.5 mm and 2 mm in diameter, are available. Action of the SAF is based on back and forth sliding motions in the vertical axis. The RDT3 head connected to an endodontic motor with the speed set to 3,000–5,000 rpm is used for the SAF. The file is connected to a VATEA (ReDent Nova, Ra’anana, Israel) pump with a silicon tube, which delivers the irrigant along the entire length of the root canal during continuous instrumentation with a speed adjustable in the 1–10 mL/min range. It is recommended to fill the pump with NaOCl or ethylenediaminetetraacetic acid (EDTA) solution. The efficacy of NaOCl and EDTA in removing the smear layer and dentin using SAF has been proved and is not dependent on the concentration of the agent, even in the apical 1/3 of the canal. It is advised to work with SAF in the canal about 4 min – 2 cycles of 2 min for 1 root canal with a short pause for recapitulation. In our study, Endostation (ReDent Nova, Israel) was used. It combines an endodontic motor with the VATEA pump in one device.

According to Tiong and Price, during ultrasonic agitation, hypochlorous acid undergoes homolysis with formation of a hydroxyl radical and a chlorine radical. The homolytic processes occur when energy is delivered to HOCl molecules. This can be achieved in many different ways, e.g., under UV irradiation as reported by Zeng et al. The resulting radicals are a species highly reactive towards a wide range of biochemical compounds – proteins and glycolipids – leading to the degradation of biological matter. The chemical equation of homolytic degradation is presented below:

\[ \text{HOCI} \rightarrow \text{HO}^+ + \text{Cl}^- \]

One of the first investigations to quantify the action of NaOCl was described by Austin and Taylor in 1918. The authors measured the amount of chlorine remaining in the solution when exposed to normal vs necrotic tissue. The amount of remaining chlorine directly corresponded to its tissue-dissolving capacity. As NaOCl dissolved the tissue, chlorine levels decreased. The amount of chlorine remaining in the solution was measured indirectly on the basis of the amount of sodium thiosulfate used to reduce chlorine in the solution.

Activation of NaOCl can be considered 2-fold: physical activation wherein the turbulence of the flow is enhanced, and chemical activation based on HOCl degradation to highly reactive radical species. The rationale for the current experiment is that a quantitative evaluation of an oxidized product will help measure the oxidative potential of NaOCl at a given time. Selective oxidation of an organic compound with NaOCl may serve as a useful method to estimate its oxidation potential. Hence, the aim of this experiment was to evaluate the chemical activation of NaOCl quantitatively by measuring the capacity of the irrigant to oxidize 9-fluorenol as a model organic compound.

Material and methods

Experimental setup

All chemicals used in this study were obtained from commercial sources and used as such, without further purification. The following chemicals were used (name, purity,
additives if present, source): NaOCl, 5.25 wt% aqueous solution (Cerkamed, Stalowa Wola, Poland); 9-fluorenol, 96% (Sigma-Aldrich, St. Louis, USA); ethyl acetate, >98% (POCH S.A., Gliwice, Poland); tetra-n-butylammonium bromide, 99% (Sigma-Aldrich); chloroform-d_3 (CDCl_3), 99.8 atom% D, with 0.5 wt% silver foil as stabilizer and 0.03% (v/v) tetramethylsilane (TMS) (Sigma-Aldrich).

^1^H NMR spectra were recorded on a Varian VNMR5 500 MHz NMR spectrometer (Varian, Atlanta, USA) in CDCl_3, using TMS as an internal standard for chemical shift calibration. All chemical shifts are reported in ppm. Armar Chemicals 5 mm NMR Tubes (5HP type) (Armar Chemicals, Döttingen, Switzerland) were used. Concentration under reduced pressure was carried out with a Heidolph rotary evaporator (Heidolph, Schwabach, Germany).

Due to the comparative character of the study, all possible measures were taken to ensure that all procedures were carried out as similarly as possible while evaluating both NaOCl activation methods. Each reading was done in triplicate and each result given as an average, along with standard deviation (SD) from the 3 readings.

Sodium hypochlorite activation

A 12 mL glass vial with a plastic cap was used to fill 10 mL samples of 5.25% (w/w) aqueous solution of NaOCl. For each activation method, 2 samples of the NaOCl stock solution were prepared as mentioned above. Activation was carried out by submerging the respective system in the sample for a 4-minute period. The SAF system - a 1.5 mm diameter and 25 mm length file – was used at a speed of 5,000 rpm, with the RDT3 head connected to a handpiece on the Endostation (ReDent Nova) with an adjustable rotary speed setting, according to the manufacturer's instructions. Constant irrigant flow was not supplied to SAF during the study. For ultrasonic activation, an ISO 20 K-file (Woodpecker, China) connected to an endodontic adapter (Endochuck 120°, Woodpecker, China) and a handpiece on an ultrasound device (Woodpecker UDS-P LED) with the endodontic function turned on (only the ‘E’ light was on and the power was at first grade, according to the user’s manual).

Reaction between sodium hypochlorite and 9-fluorenol

Each reaction was carried out in a 12 mL glass vial equipped with a plastic cap and a magnetic stirring bar. Following a procedure modified from Mirafzal et al.,\textsuperscript{23} 9-fluorenol (185 mg, 1.02 mmol) was dissolved in ethyl acetate (1.5 mL). Next, tetra-n-butylammonium bromide (30 mg, 0.0931 mmol, 9 mol%) was added as a solid in one portion. The tested sample of NaOCl solution (1.5 mL) was added to the resulting solution in one portion. The reaction mixture was stirred for 30 min. Using a syringe, a portion of the organic layer (1 mL) was transferred from the vial to a 10 mL glass round-bottom flask. The solution was concentrated under reduced pressure to dryness, using a rotary evaporator. CDCl_3 (0.7 mL) was added to the so-obtained residue, resulting in the formation of a clear solution. The absolute concentration of the NMR sample, given with respect to the organic compound used, was equal to 1.45 mol/dm.\textsuperscript{3} To prepare for ^1^H NMR analysis, the solution was then transferred to an NMR tube using a plastic syringe, and the tube was closed with a plastic cap.

Statistical analysis

The statistical analysis was made using an open source statistical environment R v. 3.5.1 (https://www.R-project.org/). The Kruskal–Wallis rank sum test was available as krustal.Test function from R package, including test statistics, p-value and number of degrees of freedom (df), follows in implementation the theoretical results from Hollander and Wolfe.\textsuperscript{25}

Results

Model for the study

The reaction under assessment is the phase-transfer-catalyzed oxidation of 9-fluorenol with NaOCl (Fig. 1).\textsuperscript{23,26} The rationale behind picking it as a model reaction is given in the Discussion section. The experiment was performed on 3 samples: commercially available non-activated NaOCl solution, NaOCl solution subjected to ultrasonic activation and NaOCl solution activated with the SAF system.

\begin{figure}
\centering
\includegraphics[width=0.5\textwidth]{figure1.png}
\caption{Equation of the model reaction used in this study: oxidation of 9-fluorenol with sodium hypochlorite and tetra-n-butylammonium bromide as a phase-transfer catalyst}
\end{figure}

^1^H NMR spectroscopy

^1^H NMR spectra were recorded in CDCl_3 at 298 K (25°C) and 499.93 MHz. A standard proton pulse sequence (s2pul) was used, with relaxation delay at 1.000 s, pulse at 45.0°, acquisition time at 2.045 s and 8,012.8 Hz spectral width. Acquisition of each spectrum comprised 16 scans. The spectra were processed using iNMR Reader software (Mestrelab Research, Santiago de Compostela, Spain). The raw NMR files (FIDs) were subjected to automatic Fourier transform (FT) with FT size equal to 64k. Automatic
baseline correction for a 1D spectrum, implemented in the iNMR Reader software, was applied to each spectrum, setting the baseline intensity to 0 and thus allowing to reliably integrate the peaks. Full view of the obtained $^1$H NMR spectra, 1 for each kind of samples examined, is presented in Fig. 2. The spectra are shown in the range between 9.0 ppm and 0.0 ppm, with TMS signal at 0.0 ppm.

The analytically useful peaks are in the 7.8 ppm to 7.0 ppm (aromatic region) and 5.6 ppm to 5.5 ppm (benzylic protons) range. The signals of chemical shifts below 5.0 ppm include peaks derived from hydrogen atoms in: ethyl acetate (solvent in the reaction), tetra-α-butylammonium bromide (catalyst), water and TMS (NMR chemical shift standard). The peaks in the region below 5.0 ppm were not relevant for the study and have been omitted in Fig. 3 for clarity. Both aromatic and benzylic signals were integrated, providing the relative ratio of the total areas of the peaks. The comparative spectroscopic analyses of reactions mixtures obtained from respective solutions are showed in Fig. 3.

### Chemical concept of the study

The chemical concept applied here is that aromatic protons can serve as an internal reference to evaluate the extent of oxidation and therefore the effectiveness of NaOCl activation. The interpretation of the changes occurring in the mixture upon oxidation is straightforward, as both the 9-fluorenol and 9-fluorenone have aromatic protons, while only the 9-fluorenol has a benzylic proton (Fig. 4). Upon oxidation, the number of aromatic protons (and thus the integration of the aromatic region) is equal both in the starting material and the product and is unchanged during the oxidation.²⁷,²⁸ The variable here is the benzylic signal, which decreases in its intensity (and integration) upon oxidation.

The ratio of integration values of signals derived from benzylic protons to aromatic protons may be correlated with the efficiency of 9-fluorenol oxidation and establishes the basis for evaluating the NaOCl activation method. The higher benzylic-to-aromatic signal ratio, the less efficient was the oxidation of 9-fluorenol to 9-fluorenone. Less efficient oxidation of the starting material means that a smaller amount of hypochlorite was left in the solution after its activation, which points to a higher extent of the decomposition of hypochlorite upon activation. This is an indication of better efficacy of NaOCl and hence a more effective method to activate the irrigant. To convey the idea more clearly, the logical sequence behind it is summarized graphically in Fig. 5.

### Numerical results of the study

The numerical results of the analyses are presented in Table 1. As each experiment was done 3-fold, 3 entries for each activation mode are provided. As aforementioned, the number of aromatic protons remain unaffected upon oxidation, so the integration of their signals can serve as an internal standard for determining the integration ratio.²⁹ The integration of the aromatic region was then set by default to 1.00. As all further discussion is based on the integration ratios, setting the aromatic region...
integration to 1.00 makes it easier to evaluate and discuss the results.

With this in mind, there is no need to reference the integration of signals against the internal standard used, i.e., TMS. The additive of TMS in the commercial deuterated solvent was used only for calibrating the chemical shift of the observed signals, with TMS having the chemical shift of 0.00 ppm. Tetramethylsilane would not have been a reliable reference for the integration due its volatility and hence tendency to change its concentration over time, making it an invalid reference for any quantitative studies.

The use of another chemical shift standard widely used in molecular medicine studies, namely 3-(trimethylsilyl)-2,2,3,3-tetadeuteropropionic acid (TSP), would not be possible due to the fact that TSP is insoluble in deuterated chloroform which was the deuterated solvent used in this study. Deuterated chloroform was picked as a deuterated solvent of choice due to its capability to dissolve the organic compounds well, both the substrate (9-fluorenol) and the product shift of 0.00 ppm. Tetramethylsilane would not have been a reliable reference for the integration due its volatility and hence tendency to change its concentration over time, making it an invalid reference for any quantitative studies.

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### Table 1. Results of the spectroscopic studies

<table>
<thead>
<tr>
<th>Activation mode</th>
<th>Benzyllic-to-aromatic signals integration ratio</th>
<th>Average</th>
<th>Standard deviation (SD)</th>
<th>Recalculated integration ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>No activation</td>
<td>0.0127 : 1.00 0.0122 : 1.00 0.0125 : 1.00</td>
<td>0.0125</td>
<td>0.0003</td>
<td>1.00</td>
</tr>
<tr>
<td>SAF</td>
<td>0.0154 : 1.00 0.0152 : 1.00 0.0143 : 1.00</td>
<td>0.0150</td>
<td>0.0006</td>
<td>1.20</td>
</tr>
<tr>
<td>Ultrasonics</td>
<td>0.0386 : 1.00 0.0382 : 1.00 0.0396 : 1.00</td>
<td>0.0388</td>
<td>0.0007</td>
<td>3.11</td>
</tr>
</tbody>
</table>

1 The benzyllic-to-aromatic signals integration ratio, recalculated with respect to the value of the “no activation” ratio equal to 1.00.
(9-fluorenone) as well as any amount of the catalyst left in the organic phase during the extraction.\textsuperscript{27,28,31} Moreover, the use of TSP could lead to misleading results as TSP tends to be unstable in the presence of oxidative agents. If any traces of NaOCl had been left after the extraction process (which is unlikely but cannot be excluded), TSP would have got oxidized, hence making its use as a standard invalid.

In all experiments, the benzylid-to-aromatic ratios were found to be between 0.0122:1.00 and 0.0396:1.00. That observation was expected, as the integration should have fallen within the 0.1250 for 9-fluoreol (equal to 1:8 ratio, with 1 benzylid proton and 8 aromatic protons) and 0 for 9-fluorenone (0:8 ratio, with no benzylid protons and 8 aromatic protons) bracket. Importantly, none of the ratios were equal to 0, as that would mean 9-fluoreol was oxidized completely which would preclude any qualitative or quantitative conclusions being drawn.

For each activation mode, an average ratio was determined from 3 experiments, along with its SD. The integration ratios were then recalculated with respect to the “no activation” ratio set to 1.00. The ratio values for the SAF and ultrasonic activation modes were found to be 1.20 and 3.11, respectively.

Differences in the results between the examined groups of samples are clearly visible in Fig. 6. The obtained integration ratios of all results in the SAF and ultrasonics groups are higher than the values for a non-activated group.

**Statistical analysis results**

Taking into account the means and SDs of benzylid-to-aromatic signals integration ratio between groups, our hypothesis states that their distribution is not equal. Due to the fact that there is no way to reliably test for normality when we have 3 observations per group, a non-parametric approach is used. The null hypothesis that the cumulative distribution function in all 3 groups is equal is tested against the alternative hypothesis that at least 2 out of 3 cumulative distribution functions in the groups are different. The hypotheses are tested using ANOVA on ranks, i.e., using the Kruskal–Wallis rank-sum test. The value of the test statistics, which under the null hypothesis has a \( \chi^2 \) distribution with 2 df, is 7.2. It follows that the p-value in this test is equal to 0.02732372, which is lower than the 0.05 significance level. The null hypothesis is rejected, thus at the 0.05 significance level there is a statistically significant difference in distributions between the groups.

It is important to note, that in this case the usual interpretation of Kruskal–Wallis rank sum test using medians is not possible. For the test to be interpreted in terms of medians, we should verify first that all 3 distributions come from the same family of location-scale distributions. This is not possible with the available data, due to the small number of observations. For the same reason, it is also impossible to perform a non-parametric post-hoc analysis using the Wilcoxon rank sum test.

**Discussion**

Stojicic et al. suggested several methods for improving efficacy of NaOCl. Changes in concentration, temperature, agitation, increase of pH, and prolonged contact time are some of them.\textsuperscript{32} Moorer and Wesselink stressed the impact of mechanical agitation to be the most significant mode of NaOCl activation. They claim fluid flow is the key for effective tissue dissolution and disinfection as well as emphasize that ultrasounds ensure the best quality of turbulent flow.\textsuperscript{33} An increase in flow and more turbulence enables the solution to reach hardly accessible areas and therefore improve efficacy. Sáinz-Pardo et al. also provide evidence that ultrasonic agitation ensures proper fluid flow throughout the working length of a root canal.\textsuperscript{34} Besides fluid dynamics, the exact mechanism responsible for the improved efficacy is not clear.\textsuperscript{35} This is the first investigation of its kind to evaluate whether a chemical activation of NaOCl also occurs during agitation.

To evaluate the methods for hypochlorite activation, we searched the chemical literature for reactions which use NaOCl solutions as a reagent. We preferred a reaction that did not take long to carry out in order to shorten the evaluation as much as possible. Other criteria that needed to be met included the following: easy reaction set-up (ideally no specialized glassware or equipment), mild conditions (preferably room temperature, atmospheric pressure, no need for the use of inert gases like argon or nitrogen), inexpensive and easily commercially available chemicals (including the starting materials, solvents and optionally catalysts) as well as the possibility to carry out the evaluation using an analytical method yielding readily interpretable
and unequivocal results. After a detailed literature survey, we picked the phase-transfer-catalyzed oxidation of 9-fluorenol with NaOCl as the model reaction.

The role of NaOCl solution is to provide hypochlorite anions (ClO\(^-\)), which serve as an oxidant towards benzylic alcohol. Importantly for the evaluation of oxidation potential, hypochlorite reacts with 9-fluorenol selectively, affording solely the oxidation product – 9-fluorenone. 9-Fluorenol is not prone to spontaneous oxidation neither in pure oxygen nor in the air, even upon prolonged storage. These properties made the quantitative evaluation of oxidation product exclusive to NaOCl, and hence valid. Moreover, it facilitates handling of 9-fluorenol, as no specialized chemical skills or sophisticated equipment is required.

Among the many variants of 9-fluorenol oxidations, the one chosen by us is preferable also because it does not require the presence of expensive, metal-based catalysts, elevated temperature or irradiation. For an effective execution of 9-fluorenol oxidation with hypochlorite, it is sufficient to make use of inexpensive and easily applicable phase-transfer catalysis (PTC).\(^{31}\) The method, originally developed in the 1960s, is particularly useful for reactions occurring in 2-phase systems, i.e., in reaction mixtures containing 2 immiscible (or miscible to a limited extent) layers. For example, such layers may be made up of a reagent aqueous solution (in our case: aqueous solution of NaOCl) and a solution of the starting organic material (here: 9-fluorenol) in an organic solvent immiscible with water (e.g., ethyl acetate). The reaction between the organic substrate and the inorganic reagent (hypochlorite anion) takes place at the interphase between those 2 layers. For the oxidation to proceed efficiently, both the substrate and the oxidant must come together in the interphase at the same time. Here a phase-transfer catalyst comes in useful. The PTC catalysts usually contain a heteroatom (particularly nitrogen), in the form of a cationic ammonium species with a general formula of R\(_2\)N\(^+\), where R is an organic group, most commonly an aliphatic chain. Such a structure has a dual nature in terms of polarity: the cationic nitrogen atom serves as the polar (hydrophilic) center, while the aliphatic hydrocarbon chains are highly non-polar (hydrophobic, but with high affinity to the organic layer). This duality makes it possible for the cation of the catalyst to be transported to both the organic and aqueous layers, efficiently delivering the hypochlorite anion from the aqueous layer to the interphase with the organic layer. In this study, tetra-n-butyrammonium bromide was used as the phase-transfer catalyst.

Having an appropriate model reaction at hand, we embarked on the search for a method for both a qualitative and a quantitative analysis of the reaction mixture composition, i.e., the identity and content of the substrate and product. We chose nuclear magnetic resonance (NMR) spectroscopy for this purpose. The NMR spectroscopy is one of the most important research techniques used to investigate the structure, content and interactions of organic compounds. The method is basically based on the phenomenon of energy transitions occurring within the nuclei of atoms when an external magnetic field is applied. By investigating the electromagnetic signals in this process and working them out mathematically, a so-called NMR spectrum can be obtained. The NMR method we decided to use was proton NMR (\(^1H\) NMR), as it allows one to obtain qualitative and quantitative information on protons (hydrogen nuclei) in the organic compounds structure present in the analyzed mixture.

Only a few reports on the use of spectroscopic methods in endodontic research have been published, all of them being relatively recent. A study by Sireesha et al. on the comparison of micro and nano substances used as temporary intracanal dressings was based on infrared spectroscopy (Fourier-transform infrared spectroscopy – FTIR) to analyze the interactions between nanoparticles.\(^{36}\) The conclusion from the study was that the tested nano-calcium hydroxide and nano-chitosan compounds are superior to ordinary calcium hydroxide and chitosan both in terms of better penetration of dentinal tubules and lower susceptibility to root fracture. One can find an example of the use of mass spectrometry (MS) in a study by Chávez de Paz et al.\(^{37}\) An evaluation of the coexistence of Enterococcus faecalis bacteria strains with other bacteria within the root canal system has shown that in case bacteria can interact synergistically as well as antagonistically. X-ray spectroscopy is another example of spectroscopy used in endodontic research, and in particular wavelength-dispersive X-ray spectroscopy (WDXS or WDS) used in the Han study to assess the calcium and silicon uptake by dentin from Biodentine and MTA.\(^{38}\) The study shows that the dentin adjacent to a given material exhibited greater influence of calcium and silicon from the Biodentine material.

By using the concept of spectroscopic data analysis described in the Results section, we were able to qualitatively and quantitatively compare the activation methods. Based on the \(^1H\) NMR integration ratios recalculated with reference to the ratio obtained for non-activated NaOCl, the occurrence of chemical activation of hypochlorite by both the SAF and ultrasonics was confirmed. Quantitatively, the ultrasonic activation was over 2 times more effective compared to the SAF activation (1.2-fold activation for SAF compared to 3.1-fold activation for ultrasonics).

All the results obtained in the SAF and ultrasonic groups predominated these in the non-activated group. What is more, all results in the ultrasonic group were higher than those measured in the SAF group. The consequence and reproducibility in each group with a low SD value lays a basis for the use of this method in other NaOCl-related studies and analyses in the future. Similar experiments on other NaOCl agitation methods would have a worthwhile contribution in understanding their clinical efficacy.

It is worth noticing that another dentistry-related study on NaOCl capacity for oxidation can be found...
in literature. However, it used a non-spectroscopic method for NaOCl analysis, namely a redox titration called iodometry. Zehnder assessed the buffered and unfurred solution of NaOCl for tissue dissolution. However, the disadvantage of that approach was that potassium iodide (KI; used as a titrating solution) is known to be active not only towards NaOCl, but also other oxidants present in the solution. This appears even more perspicuous when the equations of NaOCl comproportionation (Equation 1) as well as non-selectivity of potassium iodide oxidation (Equations 2 and 3) are taken into account.

\[
\begin{align*}
(1) & \quad 3 \text{NaOCl} & & \rightarrow 2\text{NaCl} + \text{NaClO}_3 \\
(2) & \quad \text{NaClO}_3 + 6\text{KI} + 3\text{H}_2\text{O} & & \rightarrow \text{NaCl} + 3\text{I}_2 + 6\text{KOH} \\
(3) & \quad \text{NaOCl} + 2\text{KI} + \text{H}_2\text{O} & & \rightarrow \text{NaCl} + \text{I}_2 + 2\text{KOH}
\end{align*}
\]

Sodium hypochlorite (NaOCl) is known to comproportionate into sodium chloride (NaClO\textsubscript{3}) and sodium chloride while in aqueous solutions, especially over prolonged storage. Importantly, both NaOCl and NaClO\textsubscript{3} are able to oxidize potassium iodide. The titration result may then be false due to lack of selectivity.

The reactions used in our study display specificity towards NaOCl, with 9-fluoreno1 being selectively oxidized by it. Despite a great number of works on chloride-based oxidants, NaClO\textsubscript{3} has never been found to oxidize any secondary alcohols like 9-fluoreno1.

In conclusion, a spectroscopy-based protocol for a quantitative evaluation of activation of NaOCl was proposed and successfully implemented. The comparative \textsuperscript{1}H NMR data, derived from the experiments on the irrigant capacity to oxidize 9-fluoreno1, unequivocally showed that both SAF and ultrasonics activate NaOCl. In addition, it was shown that ultrasonic agitation provided better chemical activation of the NaOCl solution than the SAF.

From a medical perspective, it is worthwhile to point out that any contact between an ultrasonic file with canal walls diminishes the activation effect.\textsuperscript{40} This is the main impediment to achieving proper ultrasonic activation on the whole working length in severely curved canals.

On another concluding note, we believe that such an interdisciplinary approach towards solving challenges on the border of medicine and science will soon become an important area for research and such results may quickly become clinically relevant. In this case, the significance of the study lies in providing a dental practitioner with specific information on choosing the best way for NaOCl activation, making canal disinfection as thorough as possible and increasing the efficiency as well as the long-term results of endodontic treatment in the clinic.

A molecular approach in exploring the dental field may give extra opportunities in future scientific projects in this area of medicine. Nowadays, microbiological and clinical studies underlie the majority of new techniques and devices being introduced into the market and, in consequence, the dental practice. However, it is molecular medicine and experimental approach that explain the fundamental aspects of biological and chemical mechanisms behind these developments and boosts the understanding of everyday activities undertaken by physicians and dentists. The described influence of the SAF motion and ultrasonic wave on changes in the molecular composition of NaOCl solution should make one aware that any step taken during a clinical procedure may hinder or improve the overall outcome. Advantageous properties of NaOCl for endodontics that are strengthened by additional activation methods like ultrasonics can be correctly understood only by means of a thorough experimental analysis. The mechanisms behind the observed macroscopic phenomena in the clinical environment are often comprehensively understandable only at the molecular level.

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