Original Research Article

Clinical evaluation of 4% hydrogen peroxide bleaching in mandibular teeth

Lais Dalmagro Peruchi1
Neimar Sartori1
Guilherme Carpena Lopes1
Andressa Ballarin1
Camila Ambrosi1
Jussara Karina Bernardon1

Corresponding author:
Universidade Federal de Santa Catarina
Campus Universitário Reitor João David Ferreira Lima – Trindade
Departamento de Odontologia
Disciplina de Dentística
CEP 88040-970 – Florianópolis – SC – Brasil
E-mail: ladape@gmail.com

1 Department of Dentistry, Federal University of Santa Catarina – Florianopolis – SC – Brazil.

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Abstract

Introduction: Bleaching agents have been constantly introduced into market; however, the efficacy of new hydrogen peroxides still needs to be evaluated. Objective: The aim of this study was to evaluate in vivo the effectiveness of 4% hydrogen peroxide bleaching agent in color change, bleaching maintenance, tooth sensitivity, and patients’ satisfaction. Material and methods: Forty subjects were selected, lower bleaching trays were constructed, and the patients were instructed to apply 4% hydrogen peroxide bleaching agent at home. Shade measurements of the mandibular anterior teeth were carried out with a spectrophotometer and Vita Classical shade guide (VITA) at the following moments: baseline, 14 days, after patients’ satisfaction, and after 90 days. In addition, tooth sensitivity was evaluated using a visual analogue scale (VAS) at the first 14 days. Shade measurements were submitted to repeated measures ANOVA with Bonferroni adjustment and level of significance set at 5%. Results: Statistically significant differences were found for mandibular tooth shade after at-home bleaching (p < 0.001). After 14 days of bleaching, 90% of the subjects were not pleased with the achieved bleaching; however, after 27 days most of the patients reported to be satisfied. Tooth sensitivity was reported to range from 1.01 in a scale from 0-10. Conclusion: At-home bleaching using 4% hydrogen peroxide is effective in lower teeth with reduced tooth sensitivity.

Keywords:
tooth bleaching;
hydrogen peroxide;
patient satisfaction.
Introduction

Tooth bleaching becomes a very popular due to be an effective, safe, low-cost, conservative treatment to obtain vital tooth whitening [16, 24]. Tooth whitening mechanism is not totally understood [21]; however, it is believed that the bleaching agent (hydrogen peroxide) is decomposed to produce oxygen ($O^+$) and hydroxyl radicals ($HO_2^-$) [20]. These free radicals attack tooth dark pigment molecules (organic macromolecules formed by aromatic rings), to obtain stability by breaking them into smaller, less complex and clearer molecules than the original ones [13].

Among bleaching techniques for vital teeth, at-home and in-office whitening are emphasized [14]. At-home bleaching main advantages are effective outcomes, technique simplicity, low cost, and to be executed out of dental office [27, 29]; the disadvantages are color regression [6], long-time treatment, which contributes to tooth sensitivity during treatment [27].

At-home bleaching treatment comprises the application of a bleaching agent gel, based on carbamide peroxide at 10% to 22% concentration or hydrogen peroxide ranging from 4% to 9% concentration placed onto a customized tray, which is daily used by the patient. The original technique advocates the bleaching agent application for six to eight-hour daily regimen, during the night [11]. Notwithstanding, Matis et al. [19] demonstrated that after two hours of bleaching, only 50% of the active agent is available; after 10 hours, bleaching agent availability decreases to 10%. Consequently, depending on bleaching agent concentration, some manufacturers recommended only 30 minutes of daily use of bleaching agent.

Scientific evidences showed that tooth bleaching can reduce tooth's microhardness, fracture toughness and calcium and phosphate concentrations [1], as well as increase enamel's superficial roughness [18, 23]. Some compounds may be added to bleaching gels, e.g. potassium nitrate and sodium fluoride, to act as desensitizing agents; sodium fluoride also allows tooth remineralization [10]. Currently, some manufacturers start to add calcium to bleaching agent gels, aiming to decrease tooth enamel's demineralization process.

Therefore, the aim of this clinical study was to assess the effects of a 4% hydrogen peroxide bleaching agent containing potassium nitrate, sodium and calcium fluoride on color change, whitening maintenance, tooth sensitivity, and patient's satisfaction.

Material and methods

This research was approved by Human Research Ethics Committee. Forty volunteers were selected according to the inclusion and exclusion criteria listed in Table I. All selected volunteers received detailed information on the research and the technique's goals and risks. Following, the participants signed a Informed Consent, meeting the Resolution number #196, from October 10, 1996, by the Brazilian Council of Health/Ministry of Health, Brasilia, DF, Brazil.

Table I - Inclusion and exclusion criteria of the research

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
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<tbody>
<tr>
<td>Patients aging from 18 to 39 years;</td>
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<td>Patients living in Florianópolis;</td>
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<td>Patients willing to participate voluntarily who agreed with the research methodology and signed the free and clarified consent form.</td>
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<tr>
<th>Exclusion criteria</th>
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<tr>
<td>Patients who presented restorations or carious lesions in the six anterior lower teeth;</td>
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<td>Patients presented any clinical history of disease which may interfere in the treatment;</td>
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<tr>
<td>Pregnant and lactating women;</td>
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<td>Presence of tooth sensitivity;</td>
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<tr>
<td>Patient with simultaneous or predicted periodontal treatment;</td>
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<tr>
<td>Patients with previous bleaching treatment;</td>
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<tr>
<td>Smokers;</td>
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<tr>
<td>Patients under analgesic and/or anti-inflammatory therapy;</td>
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<tr>
<td>Patients with serious systemic and psychological diseases;</td>
</tr>
<tr>
<td>Patients under orthodontic treatment, using fluoride supplementation or any desensitizing agent;</td>
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<tr>
<td>Patients with unavailable time to attend follow-up appointments.</td>
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</table>

At first appointment, a professional dental prophylaxis was executed, followed by filling in the specific tooth bleaching file and upper and lower arch impressions with alginate (Avagel, São Paulo, Brazil). The impressions were poured-up with die stone (Vigodent S.A. Indústria e Comércio Bom Sucesso, Rio de Janeiro). After the stone setting, working casts were adequately cut. Lower and upper trays were made and cut 2 mm above the gingival margin.
Prior to bleaching onset, color measurements of tooth #33, #32, #42, and #43 were executed by using a shade guide (Vita Classical, Vita Zahnfabrik, Bad Säckingen, Germany) and spectrophotometer (Vita Easyshade, Vident, Brea, USA) and recorded on a specific file. A demonstration of how the bleaching agent should be placed into the tray was performed patients were instructed to use the bleaching gel for two hours per day, according to the manufacturer's instructions. At that moment, patients received one syringe of 4% hydrogen peroxide bleaching agent gel (White Class, FGM, Joinville, Brazil) and the customized tray for lower arch whitening. Also a visual analogue scale (VAS) was given to the patient to register tooth sensitivity during the first 14 days of treatment. Patients were instructed to classify tooth sensitivity in a scale from 0 to 10, varying from “no discomfort at all” to “extremely unpleasant” [25].

The first follow-up appointment was performed 14 days after the treatment onset. At this evaluation, the shade measurements by using the shade guide and spectrophotometer were executed, following the same parameters of the baseline assessment. After shade measuring, the participants answered a questionnaire aiming to reveal their satisfaction degree with tooth whitening at 14 days of treatment [7]. If patient were satisfied with 14-day treatment and would not want to continue it, the bleaching agent syringe was returned. On the other hand, if patient were not satisfied, we oriented to continue treatment and a new appointment was scheduled.

For patients who wanted to continue treatment, a new follow-up appointment was performed after patient satisfaction with tooth whitening outcome. This assessment followed the same pattern of the previous ones. Twelve weeks after bleaching treatment onset, a new assessment was scheduled to verify tooth whitening maintenance over time. At this appointment, patient received the customized upper tray to execution of upper arch bleaching.

Visual shade measurement was standardized through the controlling of light environment. For this purpose, we used a light temperature of 6,500 K which corresponding to day light, at a distance of 25 cm from patient's face. Additionally, all participants wore a neutral gray apron over their clothes to avoid interference during the assessment. The shade measurement was always carried out at tooth's medium third. To assure an accurate measurement, the shade guide was ordered by value, as follows: B1, A1, B2, D2, A2, C1, C2, D4, A3, D3, B3, A3.5, B4, C3, A4 and C4 (in which B1 corresponded to color 1 and C4 to color 16). Shade value was covered by an adhesive tape to decrease examiner's error.

Objective measurement of shade was performed with a spectrophotometer following the manufacturer's instructions, with the device's tip placed onto labial surface of the medium third of the evaluated lower teeth. To standardize the tooth area which would be evaluated with the spectrophotometer, a device made out of orthodontic wire was constructed. This device was linked into the tooth to be measured and into the spectrophotometer, allowing that this latter record always the same tooth area, in all performed measurements. The spectrophotometer provides the values $L^*$, $a^*$ and $b^*$ coordinates CIE Lab, for each analyzed tooth. $L^*$ indicates the luminosity; $a^*$ and $b^*$ indicates the shade, where $a^*$ represents the color and saturation in the red-green axis and $b^*$ represents the blue-yellow axis. Shade comparison before and after tooth bleaching, for each time period, was obtained by the color difference ($\Delta E$), by the following formula: $\Delta E = (\Delta L)^2 + (\Delta a)^2 + (\Delta b)^2)^{1/2}$, in which $\Delta L = \text{final } L - \text{initial } L$; $\Delta a = \text{final } a - \text{initial } a$, and $\Delta b = \text{final } b - \text{initial } b$. Data statistical comparison was performed by ANOVA and Bonferroni tests.

### Results

ANOVA for repeated measurements showed that there was a statistically significant difference in color change of teeth, over time, in both used methods of assessment (p-value < 0.001). Table II and graph 1 showed that 14-day $\Delta E$ was smaller and statistically different of 90-day and patient satisfaction $\Delta E$.

<table>
<thead>
<tr>
<th>Periods</th>
<th>Meas±SD</th>
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<tbody>
<tr>
<td>$\Delta E$ (14 days)</td>
<td>6.03±2.59a</td>
</tr>
<tr>
<td>$\Delta E$ (patient's satisfaction)</td>
<td>7.61±3.32b</td>
</tr>
<tr>
<td>$\Delta E$ (90 days)</td>
<td>7.44±2.89b</td>
</tr>
</tbody>
</table>

Means followed by same letters are not statistically different by Bonferroni test (p > 0.05)
In table III and graph 2, we noted that tooth’s initial color mean was 7.17, which means a color between C2 and D4 when Vita Classic Scale was ordered by value. After two weeks of bleaching treatment, there was a reduction of about 3 shades, resulting in a shade close to D2, which is statistically different from baseline values. Notwithstanding, patients were satisfied after 27 days of treatment, in average, when teeth reached a shade close to B2, which was statistically different from two-week values.

Table III – Mean, standard deviation (SD) and statistical tests’ results of tooth’s color change over time evaluated by subjective method (shade guide)

<table>
<thead>
<tr>
<th>Periods</th>
<th>Mean±SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial</td>
<td>7.17±2.99a</td>
</tr>
<tr>
<td>14 days</td>
<td>3.72±1.98b</td>
</tr>
<tr>
<td>Patient’s satisfaction</td>
<td>2.89±1.33c</td>
</tr>
<tr>
<td>90 days</td>
<td>2.94±1.33c</td>
</tr>
</tbody>
</table>

Means followed by same letters are not statistically different by Bonferroni test (p > 0.05)

Graph 3 shows the results of tooth sensitivity reported by the patients during the first two weeks of treatment. Tooth sensitivity was slow; general mean of all results was equal to 1.01, in a scale ranging from 0 to 10.

Discussion

Clinical evaluations were the best way to verify bleaching agents’ effectiveness because they include patients’ individual factors (habits, food, hygiene, among others) which can influence on the final tooth shade. Therefore, because the research is conducted in a population with distinct life-style characteristics, the found bleaching degree tend to be more likely to the one obtained in clinical practice.

Among the methodologies used for evaluating tooth bleaching effectiveness in literature [15], the methods most commonly employed in clinical evaluations have been: comparison with visual standardized shade guide and instrumental measurement with spectrophotometers [22]. The main advantage of visual scale is the low cost; however, because it is a subjective method, it could be influenced by patient’s clothes and skin color as well as illumination [8], examiner experience, and visual acuity. To minimize such factors, shade assessment was performed by one single examiner under 6,500 K light. Spectrophotometer measurements has a higher cost and is more complex; on the other hand, it is objective, precise, and allows the numerical measurement of tooth color change in a tridimensional scale (CIELab) [7]. This study results corroborate literature and demonstrate that both methodologies are effective in assess tooth color changes [7]. Accordingly, both researchers and dentists can use standardized shade guides to assess bleaching treatment effectiveness when a spectrophotometer is not available.
Our results corroborate literature [3, 22, 24] and proved that at-home bleaching is effective for vital bleeding teeth. Among the bleaching materials available for at-home bleaching technique, hydrogen and carbamide peroxide is highlighted. Regardless of the material type, hydrogen peroxide is the active agent which promotes tooth bleaching acting as oxidizing agent and reacting with the aromatic rings, breaking them into smaller and clearer molecules [13]. Consequently, bleaching degree depends on hydrogen peroxide concentration, the ability of breaking aromatic ring chains, and frequency and time period that bleaching agent is in contact with macromolecules [7].

In our study, patient's satisfaction was reached at 27 days of treatment, in average. Two-week bleaching treatment promoted a clinically visible color change of the teeth (ΔE greater than 3.3 units) [28]; however, this value was not sufficient for pleasing 90% of the patients. Data of patient's satisfaction found by our study were in agreement with another clinical study: according to Bernardon et al. [2], patient's satisfaction was reached between 4 and 6 weeks of treatment, regardless of the bleaching agent or technique employed. Most of the studies used a pre-established time period to bleaching execution; however, in clinical practice is important to consider patient's opinion for establishing treatment length [7]. In this study, we observed that there was a statistically significant difference between 14-day treatment and patient's satisfaction assessment, showing that pre-established periods can be used as reference both for the patient and the clinician. Notwithstanding, ideal treatment length should be discussed with the patient during follow-up appointments.

It is also noted that after 12 weeks tooth color was stable in comparison with the color obtained at the treatment ending (patient's satisfaction). This information indicates that bleaching treatment was effective and corroborates other clinical studies, which demonstrated that there is no significant regression of tooth color after bleaching treatment [3, 7].

Tooth sensitivity has been reported as the main side effect of tooth bleaching, resulting in treatment abandonment [12]. Literature has demonstrate that 15% to 65% of patients show tooth sensitivity increase during bleaching treatment [17]. This increase occurs because hydrogen peroxide propagates through enamel and dentin, reaching the pulp [9, 27]. In our study, tooth sensitivity reported by the patients was low: 1.01 in a scale from 0 to 10. Mild tooth sensibility may be related to the presence of potassium nitrate, calcium and sodium fluoride in bleaching agent composition. According to Browning et al. [4], patients undergoing tooth bleaching with agents containing desensitizing showed a smaller sensitivity rate.

Since bleaching treatment should be conducted both in upper and lower arch, researches evaluating tooth sensitivity and bleaching agent effectiveness in both arches are of extreme importance because they help the dentist to execute clinical procedures based on scientific evidences. Therefore, further studies are necessary to compare upper and lower arch whitening degree, as well as the effects of different techniques, application time periods, and bleaching agents.

Conclusion
Our results showed that at-home bleaching technique with 4% hydrogen peroxide for 2 hours per day presents low sensitivity and it is effective. Patient's satisfaction was reached after 27 days of treatment, in average.

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References


