The aim of the present study was to evaluate the effect of prior microabrasion on the teeth color change and tooth bleaching effectiveness. Eight sound molars were mesio-distally sectioned and the halves were randomly allocated to receive enamel microabrasion or non-abrasion (control) in one of surfaces (buccal or lingual), while the remaining surface received the other treatment. The tooth color on baseline was evaluated by spectrophotometer (CieL*a*b system). After the microabrasion procedure, the tooth color was measured again. Following, the specimens were bleached with 35% hydrogen peroxide for two sessions with one-week interval. The color was re-evaluated 7 days after each section and 30 days after the second session. The effect of enamel microabrasion on color changes was evaluated by paired T-test. Deltas L*, a*, b*, and E were calculated and data submitted to 2-way repeated measure ANOVA followed by Tukey’s test. Paired T-test was also used to assess possible differences on the ultimate color achieved after tooth bleaching. Enamel microabrasion reduced the lightness and increased the redness of specimens. Specimens that received microabrasion presented higher values of ∆L* than control after each bleaching procedure; and higher ∆a* after the 2nd bleaching session. However, the prior enamel microabrasion did not affect the ultimate values of color parameters. Despite enamel microabrasion have modified the tooth color, this procedure did not affect the ultimate results achieved with tooth bleaching using a high-concentrated hydrogen peroxide.

Introduction

Tooth discoloration is usually treated with restorative procedures or using more conservative approaches such as tooth bleaching approaches (1,2). The tooth bleaching techniques performed either in office or by patients at home are carried out using bleaching agents based on hydrogen or carbamide peroxides (3-5). The bleaching effect is achieved due to oxidative action of peroxides and its sub-products on the organic matrix of the tooth tissues modifying its optical properties (6,7). Despite the high success rates observed for both in-office and at-home bleaching procedures (4,5), using oxidative agents are unable to solve aesthetic alterations of teeth when the discoloration is intrinsic in the enamel due to fluorosis staining and other enamel hypoplasia, or white lesion spot caused by dental caries (8,9).

In the presence of the enamel alterations, the superficial enamel removal by abrasive product associated to acid etching using the so-called enamel microabrasion is frequently carried out by clinicians to solve tooth discolorations limited to enamel (10-13). The enamel microabrasion procedure is performed by active application of a paste containing an acid (usually phosphoric or hydrochloric), which demineralizes the superficial layers of enamel, and an abrasive such as silicon carbide or pumice to remove the demineralized enamel (11,12). The rationale is to remove the affected outer layer, preserving the unaltered inner enamel. Therefore, the success of enamel microabrasion depends on the depth of enamel alteration. Moreover, the partial removal of enamel within reduction on its thickness increases the light transmission through of enamel, allowing an increased perception of dentin (12,14). Thus, it is expected darker color of teeth submitted to microabrasion due to thinner enamel, similarly to darkening observed on cervical area or by ageing of teeth.

Despite the possible effect the enamel microabrasion on color modification, few studies are found evaluating this outcome following microabrasion procedures. Furthermore, considering a possible tooth darkening due to microabrasion, tooth bleaching procedures using peroxides might be required following the abrasive procedure to reach proper esthetic (15,16). Considering that changes in the enamel caused by tooth bleaching have important role on the ultimate color achieve with this esthetic procedure (17,18), it is important to know about any possible effect of alteration caused in the enamel by microabrasion on efficacy of posterior tooth bleaching. However, to the best of our knowledge, there is no study evaluating the effect of prior enamel microabrasion on effectiveness of following tooth bleaching. Thus, this study aimed to evaluate the color alteration on tooth caused by enamel microabrasion as well its effect on effectiveness of following
tooth bleaching. The null hypotheses tested were that 1) the enamel microabrasion does not intervene on the tooth color; and 2) the tooth bleaching was not affected by prior enamel microabrasion.

Material and Methods

Experimental Design

Hemi-sections of sound molars were submitted to microabrasion, while the opposite hemi-section of the same tooth was used as control. Following, all hemi-sections were submitted to two sessions of tooth bleaching with 35% hydrogen peroxide. The color changes were evaluated using the parameters L*, a*, and b*. The effect of microabrasion on these parameters was analyzed as well as the bleaching effects on color changes regarding to time of assessment in three-levels (after each session and 30 days after the last session) and prior enamel microabrasion in two levels (absent or present).

Sample Preparation and Randomization

Eight sound third molars without cracks or fractures were selected for the study. The molars were sectioned using a diamond saw coupled to a hand-piece on cement-enamel junction; and the roots were excluded. Following, the crowns were mesio-distally sectioned in two halves containing the buccal or lingual enamel surface. For each molar, one hemi-section was randomly allocated to receive enamel microabrasion, while the other one was used as control for this procedure. The randomization was carried out using a randomized list created using a webpage www.sealedenvelope.com.

Baseline Measurements

Baseline color measurements of all specimens were carried-out using a spectrophotometer SP60 (X-Rite, Grand Rapids, MI, USA) based on the Commission Internationale de l’Eclairage (CIE) L*a*b system. The readings were performed under a white background during the entire experiment, while the values of parameters L*, a* and b* were recorded.

Microabrasion Procedures

For the hemi-sections allocated to be submitted to enamel microabrasion, the procedure was performed using 6.6% HCl slurry with silicone carbide micro-particles (Opalustre, Ultradent Products, Inc., South Jordan, UT, USA). The buccal/lingual enamel was air-dried and a small amount of abrasive slurry was applied over the enamel with a rubber cup attached to hand-piece for 60 seconds under low rotation. After remove the abrasive and dry the enamel, the procedure was repeated until to complete 10 applications. After the enamel microabrasion, the color measurements were performed again and the specimens were stored for 24 h under 100% of humidity.

Tooth Bleaching

The bleaching agent based on 35% hydrogen peroxide (HP Whiteness Maxx, FGM, Joinville, SC, Brazil) was manipulated according to manufacturer’s recommendation and applied over the enamel surface of all specimens, remaining for 15 minutes. After this time, the bleaching agent over the surface was replaced to totalize 3 applications (45 minutes). Then, the bleaching agent was completely removed using water-stream, and the specimens stored in distilled water for one week. After this period, the color of the specimens was again measured and a second bleaching session performed, following the procedures described previously. The bleached specimens were stored in distilled water and further color measurements carried-out after 7 and 30 days from the second bleaching session. For each time of assessment, color changes were measured by ∆L*, ∆a* and ∆b* calculation based on data measured at baseline or after the microabrasion for teeth from control and abraded, respectively. Delta E was calculated by following equation: \(\Delta E = \sqrt{(\Delta L)^2 + (\Delta a)^2 + (\Delta b)^2}\).

Statistical Analysis

In order to evaluate the effect of the microabrasion on color changes, data (before and after) of L*, a* and b* from specimens submitted to paired T-test. In order to assess the bleaching effectiveness, data from color changes (∆L*, ∆a*, ∆b* and ΔE) were individually submitted to 2-way repeated measure ANOVA ("time of assessment" as repetition factor) followed by Tukey’s post hoc test. Paired T-tests were used to assess differences on color parameters at baseline and final between the specimens allocated to receive or not enamel microabrasion. A significance level of 95% was set for all statistical analyses.

Results

Table 1 shows the changes on color parameters caused by enamel microabrasion. The microabrasion reduced the lightness and increased the redness of teeth, did not affecting the parameter b*. Table 2 presents the color changes following the bleaching procedures. Regarding ∆L*, 2-way Repeated measured ANOVA showed that both the "moment of assessment" (p < 0.001) and "prior microabrasion" (p = 0.044) affected the tooth lightness, whereas the interaction was not significant (p = 0.491). Reduced alteration on lightness was observed after the 1st session and the highest lightness changes were found 30 days after the second session, irrespective to perform or not prior microabrasion. For all times of assessment, specimens submitted to enamel microabrasion presented higher lightness changes. Data of ∆a* were affected by the "time of assessment" (p < 0.001), whereas the "prior
Discussion

The enamel microabrasion is a usually conservative aesthetic procedure used to treat spot lesions restricted to enamel tissue such as hypoplasia or white spot lesion caused by dental caries. Corroborating with the clinical observations, the enamel removal by microabrasion affected the tooth color, rejecting the first null hypothesis of study. The lightness (L*) of teeth was reduced and redness (a*) increased by the abrasive procedure. This color changes due to erosive ability of the slurry rubbed over the enamel. Despite the thickness modification of enamel was not measured in the present study, it is expected an enamel thickness reduction ranging to 25 to 200 µm due to microabrasion. A thinner enamel allows higher light transmission through of this tissue, increasing the visualization of underlay dentin and reducing the lightness and increasing the redness.

Once that the patients are frequently requiring whiter teeth, the reduction on lightness and increased redness caused by enamel microabrasion might require an additional aesthetic intervention, while the tooth bleaching is the more conservative approach. In the present study a high-concentrated hydrogen peroxide based bleaching agent was used simulating an in-office tooth bleaching procedure. Despite the at-home bleaching agents are largely used by patients with successful, the in-office bleaching still recommended for some clinical situations including patients presenting gingival recession, gastric disorders, or reduced salivary flowing. Moreover, using an in-office bleaching agent with hydrogen peroxide activation by its mixing with the thickener facilitates the experimental protocol reducing bias when compared to at-home products, while the peroxide activation depends on pH buffering by saliva. The action mechanism of bleaching agents is related to oxy-reduction reaction of peroxides within stain molecules located in the organic matrix, while these molecules become simpler and whiter. The results of the presents study found that the bleaching effect was mainly related to changes on lightness (L*) and redness (a*), while slight alteration on yellowness was observed. Teeth submitted to prior microabrasion presented higher values of ∆L* than those observed for specimens from control condition at all times of assessment. These results can be explained due to differences at color presented by specimens prior to tooth bleaching, while the teeth submitted to microabrasion presented reduced lightness. It has been demonstrated that the bleaching effect is

Table 1. Means (standard deviation) for each color parameter changes following the enamel microabrasion procedure (n=8)

<table>
<thead>
<tr>
<th>Time of assessment</th>
<th>L*</th>
<th>a*</th>
<th>b*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>87.5  (1.8)</td>
<td>2.5   (0.8)</td>
<td>10.3  (2.8)</td>
</tr>
<tr>
<td>After microabrasion</td>
<td>85.8  (1.7)</td>
<td>3.1   (1.0)</td>
<td>10.8  (4.3)</td>
</tr>
<tr>
<td>p-value*</td>
<td>0.001</td>
<td>0.032</td>
<td>0.706</td>
</tr>
</tbody>
</table>

*Calculated by paired t-test.

Table 2. Means (standard deviation) of changes on color parameters following the bleaching procedures (n = 8)

<table>
<thead>
<tr>
<th>Time of assessment</th>
<th>L*</th>
<th>a*</th>
<th>b*</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abraded</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>After 1st session</td>
<td>1.3   (1.0)</td>
<td>-1.1   (0.4)</td>
<td>-2.0   (2.0)</td>
<td>3.1   (1.5)</td>
</tr>
<tr>
<td></td>
<td>b&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Ab</td>
<td>Ab</td>
<td>Ac</td>
</tr>
<tr>
<td>After 2nd session</td>
<td>2.9   (1.3)</td>
<td>-1.7   (0.7)</td>
<td>-1.3   (3.0)</td>
<td>4.7   (1.0)</td>
</tr>
<tr>
<td></td>
<td>b&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Ab</td>
<td>Ab</td>
<td>Ab</td>
</tr>
<tr>
<td>Final</td>
<td>4.4   (1.7)</td>
<td>-2.0   (0.5)</td>
<td>-4.5   (2.2)</td>
<td>6.8   (1.9)</td>
</tr>
<tr>
<td></td>
<td>b&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Ab</td>
<td>Ab</td>
<td>Ac</td>
</tr>
</tbody>
</table>

Different letters (uppercase for rows, lowercase for columns) indicate statistically significant difference (p<0.05)
limited and usually more pronounced when carried out in darker teeth (4). This fact also explains the higher changes on parameter a* observed for abraded teeth following the 2nd session of tooth bleaching.

Beside to the differences on color change following the bleaching procedures, the results of the present study

Figure 1. Behavior of color parameters (L* - A; a* - C; and b* - E) during the experiment according to presence or absence of prior microabrasion; and comparison between the color parameters (L* - B; a* - D; and b* - F) at baseline and at final of experiment from teeth allocated to receive or not microabrasion. Bars indicate standard error.
demonstrated that the prior microabrasion did not affect the ultimate color achieved by tooth bleaching with hydrogen peroxide. Thus, the second hypothesis was accepted. In fact, the values of all color parameters were similar at the last assessment. It is reasonable to think that slower bleaching could occur for the abraded teeth due to darker color due to prior microabrasion. However, the results demonstrated similar color (absence on difference for all color parameter) between the abraded and control specimens after the 1st session of tooth bleaching. Moreover, additional bleaching effect was achieved with the 2nd session and within the 30 days following this last peroxide application. This last change color was mainly related to increasing on lightness and reduction on yellowness, whereas the no significant modification on redness was observed. The measurement of color after 30 days was chosen to determine any residual effect of tooth bleaching and to simulate clinical conditions when a rebound effect has been demonstrated (3,24). The rebound effect tends to slight reduce the effect reached by tooth bleaching and is mainly due to physiological response to pulpal damage. On the other hand, this effect is not expected using ex-vivo teeth and the color changes following the storage time might be associated to any possible residual effect of peroxide remaining in the hard tooth tissue (25,26). It is hypothesized that the absence of pulp pressure in extracted teeth can reduce the removal of peroxide and its products, mainly than located inside the dentinal tubules; while these components can produce additional bleaching effect (27).

The outcomes of the present study demonstrated that the enamel microabrasion did not affect the bleaching effectiveness, despite to reduce the lightness and increase the redness of teeth. One important limitation of method used was that teeth without any enamel discoloration (i.e. fluorosis, white spot lesion) were used. Further to facility to obtain specimens, using teeth without any enamel discoloration helps to standardize the baseline measurement. However, it is difficult to determine if the enamel removal by microabrasion techniques performed would be enough to eliminate any kind of enamel alteration. Moreover, measuring the enamel thickness and its reduction following the microabrasion procedure can help to associate this alteration with the color changes. Another important limitation of study was that the specimens were stored in distilled water instead artificial saliva after the enamel microabrasion and tooth bleaching procedures. Both enamel microabrasion and tooth bleaching are able to increase the surface roughness due to acid pH of slurry (pH of Opalustre is around 0.2) and bleaching agent (pH of Whitenees HP Maxx range from 5.6 to 6) used in the present study (19,28). Considering that changes on surface roughness can modify the light reflection and color measurement, mainly related to lightness parameter, enamel remineralizing by specimens storage in artificial saliva could to reduce any possible bias related to methodology. Thus, further studies seeking out to turn these reported limitations might be required to confirm the findings of the present study.

**References**


