Toothbrushing abrasion susceptibility of enamel and dentin bleached with calcium-supplemented hydrogen peroxide gel


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ABSTRACT

The objective of this study was to evaluate enamel and dentin susceptibility to toothbrushing abrasion, after bleaching with 7.5% hydrogen peroxide (HP) gel supplemented or not with 0.5% calcium gluconate (Ca). Toothbrushing was performed immediately and 1 h after bleaching, with two suspensions (high and low abrasiveness). Bovine enamel and dentin specimens were divided into 12 groups (n=10) according to the bleaching gel (with and without Ca), slurry abrasivity (high or low) and elapsed time after bleaching (immediately and after 1 h). As control, a group was not bleached, but abraded. The treatment cycle (7 d) consisted of bleaching (1 h) and toothbrushing (135 strokes/day) immediately or after 1 h of artificial saliva exposure. Surface roughness and surface loss (μm) were measured by profilometry and analysed by three-way ANOVA (5%). Surface roughness means were significantly influenced by slurry abrasivity (p<0.0001). For enamel loss, significant triple interaction was observed (p<0.0001). HP-bleached groups and immediately brushed with high-abrasive slurry exhibited increased loss (1.41±0.14) compared to other groups (μm). Control and HP + Ca-bleached groups brushed after 1 h with low-abrasive slurry presented the lowest loss (0.21±0.03/0.27±0.02). For dentin loss, significant interaction was observed for bleaching and interval factors (p<0.001). 7.5%HP-bleached groups and immediately brushed showed significantly higher loss (8.71±2.45) than the other groups. It was concluded that surface roughness increased when high-abrasive was used, independently of bleaching. 7.5%HP increased enamel and dentin loss, mainly with high-abrasive slurries. Calcium supplementation of bleaching gel reduced surface loss. Additionally, in order to minimize tooth wear susceptibility, it is recommended to delay brushing after bleaching.

Clinical relevance: After bleaching gel application, postponing toothbrushing is recommended, as well as brushing with low-abrasive dentifrices. Additionally, supplementation of hydrogen peroxide gel with calcium-based remineralizing agent potentially reduces tooth loss after abrasion.

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1. Introduction

Hydrogen peroxide has been widely used as a treatment agent for discolored teeth. Due to its low molecular weight and high instability, hydrogen peroxide is able to diffuse through enamel and dentin and decompose, releasing free radicals [1,2]. Its mechanism of action is based on the oxidative destruction of chromophores. This occurs by chemical degradation of molecular moieties responsible for absorbing visible electromagnetic radiation, resulting in the increase of total reflectance of the substrate, and consequently, in its brighter appearance [3].

Although studies have proven the whitening efficacy of bleaching agents, adverse effects on dental tissues are also reported and must be carefully evaluated so that their use can be considered safe [4,5].

Dental bleaching has been previously related to microstructural tooth alterations, such as microhardness reduction [6] and changes in chemical composition of tooth [7]. Morphology defects, such as irregularities, depressions and porosity formation have also been reported [8]. Although these changes were usually assigned to mineral loss due to the low pH of bleaching gels [4], degradation of
the organic matrix by oxidation reaction was reported [9], with the
described alterations also observed when near neutral agents were
used [7,8]. In order to reduce the bleaching gel demineralization
potential, the addition of remineralizing agents such as calcium
and fluoride in the bleaching agents has been proposed [10–12].

Daily oral hygiene (brushing with regular toothpaste) is usually
considered safe and not related to enamel potential harm [13].
However, individuals undergoing bleaching treatment may asso-
ciate it with high abrasivity whitening toothpastes, potentially
increasing its harmful effects on tooth surface [14]. Although
abrasivity represents the predominantly mode of action of
whitening toothpastes [15], it is known that the degree of
dentifrice abrasivity may be related to potential wear, mainly on
dentin [16]. Previous studies have reported a higher wear
susceptibility of bleached enamel [17] and dentin after brushing
[18–20].

Additionally, the effect of toothbrushing abrasion after
bleaching on tooth surface roughness has been an issue of
concern, since alterations on surface texture can lead to increased
susceptibility to staining and bacterial adhesion, and consequent-
ly, further discoloration [21,22].

It has been shown that saliva is able to reharden demineralized
bleached enamel [11]. Additionally, the supplementation of
bleaching agents with remineralizing agents could contribute to
reduce any potential harmful effect on bleached tooth [8]. Since
bleaching may be related to tooth surface microstructural
alterations and its association with toothbrushing can potentially
increase surface loss, there is a concern regarding the optimal
interval between the removal of the whitening gel and brushing,
in order to reduce the possible interaction between the bleaching
agent and the abrasive process on dental tissues.

Thus, the aim of this in vitro study was to investigate if the
elapsed time between bleaching with 7.5% hydrogen peroxide
(associated or not with calcium) and brushing (with high and low
abrasivity slurries) would affect the enamel and dentin roughness
and wear. The null hypotheses tested were that: a) 7.5% hydrogen
peroxide, associated or not with calcium would not affect the
substrate surface roughness and its susceptibility to abrasive wear;
b) the elapsed time between bleaching and brushing would not
interfere with roughness and wear, and; c) the slurry abrasivity
would not influence roughness and wear.

2. Materials and methods

2.1. Experimental design

This study followed the complete factorial $3 \times 2 \times 2$ randomized
design, with three experimental factors: 1. bleaching at 3 levels (no
bleach, 7.5% hydrogen peroxide-HP, and HP with the addition of
0.5% calcium gluconate); 2. slurry abrasivity at 2 levels according to
the RDA values (high and low); and 3. elapsed time between
bleaching and abrasion at 2 levels (immediately and 1 h after
bleaching), in a bleaching- abrasion cycling model using bovine
enamel and dentin specimens. The specimens were randomly
assigned into 12 groups ($n = 10$). The model was conducted for a
total of seven consecutive days, and response variables were
arithmetic mean surface roughness (Ra) and surface loss (in $\mu m$)
measured by contact profilometry.

2.2. Enamel and dentin specimens preparation

Freshly extracted and intact bovine incisors were stored until
required in 0.1% thymol solution, refrigerated at 4 °C. Cylindrical
enamel and dentin specimens (3 mm diameter) were prepared
from the labial surfaces of crowns and roots, respectively, using a
custom-made diamond trephine mill. The specimens were then
embedded in auto-polymerizable acrylic resin using cylindrical
silicone molds (6 mm diameter, 3 mm depth), with the labial
surface exposed for treatments, as previously described [23].

Embedded specimens were ground flat and polished with
water-cooled sequential aluminum oxide abrasive papers (1200,
2400 and 4000 grit FEPA P; Struers, Ballerup, Denmark) in a
polishing device (DP 10, Panambara, Sao Paulo, SP, Brazil). After
each grindpaper, specimens were sonicated in deionized water for
5 min. The prepared specimens were examined in stereomicro-
scope (20X–Carl Zeiss, Stemi 2000, Tokyo, Japan) to verify
the absence of cracks or other surface defects and then stored in
ultrapure water to prevent dehydration.

The baseline profiles of the enamel and dentin surfaces
were measured using a contact profilometer (MarSurf GD 25, Mahr,
Göttingen, Germany). In order to maintain the reference surfaces
for lesion-depth determination, and allow the exact superimpo-
sition of the baseline and post-treatment profiles, two parallel
grooves were marked as guides on the resin at the sides of the
embedded tooth structure. The specimens were positioned into a
custom-made specimen holder attached to the profilometer,
which allows the exact repositioning of the sample after the
treatments. The diamond stylus moved from the first reference
(resin) to the enamel or dentin area and then over to the other
reference area (4.2 mm long). Three profile measurements
were performed for each specimen at intervals of 0.25 mm.

For baseline superficial roughness analysis, the mean surface
roughness values (Ra) were determined with a cut-off value of
0.8 mm, a transverse length of 0.8 mm, and a stylus speed of
0.1 mm/s, in the previously described profiles.

2.3. Bleaching and abrasive procedures

The specimens were randomly allocated into 12 groups ($n = 10$).
The first group division was according to the bleaching gel: NB- no
bleach; HP- 7.5% hydrogen peroxide gel (pH 5.62); HP + Ca- 7.5% HP
gel with the addition of 0.5% calcium gluconate (pH 5.60). The
experimental gels were modified by the manufacturer (FGM,
Joinville, SC, Brazil), by adding or not the calcium compound in a
7.5% HP-based bleaching gel.

A 2 mm thick layer of the bleaching gel was daily applied to the
specimens’ surface and remained for 1 h. After this period, the gel
was removed with a suction tip and the surface rinsed with
ultrapure water for 20 s. In the non-bleached groups, the speci-
mens remained in ultrapure water during the period corresponding
to bleaching procedure.

After the described procedures, the specimens of each described
group were divided according to the elapsed time after bleaching
into two subgroups: immediately and one hour after bleaching for
performing toothbrushing. The specimens of the group 1 h were
kept in artificial saliva (1.5 mM CaCl$_2$ $\times$ $2$H$_2$O; 0.9 mM K$_2$HPO$_4$;
130 mM KCl; 20 mM of HEPES; pH adjusted to 7.0 with 1 M KOH
solution) [24].

The specimens were finally divided into two subgroups according
to the slurry abrasivity: H- high and L- low. The abrasive
callenge was performed using an automatic toothbrushing
machine (SEM-2T, Odeme Dental Research, Luzerna, SC, Brazil),
which imparted reciprocating motion to standard medium bristle
toothbrush stiffness (Sanfill Ultraprofissional, Sao Paulo, Brazil).
The brushes were angled 15° in relation to the specimen surface to
minimize grooves formation. During brushing, the right and left
sides of the specimens, corresponding to acrylic resin with the
reference groves, were protected with a stainless steel mask (0.1-
mm thick), with an opened window of 2-mm wide, leaving an
exposed area in the center of each specimen, preventing the
abrasion of reference areas for the profilometric analysis.
Daily, 135 brushing strokes under a vertical load of 200 g with abrasive slurries were administered, simulating 45 strokes each time, three times a day [25]. The slurries were prepared by mixing the silica abrasives Zeodont 113 [low abrasive, with RDA standard-error 86.92 [1.54] and REA standard-error 4.45 [0.46], J.M. Huber, Etowah, TN, USA] or Zeodont 103 [high abrasive, RDA standard-error 204.07 [5.21] and REA standard-error 5.49 [0.43], J.M. Huber] with a 5% carboxymethylcellulose solution in glycerol. An abrasive concentration of 10% was used [25]. The bleaching/abrasion procedures were repeated for 7 days.

### 2.4. Surface loss and roughness analysis

After the end of treatment, the final profiles of the surfaces were measured using the contact profilometer following the same parameters previously described. Final surface roughness was determined with the same specifications described before. The surface loss was calculated after superimposing the baseline and post-treatment profiles, using the previously described grooves as guides. The depth of the treated area for each specimen was calculated based on the subtraction of the two profiles, using a dedicated software (MarSurf–XCR 20 4.50-07 SP3, 2011). The background noise, used for calculation of the detection limit of the equipment, was recorded by measuring the vertical displacement of the stationary stylus for 20 s, according to the methodology described by Attin et al. [26]. This determined 0.132 μm as the lower limit of measurement. Therefore, readings below this value are considered “below detection limit”.

### 2.5. Statistical analysis

The difference between final and initial enamel and dentin values of arithmetic mean surface roughness (Ra) and surface loss (μm) from the surface of the enamel and dentin at the end of the experiment served as evaluation parameters. ANOVA three-way and post-hoc Tukey tests were used for both data. The level of significance was 5%.

### 3. Results

The mean values of initial and final surface roughness for both enamel and dentin substrates, as well as the roughness change are shown in Table 1.

Three-way ANOVA was applied both on enamel and dentin Ra alteration data (final-initial) and showed differences only for the abrasive factor (p < 0.001). The factors bleaching treatment (p = 0.41 enamel/p = 0.15 dentin) and interval post-bleaching (p = 0.18 enamel/p = 0.50 dentin) did not present significant differences, as well as the three-factors interaction (p = 0.88 enamel/p = 0.20 dentin). Brushing with the high abrasive slurry resulted in enamel and dentin Ra means significantly higher than with the low abrasive one.

The enamel surface loss analysis was also performed using three-way ANOVA, which showed significant differences for all factors (p < 0.001), and for the triple interaction (p < 0.001). Tukey’s test results are presented in Fig. 1 and showed that bleaching with 7.5% HP resulted in higher enamel loss lost compared to 7.5% HP + Ca and control groups, for both immediate and after 1 h interval. The means obtained with high abrasive were higher than the same groups brushed with low abrasive slurries. The immersion of specimens in artificial saliva for 1 h after bleaching resulted in lower enamel loss compared to the immediate groups.

For dentin, there were significant differences for the three distinct factors (p < 0.001) and for interaction between bleaching and interval factors (p < 0.001), shown in Fig. 1. The results of Tukey test showed that the groups bleached with 7.5% HP and immediately brushed showed significantly higher loss than the other groups tested (asterisks in Fig. 2).

### 4. Discussion

As the bleaching procedure is an esthetic treatment, all consequences that could affect the outcome are relevant. Surface loss potentially caused by brushing can cause damage to enamel and dentin during the years, weakening the tooth structure. Additionally, the effect of the bleaching treatment over the surface

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**Table 1**

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Interval</th>
<th>Abrasive</th>
<th>Enamel</th>
<th>Dentin</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Ra initial</td>
<td>Ra final</td>
</tr>
<tr>
<td>No bleaching</td>
<td>Immed.</td>
<td>High</td>
<td>0.025 (0.006)</td>
<td>0.088 (0.042)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low</td>
<td>0.024 (0.005)</td>
<td>0.043 (0.018)</td>
</tr>
<tr>
<td></td>
<td>After 1 h</td>
<td>High</td>
<td>0.022 (0.005)</td>
<td>0.078 (0.034)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low</td>
<td>0.022 (0.007)</td>
<td>0.040 (0.032)</td>
</tr>
<tr>
<td>7.5% HP</td>
<td>Immed.</td>
<td>High</td>
<td>0.021 (0.007)</td>
<td>0.090 (0.035)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low</td>
<td>0.021 (0.006)</td>
<td>0.050 (0.025)</td>
</tr>
<tr>
<td>7.5% HP + Ca</td>
<td>Immed.</td>
<td>High</td>
<td>0.024 (0.006)</td>
<td>0.088 (0.017)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Low</td>
<td>0.029 (0.004)</td>
<td>0.043 (0.038)</td>
</tr>
<tr>
<td></td>
<td>After 1 h</td>
<td>High</td>
<td>0.023 (0.008)</td>
<td>0.070 (0.030)</td>
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<td></td>
<td></td>
<td>Low</td>
<td>0.026 (0.005)</td>
<td>0.041 (0.027)</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>(0.007)</td>
<td>(0.017)</td>
</tr>
</tbody>
</table>
roughness can produce a more immediate negative esthetic effect, such as the formation of a scratchy surface, with increased susceptibility to extrinsic stains deposition [21]. Besides, roughness values as low as 0.25 μm can be perceived by the tongue, potentially causing discomfort to the patient [27]. Another consequence would be the biofilm deposition over rough surfaces [22]. A threshold surface roughness for bacterial retention of 0.2 μm has been reported [28].

In this in vitro study, the overall enamel and dentin surface roughness after abrasion associated or not with bleaching significantly increased compared to baseline polished specimens data. Additionally, the roughness alteration caused by brushing with high abrasive slurry was higher when a low abrasive dentifrice simulation was tested. Nevertheless, the bleaching procedures did not exacerbate the brushing effects on enamel and dentin surface roughness. Thus, only the second null hypothesis tested regarding abrasivity was rejected.

The deleterious effect of bleaching gels on tooth structures is an issue of concern, since it is known that hydrogen peroxide agents are able to penetrate enamel structure through the boundaries between nanocrystals and decompose, releasing free radicals [1]. These radicals are unspecific and act not only on cromogens, but can cause the breakage of the polypeptide chain by means of destruction of aminoacids present in tooth organic matrix [9,29]. As a consequence, several authors reported alterations on both enamel and dentin morphology and chemical composition [7,8]. Since the organic content participates on enamel integrity, these changes may result in mineral content changes [9].

Nevertheless, the effect of peroxide-based gels on enamel and dentin surface roughness is controversial, as some authors reported increased roughness after bleaching [30] and others did not observe significant differences [31,32]. Comparison among the studies is difficult, since bleaching agents, concentrations and protocols are different. It is important to note that the remineralization action of saliva is able to minimize the previously reported deleterious effect of bleaching procedures on tooth structure [4,5,11].

When bleaching was associated with toothbrushing abrasion, increased enamel roughness has been reported [33–35]. Nevertheless, we found that, regardless the bleaching treatment, the roughness alteration was determined by the toothbrushing procedure, modulated by the slurry abrasivity. Similar results were also obtained previously for bleached/abraded enamel [36]. Toothpaste abrasivity is mainly based on a radiotracer method, which provides relative abrasivity of dentin and enamel (RDA and REA), comparing them to a reference abrasive [37]. It has to be highlighted that although we classified as low and high abrasive, both slurries abrasivity were within the recommended range of ISO.
(bellow RDA 250) [38]. On the other hand, while the toothpastes usually contain fluoride, in this study the slurries were not fluoridated, as this could represent a confounding factor for the calcium-supplemented bleaching agent, potentially interfering with the remineralization action. Additionally, it was previously demonstrated that fluoridation of dentifrice did not alter its potential to affect enamel roughness [33].

Since abrasivity was the determining factor for surface roughness, postponing the brushing procedure 1 h after bleaching did not exert a protective effect for minimizing the surface roughness effects, contrarily as reported previously [39]. The irregularities caused by brushing probably were coarser than the supposed repairing action of saliva over the surface morphologic irregularities of bleached enamel and dentin.

Although the alterations in surface roughness can influence esthetic results, it is probably less clinically relevant than the potential wear process that can be related to toothbrushing associated to hydrogen peroxide use. In this study, it was overall observed that bleaching with 7.5% hydrogen peroxide associated with abrasive challenge for seven days resulted in significantly higher enamel and dentin loss than the non-bleached control group and the 7.5% + Ca bleached group. Therefore, the first null hypothesis was rejected for both substrates.

Although the effect of bleaching agents did not significantly affect surface roughness, probably because the slurry abrasivity effect overcomes the demineralization action of bleaching agents, it increased surface loss. It is speculated that the potentially demineralized surface may have been more easily removed by toothbrushing abrasion. Increased abrasive wear has been associated to bleached enamel [17] and especially to bleached dentin [19,20]. Similarly, the abrasion resistance of dental hard tissues was shown to be reduced in demineralized surfaces [38] and bleached enamel abrasive susceptibility has been related to peroxide concentration [17]. During bleaching procedures, enamel can lose calcium and phosphate ions to the oral environment [7,30]. Although demineralization potential of bleaching agents is mainly related to its pH [4,6], it was already demonstrated that its oxidative effect is also related to subsequent mineral loss [9].

The addition of remineralizing agents in a bleaching gel can minimize its demineralization potential and possibly promote ions incorporation in the demineralized enamel [10,12]. Although calcium and phosphate ions needed for remineralization process are naturally present in saliva, the presence of calcium in bleaching gel aims to act as a supplementary source of these ions in the tooth surface, maintaining its saturation during bleaching [10,12]. Calcium gluconate was added in order to minimize the demineralization potential of the bleaching gel and eventually contribute to the desensitizing effect. The salt is soluble at the conditions needed for the bleaching gel production and is easily found in adequate chemical purity [40]. According to the manufacturer (FGM), the concentration of 0.5% was determined during the product development process, in order to maintain its adequate viscosity and stability. The protective effect of calcium supplementation in bleaching gel was also demonstrated for bleached enamel further exposed to acidic solutions [23,41]. Similarly, different calcium compounds have also been added to acidic beverages in order to reduce enamel wear when subjected to erosion [42].

The remineralization potential of saliva over demineralized dental hard tissues could eventually reduce the abrasive wear, although this is a very controversial issue and a definite protective time (if really relevant) has not yet been established. Waiting periods of at least 1 h after demineralization by erosion has shown to slightly reduce enamel abrasion [43,44], and therefore used in this study. Our results showed that bleached substrates abrased 1 h post-bleaching generally exhibited reduced loss compared to immediately brushed. It has to be pointed out that, when present, the demineralization pattern promoted by bleaching product are supposed to be milder, since it has a different origin compared to acid-exposed eroded substrate. The artificial saliva may have restored the potentially surface-damaged bleached structure, reducing substrate abrasion susceptibility. On the other hand, the absence of presence of proteins in artificial saliva may have favored saliva-remineralizing properties, limiting the comparison with in situ studies [45].

In relation to slurry abrasivity, increased enamel and dentin loss was observed when higher abrasive slurry was used. The in vitro correlation of tooth wear with dentifrice abrasivity has been previously demonstrated [46]. Nevertheless, only dentin-increased abrasion has been related with toothpaste abrasivity, but associated with erosion [47] or with bleaching/erosion [18]. It has to be noted that enamel loss observed was negligible compared to dentin loss, and may not be clinically worrying. Although bleaching agents should not be applied over exposed dentin, occasionally it can reach the partially exposed root dentin when gingival retraction is present.

The in vitro nature of the present study should be taken into account, since it does not consider the presence of human saliva and acquired pellicle formation. It was demonstrated that the calcium, phosphate and fluoride content of human saliva may be responsible for maintaining mineral content and inhibit enamel demineralization during bleaching treatment [4] or promote remineralization due to the saturation of mineral components [5,11]. Additionally, it must be highlighted that the profilometry methodology demands polished specimens. The removal of native surface may favor demineralization process and exacerbate test results [48]. Thus, extrapolation of the results to clinical reality must be carefully performed.

5. Conclusion

Considering the limitations of this in vitro study, it can be concluded that surface roughness was only influenced by slurry abrasivity. 7.5% hydrogen peroxide bleaching may increase enamel and dentin susceptibility to abrasion, mainly with high abrasive slurries. Supplementation of bleaching gel with calcium and delaying brushing for one hour after bleaching reduced surface loss.

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