Dental Pulp Vascular Permeability Changes Induced by Dental Bleaching

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ABSTRACT

Aiming to compare the effect of different light sources for dental bleaching on vascular permeability of dental pulps, forty-eight incisors were used. The bleaching agent (35 % hydrogen peroxide) was activated by halogen light; LED (Light Emitting Diode) or LED, followed by laser phototherapy (LPT) (λ = 780 nm; 3 J/cm²). After the bleaching procedures, the animals received an intra-arterial dye injection and one hour later were sacrificed. The teeth were diaphanized and photographed. The amount of blue stain content of each dental pulp was quantified using a computer imaging program. The data was statistically compared (p ≤ 0.05). The results showed a significant higher (p ≤ 0.01) dye content in the groups bleached with halogen light, compared with the control, LED and LED plus LPT groups. Thus, tooth bleaching activated by LED or LED plus LPT induces lesser resulted in increased vascular permeability than halogen light.

Key words: dental pulp, laser, LED, tooth bleaching, vascular permeability

INTRODUCTION

The concern with aesthetics has made tooth whitening is one of the most required procedures in the dental clinic. The tooth bleaching can be performed internally on nonvital teeth or externally on vital teeth. Hydrogen peroxide, sodium perborate, and cabamide peroxide are generally used for bleaching (Auschill et al. 2005; Zalkind et al. 1996). The application of light sources such as halogen light, LED (Light Emitting Diode) and lasers (Dederich and Bushick 2004; Stabholz et al. 2003) can increase the rate of bleaching agent degradation and the speed of the bleaching process. These procedures have high effectiveness in bleaching, but may have undesirable effects, such as change in tooth structure (Rotstein et al. 1996), external root resorption (Crim 1992), dental pulp inflammation (Bowles and Thompson, 1986; Robertson and Melfi 1980; Seale et al. 1981) and dentin sensitivity (Dahl and Pallesen 2003; Sulieman 2004).

High incidence of tooth sensitivity (from 67 to 78%) has being reported after in-office bleaching with hydrogen peroxide in combination with heat (Nathanson and Parra 1987). Thus, heating along with penetration of bleaching agent into the dental pulp can trigger reversible inflammation (Robertson and Melfi 1980; Seale and Wilson 1985). Phototherapy with low intensity laser (LPT) is used in various areas of biological sciences to promote the tissue regeneration of injured tissues.
This therapy results in analgesic, antiinflammatory and biomodulatory effects (Ladarlado et al. 2004; Reddy 2004; Silveira et al. 2007). The laser light within the red visible and near infrared wavelengths corresponds to the energy absorption spectrum of the respiratory chain components, increasing the cellular metabolism of organisms under stress conditions (Robertson and Melfi 1980; Silveira et al. 2007). Moreover, the phototherapy with low intensity laser using infrared laser is able to compensate the cytotoxic effects of substances released by 35 % hydrogen peroxide bleaching gel (Dantas et al. 2010).

Light sources are used for speeding the in-office dental bleaching and can cause tooth sensitivity probably due to reversible inflammation in the dental pulp. Knowing that LPT can modulate the inflammatory process, it would be interesting to analyze the effect of these light sources on the intrapulpal initial signs of inflammation after in-office dental bleaching. Then, this study aimed to analyze the vascular permeability of dental pulps after in-office dental bleaching using different light sources.

**MATERIALS AND METHODS**

For the *in vivo* study, 12 adult male Wistar rats (*Rattus norvegicus albinus*) weighing an average of 180 to 200 g were used. All the animals were kept under constant temperature and with free access to food and water. The rats were anesthetized prior to the dental bleaching procedures using a mixture of 2% Xylazine (Roncum®, Bayer do Brasil, SP, Brazil) and 5% Ketamine (Ketalar®, Cristália, SP, Brazil) (0.15 mL per 100 g of rat bodyweight). Four incisors of each rat were used for the experiments. Thus, 48 incisors were divided into four experimental groups (n=12 per group), as follows: 1) Conventional method (n=12): Halogen light 2) LED (n=12) 3) LED plus LPT (n=12) 4) Control (n=12): Non-treated

The bleaching procedures were performed using 35 % hydrogen peroxide agent in 1, 2 or 3 sessions, as shown in Table 1.

| Table 1 - Distribution of the experimental groups in function of the sessions: |
|-----------------|-----------------|-----------------|-----------------|
| Groups          | 1 Session        | 2 Sessions      | 3 Sessions      |
| Conventional method (Halogen light) | G1 (n=4) | G4 (n=4) | G7 (n=4) |
|                 | 3 minutes        | 3 minutes (each session) | 3 minutes (each session) |
| LED             | G2 (n=4)        | G5 (n=4) | G8 (n=4) |
|                 | 3 minutes        | 3 minutes (each session) | 3 minutes (each session) |
| LED plus laser  | G3 (n=4) | G6 (n=4) | G9 (n=4) |
|                 | LED + LPT (3 seconds) | LED + LPT (3 seconds) | LED + LPT (3 seconds) |
|                 | (each session) with 2 days of interval | (each session) with 2 days of interval | (each session) with 2 days of interval |
| Control         | G10 (n=12)       | Non-treated: not submitted to the bleaching gels neither to the light sources |

**Experiments**

The anesthetized animals were fixed on an operational table. Dental isolation and prophylaxis was performed previously to the bleaching process using 35% hydrogen peroxide associated to the different light sources tested. For the conventional treatment (G1, G4 and G7), a 600 mW/cm² halogen light (Dabi Atlante, Ribeirão Preto, SP, Brazil) was used for three minutes. For the LED group (G2, G5 and G8), a 470 nm LED, (Kondortech, São Carlos, SP, Brazil) was used for three minutes. The LPT after the LED application was performed at the groups LED plus LPT (G3, G6 and G9) using a GaAsAl diode laser at the wavelength of 780 nm and energy density of 3 J/cm², for three seconds. The time of bleaching was determined in a pilot study. Immediately after the last tooth bleaching session, the animals were submitted to the dye application as previously described (Duval-Araújo et al. 1994). Briefly, the dye (0.6 ml of Nankin ink) was injected in the rat blood circulation using the lateral caudal artery.
After one hour, time enough for the complete excretion of the dye from the vascular system, the animals were sacrificed by anesthetic overdose and the jaw and the maxilla were removed. The specimens were processed to obtain the diaphanized samples. Briefly, the samples were immersed into 2.5 % sodium hypochlorite solution for seven days, followed for washing in running water for 24 h. Then, the samples were immersed into 5 % chloride acid solution for five days with daily change. After this procedure, they were washed in the running water for 24 h and then immersed into a crescent series of ethanol (70, 80 and 96 %) during 24 h in each solution. At the end, they were immersed into methyl salicilate and the incisor crowns were removed. After the diaphanization, the optical density of each crown was recorded using an imaging program (Imagelab 2000, São Paulo, SP, Brazil). This data indicated the amount of dye leaked from the vascular system that remained in the dental pulp tissue after the ink had been physiologically removed from the vascular system of the rats. Thus, this indirectly indicated the changes in the vascular permeability of the dental pulps of the teeth submitted or not to the bleaching procedures.

Statistical analysis
Each data point corresponded to the mean ± SEM (standard error of the mean) of the optical densities of four teeth per experimental group. The data were compared by ANOVA complemented with the Tukey’s test. The level of significance was defined as 5 % (p ≤ 0.05).

RESULTS
The results are illustrated in Figure 1A and represented in the graph of Figure 1B. The highest dye content values were observed in the group of dental bleaching using the conventional technique (p<0.01). The groups treated with LED and LED plus LPT presented similar amounts of dye.
Independently of the light sources applied during the dental bleaching, the amount of dye in the crowns was significantly higher in the samples bleached in the three sessions. In the group treated with the conventional bleaching procedure using the halogen light source, the dye increased proportionally in function of the number of bleaching sessions; however, when using LED and LED plus LPT, there was no difference between the data obtained after 1 and 2 sessions.

**DISCUSSION**

The undesirable side effects of in-office dental bleaching using the light sources for accelerating the dental bleaching process is still a matter of concern, especially due to the increased request of the general population for this dental procedure. The most common side effects of the in-office dental bleaching is tooth sensitivity, which could be due to the penetration of bleaching agent into the dental pulp and/or the increase of the pulp temperature. Thus, this study analyzed the effect of an in-office dental bleaching using 35 % hydrogen peroxide and different light sources [halogen light, LED and LED plus laser phototherapy (LPT)] on the pulp vascular permeability that is one of the first signs of inflammation. As expected, the teeth bleached with halogen light, that caused the highest temperature increase (Carrasco et al. 2008; Cobb et al. 2000), presented the more expressive increase in the dental pulp vascular permeability, whereas LED and LED plus LPT caused only small vascular permeability increases.

In order to measure the changes in the vascular permeability in the dental pulp of the teeth subjected to in-office dental bleaching with different light sources, a circulating dye was used (Duval-Araújo et al. 1994). By means of this methodology, it was possible to observe the presence of the dye particles on the dental pulp tissue where alterations of the vascular permeability occurred. The dye run inside the vascular system of the rats and as effect of vascular flow, one hour after the dye application, all dye was excluded of the vascular stream. However, in the tissues where the vascular permeability increased, the passage of the dye through the vascular walls occurred. Then, the dye reached the extra vascular environment and the tissue was permanently stained. Next, the teeth were subjected to diaphanization for revealing the dye at the dental pulp. The amount of dye in the dental pulp tissue indicated the level of edema caused by the in-office dental bleaching procedure applied. In this study, different amount of dye were observed in all the bleached teeth, showing that there was increases in the vascular permeability in all the groups.

The 35% hydrogen peroxide is one of the more usual active agents used for bleaching. This product has a thermally unstable power oxidative, dissociating in the water, oxygen and free radicals. These low molecular weight radicals are able to increase the porosity of the enamel and degrade the complex organic molecules responsible for the change in tooth color, reducing and eliminating the pigment responsible for darkening the tooth (Goldstein et al. 1989; Goodis et al. 1989; Hanning and Bott 1999; Hanks et al. 1993; Kawamoto and Tsujimoto 2004; Kurgel and Ferreira 2005). According to Hein et al. (2003), there are two tools to accelerate the chemical process of dental bleaching: the thermocatalysis (heat) or photocatalysis (light). Currently, the preferred method of speeding up the bleaching process is photocatalysis with halogen light, LED or laser. However, even photocatalysis can induce increase of intrapulpal temperature with expansion of the liquid inside the dentin tubules resulting in pulp hyperemia and pain (Goodis et al. 1989; Hanning and Bott 1999; Michida et al. 2009). In fact, light accelerates the release of free radicals such as OH from $\text{H}_2\text{O}_2$. Part of the energy is converted to heat to degrade the molecules responsible for tooth staining (Buchalla and Attin 2007) and the other part is absorbed, and may cause the intrapulpal temperature increase (Eldeniz et al. 2005).

The increase of the intrapulpal temperature is a cause for concern. The dental pulp is a highly vascularized tissue and its vitality may be compromised by an increase in the temperature through the enamel and dentin (Nyborg and Brännström 1968). Actually, Zach and Cohen (1965) observed that the increase of the intrapulpal temperature of 3.3 °C could promote reversible histological changes in this tissue. When the temperature reaches 5.5 °C, it can be threatening to the pulp vitality. Increases of more than 10 °C induce the periodontal injury, and increases more than 16 °C can result in complete pulpal necrosis. The halogen light promotes the increase of the intrapulpal temperature (Carrasco et al. 2008). This increase can be higher than those presented by argon laser (Cobb et al. 2000). Aiming at
reducing the sensitivity symptom new light sources have been introduced at the dental bleaching procedure, such as light emitting diodes (LED) associated or not to low power lasers (Dederich and Bushick 2004; Stabholz et al. 2003). These authors were looking for light sources able to accelerate the bleaching procedure without causing great increases in the intrapulpal temperature. In fact, these authors were right, since later on, in 2008, Carrasco et al. showed in an in vitro study that the halogen light promoted the highest temperature rise in the pulp when coupled with the 35 % hydrogen peroxide (1.41 ± 0.64 °C), while the lowest elevations in the temperature were observed when bleaching was accelerated with the LED light (0.44 ± 0.11 °C) and the LED associated with the laser (0.33 ± 0.12 °C).

Similar to halogen light, the LED represents a not divergent and coherent source of light. Additionally, it has a thinner emission spectral, a higher output and does not produce heat, which allows its use in the therapies without significant risks. These characteristics validate this light source as an alternative to the conventional light-curing units (Whelan et al. 2001). Moreover, these light sources are no exothermic and able to reduce the tooth sensitivity (Cohen and Chase 1979; Stabholz et al. 2003).

At the control teeth that were not bleached, there was no stain, showing that the presence of dye at the bleached teeth occurred specifically in the function of the dental bleaching procedure. Then, the vascular permeability increase could occur either due to the metabolites of 35 % hydrogen hydroxide agent breakdown or due to the changes in dental pulp temperature elicited by the light source.

All the bleached samples presented increases in the vascular permeability, independently of the light source used. Then, this result indicated that the bleaching active agent or metabolites of its breakdown could have entered in the dental pulp chamber and elicited the acute inflammation represented by vascular permeability increase. Corroborating with this hypothesis, Dantas et al. (2010), showed increase of cell death in dental pulp fibroblasts cultured in medium conditioned by 35 % hydrogen peroxide.

The source and concentration of the bleaching active agent used in this study was the same for all the bleached samples. Thus, differences in the level of edema observed amongst the experimental groups should be due to the different light sources used. The increase in dental pulp vascular permeability was significantly smaller in the samples treated with LED and/or LED, followed by LPT. As these light sources were not exothermic (Cohen and Chase 1979; Stabholz et al. 2003), the results of this study corroborated with the hypothesis that temperature was a key modulator of the acute inflammation elicited by the in-office dental bleaching. Other important observation in this study was the fact of one session of bleaching using the halogen light led to an increase in the pulp vascular permeability significantly higher than the three sessions of the other bleaching groups. This result confirmed the injury of the high temperature can cause.

There were no differences between the groups treated with LED and LED, followed by LPT. In fact, it was expected to have smaller edema in these groups than in the group of halogen light; however, the LPT was supposed to improve the dental pulp response to the bleaching procedure. This result could be due to the fact that LEDs could have similar phototherapeutic effect on the dental pulp than the low intensity laser. Although, less evidence has been gathered on the bio-stimulative effect of noncoherent light (LED), some in vitro studies suggested macrophage activation as the first target of this kind of emission (Young et al. 1989). Recent studies have shown that it can exert a laser-like curative effect on skin and mucosal wounds (Whelan et al. 2001). Independently of the number of sessions used in the treatment, the dental bleaching with 35 % hydrogen peroxide agent and LED followed or not by LPT caused lesser increase in the dental pulp vascular permeability than the conventional method using halogen light. These results and the observations of clinical trials of the absence of the sensitivity when dental bleaching was speed up by LEDs (Dederich and Bushick 2004; Stabholz et al. 2003) encouraged that for vital teeth, the dental bleaching using hydrogen peroxide should be done in association with non-exothermic light sources, especially the LEDs. Further studies should be carried out in order to clarify the processes involved in the dental pulp reaction to dental bleaching procedures.

REFERENCES
