Clinical use of photodynamic antimicrobial chemotherapy for the treatment of deep carious lesions

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Department of Restorative Dentistry, School of Dentistry, Av Lineu Prestes, 2227-05508-000, São Paulo, SP, Brazil

Abstract. The purpose of this study was to assess photodynamic antimicrobial chemotherapy (PACT) via irradiation, using a low power laser associated with a photosensitizing dye, as an alternative to remove cariogenic microorganisms by drilling. Remaining dental samples in deep carious lesions on permanent molars (n = 26) were treated with 0.01% methylene blue dye and irradiated with a low power laser (InGaAP – indium gallium aluminium phosphide; λ = 660 nm; 100 mW; 320 J cm^{-2}; 90 s; 9J). Samples of dentin from the pulpal wall region were collected with a micropunch before and immediately after PACT and kept in a transport medium for microbiological analysis. Samples were cultured in plates of Brucella blood agar, Mitis Salivarius Bacitracin agar and Rogosa SL agar to determine the total viable bacteria, mutans streptococci and Lactobacillus spp. counts, respectively. After incubation, colony-forming units were counted and microbial reduction was calculated for each group of bacteria. PACT led to statistically significant reductions in mutans streptococci (1.38 log), Lactobacillus spp. (0.93 log), and total viable bacteria (0.91 log). This therapy may be an appropriate approach for the treatment of deep carious lesions using minimally invasive procedures.

Keywords: dental caries; lasers; photochemotherapy.

1 Introduction

Caries is an infectious bacterial disease that results in the localized destruction of dental hard tissues. The disease starts with an imbalance between the processes of remineralization and demineralization on the enamel surface. The reasons are an enhance in bacterial metabolism and the consequent increase in acid production, primarily mediated by the ingestion of sucrose. The traditional treatment of advanced carious lesions commonly involves the removal of all demineralized dentin using rotary burs until sound dentin forms the entire pulpal floor. The intention is to ensure the elimination of all remaining micro-organisms, to prevent a possible recurrence of caries and pulpal injury. However, this procedure is not always successful and some micro-organisms may remain even after the removal of all softened dentin. The main risk of this approach is the unnecessary wear of dental tissues and the possible exposure of the pulp, particularly in young patients.

The inner layer of the carious lesion, adjacent to sound dentin, is partially decalcified and still contains cariogenic species, although it also leaves the original dentinal tubule structure and sound collagen fibers partially preserved. It has been shown that dentin can even remineralize to a certain degree. Thus, an alternative to the removal of decalcified dentinal tissues is to kill the bacteria and preserve the affected dentin, so that the remineralization can occur.

Photodynamic antimicrobial chemotherapy (PACT) has been shown to be a promising antibacterial treatment. The therapy involves the association of a photosensitizer agent and light of a specific wavelength, resulting in the generation of cytotoxic species in the presence of oxygen such as singlet oxygen and free radicals. Already established as a potential treatment for cancer, photodynamic therapy has once again aroused interest in the field of antimicrobial chemotherapy, with new studies demonstrating the considerable advantages of its use.

Previous studies have revealed that PACT is capable of killing oral bacteria in cultures, plaque scrapings, and biofilms. Some in vitro studies have already demonstrated that PACT may be effective against bacteria involved in caries development. Burns et al. showed that PACT could eliminate cariogenic species when a bacterial suspension was treated with the photosensitizer toluidine blue (TB) before irradiation with red laser light. Later, it was demonstrated that the lethal photosensitization of S. mutans is possible even when the bacteria is embedded on dentin.
Fig. 1 Radiograph showing the limits of carious lesion included in the study. EDJ indicates the enamel-dentin junction and PC indicates the pulp chamber. Lesion should be limited to the occlusal surface and extend beyond the inner half of dentin (interrupted line).

in a collagen matrix or when the laser light passes through the demineralized dentin slices, simulating the carious lesion. Additional studies have sought to reproduce clinical conditions and obtained favorable results with regard to PACT and its effects on the decontamination of ex vivo carious dentin or caries lesions produced both in vitro and in situ. PACT may represent an efficient and less invasive approach to the treatment of deep carious lesions. Since no previous clinical study has been reported and the antimicrobial effects of PACT are usually more efficient in vitro than in vivo, the real effect of this type of therapy needs to be tested in a clinical setting. The aim of this investigation was to evaluate the effectiveness of PACT in bacterial reduction for the treatment of deep dentinal carious lesions.

2 Subjects and Methods
2.1 Subject Selection
Healthy patients of both genders were selected from the dental clinic of the School of Dentistry, University of São Paulo, Brazil, with ages ranging from 8 to 25 years. Calculation of statistical significance was performed using the paired t-test for the paired samples (Biostat 4.0, \( \alpha = 0.05\) and \( \beta = 0.15\)%), and the minimum sample size was estimated in 22 children.

Written informed consent was obtained from each patient or from the responsible parent or guardian in cases involving minors 18 years or younger. The possible discomforts, risks, and benefits were fully explained to the patients or to their guardians. The research protocol was conducted in accordance with the Declaration of Helsinki and Brazilian Human Research Law (Res 196/96), and it was approved by the Ethics Committee of University of São Paulo, Brazil (#01/08).

Patients included in the study were required to present at least one permanent molar with an active deep carious lesion limited to the occlusal surface and without pulpal involvement. Carious lesions were standardized by means of clinical and radiographic examinations performed before the procedures, with the aid of a measuring probe calibrated in millimeters. Lesion depths had to extend beyond the inner half of the dentin (i.e., between the enamel-dentin junction and the pulp chamber, as shown in Fig. 1). Exclusion criteria were 1. the use of any antibiotics during the study period or within six months prior to its initiation and 2. signs or symptoms of irreversible pulp inflammation. After clinical and radiographic examinations, 26 total teeth were selected from 23 patients. A sample of carious dentin was collected from each tooth both before and after PACT comprising baseline and experimental samples, respectively, for the microbiological analysis. The teeth were then temporarily restored with glass ionomer cement (Maxxion, FMG, Joinville, Brazil).

2.2 Clinical Procedures
After local anesthesia with 2% lidocaine with a vasoconstrictor (DFL, Petropolis, Brazil) and after isolation with a rubber dam, demineralized dentin, and enamel were removed from only the lateral walls of the carious lesions to promote the satisfactory adhesion of the restorative material [Figs. 2(a) and 2(b)]. These procedures were performed with a sharp excavator (N 19) and a spherical carbide bur (KG Sorensen, São Paulo, Brazil).
**Fig. 2** Description of clinical procedures for PACT in deep carious lesion.
at slow rotation. Thus, carious dentinal layers were maintained
over the pulpal wall of the cavities [Fig. 2(C)].

The collection of baseline carious dentin samples from the
pulpal wall was carried out using a sterile 1 mm diameter mi-
cropunch [Fig. 2(D)] (Richter, São Paulo, Brazil) on its active
segment. The penetration depth of the micropunch in the cari-
ous lesions was calibrated between measurements in different
patients and between repeated measurements in the same pa-
tient by means of a 0.5 mm mark on its active point. Thus, the
collected samples had diameters of 1 mm, depths of 0.5 mm and
average weights of 0.059 mg. The samples were immediately
transferred to the transport medium VMGA III (Viable Medium
of Göteborg Anaerobic) containing glass beads.

2.3 Photosensitizer and Light Source

Methylene blue (MB) was used as the photosensitizer agent,
which was prepared with distilled water to obtain a final con-
centration of 0.01% (100 mg/L, 268 μM) (Fórmula & Açao,
São Paulo, Brazil). The low power laser used was a diode laser
(InGaAlP – Indium Gallium Aluminum Phosphide) with 660
nm wavelength, spot size of 0.028 cm², and fixed output power
of 100 mW. The parameters adopted were: energy density of
320 J cm⁻², time exposure of 90 s, and total energy of 9 J.
The output power was quantified with a power meter (Newport
Corp., California) before the irradiations.

2.4 Photodynamic Antimicrobial Therapy

After the baseline dentin collection, the photosensitizer MB
was applied on carious tissue with an insulin injection syringe
[Fig. 2(C)]. A volume of approximately three mL was used to fill
each lesion. The carious tissue was maintained in contact with
MB during a pre-irradiation time of five minutes. Laser beam
was then perpendicularly positioned to the occlusal surfaces of
the teeth and irradiations proceeded in one single point on the
center of each cavity [Fig. 2(E)]. After the irradiation, experi-
mental samples were collected from an area adjacent to that of
the baseline sample collection [Fig. 2(F)]. This precaution was
taken to minimize regional variations in the microbial popula-
tion. The dentinal samples were then inserted in the transport
medium VMGA III. Clinical procedures were performed by
only one trained researcher to standardize the data collection.

Teeth were then cleaned with a cotton swab and thoroughly
washed with distilled water until the photosensitizer was com-
pletely removed [Fig. 2(B)]. Glass ionomer cement (Maxxion,
FGM, Joinville, Brazil) was used for temporary restorations
[Fig. 2(T)].

2.5 Microbiological Procedures

The flasks containing the samples in the VMGA III media were
homogenized for two minutes in a vortex (Fisher Scientific,
New York, USA) to break out the aggregates of bacteria. Immediately
after homogenization, six decimal dilutions were carried out
(1:10, 1:100, 1:1,000, 1:10,000, 1:100,000, and 1:1,000,000). In
each dilution, three aliquots of 25 μl were plated onto Brucella
blood agar, Mitis Salivarius agar supplemented with 15% su-
crose and 0.2 units of Bacitracin ml⁻¹ (MSSB), and Rogosa SL
agar containing 0.13% glacial acetic acid (RSL). Brucella blood
agar was used to determine total viable counts and plates were
incubated in an anaerobic cabinet at 35 °C for seven days, in an
atmosphere of 85% N₂, 10% CO₂, and 5% H₂. Mitis Salivar-
nius agar supplemented with sucrose and bacitracin (MSSB) was
used to count mutans streptococci and the plates were incubated
at 37°C for 48 h in a candle jar. Rogosa SL agar (RSL) was
used to count Lactobacillus spp. and the plates were aerobically
incubated at 37°C for 48 h. After the incubation period, colony-
forming units (CFU) per plate were counted, and reduction was
calculated between the samples taken before and after the appli-
cation of photodynamic therapy for each medium studied.

2.6 Statistical Analysis

First, the data distribution was assessed using D’Agostino’s K-
squared test. Statistical analyses comparing the baseline data
and the experimental sample data were conducted using the
Wilcoxon signed-rank test and comparisons between the groups
were analyzed via the Mann-Whitney test at 5% significance
level.

3 Results

From March 2008 to May 2009, 165 patients were analyzed in
the dental clinic of the School of Dentistry, University of Sáo
Paulo. Among them, 23 presented with active dentin carious
lesions matching the inclusion criteria, resulting in 26 lesions
for the study. Patients were followed up from the beginning of
the study and no patient reported any pain or sensitivity after the
clinical procedures.

Statistical analyses showed significant differences in the
count of CFU before and after PACT for the three groups of
bacteria tested. The CFU reductions found after PACT on the
tested micro-organisms are presented in Table I. The therapy
promoted a mean log reduction of 1.38 (p < 0.0001) for mutans
streptococci, 0.93 (p < 0.0001) for Lactobacillus spp., and 0.91
(p < 0.0001) for total viable bacteria. No statistically signifi-
cant differences were observed regarding the level of bacterial
reduction among the test micro-organisms (p > 0.05) (Table I).
PACT was seen to promote similar levels of reduction in all
tested bacteria.

4 Discussion

Even though controversies persist regarding how much tissue
must be removed to arrest the caries process, the literature
appears to discourage the excessive removal of dentin over
the pulpal surface, supporting the idea of minimally invasive
procedures. According to this concept, it is favorable to
maintain a layer of partly demineralized dentin underneath a
filling material to preserve pulpal tissue vitality, especially to
encourage the reparative process of tubular sclerosis and ter-
iary dentin formation.

It has already been demonstrated that oral bacteria orga-
nized in biofilms can be susceptible to PACT. Wilson et al.
verified that a substantial reduction in the bacterial count
was achieved when plaque samples obtained from volunteers
were treated with toluidine blue O (TBO) or phthalocyanine
and exposed to red light. Analyses through confocal laser scan-
ing microscopy of multi-species biofilm cultured from saliva
samples and treated with TBO showed bacterial reductions of 97.4% after irradiation with a low power laser. Moreover, transmission electron microscopy has confirmed that a photosensitizer can be absorbed by the biomass found in natural oral plaque biofilms.

However, it remains unclear as to the depths to which the photosensitizer can penetrate carious dentin. Lethal photosensitization probably occurs predominantly in the outer layers of biofilm and carious tissue. This fact could occur due to the inability of the photosensitizer to spread into the inner layers or the inability of the light to be totally transmitted. These factors may explain the results obtained in the present study, because significant bacterial reductions of mutants streptococci (78.07%), Lactobacillus spp. (78.0%), and total viable count (76.03%) occurred, although the reductions were lower than those generally observed with in vitro studies in which the substrate is less complex.

Pre-irradiation exposure time seems to be an important factor for photosensitizer diffusion through the tissue. In our study, the photosensitizer was left in contact with the dentin for five minutes before laser irradiation. Muller et al. already demonstrated in vitro the minimal effect of PACT in the viability of microorganisms when MB was applied for only 60 s in contact with the biofilm and removed prior to the laser irradiation. It is known that high concentrations of dyes can induce the phenomenon of self-quenching, reducing the amount of light that actually reaches the bacteria and induces the generation of reactive oxygen species. This effect may have interfered in the effectiveness of PACT in our study, which warrants new studies with lower dye concentrations. However, because the degree of photodamage is dependent upon the dye concentration and the intensity and fluence of the laser light, a higher concentration was chosen mainly because of the complexity of the substrate. This concentration of MB has been clinically used for the treatment of herpes simplex labialis, accelerating the healing process.

Even though many studies have shown that PACT is an effective antimicrobial technique, most were performed with bacteria in an aqueous suspension, which is different from those conditions found in the oral cavity. It has been demonstrated that interposing 150-μm dentin slices between the laser light source and the bacteria leads to a reduction of 50% in the power density of the light source, although substantial kills were obtained. This effect may have occurred in our study, restricting the penetration of the photosensitizer and light transmission through the dentinal substrate. For that reason, increased energy density was chosen in order to overcome such issues in the present study. In addition, it is important and clinically convenient to have short exposure times and, therefore, the use of greater power density may represent an advantage.

Several previous reports have demonstrated that different light sources and photosensitizers can be combined to promote the bactericidal effect. The dental plaque disclosing agent erythrosin was considered a potential photosensitizer for the treatment of S. mutans biofilms grown in vitro when combined with a white light source. The complete elimination of S. mutans in a planktonic culture was as well demonstrated when it was previously treated with different concentrations of rose bengal combined with a held photopolymerizer (400 to 500 nm) or TBO combined with a light emitting diode (LED) (600 to 670 nm). Considering that longer wavelengths enable the deeper penetration of light into the tissues, the association between a blue dye and a red light source was preferred. Methylene blue has shown significant phototoxicity in different types of oral bacteria involved in periodontal diseases and endodontic infections, among others. Because this photosensitizer has an intrinsic positive charge, it can efficiently bind to both Gram-positive and -negative bacteria.

Comparing the effectiveness of MB and TBO as lethal photosensitizers for Gram-positive and Gram-negative microorganisms, it has been verified that both are capable of eradicating all microorganisms to some extent under red laser light. Nevertheless, the efficiency of the therapy can vary according to the genus of bacteria. Gram-negative bacteria were shown to be more resistant to the therapy due to the greater complexity of their cytoplasmic membranes. Although a variety of studies have found S. mutans and Lactobacillus spp. to be the major microorganisms involved in caries development, there may be large variations and changes within the lesion environment at advanced stages of caries and Gram-negative species may also occur. For that reason, this study analyzed the effect of PACT on total viable bacteria, involving both Gram-negative and -positive strains, which was demonstrated to be equally successful. It is possible that this result is related to the high dose of energy that was applied. Statistical analysis (Table 1) demonstrated a wide standard deviation for all of the groups, although this variation is expected when working with bacteria and particularly involving clinical experiments.

### Table 1 Bacterial count after PACT (qualitative and log results) – averages and standard deviations.

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>Mutans streptococi</th>
<th>Lactobacillus spp.</th>
<th>Total viable bacteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample CFU (x 10⁵)</td>
<td>baseline</td>
<td>experimental</td>
<td>baseline</td>
</tr>
<tr>
<td></td>
<td>41.59(90.88)a</td>
<td>8.51(32.35)b</td>
<td>107.52(131.77)a</td>
</tr>
<tr>
<td>Log (10)</td>
<td>5.82(1.2)</td>
<td>4.44(2.3)</td>
<td>6.71(0.59)</td>
</tr>
<tr>
<td>Reduction (%)</td>
<td>78.07 [24.03]a</td>
<td>78.00 [25.19]a</td>
<td>76.03 [21.66]a</td>
</tr>
<tr>
<td>Significance</td>
<td>p = 0.0176</td>
<td>p &lt; 0.0001</td>
<td>p &lt; 0.0001</td>
</tr>
</tbody>
</table>

*Different letters correspond to a significant statistic difference (p < 0.05).*
The present study was conducted in the clinical setting using the photosensitizer MB and a red light laser to promote dentin decontamination. The obtained results appear to be relevant for further clinical studies. The findings are in accordance with Giust et al., who demonstrated that PACT was effective in the decontamination of carious bovine dentin that was artificially induced, using a light-emitting diode light source and two different photosensitizers. The association of TBO and LED also resulted in a significant decrease in the viability of total streptococci, mutans streptococci, lactobacilli, and total micro-organisms on dentinal caries produced in situ. These data are in agreement with the present study. Although the achieved antimicrobial effect appears to be limited, it may still be considered a clinically relevant outcome and agrees with others clinical studies.

The literature encourages the maintenance of a layer of “affected dentin” over the pulpal wall in order to avoid pulp exposure. In this way, any immediate bacterial reduction obtained for dentin decontamination would increase the chances of treatment success and it is expected that further reduction occurs over time if a filling material is properly placed.

The aim of this study was to treat teeth with deep carious lesion while avoiding the risk of pulpal exposure and to arrest the carious process by favoring the repair of involved tissues. The results of this first clinical study demonstrate the potential clinical use of PACT for the treatment of caries. Also, our findings might encourage new research efforts to develop clinical protocols to make PACT feasible in clinical practice. New studies could be conducted with the goal of assessing different parameters for PACT (e.g., dye concentrations and exposure times) to improve its effectiveness.

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