Photodynamic therapy in oral potentially malignant disorders—Critical literature review of existing protocols

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Abstract

Introduction: Oral cancer is a serious public health issue. Apart from its high rate of prevalence, incidence and mortality, it can often result in more complex and expensive treatment when diagnosed late. Potentially malignant disorders (PMDs) can precede oral cancer, and are usually treated by surgical excision. However, in many cases patients are elderly and multiple interventions may be required. Photodynamic therapy (PDT) is a simple alternative, which has been successfully used in the treatment of oral PMDs.

Objective: Due to the lack of standardization regarding photosensitizers (PTSs), types of irradiation, and methods of application, the objective of this study was to analyze existing PDT protocols in an attempt to identify the one that demonstrates greater efficiency, reliability and feasibility in the treatment of oral PMDs for both researchers and clinicians.

Methods: Original clinical studies published only in English between 1993 and 2016 were searched in Pubmed/Medline database using the following keywords: photodynamic therapy; oral potentially malignant disorder; oral premalignant lesions. Review articles; experimental studies; case-reports; commentaries; and letters to the Editor were excluded from the selection.

Results and conclusion: Based on the 16 studies selected, the topical 5-ALA-20% PTS, associated to a LED light applied for 15 min with a 7-day interval between sessions emerged as the most frequently used PDT protocol, with satisfactory results. Due to its low rate of side effects, this PDT protocol presents good potential for the treatment of oral PMDs. Further clinical studies are required to ascertain its long-term validity in preventing oral cancer.

1. Introduction

Potentially malignant disorders (PMDs) refer to benign, but morphologically altered tissues, which present a greater risk of undergoing malignant transformations [1]. The World Health Organization (WHO), in 2005, changed the previous terminology “potentially malignant lesions” and “potentially malignant conditions” to PMD [2].

The term dysplasia is characterized by the presence of abnormal epithelial architecture, disordered cell growth, changes in the nuclear-cytoplasmatic ratio, atypical mitosis, and other alterations. Dysplasia can be classified into mild, moderate, and intense, according to subjective microscopic analysis [3]. It is believed that the more severe the epithelial dysplasia, the greater the risk of malignization [3].

Oral PMD management still requires clearer definition. Surgical excision, laser surgery, and cryotherapy associated with the reduction of risk factors, as well as follow up examinations for different periods of time have been suggested [2,4,5]. Photodynamic therapy (PDT) has recently emerged as a potential alternative in the treatment of PMDs, since it is capable of promoting total or partial regression of these conditions with few side effects [2,4,5].

1.1. Photodynamic therapy

PDT is a minimally invasive treatment, successfully applied in head and neck PMDs and malignant disorders. The technique is simple and can usually be performed on an outpatient basis [4,5].

A topical or systemic photosensitizer (PTS) drug is administered, which selectively binds to atypical cells. After an incubation period, a source of light of suitable wavelength is applied on the target tissue. It promotes the absorption of photons by the PTS, which turns into an extremely unstable molecule [6]. This molecule, while searching for stability, transfers the light energy to the oxygen in the cellular environment. This reaction generates oxygen reactive species, which has a
cytotoxic effect on the tissue, leading to cell death [6].

When compared to more invasive techniques, PDT systemic effects are insignificant, it is minimally toxic to normal tissue, and presents reduced morbidity and excellent aesthetic results. It can be applied in association with any other conventional treatment and repeated as often as necessary, without generating cumulative toxicity [4–6]. Sensitivity, pain, swelling, burning sensation, taste alterations, ulcerations, and loss of local sensation have been reported with PDT, but often with a low magnitude [7,8].

1.2. Photosensitizer

Photosensitizers (PTSs) are drugs that promote the transfer of light energy to the cellular environment, resulting in the formation of highly reactive chemical species, which act in the destruction of target cells.

PTSs tend to accumulate in atypical cells, but the mechanisms responsible for this process are not completely understood [6]. Several hypotheses have been proposed. One of them concerns the predominance of blood vessels with morphological changes in tumors as a result of neovascularization and the lack of lymphatic drainage, creating greater permeability and drug retention [6]. Another hypothesis states that some drugs preferentially bind to low-density-lipoprotein (LDL) receptors, which are over-expressed in atypical cells [6]. There is also the perception that low pH found in atypical cells cytoplasm promotes drug ionization, making it more hydrophilic, increasing PTS retention inside the cell [9].

To select the appropriate PTS for each clinical situation, features such as toxicity, selectivity, wavelength necessary for its activation, effectiveness, side effects, route of administration and cost have to be taken into consideration.

There are three generations of drugs, according to their chronological order of development. The first generation includes hematoporphyrin derivative (HpD) and porphyrin sodium (Photofrin®) [6]. To improve some of the drawbacks presented by the first generation of PTSs (relatively low absorption and skin photosensitivity), the second generation of PTSs was developed, with excellent results [6]. It includes 5-aminolevulinic acid (5-ALA); hypericin; phthalocyanine; benzoporphyrin derivatives, such as Vertepor®; and meta tetrahydroxyphenyl chlorin (mTHPC) derivatives, such as Foscan® [6]. A new, third generation of PTSs is now being developed to improve atypical cell selectivity, and represents an area of active research [6].

1.2.1. Light sources

The application of light in an appropriate wavelength (600–800 nm), also known as the “therapeutic window”, excites the PTS, generating a reaction with the oxygen present in the cells. Shorter wavelengths have low tissue penetration, while irradiation with longer wavelengths may not have enough power to generate reactive oxygen [6]. The blue light (450–495 nm), for example, penetrates less efficiently through the tissue when compared to the red (620–750 nm) and infrared light (> 750 nm), which can penetrate deeper in the tissue [6].

The choice of a light source is dependent on the lesion (tissue feature, size, location and accessibility), type of PTS (spectrum of absorption and administration), cost and availability of light systems [6].

Theoretically, any light source may be used in PDT [10]. Conventional lamps were the first to be used. Because they have a wide range of wavelengths, filters are required to obtain the desired wavelength [10]. With the introduction of lasers, the use of conventional lamps in PDT has decreased considerably [11]. Laser produces monochromatic wavelengths, which allows easy calculation of light dosimetry and the ideal wavelength for a specific PTS [11]. The light-emitting diode (LED) is also an alternative source of light that has been increasingly used [6].

1.2.2. Photochemistry and photophysics

The basic principle of PDT is the formation of cytotoxic agents from the interaction of the oxygen within the cells and the PTS excited by light of an appropriate wavelength.

After activation by light photons, the PTS changes from its primary state (PTS) to a singlet excited-state (PTS*) [12]. The excited PTS is very unstable and tends to return to its original and energetically more favorable state, which takes place through the emission of heat or fluorescence [12]. In this particular case, photodynamic reactions with therapeutic outcomes do not occur, but have the potential to be used as a diagnostic tool [12]. Another possibility is the passage of PTS* to another excitation state, less unstable, called triplet state (PTS*), which is capable of promoting two different photodynamic reactions [12]:

Type I reaction – PTS in its excited triplet state (PTS*) interacts with cellular substrates [6,12]. Through the transfer of electrons, ions are generated. These react with molecular oxygen, making reactive oxygen species (ROS), such as superoxide, hydrogen peroxide and hydroxyl radicals [6,12].

Type II reaction – Direct transfer of energy between PTS* and molecular oxygen occurs, exciting O₂ and leading to the formation of singlet oxygen (O2), which is also a ROS [6,12]. Reactive oxygen species are highly reactive free radicals, which have the ability to interact with different molecules and damage their normal function [13]. This is a key factor for PDT cytotoxicity [13]. ROS formation can lead to tumor destruction through three main biological mechanisms: 1) direct destruction by activating cell death pathways, such as apoptosis, necrosis and autophagy; 2) damage to the tumor vasculature, by suppressing the delivery of oxygen and nutrients; and 3) stimulation of inflammatory and/or immune response [13].

2. Methodology

Original clinical studies, published only in English between 1993 and 2016 were searched in Pubmed/Medline database using different combinations of the following keywords: photodynamic therapy; oral potentially malignant disorder; oral premalignant lesion. Sixteen clinical studies using PDT to treat oral PML were selected. The objective was to compare the PDT protocols used and their respective results. Review articles, experimental studies (in vitro or animal studies), case-reports, commentaries, and letters to the Editor were excluded from the selection.

3. Results

The selected articles are displayed in Table 1.

Based on the selected studies, a comparative analysis of the protocols and their respective results was performed. Absence or incomplete data were disregarded.

Methodological difficulties were found during table organization. In Kühler et al. [16], Tsai et al. [17], Rigual et al. [22] and Shafirstein et al. [8], for example, the number of sessions, time of light activation and interval among sessions were absent. In Yu et al. [21], Lin et al. [23], Pietruska et al. [24] the type of laser was not described, and in Fan et al. [15], Tsai et al. [17], Pietruska et al. [24] and Maloth et al. [26] the number of recurrences was not disclosed.

Number of sessions ranged from 1 to 10; light application time from 10 to 16.6 min; and the interval among sessions ranged from 3.5 to 14 days, with the 7-day interval being the most frequent [4.7,14–16]. The time from PTS administration and light activation was between 48 and 96 h when used intravenously; 1.5–4 h when applied topically; 0.5–4 h orally; and 1.5 h for intraleisional application [4.7,14–16].

Light dose administered ranged from 50 to 200 J/cm². The most frequent was 100 J/cm², used in 10 of the 16 studies [7,8,16–21,23,25]. In four studies, the light dose administered was variable [4,14,15,22]. Grant [14] used 50–100 J/cm²; Fan et al. [15], 100–200 J/cm²; Rigual et al. [22], 50–75 J/cm² and Jerjes et al. [4] used 100–200 J/cm² [4,14,15,22].

Regarding the PTS, the most frequently used drug was the 5-
Table 1
Protocols of PDT application.

<table>
<thead>
<tr>
<th>Author</th>
<th>No Patients</th>
<th>Lesion</th>
<th>N° Lesions</th>
<th>PTS Route of Administ.</th>
<th>Pre activation Time (hours)</th>
<th>Ligth (nm/l/cm²) N° sessions (weeks)/Time of light activation (minutes)/Interval between sessions (days)</th>
<th>N° Complete response/Partial response/No response</th>
<th>Follow-up (Months)</th>
<th>Recurrence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grant et al.</td>
<td>11</td>
<td>Leukoplakia</td>
<td>–</td>
<td>Purifier sodium 2 mg/kg</td>
<td>Intravenous</td>
<td>Copper vapor laser (630/50-100) 6-8 weeks at the total</td>
<td>10 CR/1 PR</td>
<td>3-19</td>
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<td>Erythroplakia</td>
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<td>Fan et al.</td>
<td>18</td>
<td>Carcinoma and PDM</td>
<td>–</td>
<td>5-ALA 60 mg/kg</td>
<td>Oral</td>
<td>Gold vapor laser (628/100-200) 3-5 weeks at the total</td>
<td>2 CR/3 PR/1 NR (CA) 12 PR (PMD)</td>
<td>28-88</td>
<td>–</td>
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<td>Kübler et al.</td>
<td>12</td>
<td>Leukoplakia</td>
<td>–</td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>Argon laser (630/100)</td>
<td>–</td>
<td>5 CR/4 PR 3 NR</td>
<td>0</td>
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<tr>
<td>Sierotzki et al.</td>
<td>12</td>
<td>Leukoplakia</td>
<td>24</td>
<td>5-ALA 10%</td>
<td>Topical</td>
<td>Argon laser (635/100) 6-8/15/14</td>
<td>10 CR 2 NR</td>
<td>4-34</td>
<td>1</td>
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<tr>
<td>Tsai et al.</td>
<td>33</td>
<td>Leukoplakia</td>
<td>–</td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>LED (635/100)</td>
<td>8 CR/13 PR/12 NR</td>
<td>6</td>
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<td>Chen et al.</td>
<td>5</td>
<td>Verruca Hyperplasia</td>
<td>–</td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>LED (635/100) 1-3/16,6/7</td>
<td>5 CR</td>
<td>3-11</td>
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<td>Chen et al.</td>
<td>8 (VH)</td>
<td>Verruca Hyperplasia</td>
<td>24 (L)</td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>LED (635/100) 2-5/16,6/7 (VH) 2-8/16,6/3,5 (L)</td>
<td>8 CR (VH) 8 CR/16 RP (L)</td>
<td>5-14 (VH) 3-16 (L)</td>
<td>0 (VH) 2 (L)</td>
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<tr>
<td>Yu et al.</td>
<td>36</td>
<td>Verruca Hyperplasia</td>
<td></td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>Diode Laser (635/100) 1-6/16,6/7 (LED)</td>
<td>36 CR</td>
<td>6-56</td>
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<td>Yu et al.</td>
<td>20 (LED) 26 (Laser)</td>
<td>Verruca Hyperplasia</td>
<td></td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>Diode Laser (635/100) Laser (635/100) 2-7/16,6/7 (LED) 2-6/16,6/7 (Laser)</td>
<td>17 CR/3 PR (LED) 25 CR/1 PR (Laser)</td>
<td>16-76 (LED) 5 (Laser)</td>
<td>3-16 (Laser)</td>
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<td>Rigual et al.</td>
<td>26</td>
<td>Dysplasia, CA in situ and T1 CA</td>
<td>–</td>
<td>Purifier sodium 2 mg/kg</td>
<td>Intravenous</td>
<td>Diode Laser or Argon laser (630/50-75)</td>
<td>24 CR/1 PR/1 NR</td>
<td>7-52</td>
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<td>Lin et al.</td>
<td>40 (VH)</td>
<td>Verruca Hyperplasia</td>
<td>40 (E)</td>
<td>5-ALA 20%</td>
<td>Topical</td>
<td>Laser (635/100) 1-6/16,6/7 (VH) 2-6/16,6/7 (E)</td>
<td>40 CR (VH) 38 CR/2 PR (E)</td>
<td>8-37 (VH) 6-30 (E)</td>
<td>0 (VH) 8 (E)</td>
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<td>Shafrstein et al.</td>
<td>18</td>
<td>Leukoplakia</td>
<td>–</td>
<td>5-ALA 20%</td>
<td>Topical e Intralesional</td>
<td>Diode Laser (585/100)</td>
<td>–</td>
<td>5 CR/7 PR (T) 3 CR/2 PR/1 not available (T)</td>
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<tr>
<td>Jerjes et al.</td>
<td>57 (5-ALA)</td>
<td>Leukoplakia and Erythroplakia</td>
<td>90 (m-THPC)</td>
<td>5-ALA 60 mg/kg</td>
<td>Topical</td>
<td>Diode Laser (628/100-200) (5-ALA) Diode Laser (632/100-200)-(m-THPC) 1-10/-4 weeks at the total</td>
<td>119 CR/12 PR/16 NR</td>
<td>Mean 7.3 years</td>
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<td>Pietruska et al.</td>
<td>23</td>
<td>Leukoplakia</td>
<td>44</td>
<td>Chlorine E6 20%</td>
<td>Topical</td>
<td>Laser (660/90)</td>
<td>10/-/</td>
<td>12 CR/22 PR/10 NR</td>
<td>–</td>
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<td>Selvam et al.</td>
<td>5</td>
<td>Leukoplakia</td>
<td>–</td>
<td>5-ALA 10%</td>
<td>Topical</td>
<td>Xenon lamp (630/100)</td>
<td>2 CR/2 PR/1 NR</td>
<td>1</td>
<td>0</td>
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<td>Maloth et al.</td>
<td>12 (L) 10 (LP)</td>
<td>Leukoplakia and Lichen Planus</td>
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<td>5-ALA 20%</td>
<td>Topical</td>
<td>LED (420/120)</td>
<td>2 CR/8 PR/2 NR</td>
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5-ALA = 5-Aminolevulinic acid, m-THPC = m-tetrahydroxyphenylchlorin, CA = Carcinoma, Administ. = Administration, CR = Complete Response, PR = Partial Response, NR = No Response, T = Topical, I = Intralesional.
aminolevulinic acid (5-ALA) 20% (Levulan®), applied topically in 8 of the 16 studies [8,16,17–21,23]. The 5-ALA was applied in a total of 256 patients, with complete remission in 78% of the lesions, partial remission in 18% and no remission in 6%. The recurrence rate was 7%.

3.1. Discussion

The 5-aminolevulinic acid (5-ALA) is a prodrug that undergoes biotransformation and only becomes active in vivo. It quickly accumulates in large quantities in tissues [27]. This feature is one of the major advantages of 5-ALA compared to other PTs, and may explain why it has been more frequently used [27]. With 5-ALA, side effects tend to be reduced, since its clearing is faster than other PTs [28]. It is used topically in the treatment of surface lesions, such as oral PMD, and is capable of penetrating 1–2 mm into the tissues [29].

LED (5 studies) and diode laser (4 studies) were the most frequently used light sources [4,8,19–22,26]. One study using 5-ALA compared LED and Laser in oral PMD, finding no significant differences between the light sources [21]. According to the authors, LED was simpler, lighter, smaller, more portable, and cheaper with the potential to replace Laser in the future. However, the authors call attention to its durability, which may be compromised by chip impairment as a result of the heat generated [21]. The Laser source (specification not provided), on the other hand, was considered more stable, more durable and was equipped with adjustable power, although it was heavier and more expensive [21]. The authors concluded that when used in erythroplasia, the choice of the light source would depend on the budget [21].

No explanation is available for the existence of varied responses using the same PDT protocol. Probably the results are influenced by specific features of PMDs such as size, color, presence of atypia and thickness of the keratin layer. Best results were achieved in minor lesions and with thinner keratin layers [8,20,21,23,25]. For example, a study reported better clinical outcome for dysplastic erythroplakia lesions than for non-dysplastic oral lesions such as leukoplakia lesions [21]. The histological, biological and structural characteristics of erythroplakia contribute to the successful clinical outcome of these lesions. Compared to non-dysplastic oral lesions, dysplastic erythroplakia lesions have less keratotic epithelial surface, as well as thinner and more permeable epithelium. Thus, the topical PTS may more easily diffuse into these dysplastic epithelia, resulting in good absorption, while the reducing effect of the thinner keratin layer on light intensity was minimal [21].

The measures adopted in case of recurrence were the repetition of the PDT protocol, lesion excision or just follow-up, depending on the lesion or patient’s wish [4,14,21–23].

PDT side effects are uncommon. Mild to moderate pain and local edema were the most reported effects [4,7,8,14,15,17,20,22,23]. Skin photosensitivity, burning sensation and discomfort were also common [4,8,15,16,20,22–24,25]. Although not frequent, erythema, ulceration and secondary infection were also present [7,8,16,17]. For pain control, nonsteroidal anti-inflammatory drugs (NSAIDs) and opiates were considered sufficient [4,5]. Systemic PTS resulted in some important side effects, when compared to topical PTS, such as residual systemic photosensitization, which can last for several days or weeks, edema, sunburn and superficial skin necrosis, when skin is exposed to bright light [4].

PDT consists of an effective, non-toxic strategy in the treatment of oral PMDs. It is a minimally invasive technique that results in less morbidity and mutilation to the oral tissues, and can be applied several times, with minimal side effects. However, standardized protocols, clinical trials with larger samples, as well as long term follow-ups are still required.

4. Conclusion

PMDs are a therapeutic challenge for dental surgeons. Based on the short-term clinical applicability and the benefits demonstrated, PDT can be a useful treatment strategy for the management of oral PMDs. Based on the literature review conducted, the protocol using 5-ALA-20% topical PTS in association with a LED light source, applied for 15 min with 7-day intervals between sessions, the number of sessions depending on the response to the treatment, emerged as a strong candidate in the treatment of PMDs. The use of systemic sensitizer (Porfimer sodium 2 mg/kg) yielded an initial complete response rate of 90%, which surely would also warrant its inclusion in any anticipated clinical trials. However, due to the side effects resulting from its use, and the small size of oral lesions compared to skin lesions, topical use seems to be best indicated in oral PMDs. It should be noted, however, that the primary aim of PDT is to prevent invasive malignancy occurring. Invasive tumors often arise in apparently normal areas of the mucosa so that it cannot be anticipated that the resolution of obvious oral PMDs will actually avoid the development of cancer. Further clinical studies with long term follow up are required to determine not only the best protocol, but also the ultimate benefits of PDT in the treatment of oral PMDs.

References


