Antimicrobial effect on *Candida albicans* biofilm by application of different wavelengths and dyes and the synthetic killer decapeptide KP

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The aim of this study was to test the application *in vitro* of different laser wavelengths at a low fluence in combination or not with proper photosensitizing dyes on *Candida albicans* biofilm with or without a synthetic killer decapeptide (KP).

*Candida albicans* SC5314 was grown on Sabouraud dextrose agar plates at 37°C for 24 h. Cells were suspended in RPMI 1640 buffered with MOPS and cultured directly on the flat bottom of 96-wells plates. The previously described killer decapeptide KP was used in this study.

Three different combinations of wavelengths and dyes were applied, laser irradiation has been performed at a fluence of 10 J/cm². The effect on *C. albicans* biofilm was evaluated by the XTT assay. Microscopic observations were realized by fluorescence optic microscopy with calcofluor white and propidium iodide.

Compared with control, no inhibition of *C. albicans* biofilm viability was obtained with application of red, blue and green lasers alone or with any combination of red diode laser, toluidine blue and KP. The combined application of blue diode laser with curcumin and/or KP showed always a very significant inhibition, as curcumin alone and the combination of curcumin and KP did, while combination of blue diode laser and KP gave a less significant inhibition, the same obtained with KP alone. The combined application of green diode laser with erythrosine and/or KP showed always a very significant inhibition, as the combination of erythrosine and KP did, but no difference was observed with respect to the treatment with erythrosine alone. Again, combination of green diode laser and KP gave a significant inhibition, although paradoxically lower than the one obtained with KP alone.

Treatment with KP alone, while reducing biofilm viability did not cause *C. albicans* death in the adopted experimental conditions. On the contrary, combined treatment with blue laser, curcumin and KP, as well as green laser, erythrosine and KP led to death most *C. albicans* cells.

The combination of laser light at a fluence of 10 J/cm² and the appropriate photosensitizing agent, together with the use of KP, proved to exert differential effects on *C. albicans* biofilm.

Introduction

Fungal infections, mainly those caused by *Candida albicans*, have an important impact on human health particularly because of the growing number of immunocompro-

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Candida albicans produces highly structured biofilms composed of multiple cell types (i.e., unicellular round budding yeasts and oval pseudohyphae, and multicellular elongated hyphae) encased in an extracellular matrix, mainly composed of proteins and glycoproteins (55%), which plays an important role in drug resistance acting as a physical barrier to drug penetration. Biofilm formation initiates when planktonic yeasts adhere to a surface and begin to aggregate and form microcolonies. This first stage is immediately followed by a proliferation of yeast cells and the beginning of hyphal development.

The ability to form hyphae or yeast cells, i.e. C. albicans dimorphism, is important for biofilm formation because of the capability of hyphae to contribute to the mechanical and architectural stability of biofilm and to support yeast cells.

In patients wearing dentures, biofilm constituted by bacteria and Candida cells can be formed both on the oral mucosa and on the surface of dentures, commonly leading to acute or chronic candidosis, including denture stomatitis.

The Antimicrobial Photodynamic Therapy (APDT) is used with different types of protocols, laser and photosensitizers. The cytotoxic effect is achieved through the local application or systemic administration (oral or intravenous) of photosensitizing agents followed by irradiation of visible light with emission spectrum appropriate to the absorption spectrum of the used photosensitizer, in the presence of oxygen. This induces oxidation phenomena with selective destruction of proteins, lipids, nucleic acids and other cellular components.

Recently the literature about C. albicans biofilms has grown with publications about the effect of APDT using different light sources and photosensitizers.

The main objective of this study was to apply different laser wavelengths at a low fluence in combination or not with proper photosensitizing dyes on in vitro C. albicans biofilm mimicking the in vivo infection. In addition, the previously described antibody-derived decapeptide KP, endowed with anticandidial activity, was also used in combination with laser and dyes application.

Materials and methods

Microbial strain and culture conditions for biofilm formation

The reference strain C. albicans SC5314, selected for its ability to form biofilm, was grown on Sabouraud dextrose agar (SDA) plates at 37°C for 24 h. Cells were suspended in RPMI 1640 medium buffered with 3-(N-morpholino) propane sulfonic acid (MOPS) at a concentration of 106 cells/ml using the McFarland turbidity standard. Yeast suspensions were cultured in flat-bottomed 96-wells plates (100 µl/well).

Plates were incubated 18 h at 37°C on an orbital shaker at 180 rpm to allow biofilm formation. After 18 hours, plates were checked for the homogeneity of biofilm growth under optical microscope (Zeiss, Model Axiosvert A1, Germany) at 4× and 10× magnification.

Dyes and laser sources

The study has been realized with three different laser diode prototypes, gently provided by the Department of Engineering and Architecture of the University of Parma, used with or without photosensitizing dyes coupled on the basis of colour affinity: red diode laser (650 nm) with toluidine blue, blue-violet diode laser (405 nm) with curcumin, and green diode laser (532 nm) with erythrosine. The final working concentration was 100 µM for erythrosine and curcumin, and 10 µM for toluidine blue; these concentrations were chosen on the basis of the effect obtained on C. albicans suspensions as described in a previous study. All the laser prototypes used have been tested with a power meter (PM-200, Thorlabs). Laser irradiation has been performed in continuous mode for the different wavelengths. On the basis of the power (30 mW) previously recorded with a power meter (Nova II, Ophir) and on the spot size (5 mm diameter), time irradiation (62 seconds) was planned for a fluence of 10 J/cm².

Killer peptide

The previously described synthetic killer decapeptide KP, endowed with candidacidal activity, was used in this study. A stock solution was prepared in DMSO (20 mg/ml) and stored at 4°C until use.

Biofilm treatment

96-wells microtitre plates incubated for 18 h for biofilm growth, as described above, were washed twice with 0.1 M phosphate-buffered saline (PBS) to remove non-adherent cells before treatment. 100 µl of a KP solution in sterile water (final concentration 20 µg/ml), dye solutions or sterile water alone (control) were added into the proper wells. After 5 minutes laser irradiation was applied to the proper wells. All the plates were left at room temperature for the same time. After laser application, associated or not with dyes and KP treatment, plates were re-incubated for 2 h at 37°C on an orbital shaker at 180 rpm. Each assay was carried out in triplicate and three independent experiments were carried out.

Evaluation of biofilm viability by XTT assay

A semiquantitative measure of biofilm viability was then assessed using a 2, 3-bis-(2-methoxy-4-nitro-5-sulfo-phenyl)-2H-tetrazolium-5-carboxanilide (XTT) reduction assay (Sigma-Aldrich, USA). Briefly, after washing 100 µl of sterile water were added to each well. Then 50 µl of
the XTT reaction mixture (activation reagent and XTT reagent) prepared according to manufacturer’s recommendations were added. The plates were incubated in the dark at 37°C for 2 h and the colorimetric change resulting from XTT reduction, directly related to biofilm metabolic activity, was measured in a microtitre plate reader (ELx800, Biotek Instruments, USA) at 490 nm. Percentage of inhibition in comparison to untreated controls was calculated by the following formula: [(control - treatment)/ control] × 100.

**Fluorescence microscopy studies**

For fluorescence microscopy studies, after laser irradiation, associated or not with dyes and KP treatment, the 96-wells plates were rinsed two times with PBS, then Calcofluor White (Invitrogen, Paisley, UK) and propidium iodide (Invitrogen,) were added at final concentrations of 25 μM and 20 μM, respectively. Plates were incubated for 30 min in the dark. Observations were performed under a fluorescent microscope (Zeiss, Model Axiovert A1, Germany) at 10 × magnification: *Candida albicans* cells in biofilms were stained by calcofluor white and visible in blue colour, *Candida albicans* dead cells were stained by propidium iodide and visible in red colour.

Pictures were taken using AxioVision 4.8 acquisition software and digital AxioCam ICc1 Rev.4 camera (Zeiss).

**Statistical analysis**

Statistical analyses were performed using GraphPad Prism 6 software.

Data are reported as the mean ± standard deviation (SD) from triplicate samples and were evaluated using one-way ANOVA. Multiple comparison analysis was realized with Tukey’s test. A value of *p* < 0.05 was considered significant.

**Results**

**Effect on *C. albicans* biofilm of treatment with different combinations of light, dyes and KP**

The effect on *C. albicans* biofilm viability of the treatment with different combinations of diode laser application, dyes and KP are shown in Figures 1-4. No difference in *C. albicans* biofilm viability has been observed with respect to the untreated control after the application of red, blue, and green diode lasers alone.

While treatment with curcumin and erythrosine alone at the adopted concentrations caused a very significant reduction in biofilm viability (*p* < 0.0001), treatment

**Figure 1:** Effects on *C. albicans* biofilm viability, as assessed by the XTT assay, of treatment with different combinations (+ with, - without) of red diode laser (650 nm, L), toluidine blue (TB) and synthetic killer decapetide (KP). Results are expressed as mean values and standard deviation (SD) of the optical density (OD) at 490 nm. There was no significant difference vs control (no treatment) nor among the different treatments. Data were analysed by one-way ANOVA and Tukey post hoc test. Data reported are from one representative experiment out of three experiments with comparable results.

**Figure 2:** Effects on *C. albicans* biofilm viability, as assessed by the XTT assay, of treatment with different combinations (+ with, - without) of blue diode laser (405 nm), curcumin (CUR) and synthetic killer decapetide (KP). Results are expressed as mean values and standard deviation (SD) of the optical density (OD) at 490 nm. Data were analysed by one-way ANOVA and Tukey post hoc test. *p* < 0.05, **** *p* < 0.0001, significant difference vs. control (no treatment); bars with the same letter represent no significant difference. Data reported are from one representative experiment out of three experiments with comparable results.
with toluidine blue alone showed no effect. Treatment with KP alone caused a reduction of biofilm viability (average inhibition of approximately 26%).

Any combined application of red diode laser with toluidine blue and/or KP gave no significant inhibition of biofilm viability (Figure 1).

In the comparison with the untreated control, the combined application of blue diode laser with curcumin and/or KP showed always a very significant inhibition, as the combination of curcumin and KP did (p < 0.0001), while combination of blue diode laser and KP gave a less significant inhibition, the same obtained with KP alone (p < 0.05). However, there was no statistically significant difference between curcumin alone and any other treatment combined with this dye, even if a light increase in % inhibition was observed when blue laser and KP were associated (90.08 vs 92.14 respectively) (Figures 2 and 4).

In the comparison with the untreated control, the combined application of green diode laser with erythrosine and/or KP showed always a very significant inhibition, as the combination of erythrosine and KP did (p < 0.0001), but no difference was observed with respect to the treatment with erythrosine alone. Again, combination of green diode laser and KP gave a significant inhibition (p < 0.01), although paradoxically lower than the one obtained with KP alone (p < 0.0001) (Figure 3).

Results obtained with XTT assay were implemented by fluorescence microscopy analysis. Staining with calcofluor white, which binds to yeast cell wall, allowed to visualizing all *C. albicans* cells in blue, while internalization of propidium iodide allowed to visualize in red dead *C. albicans* cells. Representative images are shown in Figures 5-6. Treatment with KP alone, while reducing biofilm viability, as assessed by evaluation of metabolic activity through XTT assay, did not cause *C. albicans* death in the adopted experimental conditions. Application of red, blue and green lasers alone did not kill *C. albicans* cells, nor combined treatment with toluidine blue, red laser and KP did. On the contrary, combined treatment with blue laser, curcumin and KP, as well as green laser, erythrosine and KP led to death most *C. albicans* cells (Figure 5-6).

**Discussion**

A number of previously published papers reported the effect of APDT on *C. albicans* biofilms; in most studies high fluences, up to 350 J/cm² [17] and a long time of ap-

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**Figure 3:** Effects on *C. albicans* biofilm viability, as assessed by the XTT assay, of treatment with different combinations (+ with, - without) of green diode laser (532 nm), erythrosine (ERY) and synthetic killer decapptide (KP). Results are expressed as mean values and standard deviation (SD) of the optical density (OD) at 490 nm. Data were analysed by one-way ANOVA and Tukey post hoc test. **p < 0.01, ****p < 0.0001, significant difference vs control (no treatment); bars with the same letter represent no significant difference. Data reported are from one representative experiment out of three experiments with comparable results.

**Figure 4:** Percentage of inhibition of biofilm viability, in comparison to untreated controls, after treatment with different combinations of laser (L) and dyes (red diode laser, 650 nm, and toluidine blue, TB; blue diode laser, 405 nm, and curcumin, CUR; green diode laser, 532 nm, and erythrosine, ERY) and synthetic killer decapptide (KP).
Figure 5: Fluorescence microscopy images of (A) *C. albicans* cells and (B) *C. albicans* dead cells in biofilm treated with blue laser alone; (C) *C. albicans* cells and (D) *C. albicans* dead cells in biofilm treated with Curcumin, blue laser and KP; scale bar of 250 µm, magnification 10x. Panel A and C, staining with calcofluor white; panel B and D, staining with propidium iodide. The absence of candidacidal effect for blue laser alone is visible in panel B as absence of propidium iodide (red) cells.

Figure 6: Fluorescence microscopy images of (A) *C. albicans* cells and (B) *C. albicans* dead cells in biofilm treated with green laser alone; (C) *C. albicans* cells and (D) *C. albicans* dead cells in biofilm treated with Erythrosine, green laser and KP; scale bar of 250 µm, magnification 10x. Panel A and C, staining with calcofluor white; panel B and D, staining with propidium iodide. The absence of candidacidal effect for green laser alone is visible in panel B as absence of propidium iodide (red) cells.
plication up to 26 minutes\textsuperscript{18} were used. We purposely choose a light fluence (10 J/cm\textsuperscript{2}), exportable in clinical practice without danger, discomfort or damage risk for patients or oral appliances (e.g. prosthetic devices).

Evaluation of \textit{C. albicans} biofilm inhibition was realized in this study by means of the XTT assay, a method for quantitative evaluation of metabolic activity that is not affected by inter-operator variability and is easier to perform than the CFU assay. Moreover, literature highlighted the limitation of CFU count for \textit{C. albicans} biofilms as multicellular fungi, while presenting a large biomass compared to single yeast cells, yet form a single colony in CFU assay\textsuperscript{19, 20}.

Previous studies reported that microorganisms organized in biofilms are less susceptible to photodynamic procedure compared with those in planktonic phase because of the heterogeneity of the biofilm, the reduced growth rate of cells, the differences in gene expression, and the limited penetration of antimicrobial agents across the extracellular matrix material\textsuperscript{21}.

In this study, a very significant inhibition of \textit{C. albicans} biofilm has been obtained with curcumin and erythrosine alone. Dyes concentrations were chosen on the basis of a previous study performed in vitro on \textit{Candida albicans} suspensions and \textit{in vivo} on a \textit{Galleria mellonella} model\textsuperscript{22}. In the previous study, dyes alone gave different results. The diverse adopted conditions, i.e. \textit{C. albicans} cells in suspensions compared to \textit{C. albicans} in biofilm structures, may likely be responsible for the observed differences.

Martins and coworkers in 2009\textsuperscript{23} reported that curcumin is 2.5 fold more potent than fluconazole at inhibiting the adhesion of \textit{C. albicans} to buccal epithelial cells. Curcumin acts by generating reactive oxygen species, triggers an early apoptosis in \textit{C. albicans} cells and affects membrane associated ATPase activity, ergosterol biosynthesis and protein secretion.

Some studies in the past suggested that curcumin alone, also at low concentrations (100 \textmu M), may have antifungal properties that may be improved and increased in combination with laser light\textsuperscript{25, 26}.

Dovigo and coworkers\textsuperscript{27} reported on \textit{C. albicans} biofilm a reduction of about 80\% with curcumin 40 M and 18 J/cm\textsuperscript{2}; our results showed a 90\% inhibition using a lower fluence (10 J/cm\textsuperscript{2}) and a higher curcumin concentration.

Sanità and coworkers reported in 2017 the same positive effects on \textit{C. dubliniensis} but using a light emitting diode (LED) source at 5.28 J/cm\textsuperscript{2} and describing better results for a longer preirradiation time (20 minutes)\textsuperscript{28}.

With regard to the combination of green 532 nm laser and erythrosine, at our knowledge there are no similar protocols in literature; available studies describe combination of erythrosine with green or blue-violet LED, requiring longer irradiation time to obtain the same fluence values\textsuperscript{29}. The results we reported in this study for erythrosine are consistent with results of a previously published study\textsuperscript{16} on an \textit{in vivo} model of \textit{Galleria mellonella} where, with the same dye concentration of this study (100 M) erythrosine alone had a candidacidal effect.

At last, with reference to the combination of toluidine blue and red laser, our results are in contradiction with the study of Pupo and coworkers\textsuperscript{27} which reported a value of viable \textit{C. albicans} cells significantly reduced after PDT with methylene blue and toluidine blue regardless of the photosensitizer used, even if with a higher fluence of 53 J/cm\textsuperscript{2}.

Moreover, literature describes different times of pre-irradiation, adfming that killing is not preirradiation time-dependent for planktonic cultures but it must be higher in biofilms cultures\textsuperscript{30}, so a prolonged time of pre-irradiation may be considered to improve the efficacy of treatments in our biofilm model.

**Conclusion**

In our study, curcumin and erythrosine at the chosen concentrations proved to be able to significantly reduce \textit{C. albicans} biofilm viability even without KP or laser association. Conversely, in these experimental conditions, no association of red laser, toluidine blue and KP was able to reduce biofilm viability. These considerations may be a key point for future studies where a reduced dye concentration or the increase of the fluence (for the red laser) could be investigated. The potential advantages of APDT in terms of reduced costs, scarce side effects, low overdose risk and unlikely resistance induction, are compatible with the optimization of APDT protocols in the future\textsuperscript{29-35}.

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**Conflicts of interest**

“There are no conflicts to declare.”