Low-Temperature Plasma: An Effective Approach Against Candida albicans Biofilm

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ABSTRACT: This study evaluated the antifungal potential of low-temperature plasma (LTP) on a 72-hour Candida albicans biofilm. A growth inhibition zone test was conducted with agar plates inoculated with C. albicans and submitted to LTP and argon application at 3 and 10 mm for 10, 30, 60, 90, and 120 seconds. The groups for biofilm assays were 60 seconds of LTP application with a tip-to-sample distance of 3 mm (LTP-3) and 10 mm (LTP-10); –application of only argon gas for 60 seconds with a tip-to-sample distance of 3 mm (Ar-3) and 10 mm (Ar-10); and no treatment. The C. albicans biofilm was grown on saliva-coated discs. The medium was replaced every 24 hours. Confocal laser scanning microscopy revealed the proportion of live and dead cells, and variable pressure scanning electron microscopy (VPSEM) showed biofilm/cell structure. No inhibition zone was observed for control and either Ar groups. For the LTP groups, a progressively increasing of inhibition zone diameter was observed for different treatment durations. The LTP-3 and LTP-10 groups presented higher proportions of dead cells compared with the Ar-3 and Ar-10 groups. VPSEM revealed cell perforations in the LTP-3 and LTP-10 groups. A short period of LTP exposure demonstrated an antifungal effect on C. albicans biofilm.

KEY WORDS: low-temperature plasma, oral biofilm, Candida albicans

I. INTRODUCTION

Oral candidosis is a common oral infection caused by Candida albicans and occurs in 10–75% of denture wearers. A major virulent attribute of C. albicans is its ability to adhere to a surface and form communities known as biofilms. Considering that cells within biofilms are protected from adverse environmental and host defense mechanisms, in comparison to planktonic cells, any method of microbial decontamination should be evaluated against a microbial biofilm structure.

In dental prostheses the mucosa and fitting surfaces of complete dentures act as main reservoirs for yeasts. In addition, C. albicans in the oral cavity might lead to systemic infections, which are difficult to treat and have been linked to several cases of mortality (40%).

Currently, products containing peroxides, hypochlorites, and chlorhexidine diglu-
conates are recommended for denture cleaning to remove *C. albicans*. Discoloring by oxygenating agents or black staining, however, have been considered some of the major side effects of its routine use. Therefore new approaches against *C. albicans* biofilms should be developed without damaging the living tissue and denture material.

Low-temperature plasma (LTP) is emerging as a physical treatment with microbicidal effectiveness on bacteria, parasites, fungi, spores, and viruses. Plasma is a partially ionized gas generated by an electrical discharge, which creates a highly reactive environment with ions, electrons, excited atoms and molecules, vacuum ultraviolet and ultraviolet (UV) irradiation, free radicals, and chemically reactive particles. The production of stable plasma at atmospheric pressure has attracted attention for treating living cells and tissues without thermal damage. It is also site specific, targeting only the infected area, and it seems to preserve the material’s bulk properties. In addition, plasma is usually produced by low-toxicity gases and elaborates its activity by producing a mixture of products that decay within a few seconds after the treatment process. Therefore, this approach has been suggested as environmentally friendly with no harmful residues.

This study aimed to evaluate the antifungal potential of LTP on a 72-hour *C. albicans* biofilm grown on polymethyl methacrylate discs. The research hypothesis assumed that LTP application has antifungal activity on *C. albicans* biofilm, regardless of the distance between the LTP tip and the sample.

### II. MATERIALS AND METHODS

#### A. Experimental Design

A 72-hour *C. albicans* (strain SC5314) biofilm was grown on polymethyl methacrylate (PMMA) resin discs covered by human saliva. The 72-hour biofilm assay was established based on a previous evaluation of biofilm development in variable-pressure scanning electron microscopy (VPSEM) (EVO-50; Carl Zeiss, Germany) from 24 to 96 hours (data not shown). According to this analysis, a dense polysaccharide matrix was progressively generated on the surface of the disc for up to 72 hours. After 96 hours, however, yeast cells started to detach from the polysaccharide matrix because of the biofilm characteristic of the cells’ dissemination. After biofilm formation, the samples were randomly divided into 5 groups (Table 1).

#### B. Specimen Preparation

PMMA (ProBase Cold Resin; Ivoclar Vivadent, Canada) discs, 9 mm in diameter and 2 mm thick, were fabricated according to the manufacturer’s instructions. The specimens then were stored in distilled water at 37°C for 7 days for residual monomer release and were ground using progressively smoother sandpaper (400-, 600-, 800-, and 1200-grit) (Buehler, Germany).
The surface roughness (Ra) of the specimens was subsequently measured using an interferometer (AXIO Imager M1m; Carl Zeiss). Three readings were made for each specimen and the Ra was standardized at 0.34 ± 0.02 μm.1

After the Ra was measured, the specimens were ultrasonically cleaned (Ultrasonic Cleaner FS140; Fisher Scientific) in ultrapure water (MilliQ system; Millipore, Billerica, MA) for 20 minutes to remove any contaminants and artifacts from the surfaces.28 Then, the samples were submitted to a single dose of 25.2-kGy gamma radiation (Gammacell 1000 Elite; Best Theratronics, Canada) for sterilization.

C. Biofilm Assay

A loopful of stock yeast culture of C. albicans (strain SC5314) was reactivated from its original culture in a Sabouraud dextrose agar plate (Becton, Dickinson and Co., Franklin Lakes, NJ) for 24 hours at 37°C.

Five colonies of C. albicans were resuspended in 10 mL of culture medium containing Yeast Nitrogen Base (Becton, Dickinson and Co.) supplemented with 50 mmol/L dextrose (Fisher Scientific). The suspension was cultured overnight at 37°C and 5% carbon dioxide (CO₂) and used to prepare the inoculum at a 1:100 ratio of yeast suspension and culture medium.

The biofilm was formed on saliva-coated PMMA discs through incubation with clarified human whole saliva for 1 hour at 37°C in a 3-dimensional rotator (LabLine; Thermo Scientific). The human saliva was collected from a healthy donor by stimulation with flexible film (Parafilm M; Pechiney Laboratory Safety Products & Apparel) in an ice-chilled polypropylene tube. Because no identifying information about the donor was collected, this research does not meet the definition of human subjects 45 CFR 46.102(f) and does not require institutional review board oversight. Adsorption buffer (50 mmol/L potassium chloride, 1.0 mmol/L potassium phosphate, 1.0 mmol/L calcium chloride, 0.1 mmol/L magnesium chloride; pH 6.5; 1:1), as well as phenylmethylsulfonyl fluoride

<table>
<thead>
<tr>
<th>Group</th>
<th>Treatment</th>
<th>Tip-to-Sample Distance* (mm)</th>
<th>Treatment Duration (seconds)</th>
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<tr>
<td>Control</td>
<td>No treatment</td>
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<tr>
<td>Ar-3</td>
<td>Argon gas application</td>
<td>3</td>
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<tr>
<td>Ar-10</td>
<td>Argon gas application</td>
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<td>LTP-3</td>
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<td>LTP-10</td>
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*Tip-to-sample distance represents the distance between the sample and the device used for argon and argon–low-temperature plasma (LTP) application.
(PMSF; 1:1000), were added to the saliva. The solution was clarified by centrifugation at 8500 rpm for 10 minutes at 4°C, filtered using a filter with a 0.22-μm pore size (Stericup; Millipore), and immediately used.

The saliva-coated discs were washed 3 times in sterile adsorption buffer, transferred to a new 24-well polystyrene cell culture plate (Becton, Dickinson and Co.), covered by 2 mL of the yeast suspension, and incubated at 37°C and 5% CO₂. The medium was changed every 24 hours up to 72 hours before treatment was accomplished. The procedure was performed in triplicate on 3 independent occasions.

**D. LTP Treatment**

An atmospheric-pressure LTP jet (Kinpen) developed by the Leibniz Institute for Plasma Science and Technology (Greifswald, Germany) was used for plasma treatment. The device consists of a hand-held unit (length = 170 mm, diameter = 20 mm, weight = 170 g) that generates a plasma jet at atmospheric pressure, a direct current power supply (system power: 8 W at 220 V, 50/60 Hz), and a gas supply unit. A pin-type electrode (1-mm diameter) is mounted at the center of a quartz or ceramic capillary (inner diameter = 1.6 mm). In the continuous working mode, a high-frequency voltage (1.82 MHz, 2–6 kVpp) is coupled to the pin-type electrode. The plasma is generated from the top of the electrode and expands to the surrounding air outside the nozzle² (Fig. 1).

The system works with all rare gases at gas flow rates ranging between 5 and 10 slm (standard liters per minute). In this study the device worked with argon (Ar) gas at a 5-slm flow rate. The temperature was measured with a thermometer for 2 minutes and resulted in 30.67 ± 0.58°C at a 3-mm distance from the plasma tip and 29.83 ± 0.29°C at

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**FIG. 1:** (a) Kinpen plasma unit. (b) Low-temperature plasma application on a polymethyl methacrylate sample
10 mm. The Ar gas flow/LTP plume was targeted to the center of the disc, and the tip-to-sample distance (3 or 10 mm) was kept constant during application using a fixed holder.

**E. Growth Inhibition Zone**

After reactivation of *C. albicans* (strain ATCC SC5314) culture, 5 colonies were resuspended in 10 mL of Yeast Nitrogen Base (Becton, Dickinson and Co.) supplemented with 50 mmol/L dextrose (Fisher Scientific) and cultured overnight at 37°C and 5% CO₂. An inoculum of yeast suspension at a 1:20 ratio was inoculated in culture medium and incubated at 37°C and 5% CO₂ for 8 hours. The incubation time was calculated to reach the exponential growth phase.

The cells were washed 3 times with sterile 1% phosphate-buffered solution (pH 7.4) (Invitrogen) by centrifugation at 8500 rpm for 10 minutes at 4°C and standardized to 1 × 10⁷ cells/mL, which was ascertained through a spectrophotometer (Genesys 6 spectrophotometer; Thermo Scientific) at an absorbance of about 0.209 at 660 nm.

Then, 50 μL of the *C. albicans* suspension was spread on Sabouraud dextrose agar (Becton, Dickinson and Co.) plates and pure Ar gas (Ar groups) or plasma (LTP groups) was applied. Both treatments were conducted at different tip-to-plate distances (3 and 10 mm) and for different durations (15, 30, 60, 90, and 120 seconds). A negative control group with no treatment also was evaluated. After treatment, the plates were incubated at 37°C and 5% CO₂ for 24 hours. Then, the diameter of the growth inhibition zone was measured with a digital caliper. The procedure was performed in triplicate in 3 independent occasions.

**F. Confocal Laser Scanning Microscopy**

The Live/Dead BacLight bacterial viability kit (Molecular Probes; Invitrogen) was used to reveal the proportion of live or active cells (fluorescent green) and dead or inactive cells (fluorescent red). The live/dead stain was prepared by diluting 1.5 μL of staining component A (SYTO 9) and 1.5 μL of staining component B (propidium iodide) in 1 mL of sterile 1% phosphate buffered solution (pH 7.4) (Invitrogen, USA). Each sample was covered with 1 mL of the reagent mixture and incubated for 15 minutes at room temperature, protected from light exposure. Fluorescence was captured with a Leica TCS SP5 II confocal microscope (Leica, Germany) with the objective Leica HCX APO L 40x/0.8 W U-V-I water dipping lens.

**G. Variable Pressure Scanning Electron Microscopy**

VPSEM was used in this study because it allows morphological analysis of the treated mature biofilms with no further preparation. The VPSEM (EVO 50; Carl Zeiss) in com-
Combination with the software SmartSEM (Carl Zeiss) was used for analysis. The images were captured at an extra high tension (EHT) of 15 Kv, probe current of 300 pA, vacuum of 100 Pa, and a working distance of 10.5 mm.

III. RESULTS

A. Growth Inhibition Zone

No growth inhibition zone was observed for either the negative control group or those cultures submitted to Ar gas application under all conditions (15, 30, 60, 90, and 120 seconds’ duration/3 and 10 mm distance) (Fig. 2).

**FIG. 2:** *Candida albicans* culture in Sabouraud dextrose agar. A growth inhibition zone was observed for groups treated with low-temperature plasma (LTP). Ar, argon. Ar-3, argon gas application for 60 seconds with a tip-to-sample distance of 3 mm; Ar-10, argon gas application for 60 seconds with a tip-to-sample distance of 10 mm; LTP-3, 60 seconds of LTP application with a tip-to-sample distance of 3 mm; LTP-10, 60 seconds of LTP application with a tip-to-sample distance of 10 mm.

**FIG. 3:** Mean diameter (millimeters) of the growth inhibition zone exhibited for plasma application.
For those cultures submitted to LTP treatment, the longer the treatment, the wider the growth inhibition zone at both tip-to-sample distances (3 and 10 mm). In addition, wider growth inhibition zones were observed when LTP application was conducted at a tip-to-sample distance of 3 mm compared with 10 mm for all treatment durations (Fig. 3).

**B. Confocal Laser Scanning Microscopy**

Figures 4 and 5 shows the proportion of live/dead cells in each group at 2× and 10×, respectively. Figure 4 visually shows a larger amount of dead cells in the LTP groups compared with the control and Ar groups, whereas Fig. 5 reveals morphological alterations of the yeasts in the LTP groups.

**FIG. 4:** Comparative analysis of *Candida albicans* biofilm in each group at 2×. Ar-3, argon gas application for 60 seconds with a tip-to-sample distance of 3 mm; Ar-10, argon gas application for 60 seconds with a tip-to-sample distance of 10 mm; LTP-3, 60 seconds of LTP application with a tip-to-sample distance of 3 mm; LTP-10, 60 seconds of LTP application with a tip-to-sample distance of 10 mm
C. Variable Pressure Scanning Electron Microscopy

The VPSEM images revealed cell perforations for the biofilm treated with LTP at both tip-to-sample distances (groups LTP-3 and LTP-10). On the other hand, no morphological alteration was observed for the biofilm of the negative control group (control group) and those submitted to only Ar gas application (groups Ar-3 and Ar-10) (Figs. 6 and 7).

IV. DISCUSSION

According to the results of this study, the research hypothesis was accepted because LTP exhibited antifungal activity on *C. albicans* biofilm for both tip-to-sample distances.

The growth inhibition zone observed only on LTP-treated samples demonstrated that the activity against planktonic cells was promoted by plasma activation and not

FIG. 5: Comparative analysis of *Candida albicans* biofilm in each group at 10×. Ar-3, argon gas application for 60 seconds with a tip-to-sample distance of 3 mm; Ar-10, argon gas application for 60 seconds with a tip-to-sample distance of 10 mm; LTP-3, 60 seconds of LTP application with a tip-to-sample distance of 3 mm; LTP-10, 60 seconds of LTP application with a tip-to-sample distance of 10 mm
by Ar gas flow. Data in the literature also show the effect of plasma on the inactivation of planktonic bacteria, yeasts, and spores.29–31 Similar to another study,10 the measured inhibition zone diameters yield important information on the effective coverage area of

**FIG. 6:** Variable pressure scanning electron microscopy of samples of all groups. Ar-3, argon gas application for 60 seconds with a tip-to-sample distance of 3 mm; Ar-10, argon gas application for 60 seconds with a tip-to-sample distance of 10 mm; LTP-3, 60 seconds of LTP application with a tip-to-sample distance of 3 mm; LTP-10, 60 seconds of LTP application with a tip-to-sample distance of 10 mm.
the active species generated by the plasma, which extended beyond the visible plasma plume.

Although the clear growth inhibition zone demonstrated the effect of plasma on planktonic cells, it has been repeatedly confirmed that *C. albicans* biofilms are more resistant to antifungal agents than planktonic structures (free-floating, nonadherent cells).\(^{10,32}\) The extracellular polysaccharides secreted by *C. albicans* after adhesion to dentures are the basis for the biofilm matrix, which protects the cells from different physical and chemical environmental influences.\(^2\)

The results of confocal laser scanning microscopy confirmed the antifungal effect of LTP on *C. albicans* biofilm; the plasma-treated samples (groups LTP-3 and LTP-10) exhibited a larger amount of dead cells than the negative control group and those samples submitted to only Ar gas application (groups Ar-3 and Ar-10). Koban et al.\(^2\) also demonstrated the sterilization promoted by LTP on *C. albicans* biofilm. However, their results corresponded to a 48-hour biofilm on titanium discs, and the treatment was longer (1–10 minutes), which could be inappropriate for clinical routine. Furthermore, in the present study, LTP showed an antimicrobial effect against a mature *C. albicans* biofilm in a short exposure period (60 seconds). Similar to another study,\(^{33}\) confocal microscopy also revealed dead cells in those samples submitted to only Ar gas application. Although this result could suggest a certain antimicrobial effect of irrigation with noninert gases\(^{34}\) even without plasma activation, dead microorganisms within a biofilm model grown under stationary conditions have been shown previously.\(^{35}\)

The inactivation of biological agents promoted by LTP may result from deconstruction of the microorganism’s genetic material (DNA) by UV radiation produced with plasma and erosion of the microorganisms through intrinsic photodesorption. The photon-induced desorption results from damage of chemical bonds in the microorganism after exposure to UV radiation, allowing its atoms to form volatile compounds.\(^{23,36}\)

**FIG. 7:** (a) Biofilm not exposed to low-temperature plasma (LTP) treatment. (b) Cell damage after LTP treatment

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In addition, the reactive oxygen species (ozone, atomic oxygen, superoxide, peroxide, hydroxyl radicals, and nitric oxide) produced by plasma react with cellular biomacromolecules, including DNA, lipids, and proteins. The electrostatic forces generated by plasma-charged particles may also result in membrane lysis.

The *C. albicans* cell wall is composed of mannoproteins (about 40% of the cell wall biomass), β-1,3-glucan (the major stress-bearing polysaccharide in the wall), β-1,6-glucan (a water-soluble component that interconnects mannoproteins to β-1,3-glucan), and chitin (a linear stress-bearing polysaccharide). Therefore, these major components are easy targets for the reactive species produced by LTP.

Koban et al. also stated that disinfection with plasma is based on short-lived active species, UV radiation, electrostatic effects, and reactive oxygen species that cause lipid peroxidation. According to Stoffels et al. and Pappas, even when no significant UV emission is present with low-temperature, atmospheric-pressure plasmas, the synergy of other species such as radicals and charged particles still plays a dominant role in the sterilization process.

The assumption about membrane damage caused by plasma was confirmed by the confocal and VPSEM images, revealing cell lyses (Figs. 5–7). Membrane perforation was previously reported; plasma reactive particles produce a general mechanical effect on the surface of living organisms called “etching.” Etching results from the reaction of highly reactive gas radicals with organic materials, generating by-products that are desorbed from the surface. For both plasma tip-to-sample distances, however, the effect was limited to the center of the disc submitted to the LTP treatment. A similar trend was observed by Koban et al., revealing that the narrow plasma afterglow damages only yeast cells at the actual touch point, and the reactive plasma species do not spread over the entire disc in a sufficient dose. Therefore, additional studies of treatment all over the disc are suggested.

Although it has been suggested that plasma does not induce bulk modifications on materials, additional studies are being conducted to assess the effect of plasma on the mechanical and physicochemical properties of PMMA samples at different scenarios (exposure time, plasma tip-to-sample distance, gas composition, and so on).

It is important to highlight that *C. albicans* frequently adheres not only to denture surfaces but also to other medical devices, such as voice prostheses, blood and urinary catheters, and heart valves. Therefore, although the applicability of this approach requires more extensive studies, LTP has a broader scope and relevance for antimicrobial treatment.

V. CONCLUSION

A short period of LTP exposure (60 seconds) demonstrated an antifungal effect on *C. albicans* biofilm formed on PMMA, thus accepting the research hypothesis. LTP is a promising treatment for *C. albicans* biofilm–related diseases.
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REFERENCES


