**Candida albicans** biofilm on titanium: effect of peroxidase precoating

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**Abstract:** The present study aimed to document **Candida albicans** biofilm development on titanium and its modulation by a peroxidase-precoated material which can generate antimicrobials, such as hypoiodite or hypothiocyanite, from hydrogen peroxide, iodide, or thiocyanate. For this purpose, titanium (powder or foil) was suspended in Sabouraud liquid medium inoculated with **C. albicans** ATCC10231. After continuous stirring for 2–21 days at room temperature, the supernatant was monitored by turbidimetry at 600 nm and titanium washed three times in sterile Sabouraud broth. Using the tetrazolium salt MTT-formazan assay, the titanium-adherent fungal biomass was measured as $7.50 \pm 0.60 \times 10^6$ blastoconidia per gram of titanium powder ($n = 30$) and $0.50 \pm 0.04 \times 10^6$ blastoconidia per cm² of titanium foil ($n = 12$). The presence of yeast on the surface of titanium was confirmed by microscopy both on fresh preparations and after calcofluor white staining. However, in the presence of peroxidase systems (lactoperoxidase with substrates such as hydrogen peroxide donor, iodide, or thiocyanate), **Candida** growth in both planktonic and attached phases appeared to be inhibited. Moreover, this study demonstrates the possible partition of peroxidase systems between titanium material (peroxidase-precoated) and liquid environment (containing peroxidase substrates) to limit **C. albicans** biofilm formation.

**Keywords:** adhesion, material, oral, yeast

**Introduction**

Yeasts can colonize titanium surfaces even after implantation. Because yeasts form biofilms on titanium surfaces, they can become the source of device-related infections with subsequent morbidity and occasionally death of the patient, all bearing high costs. Yeast colonization is frequently found in catheters and indwelling devices in cancer patients, for whom parenteral nutrition and immunosuppression are aggravating factors.1–4 Fungal biofilms are difficult to treat on account of their increased resistance to antimicrobial agents. This is the reason why fungemia associated with catheterization or prosthetic devices should prompt the removal of these implanted materials, because antifungal drug treatment remains an uncertain alternative.5,6

Prophylactic antibiotic therapy to prevent bacterial device-associated infections occasionally leads to increased mycotic complications.4 In dentistry, **Candida** species and periodontal pathogens have been isolated around titanium implants.7,8 Titanium-coated silicone was thought to prevent **Candida** biofilm formation on voice prostheses by increasing the smoothness of the material surface.9 Titanium material does not possess antifungal properties by itself, but TiO₂ can act as a photocatalyst for the decomposition of organic compounds under ultraviolet light, thereby reducing the viability of **Candida albicans.**10,11 Bioactive coating, used as an osseointegration
facilitator, is another option for controlling biofilm formation. In order to reduce bacterial and fungal infections, some groups have proposed the precoating of orthopedic devices with antimicrobial agents (chlorhexidine/chloroxylenol). However, bioactive protein coating is not ideal for in vivo use because it can induce adverse immunologic reactions, yet remains an interesting area for studying biofilms in vitro. Adsorption of lactoperoxidase (LP) onto titanium surfaces, which has antimicrobial properties similar to those of oral peroxidase, has already been described. LP produces hypo-thiocyanite (OSCN⁻) and hypohydrocyanic acid (HO SCN⁻) compounds from thiocyanate (SCN⁻) in the presence of hydrogen peroxide (H⁡₂O₂). In vitro, iodide (I⁻) has also been used as substrate, because it is oxidized into hypooiodite (OI⁻) or hypoiodous acid (OIH) by peroxidase activity.

Nevertheless, attempts to understand interface biology better and to improve material surface treatment need a model of biofilm on titanium pieces that are easy to manipulate in the laboratory. Few studies have developed a titanium model in order to investigate procedures for limiting yeast adherence and biofilm formation. The present study aimed at demonstrating and quantifying both Candida adherence to titanium (powder and foil) and its modulation in the presence of peroxidase systems.

**Materials and methods**

**Materials**

*Candida* adherence to titanium (powder or foil) was evaluated in comparison with its adherence to other materials, such as acrylic resins (hard or soft) and polystyrene, which are commonly used oral biomaterials and laboratory assay tube manufacture materials, respectively. Aliquots (0.5 g) of titanium powder (Alfa Aesar®, Karlsruhe, Germany; 99.5% purity; ∼325 mesh) were transferred into sterile polystyrene cell culture tubes (BD Falcon®, Franklin Lakes, NJ; size 125 × 16 mm, round bottom). Titanium pieces were cut from native foil (Advent®, Oxford; 99.6% purity; thickness 25 μm, size 25 mm × 5 mm). As previously described, each piece was weighed and the total surface was calculated using the following algorithm: 

\[
S = 2 \left( \frac{m}{p} + h(L + 1) \right)
\]

where 

- \( S \) = total surface (mm²), 
- \( m \) = foil mass (mg), 
- \( p \) = volumic mass (4.544 mg/mm³), 
- \( h \) = foil thickness (0.025 mm), 
- \( L \) = foil length (mm), 
- \( l \) = foil width (mm).

Total surface averaged 246 ± 4 mm² (n = 16) with a variation coefficient of 6.5%.

Two different resins for dentures (Vertex soft® and Vertex Rapid Simplified®; Vertex Dental, Zeist, Netherlands) were investigated with and without varnishing (Palaseal®; Heraeus Kulzer, Wehrheim, Germany). Acrylic resin pieces (thickness ∼2 mm, size 25 mm × 5 mm) were processed according to manufacturer’s instructions. The mass of each resin piece averaged 222 ± 19 mg (±SD, n = 96) with a coefficient of variation of 8.6%.

**Cultures of Candida albicans**

*C. albicans* ATCC10231 (Culti-Loops®, Oxoid®, Basing-stoke, UK) was grown at 37°C on Sabouraud-gentamicin-chloramphenicol agar (Becton-Dickinson®, Belgium). Yeasts were harvested from the third subculture and used to inoculate Sabouraud broth adjusted to an absorbance of 0.800 at 600 nm. The *Candida* count was conducted in the adjusted suspension after culture (37°C for two days) on Sabouraud agar of 20 μL aliquots from serial dilutions (until 10⁴-fold) in Sabouraud broth. The number of colony-forming units (CFU) in the initial suspension yielded 16 ± 3 × 10⁶ CFU/mL (n = 6).

**Candida albicans adherence**

Titanium powder (0.5 g) was suspended in 3.6 mL Sabouraud broth and yeast cells (0.4 mL at included concentration) were added. Negative controls consisted of titanium powder incubated in 4 mL sterile Sabouraud broth. After continuous stirring (three revolutions per minute, Stuart® SB3 rotator, Barloworld Scientific, Stone, UK) for two days at room temperature, the titanium powder was separated from the liquid medium by sedimentation (10 minutes) and washed three times with 4 mL sterile Sabouraud broth, changing the assay tube for each washing cycle. From some negative controls, after two days of incubation and removal of sterile Sabouraud, titanium powder was contaminated by 4 mL of the first supernatant from the contaminated assays in order to evaluate washing process efficiency. The *Candida*/titanium contact time was then reduced to sedimentation time (10 minutes). All the supernatants were monitored by turbidimetry at 600 nm to evaluate the planktonic yeast phase. Titanium and resin foils were similarly treated. Material-adherent yeast biomass (anchored yeast phase) was evaluated by the tetrazolium salt MTT (1.7 mg/mL in 2:1 Sabouraud phosphate-buffered solution at pH 7.40) which is enzymatically reduced by viable fungi, with resultant formation of MTT-formazan absorbing at 570 nm. After MTT-formazan extraction with isopropanol, absorbance was immediately measured in a spectrophotometer at 570 nm. The main characteristics of the method were: molar absorption coefficient ε = 570 of 17,000 M⁻¹ × cm⁻¹, sensitivity of 0.5 × 10⁶ blastoconidia/mL, coefficient of variation ≤12.0%, analytic
range $0.5–10 \times 10^6$ blastoconidia/mL. Figure 1 illustrates the rate of tetrazolium salt MTT reduction into formazan in the presence of a fixed number of blastoconidia as a function of the tetrazolium salt concentration.

**Microscopic inspection**

*C. albicans* adherent to titanium powder were observed using a light microscope (fresh preparation) equipped with epifluorescence (after calcofluor white staining). Yeast-contaminated titanium was examined microscopically in the presence of 1 mg/mL calcofluor white M2R stain (fluorescent brightener 28; Sigma®, St Louis, MO) known as a marker of cell wall chitin.

**Lactoperoxidase adsorption on titanium**

LP (Biopole®, Belgium) was dissolved in demineralized water. Pieces of titanium foil were immersed for 30 minutes at 37°C in 5 mL of a 0.8 mg/mL LP solution or in demineralized water alone (control). After incubation, the liquid phase was removed and titanium foils were washed three times with 5 mL of demineralized water and then transferred into another tube with the last washing. This procedure was repeated five times prior to a last removal of the liquid phase.

**Lactoperoxidase systems**

Two different peroxidase systems were tested in this model of yeast biofilm. Firstly, a hydrogen peroxide (H$_2$O$_2$)/LP (1 U/mL)/iodide (I$^-$, 1.2 mM) system generating hypoiodite (OI$^-$) and secondly a H$_2$O$_2$/LP (1 U/mL)/thiocyanate (SCN$^-$, 1.2 mM) system producing hypoiodite (OSCN$^-$). H$_2$O$_2$ was enzymatically produced by glucose-oxidase (GOD, 1.3 U/mL or 0.2 U/mL) and glucose (G) present in the culture medium (115 mM).

**Statistics**

Data were analyzed using the GraphPad Prism version 5.00 (GraphPad Software®, San Diego, CA). ANOVA completed by a Dunnett’s or Bonferroni’s post-test were performed for statistical comparisons using the same software. Mean values were expressed with their related standard error of the mean unless otherwise indicated.

**Results**

*Candida albicans* growth in planktonic phase and adhesion to tested materials

The growing yeast phase in supernatant was evaluated by measuring the 600 nm spectrophotometer absorbance ($A_{600\text{nm}}$) after different periods of incubation (2–21 days). After removal of the supernatant, the immersed materials were washed with sterile broth and the turbidity of the subsequent supernatant after sedimentation was monitored using the same method. Figure 2 illustrates the evolution of $A_{600\text{nm}}$ during the washing process of different materials (titanium powder and foil, resin foil) after a two-day incubation in a contaminated environment. When titanium powder was contaminated by a yeast liquid culture ($A_{600\text{nm}}$: 0.653 ± 0.074, n = 21) just before the washing process, the sampling of the supernatant after a 10-minute sedimentation enabled the recovery of only 86.7% ± 2.9% of the initial absorbance at 600 nm (n = 16). This discrepancy (~13.3%) from the theoretic value of 100% is statistically significant (one-sample t-test, $P = 0.0004$). The three subsequent washing-mixing-sedimentation cycles decreased supernatant turbidity to 7.0% ± 1.0% of the initial value (n = 18) according to a one-phase exponential decay model ($y = 100 e^{-1.26t}$, $r = 0.9940$).

By contrast, when titanium powder was incubated for two days with yeast cells, the liquid-phase turbidity of the third washing in the assays still yielded 110.1% ± 15.0% of...
and resin foils also showed a linear decrease of turbidity in the liquid broth used for the washing solution ($y = -49x + 97$, $r = 0.9948$ and $y = -45x + 1030$, $r = 0.9919$, respectively) rather than a one-phase exponential decay. In the case of titanium foils, absorbance at 600 nm yielded $42.3\% \pm 8.6\%$ ($n = 12$) and $2.0\% \pm 0.5\%$ ($n = 12$) of the initial value in the first and second washing medium, respectively. In the case of the resin foils, it yielded $64.8\% \pm 11.1\%$ ($n = 24$) and $9.5\% \pm 2.0\%$ ($n = 24$) of the initial value in the first and second washing medium, respectively. When contaminated titanium powder was washed immediately, the anchored blastoconidia amount was evaluated as $1.1 \pm 0.1 \times 10^6$ blastoconidia per g of titanium powder.

As shown in Table 1, the MTT-formazan assay conducted in the titanium powder phase after a two-day incubation and three washings showed the presence of a residual adherent biomass equivalent to $7.5 \pm 0.6 \times 10^6$ blastoconidia ($n = 30$) per g of titanium powder instead of $494.9 \pm 40.2 \times 10^6$ blastoconidia per cm$^2$ ($n = 12$) on titanium foil ($13.9 \pm 0.2$ mg, $n = 12$). Moreover, the presence of anchored Candida biofilm was also demonstrated on two different resins (hard and soft, used for denture manufacturing) and Falcon tube polystyrene wall. This amount represented $20.5 \pm 3.9 \times 10^4$/cm$^2$ ($n = 12$), $13.7 \pm 3.2 \times 10^4$/cm$^2$ ($n = 12$) and $36.7 \pm 3.7 \times 10^3$/cm$^2$ ($n = 4$) for soft resin, hard resin, and polystyrene assay tube wall, respectively. Microscopic examination confirmed the presence of yeast on the surface of titanium powder.

Figure 3 demonstrates different morphologic aspects of Candida biofilm onto titanium. Figure 3a shows one titanium grain surrounded by a layer of blastoconidia (magnification $\times$400) after one week incubation and three washes. After a longer incubation (three weeks), filamentous structures (hyphae and pseudohyphae) can be observed (Figure 3b). The lower panel shows titanium particles after three weeks’ incubation and four washings. In this condition, some blastoconidia were still present with a trabecular structure (Figure 3c) which can be stained by the fluorescent Brightener 28, a polysaccharide structure marker for fluorescent light.

**Time course of Candida biofilm production on titanium powder**

Development of yeast biomass was monitored weekly for three weeks in the planktonic phase and in the anchored phase by the procedures mentioned above. Figure 4 (upper panel) reports the time course of the supernatant absorbance at 600 nm, reflecting the planktonic yeast mass. Turbidity reached $0.653 \pm 0.074$ ($n = 21$) after a two-day incubation at room temperature; then it slightly increased (to $0.781 \pm 0.095$, $n = 4$, on day 7; and to $0.929 \pm 0.142$, $n = 4$, on day 14). After the second week, it
Candida adhesion to titanium decreased and reached 0.161 ± 0.058 (n = 4) on day 21, which is less than the values previously obtained (corresponding to 24.7% ± 9.3% of those found on day 2 and 17.3% ± 6.8% of those on day 14). An ANOVA test completed by a Bonferroni’s multiple comparison post-test did not show any difference between days 2, 7, and 14 (P > 0.5); but the turbidity on day 21 was significantly decreased in comparison with values obtained earlier (ANOVA, P = 0.0078). The time course of the anchored yeast mass was rather different (Figure 4, lower panel), reaching steady state on day 2 (0.924 ± 0.060, n = 30) and persisting until day 21 (0.887 ± 0.044, n = 4). Intermediate values were 0.829 ± 0.206 (n = 4) on day 7 and 1.280 ± 0.112 (n = 4) on day 14. An ANOVA test completed by a Bonferroni’s multiple comparison post-test did not show any difference between the anchored yeast mass obtained for different periods of incubation (ANOVA, P = 0.1717).

Modulation of Candida biofilm production on titanium powder by peroxidase systems

In the presence of 1.3 U/mL GOD and 115 mM G, both G/GOD/1/-LP and G/GOD/SCN/-LP systems prevented Candida growth in the planktonic as well as the attached phase for at least 21 days. Table 2 indicated the absorbance values at 600 nm (for the planktonic evaluation) and at 570 nm (for the attached phase evaluation) expressed as the percentage of the time-paired control in the absence of oxidant. Both peroxidase systems maintained the titanium powder free of yeast cells for three weeks, with absorbance at 600 nm equal to zero. Repeated measurements (n = 4) at one, two, and three weeks showed persisting growth inhibition in the broth. By contrast, a limited but existing anchored phase was demonstrated on titanium powder by the MTT procedure.

Table 1 Number of blastoconidia anchored onto different materials (titanium, resin, and polystyrene)*

<table>
<thead>
<tr>
<th>Material</th>
<th>Incubation</th>
<th>Adherent blastoconidia number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Powder</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Titanium</td>
<td>Immediate treatment</td>
<td>1.1 ± 0.1 × 10^6/g (n = 3)</td>
</tr>
<tr>
<td></td>
<td>Two-day</td>
<td>7.5 ± 0.6 × 10^6/g (n = 30)</td>
</tr>
<tr>
<td>Foil</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Titanium</td>
<td>Two-day</td>
<td>494.9 ± 40.2 × 10^3/cm² (n = 12)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(equivalent to 35.6 ± 2.9 × 10^6/g)</td>
</tr>
<tr>
<td>Soft resin</td>
<td>Two-day</td>
<td>20.5 ± 3.9 × 10^3/cm² (n = 12)</td>
</tr>
<tr>
<td>Hard resin</td>
<td>Two-day</td>
<td>13.7 ± 3.2 × 10^3/cm² (n = 12)</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>Two-day</td>
<td>36.7 ± 3.7 × 10^3/cm² (n = 4)</td>
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*Incubation was at room temperature. Material-adherent blastoconidia were evaluated by the MTT procedure. Data are expressed per mass unit for powder or per surface unit for foils.

In the presence of the SCN^- substrate (n = 4), a small anchored phase was observed to evolve from 11.3% ± 12.3%, to 24.9% ± 8.3%, then finally to 18.7% ± 3.5% of the control on days 7, 14, and 21, respectively. In the presence of I^- substrate (n = 4), no anchored biomass could be shown (zero

**Figure 3A** Candida albicans blastoconidia adherent to titanium granule (magnification X400 in the absence of any stain procedure) after a one-week incubation and three washings. **B** Titanium grains surrounded by filamentous structures (hyphae and pseudohyphae) after a three-week incubation. **C** Titanium particles (after three weeks of incubation and four washings) with some remaining blastoconidia and trabecular structure.
absorbance at 570 nm) after one week of incubation at room temperature, and only appeared after 14 days’ incubation (14.5% ± 3.7% of the paired control) and slightly progressed to 20.2% ± 3.7% of the control on day 21. At any incubation time, data from both peroxidase system conditions were statistically lower than their paired controls (ANOVA completed by a Dunnett’s multiple comparison test).

Yeast recovery from deposits onto titanium foil
Droplets (20 µL of sterile water containing from 10 to 200 blastoconidia in five independent experiments) were deposited onto titanium foil and incubated at 37°C for 30 minutes. Foils were then rubbed on Sabouraud solid medium for cultures at 37°C for two days. Yeast recovery (81.9% ± 7.0%, n = 5) was statistically independent (ANOVA, P = 0.6167) of the numbers of blastoconidia deposited onto the foil. Figure 4 reports Candida recovery after rubbing the foil prepared in the following conditions: a) blastoconidia droplets were directly deposited onto foil; b) blastoconidia droplets were deposited onto lactoperoxidase-coated titanium; and c) blastoconidia droplets with added G/GOD were deposited onto lactoperoxidase-coated titanium, d) blastoconidia droplets with added G/GOD/KI were deposited onto lactoperoxidase-coated titanium (n = 5 for each condition). Data were analyzed by an ANOVA test (P = 0.0009) completed by a Dunnett’s multiple comparison test (***P < 0.001).

Figure 4 Time course of yeast biomass in the planktonic phase (top) and in the anchored phase (bottom).
Note: Development of yeast biomass was weekly monitored for three weeks in the planktonic phase (absorbance at 600 nm) and in the anchored phase (absorbance at 570 nm after MTT procedure).

Abbreviations: G, glucose; GOD, glucose oxidase; LP, lactoperoxidase; I, iodide; SCN, thiosulfate.

Discussion
Frequent occurrence of C. albicans at peri-implantitis sites,7 inside port reservoirs of totally implantable vascular access devices,1 and other titanium implants justifies laboratory attempts to devise a reproducible model in order to investigate in vitro yeast biofilm formation and its modulation. This paper shows the reliability of a model using titanium powder and foil as support for C. albicans adhesion and biofilm formation with the aim to study yeast growth inhibitors on biomaterial surfaces, in particular titanium implants. Numerous in vitro biofilm models have already been described on various material surfaces in order to investigate new antimicrobial strategies. However, only a few studies have evaluated titanium support in the yeast biofilm area. One paper15 reported the efficiency of a laser decontamination procedure in dentistry applied to a titanium model derived from previous studies19 of yeast biofilms on denture polymethylmethacrylate surfaces.
The present study uses similar inoculum sizes and procedures for the quantification of yeast biomass anchored to a titanium support, as in the two last quoted papers. Nevertheless, the present model differs in several aspects, ie, the use of material powder increases support surfaces in the same manner as cell cultures on beads. Moreover, adhesion of *C. albicans* to the support precedes the formation of a biofilm. These two steps are not set apart in the model described here, which enables the simultaneous study of yeast growth both in the supernatant (planktonic phase) and in the biofilm (anchored phase). Within a very short time, anchored blastoconidia progressively covered the available surface on the biomaterial. The small but significant decrease of culture liquid turbidity immediately after addition of titanium powder should reflect attachment of *Candida* to titanium, starting as early as the material has been immersed. The anchored blastoconidia amount was then evaluated by using the MTT method as 1.1 ± 0.1 × 10^6 blastoconidia per g of titanium powder, demonstrating a rapid attachment. After covering the entire available surface, biofilm grew following a multilayer pattern (as seen in Figure 3a), susceptible to liberation of numerous blastoconidia during the washing process. This desorption resulted in modifying the decay model of turbidity during the washings. After the longest incubation (three weeks), titanium powder was observed embedded in a network of hyphae. To investigate the time course of events, planktonic and biofilm growths were evaluated between two days and three weeks. As shown in the results section, absorbance at 600 nm of the liquid medium rapidly increased to reach a steady state at two days, then decreased after two weeks, in agreement with theoretic microbiologic growth curves, the decline being attributed to modifications of the culture liquid (eg, nutrient exhaust, toxic catabolites, and pH change). MTT reduction assay data revealed that the *C. albicans* biofilm developed rapidly (at least in two days), while the growth decline stage was delayed (albeit this was not part of this investigation).

Photonic and fluorescent microscopy was performed at all stages in order to visualize the biofilm structure, composed of a dense blastoconidia layer and hyphal forms. At 21 days, added extracellular polymeric material could be observed, with an amorphous reticular appearance. Initially, *C. albicans* cells were observed as blastoconidia, and (pseudo)hyphae appeared later. Production of extracellular polysaccharide material by *Candida* cells and filamentation of yeast cells contribute to modify biofilm architecture into a heterogeneous structure. Further studies should describe more precisely the kinetics of *Candida* biofilm formation on titanium.

Titanium is widely employed for implant manufacturing due to its good biocompatibility and mechanical properties, but infection remains a cause of failure leading to removal. The titanium surface is not antimicrobial by itself, so it could be used as support for a *Candida* biofilm in this investigation. Making implant surfaces resistant to microbial colonization should reduce infectious complications. Such developments need an *in vitro* model which allows investigation of the effect of surface modification and coatings on biofilm production. Yeast biofilms are characterized by a significantly decreased susceptibility to usual antifungal drugs, encouraging the search for other inhibition methods, such as peroxidase systems. Previous studies have shown that a 30-minute exposure to OI^-1^ was sufficient to inhibit planktonic growth *in vitro*. The present investigation demonstrated that *Candida* biofilm development on material surfaces could be reduced or even suppressed by LP-generated OI^-1^ and OSCN^-\_, not only when peroxidase and substrates system were dissolved in the liquid phase into material and immersed, but also when material-precoated peroxidase was activated by simple addition of the substrates to the liquid surrounding the precoated material. Those data demonstrate...
the efficiency of peroxidase systems against a Candida strain, and concomitantly show an application of the titanium powder model described supra. This investigation did not demonstrate that enzymes remain active after a three-week incubation, but clearly showed that their inhibition effect persisted during this time. Previous investigations demonstrated, firstly, that coating onto titanium did not modify LP activity, and secondly, that LP incorporated into oral gel maintained its activity for at least one year and thirdly, that the substrate exhaust (namely H2O2, I−) is the actual limiting factor.20 Transfer of this finding from bench to clinic is questionable. Indeed, immunogenicity of coated LP should limit the applications of this system to ex vivo conditions. A previous investigation indicated an antibacterial effect with Gram-positive and Gram-negative bacteria, suggesting a nonspecific inhibitory effect of hypoiodite on microbial metabolism and growth.22

Conclusion

In conclusion, C. albicans biofilms on titanium powder could offer a simple and reliable model for further investigating new antimicrobial strategies. Moreover, the model could be used in other studies of contaminating microorganisms on implanted materials.

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Disclosure

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