Killing of *Candida auris* by UV-C: Importance of exposure time and distance

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**INTRODUCTION**

*Candida auris* is a globally emerging yeast, only recognised in the last 10 years causing severe infections. This yeast has been reported to cause hospital outbreaks in various healthcare facilities across the globe. Skin colonisation and inanimate surface contamination in close vicinity of infected and colonised patients is likely an important factor in patient to patient transmission. Rapid identification of *C. auris*, skin decolonisation of patients, and decontamination of hospital surfaces are essential steps in controlling *C. auris* outbreaks.

Although there are no established guidelines for decontaminating surfaces contaminated by *C. auris*, healthcare organisations have issued different recommendations. The Center for Disease Control and Prevention (CDC) suggests the use of Environmental Protection Agency (EPA)-registered hospital grade disinfectant effective against *Clostridium difficile* spores, while Public Health England...
(PHE) suggests the use of hypochloride, possibly in conjunction with other products. These recommendations have been based on an increasing number of studies, which demonstrated that such agents are indeed effective in vitro as well as during clinical care. In addition, in vitro studies demonstrated the effectiveness of hydrogen peroxide vapour.

Another approach in the decontamination of hospital surfaces is the use of mobile ultraviolet-C (UV-C) devices, which are broadly applied in some countries. PHE also recommends UV-C as a potential adjunct in controlling the spread of C. auris, although there is no clinical evidence for its efficacy. A single in vitro study by Cadnum et al. demonstrated that 10 or 30 minutes UV-C led to a 40 to 1 x 10^6-fold reduction, respectively, in colony-forming units, demonstrating the importance of time as an essential factor in UV-C efficacy. The samples were placed 1.5 m from the UV-C device. In most hospital rooms, the radius is larger than 1.5 m, requiring the UV-C device to be relocated after a first exposure cycle in order to cover the whole room. To determine the effect of distance on C. auris decontamination with UV-C, we implemented a sample distance of 2 and 4 m, which better reflects the clinical situation. In addition, we tested the effect of UV-C exposure time and the UV-C sensitivity of C. auris strains from different worldwide clades and origins.

2 | MATERIALS AND METHODS

2.1 | Candida auris and Candida albicans strains

In the present study, clinical C. auris strains originating from Venezuela (n = 3), Spain (n = 3), India (n = 3) and Japan/Korea (KCTC 17809, KCTC 17810 and JCM 15448) were used, along with C. albicans strains ATCC 90028, ATCC 10231 and ATCC 24433.

2.2 | UV-C decontamination device

The UV-360 Room Sanitiser (UltraViolet Devices, Inc., Valencia, CA) device was used. It is a four-wheeled unit containing four vertically placed maximum output UV Germicidal lamps that are 158 cm tall and emit light of predominantly 254 nm in 360° (Figure 1). The system also contains four motion sensors, which abort the cycle if someone enters the room during use.

2.3 | Killing Candida yeasts with UV-C

Strains were suspended in 500 μL phosphate-buffered saline (PBS, Diasorin Molecular LCC, CA), and yeast quantity was determined using the Genesys 20 Spectrophotometer (ThermoSpectronic, UK). Subsequently, 10 μL of suspension with 1 x 10^5 or 1 x 10^6 colony-forming units (CFU) was spread to cover 50 mm² wells of multistest glass slides (MP Biomedicals, LCC, Illkirch, France). These were placed in petri dishes and allowed to air dry for 30-40 minutes in incubator at 35°C. Subsequently, the multistest slides were transferred to a standard laboratory room (3 x 5 m) and placed on a table with a standard height of 90 cm at a distance of 2 or 4 m from the UV-360 Room Sanitiser. Thus, the samples were perpendicular (horizontal) to the UV-C source. The glass slides with Candida cells were directly exposed to a UV-C cycle of 5, 10, 20 or 30 minutes incubated for the same time without exposure to UV-C. After exposure, yeast were suspended in 100 μL PBS for 1 minute and the suspension was further diluted in PBS. Finally, 100 μL of the diluted suspension was transferred to Sabouraud dextrose agar plates (Tritium Microbiologie BV, Eindhoven, The Netherlands) and incubated at 35°C for 24-48 hours for colony count determination. Log reductions were calculated in comparison with unexposed samples. Experiments were performed in triplicate.

2.4 | Statistical analysis

A 1- or 2-way ANOVA with Bonferroni's multiple comparisons test was used to compare CFU of non-exposed isolates and mean log
3 | RESULTS

To determine the effect of UV-C on the decontamination of surfaces with *C. auris*, we first investigated whether UV-C exposure time and *C. auris* seeding density affected UV-C efficacy using a single *C. auris* strain. After seeding *C. auris* on glass well plates at two different densities, these were placed at 2 m exposure distance from the UV-C device and exposed for 5, 10, 20 and 30 minutes. A strong time-dependent reduction of *C. auris* CFU was observed, which was significantly different (*P* < 0.05) for all time points for each density (Figure 2A). At 10, 20 and 30 minutes, CFU reduction was significantly higher in *C. auris* inoculated at a density of 1 x 10^5 as compared to those seeded at 1 x 10^6 CFU (Figure 2A). Subsequently, the effect of distance was studied. Increasing the distance to the UV-C source from 2 to 4 m strongly reduced the efficacy of UV-C to kill *C. auris* (Figure 2B).

Additionally, we investigated the sensitivity of *C. auris* strains originating from different countries to UV-C and compared this to the sensitivity of *C. albicans* reference strains. For this experiment, we used the low density of 1 x 10^5, as this count better reflects the clinical situation. With a UV-C exposure time of 10 minutes at 2-metre distance, we found a decreased efficacy of UV-C to kill *C. auris* strains originating from Venezuela, Spain and India as compared to the *C. auris* strains from Japan/Korea. The latter *C. auris* strains were similar to *C. albicans* (Figure 3). Similar results were obtained for the UV-C exposure time of 30 minutes at a distance of 4 m, while for the other exposure conditions there were no significant differences among the strains (Figure 3).

![Figure 2](image-url)  
**FIGURE 2** UV-C efficacy in killing *Candida auris* strains from different countries. *Candida albicans* and *C. auris* isolates from different countries were seeded on multitest slide glasses at a density of 1 x 10^5 CFU per well. Glasses were exposed for 10 or 30 min to UV-C at a distance of 2 or 4 m. Significant differences (*P* < 0.05) as compared to *C. auris* reference strains are indicated with an asterisk. Ref, reference; Ven, Venezuela; Spa, Spain; Ind, India

4 | DISCUSSION

The present study demonstrates that *C. auris* can be effectively killed by UV-C, although the density of *C. auris*, the time of UV-C exposure and the distance to the UV source strongly influenced the effectiveness of UV-C treatment. With a 10-fold higher concentration of *C. auris*, the effectiveness of UV-C treatment diminished around 10-fold when exposed for 20 or 30 minutes UV-C radiation. Time and distance were even more important parameters, as with a 2-fold increase in time or decrease in distance, respectively, and ~10- and ~50-fold reductions in CFU were found. A maximal effect of UV-C was reached with 30 minutes exposures with UV-C and when surfaces were at 2 m of the UV-C device. Cadnum et al also demonstrated that 30 minutes UV-C exposure is required to reach a maximal effect of *C. auris* reduction while placing the specimen at 1.5 m from the UV-C device. A recent report from Ponnachan et al found that 15 minutes of exposure killed all *C. auris*. It has
to be noted though that in the latter study the standard sample distance to UV-C light was only 1 m, while the samples were directly facing the UV-C lamp. Facing samples in parallel directly to the UV-C source led to a >10-fold higher effectiveness in killing methicillin-resistant *Staphylococcus aureus* (MRSA) and a >5-fold higher effectiveness to kill *Clostridium difficile*, as compared to placing them horizontally to the UV-C lamp.18 As the minimum distance of sample to UV-C source in our study was 2 m, while the samples were placed horizontally to the UV-C source, the previous studies14,17 are largely in agreement with our findings regarding the effect of time and distance in killing *C auris* with UV-C.

The report of Cadnum et al14 found that the sensitivity of *C auris* for UV-C in a laboratory setting appeared to be similar to *C difficile*. A 10 minutes UV-C cycle at 1.5 m was not sufficient to kill all *C auris* nor all *C difficile*, which was in contrast to MRSA that was effectively killed under these conditions. The effect of UV-C on *C auris* killing has—to our knowledge—not yet been investigated in the clinical setting, while there have been different hospital and patient studies on the effect of UV-C on *C difficile* surface contamination and hospital-acquired infection rates. In agreement with the relative resistance for UV-C in the laboratory setting, the contamination of high-touch sites14,17 are largely in agreement with our findings regarding the effect of time and distance in killing *C auris* with UV-C. Thus, despite the fact that the laboratory settings suggest that a total UV-C exposure time of 10 minutes is not sufficient to kill all *C difficile*, in the clinical setting a total UV-C exposure time of 10-12 minutes leads to reduced *C difficile* infection rates. Moreover, the effect of UV-C on hospital-acquired infection rates for MRSA, which is much more sensitive to UV-C than *C difficile*, seemed equally or even less efficient in several studies.20-22 Thus, despite differences in UV-C sensitivity to kill pathogens in the laboratory or clinical setting, the actual hospital-acquired infection rate might differ, demonstrating the importance to identify the effect of UV-C on hospital-acquired *C auris* infection rates.

Altogether, our findings demonstrate that, with a longer cycle time, and at an optimised distance, *C auris* can successfully be decontaminated by UV-C. Future studies in hospitals struggling with endemic *C auris* presence should further investigate the effect of UV-C under routine conditions in clinical practice.

**CONFLICT OF INTEREST**

The authors declare that they have no competing interests.

**AUTHOR CONTRIBUTIONS**

Acquisition, analysis and interpretation of the data: TG, AC, JM, AV.

Statistical Analysis: TG. Drafted the manuscript: TG. Reviewed and modified the manuscript: TG, AC, JM, AV.

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