Investigation of Microbial Contamination in the Clinic and Laboratory of the Prosthodontics Department of Dental School

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ABSTRACT

Objective: To determine the level of clinical contamination in the clinic and laboratory of the prosthodontics department of Kerman Dental School. Material and Methods: Clinical surfaces of the dental units, the laboratory, and the professors’ lounge of the prosthodontics department were randomly sampled. The sampled surfaces included the dental units’ console, light switch, light handle, headrest, and air-water spray syringe in the clinic, plastering tables, buttons of the vibrator, polishing, and trimmer machines, acryl tables, handles of pressure pot and press machine, handpiece holders, work desks, and drawer handles in the laboratory, and desks, computer mouse and keyboard, telephone sets, and doorknob in the professor’s lounge. The samples were examined for the type and growth of microorganisms. The data were entered into SPSS, where they were analyzed using the chi-square test at the 0.05 significance level.

Results: Of all the samples taken, 89.9% showed microbial contamination. The most common type of contamination was fungus (34.8%) and the least common types were Enterococcus faecalis and Staphylococcus epidermidis (1.1%). The second and third most common types of bacteria in the samples were Staphylococcus aureus (18%) and Pseudomonas aeruginosa (12.4%), respectively. There was no significant difference between the frequencies of microbial contamination in the clinic, the laboratory, and the professors’ lounge.

Conclusion: Given the strong chance of cross-infection in the examined department and laboratory, it is necessary to enforce protocols for proper disinfection of surfaces before, between and after treatments.

Keywords: Cross Infection; Microbiology; Bacteria.
Introduction

Infection prevention and control are essential for creating a safe environment for all patients and health care professionals, particularly in dentistry. Dental students, dentists, and dental assistants are at risk of diseases such as HBV, HCV, HSV-1, HIV, influenza, and rubella through exposure to patients and the water of dental units and handpieces that can cause cross-infection in dental clinics [1-3]. In dental settings, infections can be transmitted directly through contact with infected saliva and blood and their droplets and indirectly through contact with contaminated instruments, equipment, or surfaces [4-6]. Important bacteria that can cause infection in dental settings include *Streptococcus pneumoniae, Mycobacterium tuberculosis, Klebsiella pneumoniae, Escherichia coli, Legionella pneumophila*, and *Pseudomonas aeruginosa* [7,8].

Cross-infections in dental settings can be caused by many pathogenic organisms present in the oral cavity and respiratory tract [6,9]. Considering the ease with which diseases can be transmitted in dental clinics, the prevention of cross-infection in these clinics is an important aspect of the dental profession [10].

Light switches and handles, control panel and console, tray, headrest, chair position control buttons, handpiece hose, and air-water spray syringe of dental units, and dental stools are examples of clinical contact surfaces that, although not directly in contact with the patient, can be easily contaminated by splattered blood or saliva particles or through exposure to contaminated dental equipment or gloves of dentist or assistants [11].

Nonclinical surfaces are surfaces that are typically not exposed to contaminated gloves or equipment [12-14]. Research has shown that dental laboratories can also be the source of contamination [15]. Infection transmission can also occur during the transfer of dental molds and other equipment from the clinic to the laboratory.

Given the importance of infection control in dental settings and in the continuing interest in dental researches [16-19], this study aimed to investigate the frequency of microbial contamination in the clinic and laboratory of the Prosthodontics Department of Kerman Dental School.

Material and Methods
Study Design

This experimental study was performed on the clinic and laboratory of the prosthodontics department of Kerman Dental School, Kerman, Iran.

Data Collection

The samples were collected from the surface of 11 operational dental units, 5 laboratory tables, 7 spots in the laboratory and 5 spots in the professors' lounge of the prosthodontics faculty. The sampled surfaces included the dental units’ console, light switch, light handle, headrest, and air-water spray syringe in the clinic, plastering tables, buttons of vibrator, polishing, and trimmer machines, acryl tables, handles of pressure pot and press machine, handpiece holders, work desks, and drawer handles in the laboratory, and desks, computer mouse and keyboard, telephone sets, and doorknob in the professor's lounge.

All samples were collected by a trained senior student wearing sterile gloves, who changed gloves after each sampling. Samples were obtained by sterile swabs during the launch hour. Before sampling, the swabs and the tubes containing the culture medium were sterilized by autoclaving (135°C, 20 bar, 45 min). For sampling, a swab was moistened with the nutrient broth, firmly placed on the surface for 1 minute, and then placed in a tube containing nutrient broth. The tubes were then transferred to the microbiology laboratory, where it was incubated at 37°C for 24-hours.
A total of 89 samples were collected in this study. To isolate gram-positive and gram-negative bacteria, after the 24 hours of incubation, the liquid (broth) samples were diluted and transferred to a solid medium (agar) to grow gram-negative and a blood agar medium for the growth of gram-positive bacteria.

After 24 hours, the growth of bacterial colonies was examined. To identify the type of bacteria in each colony, gram staining was performed to determine whether the colony is gram-positive or gram-negative and whether it is coccci or bacilli. After this examination, differentiation tests were performed with catalase, mannitol, 6.5% sodium chloride, NaCl, and coagulase (to detect gram-positive bacteria) and oxidase, citrate, lysine decarboxylase, TSI, SIM, and OFtest (to detect gram-negative bacteria). All media were incubated for 24 hours before the test to ensure sterility.

Data Analysis

The results were recorded in a checklist and then entered into SPSS, which they were analyzed using ANOVA and T-test at the 0.05 significance level.

Ethical Aspects

This study's proposal was approved by the ethics committee of Kerman University of Medical Sciences with the ethics code IR.KMU.REC.1397.120.

Results

Overall, 89.9% of the 89 collected samples showed microbial growth. The most common type of microbial contamination in the samples was fungus with 31 cases (34.8%), and the least common were Enterococcus faecalis and Staphylococcus epidermidis with one case each (1.1%) (Figure 1).

![Figure 1. Frequency distribution of microbial contamination by type.](image)

There was no statistically significant difference between the sampled spots in microbial contamination (p=0.143) (Table 1). No contamination was observed in the dental units’ light handles, air-water syringes, and chair control panels. However, microbial contaminations were observed in all headrests.

The samples taken from the lounge showed no sign of Streptococcus viridans, Bacillus subtilis, Klebsiella pneumoniae, Enterococcus faecalis, Staphylococcus epidermidis, Escherichia coli, and fungi. The samples taken from the
laboratory did have *Bacillus subtilis*, *Klebsiella pneumoniae*, *Enterococcus faecalis*, and *Staphylococcus epidermidis* (Figure 2).

Table 1. Relationship between the presence of microorganisms in the dental units, the laboratory, and the lounge.

<table>
<thead>
<tr>
<th>Location</th>
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<th>%</th>
<th>Yes</th>
<th>%</th>
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<td>Different Parts of Dental Unit</td>
<td>48</td>
<td>85.7</td>
<td>8</td>
<td>14.3</td>
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<td>Different Parts of Prosthodontics Laboratory</td>
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<td>96.2</td>
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<td>Different Parts of Lounge</td>
<td>7</td>
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<td>0</td>
<td>0.0</td>
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<tr>
<td>Total</td>
<td>80</td>
<td>89.9</td>
<td>9</td>
<td>10.1</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2. Frequency distribution of the types of microorganisms found in the dental units, the laboratory, and the lounge.

A significant difference between the types of microorganisms was found in different rooms. *Fungi* were significantly more common in the dental units (p=0.0001). In the laboratory, microbial contaminations were detected in handpieces, desks, drawer handles, buttons of the vibrator, polishing and trimmer machines, plastering tables, acryl tables, and press machine and its handle. In the professors’ lounge, contaminations were observed in the keyboard, mouse, telephone set, desk, and doorknob.

Discussion

The highest concentration of microorganisms in dental clinics is in the mouth of patients. However, when infected with a patient's saliva, gingival fluid, or blood, dental practitioners themselves can become a major source of surface contamination in the clinic [20].

In the present study, microbial contamination was observed in 85.7% of the samples taken from different parts of dental units. In a study by Valian et al. [11] on microbial contamination in the units of periodontics and restorative dentistry departments of the Beheshti Dental School, 63.3% of the units were contaminated at the end of the work. This discrepancy may be due to differences between this study and Valian et al. [11] in terms of the method of work and the type of clinic. However, it should be noted that some degree of microbial contamination after dental work is unavoidable.
In this study, all but two parts of the dental units (in the chair control panel) were found to be contaminated. The lower contamination of the chair control panel can be attributed to the fact that students rarely use this control panel after the initial adjustment of the chair position. The results of a previous study on the contamination of dental office surfaces, including light handle, floor, and sink, showed that the bacterial count was higher at the end of the workday than at the beginning [21].

It appears that the prosthodontics students in the studied dental school are not as committed to infection control as necessary. Working with high-speed spinning dental handpieces such as turbines during treatment produces particulates containing oral cavity microbes (aerosols), which have a significant contamination potential and a large dispersal radius; features that make them a growing concern in dentistry [22,23]. It has been shown that there is likely to be four times more contamination in the room air during dental treatment than during other times [7].

In the present study, microbial contamination was observed in all sampled spots in the professors' lounge. This appears to be caused by the students walking into the lounge to show their molds or finished works or to bring patient records and radiographs from the ward. Therefore, to reduce cross-infection, necessary measures should be taken to prevent students and assistants from bringing these items to the professors' lounge.

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In the present study, 96.2% of the sample taken from the laboratory showed microbial contamination signs. A previous research has reported that dental polishing is a source of infection transmission [24]. In a study on cross-infection in prosthodontics laboratories, it was found that they were substandard in this respect and there was no protocol or training program to limit cross-infection in these laboratories [25].

In the present study, the most common types of microorganisms in the sampled areas were fungi, followed by Staphylococcus aureus. A previous study found that 38% of contaminations were of fungal type [15]. Valian et al. study on microbial contamination in the periodontics and restorative dentistry departments reported that the most common microorganism was Staphylococcus aureus [11]. In a study on the possibility of transmission of methicillin-resistant Staphylococcus aureus through dental surfaces, Kurita et al. [26] reported infections with this bacterium in 8 out of the 140 examined dental patients.

A study by Qavam and Aligholi [27] on the contamination of commonly used dental equipment reported detecting Staphylococci, Enterococcus faecalis, and bacillus species in these pieces of equipment. In this study, the least common type of microorganism was Enterococcus, which is consistent with the study by Esfahani et al. [28]. In contrast, the most common type of microorganism in the present study was fungus, which could be because it was performed in the prosthodontics clinic and laboratory.

This study's sample also contained other microorganisms, including Streptococcus viridans and Bacillus subtilis, Klebsiella pneumoniae, Enterococcus faecalis, Staphylococcus epidermidis, and Escherichia coli. Indeed, Bacillus subtilis is the microorganism used to measure autoclaving efficiency [29]. Streptococcal virulence is the cause of subacute bacterial endocarditis. Also, 1.1% of the samples taken in this study had Pseudomonas spp., most of which have multiple resistant genes [30]. Enterobacteria have also been reported as the most important gram-negative bacteria causing resistant nosocomial infections [31].

Most of these bacterial species are normal flora of the human body and generally non-pathogenic but can become pathogenic under certain conditions, including a weakened immune system. Given the association of transmission of contagious diseases with the infection control measures taken during both invasive and non-invasive dental treatments, this issue is of great importance for mitigating the risk of disease transmission during treatment for dentists and patients [32]. Dental professionals should be aware of the risks involved in
this profession and be trained and encouraged to commit to minimizing the spread of microorganisms in their work environment.

**Conclusion**

This study found 89.9% microbial contamination in the samples taken from the examined prosthodontics clinic and laboratory. The most common type of microorganism observed in these dental settings was fungi, followed by Staphylococcus aureus. Further actions are needed to ensure proper disinfection of clinical surfaces in the examined prosthodontics clinic and laboratory.

**Authors’ Contributions**

<table>
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<td>Methodology, Investigation and Writing - Review and Editing.</td>
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All authors declare that they contributed to critical review of intellectual content and approval of the final version to be published.

**Financial Support**

None.

**Conflict of Interest**

The authors declare no conflicts of interest.

**Data Availability**

The data used to support the findings of this study can be made available upon request to the corresponding author.

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**References**


