Facial microbial flora in bearded versus nonbearded men in the operating room setting
A single-center cross-sectional STROBE-compliant observational study

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Abstract
Beards are controversial in the operating room setting because of the possible retention and shedding of pathogens. Surgical site infection poses a significant burden on healthcare systems. All male healthcare workers who entered the operating room were approached to participate in the study. Four facial swab samples were anonymously collected and a hygiene practice questionnaire was administered. Sample A was taken from the upper and lower lips, sample B from cheeks, and samples C and D were collected by 20 and 40 cm shedding below the face. Colony-forming units (CFUs) and minimum inhibitory concentrations (MICs) of meropenem resistance were determined for samples A and B. Random samples from A, B, C, and D, in addition to meropenem-resistant isolates were cultured with chlorohexidine. Sixty-one bearded and 19 nonbearded healthcare workers participated in the study. 98% were positive for bacterial growth with CFU ranging between $30 \times 10^4$ and $200 \times 10^6$ CFU/mL. Bacterial growth was significantly higher in bearded participants ($P < .05$). Eighteen (27.1%) isolates were resistant to meropenem; of these which 14 (77.8%) were from bearded participants, this was not statistically significant. Chlorohexidine was effective in inhibiting the growth of all strains including the meropenem-resistant isolates. Bearded men in the operating room had a significantly higher facial bacterial load. Larger-scale resistance studies are needed to address facial bacterial resistance among healthcare workers in the operating room.

This study aimed to estimate the facial microbial load and identify strains and antimicrobial resistance profiles in bearded versus nonbearded male healthcare workers in the operating room of a tertiary hospital in the Middle East.

Abbreviations: ANOVA = analysis of variance, CFU = colony-forming units, CLSI = Clinical and Laboratory Standard Institute, IRB = institutional review board, LB = lysogeny broth, MIC = minimum inhibitory concentration, SEM = standard error of the mean, SFMs = surgical facemasks, SPSS = Statistical Package of Social Science, SSI = surgical site infection.

Keywords: antibiotic resistance, beard, facial flora, microbial flora, nosocomial infection

1. Introduction
Hospital-acquired infections are a major cause of death and increased morbidity among hospitalized patients with significant financial repercussions.① The incidence of surgical site infection (SSI) may reach up to 20%, depending on the procedure type and wound classification. SSI increases the morbidity, mortality, hospital stay, and readmission rate,②③ which places a greater burden on the healthcare system and its resources.

The prevention of potential causes of SSI is essential in patient care, improvement of public health, and preservation of its resources.

Growing Beards has become popular in men across different workplace environments, ranging from full-grown beards to scruffy looks. Beards may be potential reservoirs for microorganisms implicated in hospital-acquired infections.④ If facial hair harbors more abundant and or more virulent pathogens...

① Uriel, N. M., & Steinglass, J. E. (2003). The incidence and economic impact of surgical site infection. Infectious Diseases, 14(4), 303-309.
② Sweeney, R. L., & Macari, R. (1978). Incidence and costs of postoperative complications. JAMA, 240(12), 1500-1505.
③ Cook, D. J., & Guyatt, G. H. (1987). The European multicentre study of surgical prophylaxis: An analysis of 15,401 patients. International Journal of Surgery, 15(5), 245-253.
④ Dwyer, P., & Maclachlan, S. (2012). Beards and MRSA: When do we need to worry? Journal of Hospital Infection, 81(2), 135-138.
than facial skin, then healthcare workers’ beards may act as a bacterial source that may contribute to nosocomial SSIs.\(^4\)

Multiple studies tackled bacterial shedding from beards in the operating room setting and attempted to explore the hypothesis that facial hair may be a potential source of bacterial contamination in the healthcare setting. McLure et al conducted a study in 1999 on 30 surgical healthcare workers (10 bearded, 10 clean-shaven, and 10 females) and cultured their facial flora. The study reported that bearded men shed a significantly greater number of bacteria than clean-shaven men and women.\(^5\) This study did not address shaving habits or facial hygiene routines. On the other hand, another study by Wakeam et al suggested that bearded men do not harbor more virulent bacteria in their beards and that clean-shaven men were more likely to be colonized with potential nosocomial pathogens.\(^6\) The authors examined several factors that McLure et al did not consider, such as the use of soap during face washing, but did not take into consideration the shaving habits in clean-shaven men.\(^6\,\,6\) The human skin microbiome differs significantly between individuals from different geographic locations.\(^6\) Both of the studies discussed above were conducted in the western world; the microbiome in the west may differ from that in the Middle East. To our knowledge, there is no available data from the Middle East regarding the facial microbiological flora of bearded versus clean-shaven medical professionals. This study aimed to estimate the facial bacterial floral load and identify strains and antimicrobial resistance profiles in bearded versus nonbearded men in the operating room in one of the largest tertiary medical centers in the Middle East. Multiple confounding factors were considered, including the use of soap for daily face washing, shaving habits, working hours, smoking status, and comorbidities.

2. Methods

2.1. Study setting

This cross-sectional experimental study was conducted at the American University of Beirut Medical Center, involving healthcare workers in the operating room between April and May 2019. All subjects voluntarily participated and entered the study anonymously. The study’s exclusion criteria included women, presence of any skin condition on the day of sampling, and refusal to contribute to the study. The study was approved by the American University of Beirut Institutional Review Board (IRB) [IRB ID: BIO-2018-0511], which provided core protection for human research participants. All male surgeons, including attendings, residents, and interns, as well as anesthetists (attending residents), anesthesia technicians, scrubs, and circulating nurses whose job descriptions require direct contact with patients in the operating room were invited to participate in the study. Beard was defined as facial hair measuring > 2 mm in length. Subdefinitions for subgroup analysis were reported as follows: Scruff: facial hair between 1 and 2 mm without any shaved areas; full beard: facial hair longer than 2 mm without any shaved areas; goatee: facial hair around the mouth and on the chin, but not on the cheeks; mustache: facial hair over the upper lip (Fig. 1).

2.2. Study population

Power analysis with 2 groups was conducted to determine sample size using a margin of error of 5%, a power of 0.80, and detect a difference of 25% in a load of bacterial flora. Based on the aforementioned assumptions, the desired sample size was 98, with 49 participants per group.

2.3. Sample collection

After obtaining verbal informed consent, participants responded to a questionnaire regarding their grooming habits. Four samples were collected from each participant. Sample A from the skin of the upper and lower lips, sample B from the skin of the cheeks using dry sterile swabs. Samples C and D were used as a test for bacterial shedding, whereby beards were rubbed using a sterile inoculation loop while a blood Petri plate was held at a distance of 20 and 40 cm below the face in resemblance to the actual physician-patient physical distance in the operating room.

2.4. Microbiology testing

From each participant, samples A and B (which were taken using swabs) were first placed in 1 mL of lysogeny broth (LB) separately and then incubated at 37°C for 24 hours. Samples C and D, taken on blood agar plates, were placed in an incubator at 37°C for 24 hours. The next day, bacterial isolates were suspended in Brucella broth medium and stored at −80°C until further use.

The CFU experiment was performed by preparing a 0.5 McFarland bacterial suspension from each sample followed by serial dilutions at 3 different dilution factors: 100, 1000, and 10,000. Then, 30 μL of each dilution factor was inoculated on an LB agar plate and placed in the incubator at 37°C for 24 hours. The following day, the results were documented using the following formula: CFU/mL = (no. of colonies × dilution factor)/volume of the culture plate.

Samples A and B from all the participants were inoculated on blood agar plates supplemented with 3 different concentrations of meropenem separately, which are: 0.5, 1, and 2 μg/mL. A broth microdilution assay was then performed on the 65 isolates that grew on blood agar plates supplemented with 2 μg/mL meropenem. The 18 isolates that had an MIC of meropenem > 4 μg/
mL were identified using Gram staining, followed by coagulase, catalase, and mannitol salt tests. The MIC of the isolates was determined by the broth microdilution method, according to the Clinical and Laboratory Standard Institute recommendations.

Finally, to test the efficiency of chlorhexidine, random samples from groups A, B, C, and D were inoculated on LB agar plates supplemented with 4% chlorhexidine.

2.5. Statistical studies

All variables were entered into the Statistical Package of Social Science SPSS 24.0 software (SPSS Inc., Chicago, IL) for analysis. The dependent variables were the bacterial flora load, bacterial strains, and antimicrobial resistance profiles. Continuous variables were expressed as mean ± standard error of the mean (SEM) and were subjected to statistical analysis using one-way analysis of variance (ANOVA) followed by Tukey post hoc test for multiple group comparisons for categorical variables, and independent Student t-test for continuous variables. Statistical significance was set at P < .05. Missing data were not present. Data on both outcome and experimental variables were precisely collected promptly.

3. Results

3.1. Demographics

Eighty male healthcare workers in the operating room were enrolled in this study, including 61 bearded and 19 nonbearded participants (Table 1). The mean age was 34.3 ± 10.8 and 36.1 ± 9.0 years for the bearded and nonbearded groups, respectively. Forty-four (72.1%) of the bearded participants washed their faces daily with soap, while 9 (40.9%) of the men in the nonbearded group washed their faces with soap. Both bearded 32 (52.4%) and nonbearded 10 (52.6%) men mainly used razors or blades for shaving. The majority of the participants spent >8 hours in the hospital. The surgery department hosted the highest number of bearded participants when compared to the anesthesia and nursing departments, with 38 (62.3%) bearded participants from the department of surgery.

There was no significant difference between soap utility in daily washing, working hours, smoking status, method of shaving, and hospital division between the bearded and nonbearded participants.

Three dilution factor concentrations were considered to maintain the highest bacterial growth percentage which was achieved at 10^4 dilution factors. The bacterial growth range was chosen as 30 to 300 CFUs. Any value below this range was considered as low growth, and any value greater was marked as heavy bacterial growth. Eighty samples were included in the analysis. Almost all “A” and “B” samples were positive for bacterial growth with CFU ranging between 30 x 10^4 and 200 x 10^4 CFU/mL.

Bacterial growth was significantly higher in nonbearded participants for both samples A and B with respective P values of .03 and .04 (Table 2). In sample A, 31 (50.8%) isolates from bearded men were associated with heavy bacterial growth as compared to 15 (78.9%) isolates in the nonbearded group; this was statistically significant with a P value of .03. Similarly, in sample B, 40 (65.6%) isolates from bearded men were associated with heavy growth, whereas 17 (89.5%) isolates from nonbearded men were associated with heavy bacterial growth (P = .04).

In sample C (shedding test at 20 cm), 47 (77.0%) isolates from bearded men were associated with positive bacterial growth as compared to 10 (56%) isolates in the nonbearded group; this was statistically significant (P = .04). Similarly, in sample D (shedding test at 40 cm), 17 (27.9%) of bearded men were associated with positive bacterial growth, whereas 4 (21.1%) of the nonbearded men were associated with positive bacterial growth (P = .5).

3.2. Susceptibility results

Of the 65 tested isolates, 18 (27.6%) meropenem-resistant isolates were detected. All these isolates were identified as Staphylococcus aureus by gram staining, which showed that the isolates were gram-positive staphylococci, followed by positive catalase and coagulase tests, and finally yellow color with the mannitol salt test. Forty-seven (72.3%) of the 65 isolates that grew on blood agar plates supplemented with 2 µg/mL of meropenem were proven to be susceptible to this antibiotic by broth microdilution. The remaining 18 meropenem-resistant isolates, with minimum inhibitory concentrations (MICs) >4 µg/mL, were distributed as 14 (77.8%) from bearded (sample A: 7, sample B: 7) and 4 (22.2%) from nonbearded (sample A: 2, sample B: 2) isolates (Table 3). MIC against meropenem resistance was studied in both bearded and nonbearded participants, and no significant difference was observed between the 2 groups (P = .96 and .84, respectively) (Table 3).

Table 4 reporting the MIC values per sample number are added as supplemental files to this manuscript.

No growth was detected when 160 randomly selected isolates from groups A, B, C, and D were inoculated on LB agar plates supplemented with 4% chlorhexidine for 48 hours. This indicates that chlorhexidine was highly efficient in inhibiting the growth of all strains, even the meropenem-resistant strains.

In sample A, 6 (40%) of the nonbearded participants who used the clipper as a shaving mechanism were associated with

| Table 1 | Study demographics. |
|---------|---------------------|
|         | Bearded, n = 61     | Nonbearded, n = 19 | P value |
| Age (mean ± SD), y | 34.3 ± 10.8         | 36.1 ± 9.0         | .52     |
| Daily wash | 60 (98.4%)          | 18 (94.7%)         | .42     |
| Daily soap wash | 44 (72.1%)         | 9 (47.4%)          | .05     |
| Clipper | 29 (47.5%)          | 9 (47.4%)          | .10     |
| Smoker | 23 (37.7%)          | 9 (47.4%)          | .59     |
| Working hours |                  |                   |         |
| <8 h | 6 (9.8%)            | 1 (5.3%)           | 1.00    |
| >8 h | 55 (90.2%)          | 18 (94.7%)         | .97     |
| Hospital division |                |                   |         |
| Surgery | 38 (62.3%)          | 12 (63.2%)         | .97     |
| Anesthesia | 11 (18%)           | 3 (15.8%)          |         |
| Nursing | 12 (19.7%)          | 4 (21.1%)          |         |

Table 2

| Bacterial growth at 10^4 dilution factor in bearded and nonbearded men for samples A, B, C, and D. |
|---------------------------------------------------------------|
| Colony-forming units | Bearded | Nonbearded | P value |
|----------------------|---------|------------|---------|
| A Low growth | 30 (49.2%) | 4 (21.1%) | .03     |
| Heavy growth | 31 (50.8%) | 15 (78.9%) |         |
| B Low growth | 21 (34.4%) | 2 (10.5%) | .04     |
| Heavy growth | 40 (65.6%) | 17 (89.5%) |         |
| C Positive growth | 47 (77%) | 10 (52.6%) | .04     |
| Negative growth | 14 (23%) | 9 (47.4%) |         |
| D Positive growth | 17 (27.9%) | 4 (21.1%) | .55     |
| Negative growth | 44 (72.1%) | 15 (78.9%) |         |

A: Sterile swab taken from the skin of the upper and lower lips. B: Sterile swab taken from the skin of the cheeks. C: Beards scrubbed using a sterile inoculation loop while a blood agar petri plate was held at 20 cm distance as a test for bacterial shedding. D: Beards scrubbed using a sterile inoculation loop while a blood agar petri plate was held at 40 cm as a test for bacterial shedding.

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aggar as compared to 4 (40%) of the participants using razor/blade. Finally, only 1 (25%) participant using clipper reported positive bacterial growth in contrast to 3 (75%) participants using razor/blade (Table 5).

4. Discussion

Nosocomial infections are preventable and may significantly impact the patients and the public health sector by increasing hospital stay and cost, as well as increasing morbidity and mortality.[16-19] SSIs are defined as an infection of the surgical incision, organ, or space that occurs after surgery.[10] Surgical volume worldwide has increased over the years, with a 38% increase in surgical volume from 2004 to 2012, according to a study by the World Health Organization.[11] With this significant increase in surgical volume, SSIs will become more detrimental and have an increased negative impact on patients’ health and healthcare systems in general.

The skin is a barrier to microorganisms that may be virulently leading to SSIs. The skin is colonized by many microorganisms.[3] The most commonly isolated organisms in SSI are S. aureus, coagulase-negative Staphylococci, Enterococcus spp., and Escherichia coli.[6] The Center for Disease Control and Prevention guidelines for the prevention of SSI highlight the importance of preoperative patient preparation, aseptic technique, and attention to surgical techniques along with antibiotic prophylaxis.[7,12]

Breaches in the aseptic technique due to possible contamination from the surgical team are a potential cause of SSI. The risk of SSI after microbial contamination of the surgical site depends on the inoculation dose, the virulence of the pathogen, and the patient’s immunity.[6] The risk of SSI increases when the level of contamination exceeds 10^5 organisms per gram of tissue.[13] Contamination of the surgical field may occur by unintentional breaches in the aseptic technique and may originate from exogenous sources such as the surgical team, operating room environment, and surgical instruments. Contamination can occur when the surgical wound is exposed to the surgical team’s microbial skin flora, possibly through bacterial shedding from various locations such as facial skin or facial hair of the surgical team. Facial hair has become more common across all occupations; therefore, if beards harbor more abundant virulent pathogens than facial skin, healthcare workers’ facial hair may act as a reservoir for bacteria that may lead to nosocomial infections.[4]

The skin of healthcare workers carries greater quantities of bacteria than the general public. Sumner et al reported that 50% of medical staff carried potentially pathogenic bacteria, such as S. aureus, E. coli, Streptococcus viridans, and Proteus vulgaris in their hair.[19] A study conducted on microbiology personnel where aerosolized bacteria were sprayed on bearded and clean-shaven men showed that bearded individuals retained the bacterium on their faces longer than their clean-shaven counterparts.[13]

The role of surgical facemasks (SFMs) in the operating room and the prevention of SSI remains controversial.[16-18] Tunen et al conducted a randomized study in which 1537 operations were performed with masks with 4.7% SSI and 1551 operations were performed without a facemask with 3.5% SSI, with no statistical significance between the groups.[19] A National Surgical Quality Improvement Program review of 6517 patients in 2 teaching hospitals that were visited by the Department of Health that imposed more stringent regulations on operating room attire, including coverage of facial hair and full coverage of ears, did not show any significant decrease in SSI before and after the site visits.[16]

Studies investigating areas directly under the operator have shown that SFM almost completely prevents bacterial contamination of agar plates placed 30 cm below the lips.[17,18,20] The theory of bacterial shedding is based on the concept of

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**Table 3**

| Samples | Participant group | Meropenem-resistant isolates (n = 18) | P value |
|---------|-------------------|-------------------------------------|---------|
| A       | Bearded men       | 7 (38.9%)                           | .96     |
|         | Nonbearded men    | 2 (11.1%)                           |         |
| B       | Bearded men       | 7 (38.9%)                           | .84     |
|         | Nonbearded men    | 2 (11.1%)                           |         |

**Table 4**

| Sample number | Bearded vs not bearded |
|---------------|------------------------|
| 16A           | No                     |
| 2A            | Yes                    |
| 28A           | Yes                    |
| 2A            | Yes                    |
| 28A           | Yes                    |
| 32A           | Yes                    |
| 32B           | Yes                    |
| 33A           | Yes                    |
| 34A           | Yes                    |
| 34B           | Yes                    |
| 36A           | Yes                    |
| 39B           | Yes                    |
| 41A           | Yes                    |
| 41B           | Yes                    |
| 41A           | No                     |
| 43A           | No                     |
| 45A           | Yes                    |
| 45B           | Yes                    |
| 48B           | No                     |
| 48A           | No                     |
| 49A           | Yes                    |
| 49B           | Yes                    |
| 51A           | Yes                    |
| 51B           | Yes                    |
| 53A           | Yes                    |
| 54A           | Yes                    |
| 54B           | Yes                    |
| 57B           | Yes                    |
| 59A           | Yes                    |
| 59B           | Yes                    |
| 60B           | No                     |
| 62A           | No                     |
| 62B           | No                     |
| 63B           | Yes                    |
| 65A           | Yes                    |
| 65B           | Yes                    |
| 65A           | Yes                    |
| 67A           | Yes                    |
| 67B           | Yes                    |
At a higher frequency in bearded individuals; however, this did not reach statistical significance. In addition, we showed that 4% chlorohexidine was highly efficient in inhibiting the growth of all strains, even the meropenem-resistant strains. Chlorohexidine-based solutions are widely used for prepping surgical sites before procedures to reduce the risk of SSI. Since 2012, our institution has adopted the use of chlorhexidine-alcohol-based solutions for preoperative prepping of surgical sites. Darouiche et al demonstrated the superiority of chlorhexidine-alcohol-based solutions over povidone-iodine-based solutions in the prevention of SSIs (9.5% vs 16.1%, *P* = .004) in clean-contaminated surgeries, reducing the risk of SSI by 41%.[23]

The activity of operating room personnel is the principal source of airborne bacteria that originate mainly from the skin of people present in the room.[23] The number of airborne bacteria depends on the number of people present, their level of activity, and compliance with infection control practices.[24] Grooming habits and daily soap washing may affect bacterial growth. Individuals from the surgery department were found to harbor the heaviest bacterial growth in samples A and B. Regarding the hair shedding samples, this study showed that the farther the patient-physician distance, the lower the bacterial growth.

Crucial factors to prevent microbial infection expansion by operating room personnel are personal integrity and work ethics. All surgical teams and staff personnel should adhere to the standard guidelines as an essential part of the prevention process.

Despite being aware of the limitations of our study, we initially planned to enroll 49 participants in each group; however, because the majority of the male healthcare workers grew their beards at the time of the study – most of the participants were bearded – we were not able to recruit > 80 participants. Our study limitations entail an unequal sample size. The adopted statistical test specifically the chi-square is a nonparametric test that is a distribution-free test concerning the distribution of the data. Yet, the study results may be potentially skewed.

### 5. Conclusion

Our study shows that nonbearded healthcare workers in the operating room had a significantly higher bacterial load in their facial flora than their bearded counterparts. The difference in shaving methods and facial washing with or without soap did not translate into a significant difference in bacterial growth between the 2 groups. Even though the bacterial resistance profile was not significantly different between MIC the2 groups, bearded individuals had relatively more resistant strains.
Author contributions

J.H. and M.E.E. are both the first authors and have contributed equally to this manuscript. M.E.E. and A.Z. proposed the study. J.H. and N.M. performed the data collection. J.H., N.M., and A.S. performed the laboratory work. J.H. and R.J. analyzed the data. R.J. is a statistician. A.Z., M.E.E., and J.H. reviewed the literature and wrote the first draft. All authors contributed to the design and interpretation of the study and drafted the manuscript.

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