ASSESSMENT OF MICROBIOLOGICAL AEROSOL
CONCENTRATION IN SELECTED HEALTHCARE
FACILITIES IN SOUTHERN POLAND

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SUMMARY

Objectives: This study was aimed to assess the concentration of microbial aerosol and species composition of airborne staphylococci in 10 healthcare facilities in southern Poland including primary healthcare units and hospital wards; and to assess whether the selected components of microbial aerosol pose a threat of severe infections to either patients or the personnel.

Methods: The study was conducted at monthly intervals over a period of one year. Air samples were collected by MAS-100 sampler. The number of mesophilic bacteria, mould fungi, actinomycetes and staphylococci was determined on general and selective media. The species identification of staphylococci was conducted using API tests for strains that were pre-selected based on macroscopic and microscopic observations.

Results: A total number of 1,584 samples were collected during the sampling period. The numbers of airborne microorganisms varied both between the examined premises and between the seasons of the year. The observed differences were statistically significant with one exception for actinomycetes and their differences between the examined premises. The concentrations of mesophilic bacteria varied from 5 to 297 CFU/m³ of air, for Staphylococcus the values ranged from 1 to 96 CFU/m³, for fungi – from 1 to 100 CFU/m³, and the number of actinomycetes ranged from 7 to 321 CFU/m³. Ten species of coagulase-negative staphylococci (CoNS) were identified among 55 isolates with S. saprophyticus and S. warneri being the most frequently detected (n = 14 and 13, respectively). S. haemolyticus, which is one of the most common causal agents of nosocomial infections was observed in four facilities (n = 5).

Conclusions: The microbial concentrations varied both between the seasons of the year and between the examined facilities. The highest bioaerosol concentrations were observed in most crowded premises. The identified species of staphylococci, although not typically associated with human infections, are common causal agents of nosocomial infections and infections in immunocompromised people.

Key words: airborne microorganisms, healthcare facilities, bioaerosol, microbial aerosol, Staphylococcus spp.

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INTRODUCTION

The concentration of airborne microorganisms in healthcare facilities can be affected by a number of different biotic and abiotic factors. Biotic, or biological factors include the health condition of both personnel and patients, as well as the presence of persons who can be transmitters of microorganisms (1). Another problem related to microbial contamination of air in healthcare units is related to the appearance of microorganisms of modified properties, mainly antibiotic resistant bacteria (2).

By using precise measuring techniques and by culturing the air samples on blood agar for 24–48 h at 35°C, Bischoff et al. (3) demonstrated that sneezing causes a highly significant increase in S. aureus, coagulase-negative staphylococci (CoNS) and other (not determined to the species level) bacteria in environmental samples. One sneeze expels, on average, mostly other bacteria, such as α-haemolytic streptococci (474 CFU/m³/min), but also 3.24 CFU of CoNS/m³/min and 2.83 CFU of S. aureus/m³/min. Also, mean airborne counts may reach up to 15, 47 and 2,522 CFU/m³/min for S. aureus, CoNS and other bacteria after sneezing. During sneezing, the infected person emits up to 40,000 droplets with the velocity of 100 m/s. Larger droplets fall to the ground very quickly, but the smaller ones evaporate and become droplet nuclei, which are typically 1–5 µm in diameter and can remain suspended in the air for long hours (4). Such small numbers of droplet nuclei allow them to pass the host defence mechanisms and the human upper respiratory tract to be deposited in the alveoli in the lungs (4). Additionally, spreading of bioaerosol is affected by the air movement caused by ventilation, differences in temperature or humidity, or even moving of hospital equipment (5). Another group of abiotic factors affecting transmission of microorganisms includes the types of carriers, such as the materials of which the equipment is made, clothes or room facilities, and equipment (1). Studies show that in hospital conditions microorganisms can be deposited on clothing of medical personnel, blankets or towels and that the survival rate depends on the microbial genus or species. For instance, the highest survival rate on towels and cotton clothes was detected for Staphylococcus spp., enterococci and mould fungi (1, 6).
While healthcare facilities face challenges common to all buildings, they meet an additional challenge of high-density populations of potentially contagious and immunocompromised people (7). This results in various problems regarding infection control, since all respiratory pathogens may cause hospital-acquired infections (8). The list of known hospital-acquired infections presented by, among others, Fernstrom and Goldblatt, (7) includes those caused by different viruses, as well as fungi such as Aspergillus spp., Mucor or Rhizopus stolonifer, or bacteria such as Mycobacterium, Nocardia, Pseudomonas aeruginosa or staphylococci.

Having in mind the significance of the problem related to the microbiological quality of air in a specific group of places such as hospitals and healthcare facilities, the study was undertaken in order to assess the concentration as well as the composition of microbial aerosol in the selected healthcare facilities (outpatient units and hospital wards) in southern Poland. Particular attention was paid to the concentration and species composition of Staphylococcus spp., as some species of these bacteria, such as S. aureus, S. epidermidis, S. saprophyticus or S. haemolyticus may be associated with particularly severe infections acquired in hospitals and healthcare units (2, 9).

MATERIALS AND METHODS

The study was conducted over a period of one year, at monthly intervals from May 2014 to April 2015 in ten selected healthcare facilities (Table 1), including four waiting rooms of outpatient specialized units (I–IV), one hospital waiting room (V), one treatment room of an infectious ward (VI), two sickrooms – one of a children’s ward (VII) and of an intensive care unit (VIII), intensive care room of a cardiology ward (IX), and isolation ward of an intensive care unit (X). Atmospheric air collected in the park outside one of the units was treated as control (C). Air samples of 100 L were taken using MAS-100 (Merck) air sampler (with the cut-off size of 1.47 µm). The measurements were conducted according to the procedure described in the PN-Z-04008-08:1989 standard (10), which means that the sampler was placed at approximately 1.5 m above the ground, i.e. in the human breathing zone and the volume of air collected was empirically adjusted to the expected concentration of microbiological aerosol. The measurements were conducted on usual working days of the tested facilities, when both the patients and the personnel were present. The approximate hours of sampling were from 10 to 13 (i.e. during the visiting hours in hospitals and during regular working hours of outpatient units). The numbers of the total mesophile bacteria (Trypticase Soy Agar – Biocorp, 48 h at 36±1°C), fungi (Malt Extract Agar – Biocorp, 3–5 days at 24°C), actinomycetes (Gauze Medium, 7 days at 24°C) and staphylococci (Chapman Agar – Biocorp, 48 h at 36±1°C) were assessed. After incubation, the numbers of colonies characteristic for different microbial groups were counted and expressed as colony forming units per cubic meter of air (CFU/m³). The actual colony count per each culture plate was corrected according to the positive hole correction table (Operator’s manual MAS-100) (11). All measurements were conducted in triplicates, which – having regard to the fact that the sample collection was conducted in eleven sites, once per month over a year, with four groups of microorganisms – gives a total number of 1,584 samples collected during the sampling period. The results from three replications were used to calculate mean values, which are shown as final results in Figures. The samples were collected one after another and between each sampling, the impactor was disinfected by using cotton balls immersed in 70% ethanol. Currently, there are no generally applicable standards that would determine the limit values for concentrations of airborne microorganisms. Therefore, the results obtained based on the analysis were compared to the proposed guidelines by the Team of Experts in Biological Factors (Polish acronym ZECB) (12) on the concentrations of airborne microorganisms in indoor air, including living spaces and public utility premises.

Moreover, after incubation, characteristic colonies grown on Chapman medium were selected for further analysis. Pure cultures were obtained by plate streaking on Chapman medium. Further identification was based on Gram staining, furazolidone sensitivity test and API Staph tests (BioMerieux).

Statistical analysis was performed using Statistica v.10 (StatSoft) – basic descriptive statistics were calculated, and a one-way ANOVA test was employed to verify the significance of seasonal differences in the number of microorganisms and the differences between individual healthcare facilities.

<table>
<thead>
<tr>
<th>Sampling sites</th>
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<tr>
<td>Symbol</td>
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<tr>
<td>----------------</td>
</tr>
<tr>
<td>Internal medicine outpatient clinic – waiting room</td>
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<tr>
<td>Specialist outpatient clinic – waiting room</td>
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<tr>
<td>Allergology outpatient clinic – waiting room</td>
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<tr>
<td>Internal medicine outpatient clinic 2 – waiting room</td>
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<tr>
<td>Hospital 1 - waiting room</td>
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<tr>
<td>Hospital 2 infectious ward – treatment room</td>
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<tr>
<td>Hospital 3 children’s ward – sickroom</td>
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<td>Hospital 4 intensive care unit – sickroom</td>
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<td>Hospital 5 cardiology ward – intensive care room</td>
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<td>Hospital 4 intensive care unit – isolation ward</td>
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<td>Control site – outdoor air, park in front of one of the units</td>
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RESULTS

The results of the conducted studies are summarized in Figures 1–5. The values presented in Figures 1–4 are means of measurements conducted in triplicates in individual months. The results presented in Fig. 5 are means of measurements conducted annually.

The analysis of variance showed that the differences between the studied locations in the numbers of mesophilic bacteria, staphylococci and fungi were statistically significant (F value of 7.67, 2.67 and 3.24, respectively, p < 0.05). Only in the case of actinomycetes these differences appeared not to be significant. Also, the analysis of the differences in the microbial numbers between the seasons of the year indicated the statistical significance of differences (F value of 3.18, 3.86, 20.94 and 9.28 for mesophilic bacteria, staphylococci, fungi and actinomycetes, respectively). The differences are significant at p < 0.05.

The number of mesophilic bacteria ranged from 5 CFU/m³ (site No. IV, winter) to 297 CFU/m³ (No. V, autumn). Also, as shown in Fig. 1, the greatest concentration of mesophilic bacteria was observed in autumn and winter – in autumn the highest values were recorded in the sites No. II, III, IV and V, while in winter – in the sites No. VI, VII and IX. The smallest mean number of these microorganisms was observed in the facility No. VIII, i.e. 23 CFU/m³ while the greatest mean number, 128 CFU/m³, in the facility No. V (Fig. 5).

The number of airborne staphylococci varied between 1 CFU/m³ (IX, spring) to 96 CFU/m³ (IX, winter). In most of the studied sites the greatest values of staphylococcal concentration were recorded in summer – sites No. I, III, IV, V, VIII and X (Fig. 2). The mean numbers of staphylococci ranged from 8 CFU/m³ in the site No. IV and 9 CFU/m³ in the site No. X to 38 CFU/m³ in the site No. IX (Fig. 5).

As shown in Fig. 3, the highest values of fungi were also detected in summer in all of the examined sites and they ranged from 1 CFU/m³ in winter at the site No. VIII to 100 CFU/m³ in summer at the site No. IV. In general, the smallest number of fungi was recorded at the facility No. VIII (6 CFU/m³) while the largest value was observed at the facility No. II (i.e. 43 CFU/m³) (Fig. 5).

Finally, the greatest numbers of actinomycetes were observed in winter – in six out of ten studied healthcare facilities, i.e. No. I, II, III, IV, VIII and IX. The concentration of airborne actinomycetes ranged from 7 CFU/m³ (site No. IV both in spring and autumn) to 231 CFU/m³ (site No. VII in winter) (Fig. 4).
and widespread coagulase-negative staphylococci (CoNS) form biofilms during the respiratory tract infections. This may contribute to, among others, ventilator-associated pneumonia (18). Nosocomial infections can also be caused by the presence of high concentrations of fungi, including the most commonly found environmental moulds, such as Aspergillus, Mucorales, Paecilomyces, Cladosporium, Fusarium and others (19).

There are no regulations concerning the acceptable levels of microbial aerosol concentration in indoor air in Poland (2). According to the “Design guidelines for general hospitals” there are three classes of hospital premises, depending on their cleanliness (20):

- Class I – premises of the highest possible aseptic – the minimum level of bacteria, including sterile boxes, highly aseptic operating rooms etc., with the permissible concentration of 70 CFU of bacteria/m³ of air.
- Class II – premises with low levels of bacteria, including septic and aseptic operating rooms, post-operating rooms and intensive care units with bed premises, with the permissible concentration of 300 CFU of bacteria/m³ of air.
- Class III – premises with normal levels of bacteria, including treatment rooms or diagnostic premises, with the maximum permissible concentration of 700 CFU of bacteria/m³ of air.

Two of the premises examined in this study can be included into class II (sites No. IX and X), while the remaining premises belong to class III. Thus, considering the number of mesophilic bacteria, the permissible level was not exceeded in any of the examined cases.

The microbial aerosol concentrations recorded in our study were also compared to the guideline proposals, given by the Polish Team of Experts in Biological Factors (12) on the concentrations of airborne microorganisms in indoor air, including living spaces and public utility premises. Among the microorganisms examined in our study, this proposal includes mesophilic bacteria and fungi, and the concentrations observed in our study did not exceed the proposed limits, which were $5 \times 10^{3}$ CFU/m³ for mesophilic bacteria and $5 \times 10^{5}$ CFU/m³ for fungi, as the maximum numbers of mesophilic bacteria was 297 CFU/m³ and for fungi +100 CFU/m³.

The recorded concentrations of mesophilic bacteria are in between the ones observed by other authors, as e.g. even the highest concentration observed in this study, i.e. 297 CFU/m³ was lower than the smallest values in the study by Karwowska et al. (2), who observed the number of mesophilic bacteria ranging from 320–560 CFU/m³ in the patients’ waiting rooms. On the other hand, the concentration of bioaerosol composed of mesophilic bacteria in the children hospital sickrooms (site No. VII – values ranging from 24 to even 286 CFU/m³ in winter) were higher than the ones observed by Li and Hou (21), where they were lower than 160 CFU/m³. In 2005, Pastuszka et al. (22) published their preliminary results of measurements conducted in Silesian hospitals, in which the level of bacterial aerosol was found to be about $10^6$ CFU/m³ in clinical outpatient rooms and ranged from $10^5$ CFU/m³ to $10^7$ CFU/m³ in hospitals, depending on the number of occupants and physical quality of the building. In our study, the level of $10^6$ CFU/m³ of bacteria was exceeded.

**DISCUSSION**

The problem of microbiological contamination of air is associated with many different types of premises, including university rooms, offices, laboratories, canteens as well as healthcare centres and hospitals (13–17). This problem becomes even more severe when it can affect the health of people dwelling in such premises, as in the case of healthcare units and hospitals. The respiratory system of human organism is a gateway for bacterial or viral infections, as well as for environmental pollutants. After colonization of mucous membranes, microorganisms can dwell asymptotically and in cases of impaired host resistance – they can cause respiratory infections and in severe cases such infection can spread throughout the body. The most important step in bacterial colonization is their adhesion to epithelial cells, where they must survive the purification by secreted mucus. Both pathogens, such as *Streptococcus pneumoniae* or *Haemophilus influenzae* and widespread coagulase-negative staphylococci (CoNS) form biofilms during the respiratory tract infections. This may contribute to, among others, ventilator-associated pneumonia (18). Nosocomial infections can also be caused by the presence of high concentrations of fungi, including the most commonly found environmental moulds, such as *Aspergillus*, *Mucorales*, *Paecilomyces*, *Cladosporium*, *Fusarium* and others (19).

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**Fig. 5.** Annual mean numbers of airborne microorganisms in studied healthcare facilities (CFU/m³).

I–X – numbers of tested facilities; C – control site located outdoors.

**Fig. 6.** Numbers and share (%) of individual staphylococci species in the total of 55 isolates.
only five times, mostly in waiting rooms, where the number of persons per day exceeded 200. The only patient room, where the level of bacterial aerosol was the highest, was in the children’s hospital in winter. Generally, in winter the numbers of microbial components of bioaerosol were very high in the children’s hospital and the highest numbers of actinomycetes were observed in these particular premises. This can be related to the fact that winter is the period of increased infections, which also affected the number of patients in this hospital. Moreover, rooms in this hospital are small with carefully closed doors and windows, and with each child patient being accompanied by at least one parent, the rooms became severely overcrowded.

The maximum concentration of *Staphylococcus* spp., i.e. 96 CFU/m³ as well as the highest mean concentration of these bacteria was recorded at the site No. IX, which is the intensive care unit of the cardiology ward. The observed concentrations of staphylococci are slightly higher than the ones observed by Klánová and Hollerová (23), who recorded the values ranging from 5 to 70 CFU/m³. On the other hand, our values are much lower than the ones recorded by Karpowska et al. (2), who observed the concentrations of *staphylococci* even up to 350 CFU/m³. However, patients of this ward are very often intubated people, whose breathing is supported with ventilators, in which even small concentrations of *staphylococci* may promote the emergence of nosocomial infections (24). Also, in the study by Pastuszka et al. (22), the numbers of *staphylococci* were mostly higher than the ones observed in our study, as they ranged from 60 CFU/m³ in an operating room to 379 CFU/m³ in one of the patient rooms. The values recorded in this study did not exceed the level of 60 CFU/m³, except for one observation, mentioned earlier, i.e. 96 CFU/m³ in the intensive care unit of the cardiology ward. It needs to be mentioned, that the study by Pastuszka et al. (22) was conducted nearly 15 years ago, therefore, the observed differences may result from the fact that the procedures used in outpatient clinics and hospitals have changed over this period, and it may provide an interesting suggestion to conduct a detailed analysis of the changes in the bioaerosol levels in hospital premises, as a possible result of the renovations and procedure changes.

As for the number of fungi, the concentrations detected in this study, ranging from 1 to 100 CFU/m³, are higher than the ones reported by Karpowska et al. (2), who observed the values of 15–35 CFU/m³ of air. They are very similar to the values observed by Augustowska and Dutkiewicz (25), which range from 10 to 96 CFU/m³ and they are smaller than those observed by Li and Hou (21), i.e. 260 CFU/m³. Still, the observed values do not exceed the limits proposed by the Team of Experts (12), i.e. 500 CFU/m³ of air for indoor and public utility premises or by Krzysztofik (26), set at 200 CFU/m³ of air for sickrooms.

The values of airborne actinomycetes ranged from 7 CFU/m³ in one of the sickrooms to 231 CFU/m³ in the children’s ward. The highest concentration was recorded in winter, similarly to the number of mesophilic bacteria – in the period when the rooms were overcrowded with patients and their parents. In addition to the fact that actinomycetes are well recognized indoor air pollutants, high concentrations of their spores in the air are related to the incidence of allergic alveolitis and asthma, as well as other health effects (27). According to Hirvonen et al. (28), the spores of *Streptomyces* spp. can stimulate lung macrophage reactions, resulting in inflammation and tissue injury.

A total of 55 isolates belonging to the genus *Staphylococcus* were subjected to further identification, which revealed the presence of 10 different species, all of which were coagulase negative (CoNS). Among them, *S. saprophyticus* (n = 14) and *S. warneri* (n = 13) were most frequently detected. Until recently, CoNS were described as ubiquitous commensals of healthy human skin and mucosa. However, they are being more frequently reported as important opportunistic pathogens, mainly associated with healthcare-acquired infections in patients with indwelling medical devices (29). The most abundant in this study, *S. saprophyticus*, which is a part of a human microflora, is also one of the most frequent agents of community-acquired urinary tract infections (30, 31). The second most frequently identified species, *S. warneri* was isolated from one of the healthcare units and four hospital wards, including cardiology, sickroom of an intensive care unit and an infectious ward, whose patients’ immune system may be compromised. This information might be important, as this species was reported to cause infections in hospitalized patients and was also suggested as a cause of, among others, ventricular shunt infections, endocarditis and even sepsis (32, 33). One of the most dangerous CoNS, *S. haemolyticus* was isolated from 4 sites (n = 5) – three hospitals, including the isolation ward of an ICU. This species is very frequently isolated nosocomial infections agent, causing various infections, including endocarditis or septicemia, as well as it is recognized as one of the most multidrug resistant CoNS (34).

**CONCLUSIONS**

Based on the presented results it can be stated that microbiological aerosol within healthcare facilities changes with seasons of the year and its concentration may affect the health of both patients and personnel. Most densely populated premises proved to be characterized by the highest concentration of microbial aerosol. All of the detected species of *Staphylococcus* are coagulase-negative and are not typically associated with human infections, however, they are very common causal agents of nosocomial infections and infections in immunocompromised people. Such infections are dangerous to both patients and medical personnel and regular examinations of airborne microbiological contamination should be one of the methods employed in prevention against such infections. Moreover, ensuring proper maintenance of ventilation and air conditioning systems is one of the most important factors affecting the air quality.

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**Conflict of Interests**

None declared.
REFERENCES

5. Tang TW, Li Y, Eames J, Chan PKS, Wlazlo A. Bacterial aerosol in Si-

13. Edmiston CE, Seabrook GR, Cambria RA, Brown KR, Lewis BD, Som-
15. Tang TW, Li Y, Eames J, Chan PKS, Wlazlo A. Bacterial aerosol in Si-

17. Pastuszka JS, Marchwińska-Wyrwal E, Wlazlo A. Bacterial aerosol in Si-

22. Pastuszka JS, Marchwińska-Wyrwal E, Wlazlo A. Bacterial aerosol in Si-

27. Reponen T, Gøtschke GV, Grishpun SA, Willeke K, Cole EC. Characteris-

33. Widerström M, Wiström J, Sjöstedt A, Mønssen T. Coagulase-negative staphylococci: update on the molecular epidemiology and clinical presen-

36. Pastuszka JS, Marchwińska-Wyrwal E, Wlazlo A. Bacterial aerosol in Si-

41. Reponen T, Gøtschke GV, Grishpun SA, Willeke K, Cole EC. Characteris-

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