Bioaerosols in an Underground Tourist Trail

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Abstract

Cultural heritage objects, often aged and vulnerable, are at a particularly high risk of microbial pollution. Enzymatic activity of bacteria and fungi might lead to the damage of precious antique objects, while microbial pathogenicity and immunogenicity can pose a threat to the health of staff and visitors. Therefore, monitoring the microbiological contamination of cultural heritage objects is necessary, especially in the case of unique objects, like tourist trails located underground. We have evaluated the concentration and diversity of microorganisms present in the air and surfaces of the Lublin Underground Trail located in Poland. We have found the prevalence of bacterial species being a part of the human microbiota in the analysed air. The average fungal contamination of the air was low (337 CFU/m³), but we have identified species of high immunogenicity and those contributing to biodeterioration. Microclimatic parameters were considered as favourable for microbial growth, which emphasizes the role of adequate heating and ventilation air conditioning systems providing sufficient air quality in cultural heritage objects.

Keywords: bioaerosol, indoor air, historic buildings, cultural heritage, biodeterioration

Introduction

Cultural heritage objects are at a particularly high risk of microbial pollution due to their location in historic buildings. Old buildings are often unadapted to the role of libraries, archives and museums, thus providing insufficient conditions to store art works vulnerable for the destruction caused by enzymatic activity of microorganisms during the process called biodeterioration. Long-term exposure to biological agents might lead to allergies, asthma, allergic rhinitis, inflammation of the sinuses and bacterial or fungal infections in immunocompromised people [1]. The role of monitoring the microbiological quality of the air is emphasized by the fact that people spend up to 90% of their time at home or at work [2].

Air constitutes a vector of bacteria, fungi, and viruses, as well as their metabolic compounds like endotoxins, mycotoxins and MVOCs (volatile organic compounds). Dust plays a key role in transmitting microorganisms by air, because it acts as a carrier of organic particles. Bioaerosols – all organic particles suspended in the air – represent even one third of the indoor air pollutants [1]. Bioaerosol particles are characterized by a large variation in size (from 0.5 μm of bacterial cells to even 25 μm of fungal
spores), as well as multiple microbial species in different concentrations [3]. The presence of a small amount of organic matter, i.e., in dust, is adequate for the growth of fungi and bacteria. The factors that favour the growth of bacteria and filamentous fungi include: high water activity, increased levels of carbon dioxide, insufficient ventilation, lack of appropriate air exchange systems, increased humidity and temperature fluctuations [1, 4]. Increased humidity along with a high level of salinity leads to the development of halotolerant and halophilic microorganisms that are responsible for a very fast corrosion process that destroys inorganic materials. The activity of this group of microorganisms is observable in a form of characteristic salt efflorescence on infected surfaces [5]. The most easily observed signal indicating biodeterioration is also surface change of material, including fiber damage, structural defects, visible growth of microorganisms and decolourisation [4]. The selection of an HVAC system has critical significance both for maintaining the proper microclimatic conditions that reduce microbial contamination of the air as well as for protecting historic objects against biodeterioration. In the case of HVAC systems, constant monitoring of the technical condition of the system and its valid operation is equally important, because even the inappropriate direction and the speed of airflow can increase the level of microbiological contamination of the indoor air. It was also established that a constant temperature significantly limits the number of bacteria in the air [6].

Monitoring the microbiological quality of the air allows us to identify factors potentially harmful to human health in specific work places, which is also an issue highlighted by other authors [7]. For those reasons, sources, concentration and the diversity of microorganisms in cultural heritage objects should be determined.

Our aim was to determine the concentration and diversity of microorganisms present in the air and on surfaces of the Lublin Underground Trail as well as to discuss the potential sources of identified microorganisms and their role in the biodeterioration of cultural heritage objects.

Material and Methods

Sampling Site

The underground tourist trail is currently a very popular attraction in Lublin (Poland). This 280-m (918.6 feet) long maze was created through a connection of restored cellars, located over a dozen meters below ground; the oldest cellars were built in the 16th century. Gravity ventilation ducts are located along the entire trail, but there is no heating and air conditioning. One of the main attractions is wooden maquettes visualizing the development of the city. In 2017 a total of 27,647 tourists visited the trail. The average number of tourists per month was 2304±1242; the lowest noted for January (576) and the highest for May (4098). The Lublin Underground Trail consists of one main corridor connecting 14 small chambers. Side chambers, however, are not separated from the corridor. Measuring points were located at equal distances from each other (approximately 50 m) to ensure the best reflection of the average concentration of microorganisms in the air of the entire trail. Because of the age of the studied place and underground location, the tourist trail creates unique environmental conditions.

Sampling and Analytical Methods

Microbiological analysis of the air was carried out using the collision method and a Microbio MB1 microprocessor air sampler (De Ville Biotechnology, France). All measurements were repeated twice. Samples of the air (50 L) were collected in seven selected points at a height of 1.5 m above the floor. To determine the total number of bacteria and fungi in the analyzed air we used tryptic soy agar and Sabouraud medium (BTL Ltd, Poland). The presence of staphylococci and Gram-negative rods was determined using mannitol salt agar and MacConkey medium (BTL Ltd, Poland), respectively. Plates were incubated for 24 h at 37°C, except Sabouraud medium, which was incubated for 5 days at 25°C. After the incubation, the number of bacterial and fungal colonies were counted. The results were corrected according to the conversion tables provided by the air sampler manufacturer and inserted into the following formula, and the concentration of microorganisms in the air were presented as colony forming units per 1 m³ of air (CFU/m³):

\[ X = \frac{(a \times 1000)}{V}, \]

where:

- \( a \) – the sum of microbial colonies grown on a given medium
- \( V \) – the volume of air sample (L)

The type of fungal contamination of surfaces was determined by swabbing wooden maquettes, walls of the corridor and cellars, and the openings of ventilation ducts with sterile cotton swabs. Swabs were inoculated on Sabouraud medium and incubated in the conditions mentioned before. The following microclimatic parameters were also measured: relative humidity (8706 Psychrometer, AZ Instrument Corp.), air movement, temperature (Hot Wire Anemometer AM-4204, Lutron Electronic) and the concentration of carbon dioxide (MX6 iBrid Six-Gas Monitor, Industrial Scientific). Species identification of bacteria was conducted based on Gram staining, production of catalase, coagulase, cytochrome oxidase and biochemical API tests (API 20E/CORYNE/STAPH/20 STREP, BioMérieux). The fungi were identified both on the basis of their macroscopic and microscopic characteristics using the key for fungi identification [8]. The correlation
between the number of microorganisms in the air and microclimatic conditions was determined using Spearman’s test and Statistica 13 software (StatSoft Inc.).

Results and Discussion

Total Number of Bacteria and Fungi

The microbial load in the air of historic old buildings depends on many factors, including humidity, type of ventilation, geographic location, the presence of antique objects and the number of visitors. We have assumed that the underground location of the tourist trail promotes the accumulation of water, therefore microbial contamination of the air would be significant. However, we found that the average number of bacteria isolated from the air of the Lublin Underground Trail – 466±247 CFU/m$^3$ – was comparable to that found in other studies (Fig. 1). Similar results were found in the air of museum rooms – 340-560 CFU/m$^3$ [9], as well as archives and library rooms – 110-470 CFU/m$^3$ [10], both located in Poland. In four out of seven measurement points, the number of bacteria were at least twice as high as the number of fungi. The prevalence of bacteria in the indoor air bioaerosol is commonly found in many studies. Skóra, Gutarowska, and Pietrzak [10] noted that bacteria accounted for 56-84% of the bioaerosol of Polish archives and library rooms. Low average fungal contamination (337±200 CFU/m$^3$) was probably caused by the lack of airflow in the vents, depriving the environment of access to the atmospheric air, which is the main source of these microorganisms (Table 1).

The number of mannitol-positive staphylococci varied (0-90 CFU/m$^3$), but in six out of seven measurement points this was higher than the number of mannitol-negative ones (Fig. 2). Genus *Staphylococcus* is one of the indicators of air quality, including contamination with potentially pathogenic species like *S. aureus* [11]. The high concentration of staphylococci in the indoor air is most often caused by emissions from the human upper respiratory tract [12], thus visitors might increase the bacterial air contamination. The presence of Gram-negative rods were noted only in three measurement points and the concentration in the Lublin Underground Trail air was low (0-30 CFU/m$^3$).

Table 1. Concentrations of CO$_2$ and microclimatic parameters in the Lublin Underground Trail.

| Air sample | CO$_2$ (% vol.) | Temperature (ºC) | Relative humidity (%) | Air movement (m/s) |
|------------|-----------------|-----------------|----------------------|-------------------|
| 1          | 0.06            | 17.1            | 63                   | 0                 |
| 2          | 0.06            | 16.2            | 63                   | 0                 |
| 3          | 0.03            | 15.3            | 59                   | 0                 |
| 4          | 0.06            | 13.5            | 66                   | 0                 |
| 5          | 0.06            | 14.5            | 85                   | 0                 |
| 6          | 0.06            | 12.0            | 62                   | 0                 |
| 7          | 0.06            | 13.5            | 58                   | 0                 |
| Mean±SD    | 0.06±0.01       | 14.6±1.8        | 65.1±8.5             | 0                 |
Bacterial and Fungal Species

The main source of microorganisms in indoor air is outdoor air, as well as people and animals. People spread bacteria and fungi by moving, coughing or talking; they can also transfer environmental microorganisms on their clothes or shoes [13]. We have found presence of multiple bacteria species in the Lublin Underground Trail that belong to the physiological human microbiota and usually inhabit the skin, mucous membranes of the oral cavity or the upper respiratory tract (Table 2). Being a component of the human microbiota, *Staphylococcus* spp., *Corynebacterium* spp., *Micrococcus* spp. and *Kocuria* spp. are often isolated from indoor air [9, 14]. It is also well known that the high density of streptococci in bioaerosols can be found in crowded rooms [15]. Part of identified commensal microorganisms act as opportunistic pathogens, and in high concentrations may pose a threat to human health, which is confirmed by the examples of diseases reported by different authors (Table 2). However, these bacterial disease-causing capabilities activate in the case of the hosts’ immune system disorders or the presence of serious comorbidities, thus the probability of infection among the staff and visitors of the Lublin Underground Trail is low. On the other hand, we observed salt efflorescence on the walls of the main corridor and side rooms as well as water condensation. Such conditions – high salinity and humidity – facilitate biodeterioration and provide an environment suitable for the development of halophilic and halotolerant microorganisms, like genus *Staphylococcus* and *Micrococcus* able to grow in 15% salinity [16-18].

All of the fungal species isolated from the analysed air are being commonly found as part of an outdoor bioaerosol, and their pathogenicity is mainly connected with the high concentration in the air, the presence of immunogenic particles and/or the

| Bacterial species | Possible sources | Pathogenicity | Biodeterioration abilities |
|-------------------|-----------------|--------------|---------------------------|
| *Propionibacterium avidum* | skin of groins, armpits and crotch [19] | NDA | none [19] |
| *Cellulomonas* spp. | soil [20] | NDA | decomposition of cellulose [4] |
| *Cellulosimicrobium cellulans* | soil [20] | NDA | biocorrosion of mineral building materials [4] |
| *Brevibacterium* spp. | water [21] | NDA | deterioration of concrete, brick, glass [4] |
| *Leifsonia aquatica* | water [22] | acute septicemia [22] | NDA; *Leifsonia poae*: deterioration of wooden altar [23] |
| *Staphylococcus haemolyticus* | skin [24, 25] | neonatal sepsis [24] | deterioration of photographic paper [26] |
| *Staphylococcus hominis* | skin of arms, legs, trunk [27, 28] | general infections, sepsis [29] | deterioration of photographic paper [26] |
| *Staphylococcus capitis* | skin [30] | prosthetic joint infections [31] | deterioration of stone [26] |
| *Staphylococcus* spp. | skin and mucous membranes [14] | dependent on the species | deterioration of stone and murals [32] |
| *Micrococcus* spp. | skin and mucous membranes [14] | NDA | deterioration of stone and murals [32] |
| *Kocuria* spp. | skin, mucosa and oropharynx [33] | endocarditis, catheter-related bacteraemia [33, 34] | NDA |
| *Pseudomonas oryzihabitans* | water and soil [35] | catheter-related nosocomial infections, skin and soft tissue infections [36, 37] | deterioration of stone and murals [32] |
| *Alcaligenes* spp. | water and soil [38] | NDA | corrosion of metals, metal alloys, plant and animal fibers, paint coatings [39] |
| *Aerococcus viridans* 2 | upper respiratory tract [40, 41] | endocarditis, osteomyelitis, meningitis, septic arthritis [40, 41] | NDA |
| *Streptococcus constellatus* sp. *constellatus* | oral cavity, gastrointestinal tract [42, 43] | potential role in chronic periodontitis [44] | NDA |

Note: NDA – no data available
production of mycotoxins. Special attention should be paid to filamentous fungi that belong to primary colonizers – fungi able to grow in the lowest water activity environment, i.e., *Penicillium expansum* [45-47]. We found that most of the isolated fungal species were theoretically capable of producing mycotoxins (Table 3). As previously shown, *Aspergillus flavus* can produce aflatoxin B1 or/and B2 and is an etiological factor of aspergillosis [48, 49]. The presence of genus *Aspergillus* was found in the air, and swab samples from the walls and ventilation ducts (*A. versicolor, A. niger, A. flavus*). For some species, i.e., *A. versicolor*, the environmental factor triggering the production of mycotoxins is the high humidity of colonized materials [47]. Other signal for the production of the fungal secondary metabolites is periodically unfavorable microclimatic conditions [50, 51], thus future studies about the pathogenicity of fungi in the indoor air should cover longer periods of time and consider the presence of mycotoxins in the air.

We found the presence of *Cladosporium* spp. in all of the air and swab samples; it was also the only species colonizing the wooden maquettes. *Cladosporium* spp. has very low toxicity, but high immunogenicity: ten different antigens have been described so far. Longitudinal exposition to *Cladosporium* spp. spores might lead to the development of allergy and asthma [52, 53]. According to Chmiel, Frączek and Grzyb [54], the prevalence of one fungal species in the bioaerosol and the longitudinal exposition might be a cause of adverse health effects. Although the fungi concentration in our studies was low, other authors highlight the role of the immunogenicity of identified species. Gutarowska [55] found that 40% of the inhabitants of mouldy buildings had increased levels of IgE antibodies. Similarly, Wizgiewska et al. [56] noticed that 30% of the employees of the National Museum in Warsaw were allergic to at least one of the filamentous fungi: 13% to *Cladosporium* spp., 11% to *Penicillium* spp., 9% to *Alternaria* spp. and 9% to *Aspergillus* spp. Fungal species present in bioaerosol of the Lublin Underground Trail were commonly found in other historical buildings in Poland [50], and were considered by some authors to be indicators of microbial air pollution (*A. niger, A. versicolor, C. macrocarpum, C. herbarumgenera, Penicillium spp. and Rhizopus nigrans*), [9].

**Microclimatic Conditions**

Lack of air flow in all measurement points testifies to the ineffectiveness of ventilation present in the tested location (Table 1). Appropriate moisture insulation of a building along with the implementation of ventilation or air conditioning systems guarantee the stability of microclimatic parameters [1, 16]. On the other hand, in the case of historical buildings the airflow rate itself could be a critical factor because rapid air movement enables the microorganisms to transfer and displace between different environments [54]. The average air temperature in the Lublin Underground Trail was 14.6°C. The temperature range 18-32°C is considered to be favourable for the growth of most filamentous fungal species in the moderate climatic zone [55]. Optimum temperature for the growth of cellulitic fungi (14-15°C) evaluated by Szostak-Kot [57] is similar to the temperature measured in our studies, which explains the intensive biodeterioration of wooden maquettes. Temperature of the air of the Lublin Underground Trail was not only sufficient for microbial growth, but had a potential to stimulate the production of mycotoxins. Also, the humidity (58-85%) allowed for the growth of moulds as well the development of a biodeterioration process, which has also been observed by other authors [54, 58]. In our studies, the consequences of dampness were visible as water condensation on the walls along with the presence of salt efflorescence. The concentrations of carbon dioxide did not exceed the permissible concentration in enclosed spaces (0.1%) as set by WHO [59].

We have not found any correlation between temperature and relative humidity of the analyzed air and microbiological parameters in all measurement points (n = 7): total number of bacteria (*r* = -0.18, *p* = 0.71 and *r* = -0.17, *p* = 0.71), total number of fungi (*r* = 0.49, *p* = 0.26 and *r* = -0.19, *p* = 0.68), number of mannitol-positive staphylococci (*r* = -0.43, *p* = 0.34 and *r* = 0.06, *p* = 0.9), number of mannitol-negative staphylococci (*r* = 0.32, *p* = 0.49 and *r* = -0.58, *p* = 0.17) and number of gram-negative bacilli (*r* = -0.59, *p* = 0.16 and *r* = -0.13, *p* = 0.79).

| Table 3. Fungi isolated from the surfaces and the air of the Lublin Underground Trail. |
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| **Fungi isolated from the surfaces** |
| **Room walls** | **Corridor walls** | **Ventilation ducts** | **Wooden maquettes** |
| *Aspergillus versicolor* | *Cladosporium* spp. | *Paecilomyces* spp. | *Cladosporium* spp. |
| *Aspergillus flavus* | *Paecilomyces* spp. | *Ulocladium* spp. | *Cladosporium* spp. |
| *Cladosporium macrocarpum* | *Fusarium* spp. | *Mucor* spp. | *Aspergillus flavus* |
| *Ulocladium* spp. | *Mucor* spp. | *Aspergillus niger* | |
| *Phoma* spp. | | | |
| *Rhizopus oryzae* | | | |
| **Fungi isolated from the air** |
| *Rhizopus oryzae*, *Penicillium expansum*, *Mucor* spp., *Aspergillus flavus*, *Penicillium rhizogenum*, *Cladosporium* spp. |
Conclusions

Monitoring the microbiological quality of the air of cultural heritage objects ensures the safety of historic buildings, workers and visitors. Introducing adequate ventilation systems provides desired relative humidity and air temperature. The appropriate microclimatic parameters of the environment not only limit the growth of potentially harmful and toxicogenic fungi, but also slow the biodeterioration process. Analysis of the air of the Lublin Underground Trail showed that people are the main reservoir of isolated bacterial species, because the air itself is an environment inimical for the growth of bacteria and can only lead to their transmission. Based on this observation, the number of visitors to historic buildings on the concentration of bioaerosol.

Conflict of Interest

The authors declare no conflict of interest.

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