The importance of cleanrooms for the treatment of haemato-oncological patients

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Introduction

Patients with haematological malignancies or after autologous or allogeneic hematopoietic stem cell transplantation (HSCT) rank among immunosuppressed individuals. Prolonged and deep neutropenia is considered a key risk factor of the occurrence of an exogenous infection. One of the possibilities of preventing an exogenous infection in these patients is to place them in a “cleanroom” for the crucial period of time. Cleanrooms are intensive care units with reverse isolation.

The final part of the general article below provides an overview of the technology and types of cleanrooms for immunosuppressed patients in compliance with the current recommendations and technical standards.

Key words: cleanroom, haemato-oncological patients, exogenous infection risk.
Cleanroom technology

The beginnings of cleanroom technology date to the first half of the 20th century. Towards the end of the 1930s the HEPA (high performance particulate air) filter was developed, to be used in the Second World War for the gas masks of the Swiss army. The second condition is to ensure a one-way air flow in order to successfully control the particles it contains. A solution to this problem was published by Willis J. Whitfield [1] in 1962.

The cleanroom technology consists of a sequence of follow-up activities designed to control and minimize contaminants in the supply/exhaust air. It needs to meet the following requirements:

- protect the process or product from becoming damaged by contaminated air,
- protect persons from exposure to airborne contaminants that could endanger the persons’ health, and
- protect the outside environment from harmful emissions generated in connection with the operation of the facility [1].

With airing and air conditioning, the filtration of the atmospheric air is the basic method of maintaining the required purity of the interior air of the given room. When creating cleanrooms, multilevel filtration forms an integral part of highly efficient filtration, where the requirements for high air purity ensue from the requirements of individual work procedures and of public health protection [2]. Three-level air filtration is recommended in order to meet the required air cleanliness class in cleanrooms in health care centres. A fine filter of class F 7 is recommended as the pre-filter of the second and third filtration level. Highly efficient HEPA filters H12-H13 are recommended as the basic filters for all cleanrooms of class 100 to 100,000 pursuant to FED-STD-209E [3], which currently corresponds to class 5 pursuant to ČSN EN ISO 14644-1 [4]. They are highly efficient for all types of dust and aerosols including viruses. The three-level filtration must also be installed in discharge systems designed for hazardous aerosols (health care, biotechnology, nuclear power) [4]. For air cleanliness classes, see Table 1.

Although the above represents only a small part of cleanroom technology, it is one of the basic prerequisites for creating the cleanrooms. Cleanrooms in health care centres are air-conditioned with the help of a sanitary air-conditioning unit working exclusively with fresh air. The supply air is humidified/dried and heated/coolied as required. At present, all required parameters of the supply air are carried out with the help of a computer program.

Another vehicle, in many cases even more hazardous than air, is water used for the standard hygiene of patients placed in a protective environment room. Although water coming from water pipes in health care centres meets the drinking water requirements of the applicable legislation, it is not suitable for these patients [5]. It can contain microbial infectious agents that are safe for healthy individuals but in immunosuppressed patients they may lead to even fatal infections [6, 7]. Apart from a number of Gram-negative non-fermenting bacteria, drinking water distribution systems in health care centres are often contaminated by various types of microscopic filamentous fungi [8–12]. Probably the greatest risk is the presence of Legionella in the hot water pipeline. Although hot water is produced from drinking water, it is termed “hot industrial water”. Methods of efficient prevention of the occurrence of this bacterium are complex and costly [13].

Patients in isolation rooms usually use specially treated water for their personal hygiene. This water is a combination of mechanical and chemical treatment of drinking water designed to minimize the risk of the presence of any infectious agent. The occurrence of Legionella in the pipeline of treated water is minimized by having the water heated to more than 64°C and placing final filters on the shower and the faucet shoulder. Membrane pores in these filters are 0.22 μm large, which prevents Legionella and other bacteria from entering.

Places recording increased humidity, e.g. drains, siphons, and sanitary facilities, are considered highly dangerous vehicles for the survival of usually conditioned pathogenic infectious agents. These as a rule involve Gram-negative non-fermenting bacteria that induce dangerous infections in immunosuppressed patients [6, 14–18]. All these places must be paid increased attention and be regularly and efficiently decontaminated. The decontamination results need to be verified using growth media.

Table 1. Selected airborne particulate cleanliness classes for cleanrooms and clean zones (ISO 14644-1:1999 E)

| ISO classification number (N) | Maximum concentration limits (particles/m³ of air) for particles equal to and larger than the considered sizes shown below |
|-------------------------------|---------------------------------------------------------------------------------------------------------------------------------|
|                               | 0.1 μm | 0.2 μm | 0.3 μm | 0.5 μm | 1 μm | 5 μm |
| ISO Class 1                   | 10     |        |        |        |      |      |
| ISO Class 2                   | 100    | 24     | 10     | 4      |      |      |
| ISO Class 3                   | 1 000  | 237    | 102    | 35     | 8    |      |
| ISO Class 4                   | 10 000 | 2 370  | 1 020  | 352    | 83   |      |
| ISO Class 5                   | 100 000| 23 700 | 10 200 | 3 520  | 832  | 29   |
| ISO Class 6                   | 1 000 000| 237 000| 102 000| 35 200 | 8 320| 293  |
| ISO Class 7                   |        |        |        | 352 000| 83 200| 2 930|
| ISO Class 8                   |        |        |        | 3 520 000| 832 000| 29 300|
| ISO Class 9                   |        |        |        | 35 200 000| 8 320 000| 293 300|
Cleanrooms

The origin of cleanrooms in health care centres dates to the time of Joseph Lister, who decontaminated the air in operating rooms by spraying Lysol solution.

The building technologies and materials of the twentieth century facilitated the construction of cleanrooms, which were supposed to solve two fundamental problems. Firstly, they aim to protect the patient from an infectious agent (today also in other departments than just the operating room). Secondly, cleanrooms are built to prevent dangerous infectious agents from leaking from an infectious patient to the ambient environment. The latter was successfully solved by the American aerospace organization NASA, which was able to isolate astronauts in a moon laboratory after they returned from space, preventing the possible leakage of a hazardous infectious agent that could put humankind at risk.

Depending on the pressure ratios of the isolators’ internal environment to the external one, cleanrooms are divided into two basic groups [19].

Airborne infection isolation rooms (AII), designed to isolate patients with hazardous infectious diseases or if there is a suspicion of such a disease. The pressure of the surrounding air should foreclose leakage of the infectious agent. Contaminated air is drained through three-level filtration; the last filtration stage is ensured with HEPA or ULPA filters. These cleanrooms form part of the NASA moon laboratory. Cleanrooms of the above type are of great importance even today. Patients with the MRSA and SARS infections, pulmonary or laryngeal tuberculosis, and HIV positive patients with high-resistance Mycobacterium tuberculosis strains are placed in airborne infection isolation rooms (Fig. 1) [20].

Health care and construction institutions in the USA, e.g. CDC and ASHRAE, issue recommendations concerning cleanrooms for patients treated for tuberculosis [21]. Figure 2 shows examples of AII rooms with an anteroom and neutral anteroom.

Protective environment rooms aim at protecting the patient for the necessary period of time from a possible infectious agent from the outside, preventing an exogenous hospital-acquired infection. They are therefore sometimes referred to as “reverse isolation” (Fig. 3).

In the 1960s, plastic isolators with regulation of the pressure parameters were used by some surgical specialists, e.g. gynaecologists. The latter would deliver children into a sterile environment when the newborn was suspected to be suffering from a serious primary immunodeficiency. The isolator was a plastic “bubble” attached to a sterile pad. Through HEPA filters, sterile air flows into the interior environment, where a drain for the contaminated air is placed at the upper part of the equipment. A slight positive pressure against the ambient environment was maintained in the interior part of the designated area. Following the introduction of these isolators, surgeons recorded a lower number of postoperative infectious complications [6], and yet the isolators failed to become widely used. Basic technical parameters of both types of cleanrooms are listed in Table 2.

In health care centres, the majority of patients need to be protected from infectious agents. New treatment methods, e.g. cytostatic treatment and radiation therapy, require the patient to be protected from an exogenous infection for varying periods.

The implementation and incorporation of cleanrooms into health care centres is, above all, connected to haematological patients.

An isolator designed for clinical purposes was launched in the USA in 1957. This equipment was associated with the name Trexler [22]. In September 1964 the National Cancer Institute (Bethesda) began using an isolator, type “Life Island” produced by Matthews, for post-chemotherapy patients. It was a plastic isolator made from a translucent foil, into which air flew through HEPA filters that eliminated over 99.97% of particles larger than 0.3 μm. The air was changed twice to 16 times an hour, as required. Inside, the staff used special gloves placed in the wall. Two UV tubes were installed at the foot of the bed. The authors provided a precise description of the procedure for transporting a patient into the above isolator, including a strict decontamination regime, which was to prevent transfer of an infectious agent and the origin of an exogenous infection. The microbial contamination of the air (fall-out method) and the surfaces (swabs from hotspots) were continuously checked. Cultivation procedures focused on demonstrating aerobic

Fig. 1. Example of positive-pressure room control for protection from airborne environmental microbes
The importance of cleanrooms for the treatment of haematological patients. In terms of size, the facility was quite small. The treatment was problematic, generating undesirable mental states in patients [23].

In 1967 James and his colleagues [24] introduced a unit that was to decrease the possibility of an exogenous infection in patients after cytotoxic therapy. Two single rooms shared a sanitary facility and all these premises were supplied with treated air (20 air changes per hour; the filtration device retained particles larger than 0.5 μm, while the air temperature was 20°C and relative humidity 50%). These units were surrounded with a “protective” gallery for communication with the attending personnel. The authors stated that it was a compromise between the sterile room system and a single bed isolator.

In 1969 Schneider and his team published their three years’ experience with the pathogen-free isolation unit [25]. Located in France, the unit contained five single rooms with facilities for the nurses and physicians, sterilisation, a kitchen for preparing sterile food, and a piping system with treated water sterilized by UV radiation. The supply air was treated by means of two-stage filtration and germicidal lamp radiation. All the above isolation units only supplied treated air and drained contaminated air.
In 1969 Bodey and his colleagues [26] reported about a stable isolation unit equipped with a horizontal laminar flow of the treated air. An entire wall (behind the patient’s head) was formed by HEPA filters with 99.97% efficiency for particles larger than 0.3 μm. The laminar flow of the intake air required a certain speed and a minimum amount of objects in the room to avoid a turbulent flow. The patient was not supposed to feel the exchange of the contaminated air as a draught. Here, too, a precise schedule of the patients’ reception and stay in the facility was specified. Here, too, a precise schedule of the patients’ reception and stay in the facility was specified.

Czechoslovakia saw the launch of an improvised reverse isolation room for the treatment of patients suffering from haematological diseases in 1972 and the introduction of the laminar air flow system “Life Island Mark 12” produced by Matthews in 1974 [27]. At present, the reverse isolation units are being installed in the majority of Czech faculty hospitals (e.g. Faculty Hospital Motol, Faculty Hospital Brno, and Faculty Hospital Olomouc).

Different opinions on whether the isolator needs to be equipped with a HEPA filter appear in the professional literature. In his study from 2001, Dykewicz states that while the filter’s importance in patients with autologous HSCT is lower or had not been demonstrated, patients with long-term neutropenia should be placed in a cleanroom equipped with a HEPA filter as it is one of the highly hazardous factors for the origin of hospital-acquired aspergillosis [29]. The increased risk of its emergence is related to the remodelling and reconstruction works carried out in health care centres. Any shifts in the old walling are always connected to the release of aspergillus spores. The only protection against this hospital-acquired infection for immunosuppressed patients is a fully functioning three-stage air filtration concluded with HEPA filters [25].

In order to ensure smooth functioning of the entire HVAC device (a defect, reconstruction), the isolation rooms must have a back-up source of energy [31].

The technology of isolation rooms protecting critically ill patients from external infectious agents has existed for over fifty years. The cleanroom technology now provides optimal conditions of a clean environment, minimizing the risk of an exogenous infection. Protective environment rooms are today used especially for patients after autologous or allogeneic HSCT and peripheral stem cells, after high doses of chemotherapy, and for those suffering from a serious form of aplastic anaemia. Infections represent the main cause of morbidity and mortality of most haematological patients. The susceptibility to infection in these patients is closely related to their primary disease, chemotherapy, intravenous catheters, and special operations. 60% of febrile neutropenic patients report a bacterial infection. Bacterial infection prevails at the beginning of neutropenia, while infections induced by microscopic filamentous fungi usually emerge later. A viral infection may appear during the entire neutropenia period. Monitoring the risk of infection in
these patients must also apply to the attending personnel and the whole environment of the relevant department.

Ideally, immunosuppressed patients would be placed in an entirely sterile environment, which can unfortunately be created only theoretically. This quality is violated every time a patient or attending personnel enter the room [28].

Patients themselves may represent the source of an infectious agent. Potential sources of infectious agents include the hair, skin, and mucous membranes of both the patients themselves (endogenous infection) and the attending personnel (exogenous infection). The risk of endogenous infection may be decreased by partial or total decontamination of the microbial population of the oral cavity, nasopharynx, the gastrointestinal tract, genitals, and skin. This problem is handled by clinical staff. Dust particles or food can serve as potential vectors [28]. The most prominent are Enterobacteriaceae, Pseudomonas aeruginosa, Staphylococcus aureus and Candida spp. [30].

Models that distinguish between infectious complications occurring during different post-transplantation phases have been put forth, based largely on a myeloablative paradigm in which phase I is the pre-engraftment phase (< 15–45 days after HSCT); phase II is the post-engraftment phase (30–100 days after HSCT); and phase III is the late phase (> 100 days after HSCT). During phase I, prolonged neutropenia and breaks in the mucocutaneous barrier result in substantial risk for bacteremia and fungal infections involving Candida species and, as neutropenia continues, Aspergillus species. In addition, herpes simplex virus (HSV) reactivation occurs during this phase. During phase II, infections relate primarily to impaired cell-mediated immunity. The scope and impact of this defect are determined by the extent of GVHD and the immunosuppressive therapy for it. Herpesviruses, particularly CMV, are common infectious agents during this period. Other dominant pathogens during this phase include Pneumocystis jiroveci and Aspergillus species. During phase III, persons with chronic GVHD and recipients of alternate-donor allogeneic transplants remain most at risk for infection. Common pathogens include CMV, VZV and infections with encapsulated bacteria (for example, Streptococcus pneumoniae). The relative risk for these infections is approximately proportional to the severity of the patient’s GVHD during phases II and III. For recipients of non-myeloablative grafts, substantial differences may be observed during phase I, but the susceptibility to infections involving phases II and III is largely similar, and driven primarily by the status of the underlying disease, a history of GVHD and/or the need for ongoing immunosuppression. The risk of disease from community-acquired respiratory viruses is elevated during all three phases; however, in phase III, the outpatient status of haematopoietic cell transplant recipients can complicate efforts to reduce exposure and provide timely intervention. Thus, the risk of infection is primarily determined by the time from transplant and the presence or absence of GVHD. Unfortunately, there is currently no definitive laboratory marker of immune reconstitution that can predict infectious risk that could be used to tailor infection prophylaxis [32].

Infection prevention and control in health care facilities where hematopoietic stem cell transplantation recipients are treated

It is necessary to meet the following requirements for efficient infection prevention and control in the cleanrooms for patients after HSCT:

- ≥ 12 air exchanges per hour;
- central or point-of-use HEPA filters with 99.97% efficiency for removing particles ≥ 0.3 μm in diameter;
- correct filtration is particularly critical in HSCT centres with ongoing construction and renovation;
- continuous pressure monitoring, especially while rooms are occupied;
- self-closing doors to maintain constant pressure differentials of anterooms should be used to ensure appropriate air balance;
- floor surfaces should be smooth, nonporous, and scrubable to minimize dust levels;
- HSCT recipients should be placed in single-patient rooms, if possible. If the availability of single-patient rooms is limited, their use should be prioritized for the most severely immunosuppressed patients;
- hand hygiene includes both use of alcohol-based hand rubs and hand washing with soap (plain or antimicrobial) and water;
- equipment and devices should be cleaned, disinfected or sterilized, and maintained as directed by established guidelines;
- it is recommend that plants and dried or fresh flowers should not be allowed in hospital rooms during conditioning or after HSCT because Aspergillus species have been isolated from the soil of potted ornamental plants, the surface of dried flower arrangements, and fresh flowers.

The above recommendations were compiled according to Guidelines for Preventing Infectious Complications among Hematopoietic Cell Transplantation Recipients [33].

Visitors and their influence on the patient’s mental state

The problem of attending personnel or visitors entering the cleanrooms remains a question. Patients are allowed visitors directly in the room only in exceptional cases, where their health and circumstances permit. Both visitors and the attending personnel are obliged to wear personal protective equipment and aids (mouthpieces, gloves, disposable lab coats, etc.) and disinfect their hands. These are precautions against spreading the infectious agent. The cleanroom units provide detailed instructions for visitors (if permitted) on how to minimize the risk of transferring an infectious agent. Some types of isolation rooms have galleries built around these rooms for visitors.

Persons with infections of the respiratory tract, flu-like illnesses, etc. are excluded from visiting. In some foreign countries, children are allowed to visit cleanrooms. They must be paid increased attention, however, as there is a possibility of the transfer of infectious diseases [28].

The role of visitors is indispensable – mainly psychologically. Patients may feel isolated and lonely and, likewise, they may believe (as they are placed in an isolation room) that
the attending personnel visit them less often than the other rooms. If this is the case, the patient needs to be explained the necessity of the minimization of the possibility of transferring an infectious agent, and, above all, informed that a lower frequency of the personnel’s entrance does not mean lower or impaired health care.

Within the current health care of immunosuppressed patients, cleanrooms hold a unique position in minimizing the risk of a hospital-acquired exogenous infection and in the recovery of these patients.

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