Mould Contamination Experimental Study in Public Building Environment

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Abstract. This paper is aimed at the experimental study on mould contamination status. Mold contamination has seriously threatened residents' physical and mental health and buildings. However, few experimental studies on mold contamination were made in the built environment under hot and humid climate. Then, we cultured the sample in 3 sampling points by the five-point sampling method in a classroom of USC. Then, we cultured the sample in 28 ℃ constant temperature condition, counted mould number with plate counting method and referred to the Omeilianski formula to calculate indoor mould concentration. The study found that the average concentration of mould spore in the typical public building was 629cfu/m², and the exceeding rate was up to 25.8%. The results show that the typical public building environment in Hengyang has serious mould contamination, thus further environmental optimization research is urgently needed.

1 Introduction

With the rapid development of China's construction industry in recent years, building environment problems have become increasingly prominent, among which mould contamination is a key problem, especially in the humid and hot climate areas in south China. The humid and hot climate in the south provides favorable conditions for mould breeding.

Mould contamination in building environment has a detrimental effect to buildings and residents. Mould can make metamorphism, degradation, decomposition, corrosion and crack to building materials, which disable their original function, produce the mildew on the wall and cause the external wall surface of the building to bulge and its insulation material to fall off[1,2]. The service life and aesthetics of the wall are seriously affected. Mould growth can cause Sick Building Syndrome (SBS), which leads to high renovation costs [3-4]. Long-term exposure to mould environment can easily cause respiratory diseases and allergic symptoms, such as bronchitis, asthma and allergic pneumonia. For the elderly and children with low immunity, mould may also cause headache, fever, inflammation of skin or mucous membrane and other diseases [5]. The mycotoxin produced by mould in the process of metabolism is more harmful to human body and damages human nervous and immune system.

According to the World Health Organization (WHO), more than 80% of the world's residences have molds in varying degrees. Due to the influence of many factors such as climatic conditions, building quality and indoor environment, mould contamination in Chinese residential buildings is particularly serious. Wei [5] found in their indoor mould investigation in Shanghai that the housing with mould accounted for 56.3%, and the housing with serious mould contamination accounted for 16.9%. Liu [6] conducted a questionnaire survey on 15,266 students and found that the incidence of visible mould and mouldy odor in Shanghai residential buildings was 7.9% and 11.9% respectively. Yuan [7] investigated the indoor mould contamination status of 16 buildings, 12 bungalow houses and 6 office buildings in Dongzhimen district, Beijing, in which indoors mould contamination was found to varying degrees, and the mould contamination in summer was higher than that in winter. Li [8] investigated the atmospheric microbial contamination in Nanning city and found that the range of atmospheric microbial content in the urban area was 1101-29094 cfu/m³, among which mould accounted for 20.8%.Lin[9]investigated the air microbial contamination in urban areas of Jiaxing city and found that the counts of bacteria and mould in the air were 568-4630cfu/m³ and 507-5347cfu/m³, accounting for 47.4% and 52.6% of the total number of airborne microorganism, respectively.

In this paper, the typical public buildings in Hengyang, which belongs to the typical hot and humid climate area, were studied by experimental test. The results have a good reference to the mould research in other humid and hot climate areas.

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2 Mould test in building environment

We selected a classroom on the first floor of the fifth teaching building in USC, Hengyang as the research object. The fifth teaching building is a brick and concrete structure, six stories high, which has existed for 20 years. It is orientated south and north, shaded by other buildings and greenery, and has good ventilation. Mould contamination in construction is positively correlated with the concentration of mould spore in the air, thus we sampled indoor mould spores to evaluate indoor mould contamination [10,11]. At the same time, we recorded the related factors of building characteristics and indoor environment. The actual measured time was December, 2018.

SAB medium was prepared with reference to Examination Methods for Public Places with Chinese standards (EMPP)(GB/T 18204.4-2013) [12]. The main components were: 0.5L distilled water, 5g peptone, 20g glucose, 10g agar, and 0.05g chloromycetin. We mixed the above ingredients together even after adjusting the pH to 5.6±0.2 and bottled under 115 ℃ high pressure sterilization for 15 minutes. After being cooled to an appropriate temperature, the culture medium was poured into a glass plate with a diameter of 90mm according to 20ml-25ml per plate. After the agar in the culture medium was solidified, it was put into a culture box for 24 hours to observe whether there were bacterial colonies. If there were bacterial colonies, it was discarded.

We used the natural sedimentation method to test the number of mould spores in indoor air. According to the five-point sampling method, we set five sampling points. Each sampling point is 1.2m to 1.5m away from the ground and more than 1m away from the wall. The mediums were exposed to the environment for 5min, and a group of medium was randomly placed with parafilm as control group. After sampling, we covered the lid, inverted the plate and sealed it with parafilm to prevent contamination of bacteria during culture. After the sealed plate was placed in a constant temperature environment of 28 ℃ for 72 hours, the mold in the plate was counted. Since temperature and humidity have a major impact on mold growth and spore release, it is necessary to record changes in indoor temperature and humidity. The temperature and humidity recorder was placed at a height close to the sampling point while sampling the mould. The automatic recorder recording interval is 30min, and the recording time is 24h after the start of sampling.

The matrices in the 6 petri dishes were taken from the same batch of prepared medium, and the contents of each petri dish were consistent. The 6 petri dishes were disinfected and sterilized simultaneously while taken out, and were randomly numbered. The counting method was in accordance with the plate counting method in GB/T 18204.4-2013 EMPP [12]. The results of mould counting in petri dishes were shown in table 1. The real-time growth of mould in each petri dish was shown in the table:

| Time   | No.1 | No.2 | No.3 | No.4 | No.5 | Control group |
|--------|------|------|------|------|------|---------------|
| Begins | 0    | 0    | 0    | 0    | 0    | 0             |
| 12 h   | 0    | 0    | 0    | 0    | 0    | 0             |
| 24 h   | 0    | 0    | 0    | 0    | 0    | 0             |
| 48 h   | 6    | 5    | 3    | 1    | 4    | 0             |
| 72h    | 10   | 12   | 7    | 6    | 13   | 0             |

The petri dish was incubated at a constant temperature for 24–48 hours. At the 48th hour, petri dish No. 1 was observed to have the most colonies, No. 3 dish and No. 4 dish had few colonies, and No. 4 dish had a single large white trichome colony, but other colonies in NO. 4 dish were small. After 72 hours, 10 colonies were observed in No.1 petri dish; 12 in No.2 petri dish; 7 in No.3 petri dish , 6 in No.4 petri dish , and 13 in No.5 petri dish. No mould growth was observed in the control group. There was no significant difference in mould in the petri dishes except that No.4 had a large colony. We suspected that the sampling points No.2 and No.5 were...
close to the corner of the wall, and the wall is seriously polluted by mould. Therefore, the results show that No.2 and No.5 have a large number of mould colonies. No.3 and No.4 sampling points are close to the corridor and ventilated, and the results show that No.3 and No.4 have a small number of mould colonies. The distribution of mould in No.5 dish was shown in figure 2.

Fig 2. Distribution of mould in petri dish No. 5

According to Omeilianski's suggestion, the total number of microbes falling into 100cm² plates within 5min is equal to the total number of microbes in 10L of air. The indoor mould concentration was calculated by the Omeilanski formula, as shown in the following formula.

\[ C = N \times \frac{100}{A} \times \frac{5}{t} \times \frac{1000}{10} = 50000N \times \frac{A}{t} \]

In this formula: \( C \) is the microbial concentration in the environment, cfu/m³; \( N \) is the average number of colonies in the dish, number per dish; \( A \) is the area of the dish, cm²; \( t \) is sampling time, min.

By substituting the experimental results into the above formula, we obtained the average mold concentration in the measured house as 629 cfu/m³. WHO suggested that 500cfu/m³ should be taken as the threshold value of indoor mould spore concentration to determine whether there is mould contamination. This experiment detected that the total number of mould colonies in typical public building environment in Hengyang exceeded the standard rate by 25.8%, indicating that there is serious mould contamination in the building environment.

4 Conclusion

The SAB medium was prepared with reference to EMPP (GB/T 18204.4-2013), and a classroom on the first floor in USC was selected as the sample room. The sampling point was set by five-point sampling method, and the sampling time was 5min. Then, the sample was plated in a 28°C temperature condition. The counting method was in accordance with the plate counting method. After 72h, we found that No.2 and No.5 medium had a small number of mould colonies. It may be that sampling points No.2 and No.5 are close to the corner of the wall, where has serious mould contamination, while sampling points No.3 and No.4 are close to the corridor and ventilated, so the number of mould colonies is small. This fully proves that there is a positive correlation between construction mould contamination and the concentration of mould spore in the air.

In this paper, mould spore concentration in the air of typical public buildings in Hengyang was tested experimentally, so as to study mould contamination in the building environment. The indoor mould concentration was calculated according to the Omeilanski formula, and the measured results show that the average mould spore concentration is 629cfu/m³, which is higher than the threshold of spore concentration. This indicates that there is serious mould contamination in the sampling site. The results can be used as a reference for the study of mould contamination in building environment in hot and humid climate.

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