Characterizations and Analysis of the Mold from Animal Specimens in Shenzhen Museum

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Abstract

Biological specimens play an important role in cultural exchange, science popularization, scientific research and economic window, but the preparation and preservation technology system of biological specimens is relatively unsafe and inefficient. Mold grows seriously on animal specimens, which is not only harmful to human beings’ health and environment, but also is one of the factors that restricts the development of the natural history museums where these specimens are kept. This paper identified the mold species of animal specimens by PCR with ITS primers, bio-microscopic observation, sequencing and phylogenetic tree analysis. The results showed the mold of animal specimens mainly belonged to Aspergillus and Neurospora. This study established the foundations of controlling and restoring the mold that infected animal specimens and guided a new methodology of preparation and environmental friendly exhibition for animal specimens.

Keywords

Animal Specimens, Mold Identification, PCR, Sequencing, Phylogenetic Tree Analysis

1. Introduction

1.1. Beautiful Appearance but Problems inside

Biological specimens display beautiful appearances [Figure 1], but there are serious potential problems hidden

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Biological specimens display the very beautiful appearance. Exposure to hot and humid environments has damaged specimens. This environment has been especially hard on animals that have been taxidermied because they have external skin, fur, internal fat, protein, connective tissue, etc. Infection by mold spores and hyphae is an acute problem to animal specimens, which have been associated with many human beings' serious health problems and environmental safety problems. It would also lead to great loss of specimens in natural history collections. Because animal specimens are more precious than any other forms of biological specimens such as plants and other specimens. They are very expensive, so it would be a huge economic loss if they were damaged. The climate of South China is hot and humid which lasts for about 8 - 10 months in one year. In Guangdong province, the annual average relative humidity is 65% - 85%. Especially during the mold rainy season, the relative humidity often reached more than 90%, sometimes even was 100%.

At present, there are many reports on the ancient cultural relics such as paper, ancient lacquer painting, silk cotton fabric, leather, ivory, bamboo, jade, pottery and porcelain, and glass [1]-[4]. Recently there have been a few reports on the mold that infected animal specimens.

So, there are great challenges and difficulties in the most important techniques for sample collection, preparation and storage of animal specimens.

1.2. Current Situation: The Harmfulness of Chemical Mold Proof Reagents Used in Preparation of Animal Specimens

At present, there are many poisonous reagents used in preparation of animal specimens, such as arsenic trioxide (As$_2$O$_3$), Camphor, Phenol, Oxymethylene, and so on. It is harmful to people’s health and the environment of exhibition, such as conservation workers, and the general public. Mold is the most important role to damage the animal specimens [Figure 2].

1.3. Mold Damage: The Harmfulness of Animal Specimens Mold to Human Beings

The incidence and prevalence of serious mycoses continue to be a public health problem.
There are many poisonous reagents used in preparation of animal specimens. Thousands of species of microscopic MOLD are common in the environment, including natural history specimens. The species involved Aspergillus, Penicillium, Geotrichum, Alternaria, Cladosporium, Mucor, Rhizopus, Trichoderma, Fusarium which are associated mostly with allergic response of different types. At first, mold is harmful to human beings. Microbiological contamination with mold, It not only can pose a significant health hazard to those working in archives or museums [5], but also could lead to many serious illness. The nose and paranasal sinuses almost always harbor numerous mold species. Amongst the species recovered, many have the potential to produce mycotoxins including Aspergillus (flavus, niger, fumigatus, versicolor), Chaetomium, Fusarium, Penicillium and Trichoderma. This group also found “mold elements (hyphae, destroyed hyphae, conidia and spores)”. Mold and eosinophilic mucin are the markers of sinus involvement in the CRS patients [6]. Chronic Rhinosinusitis (CRS), A primary concern was that exposure to high concentrations of Aspergillus spores may cause respiratory illness, particularly in individuals that are immunocompromised, it also led to endocrinopathy with growth hormone deficiency; Dennis-Robertson syndrome [7]-[9]. Molds and mycotoxins cause neurobehavioral and pulmonary impairment as well as adverse health effects especially with allergies and asthma [10]-[19]. So, it is very important to characterize and analyze the mold of animal specimens.

Shenzhen Museum was established in 17th Oct, 1981. The area of structure is 32,000 square meter. By the end of 2015. There are collected more than 35,000 different kinds of cultural relics, including about 25,000 of marine animals and wild animal specimens and fossils collected. The specimens of biological fossils dated from 100 million years, ancient history and art treasures, showing 5000 years of civilization of China and important historic materials about the development history of modern and contemporary Shenzhen.

2. Materials and Methods

2.1. Experimental Materials

The mold samples were collected from animal specimens of Shenzhen museum during December 2013-March 2014. These specimens include porcupine, birds, turtles, antelope and deer. Because of the mould rainy day, there are plenty of mold exposed on them (Figure 3).

2.2. Methods

2.2.1. Mold Cultured and Isolated and Microscopes Observation

PDA medium was used to culture and isolate the collected mold. Identification of the mold was confirmed by culturing swabs from infected specimens and examining mold hyphae with both light bio-microscopes. The swabs were immersed and homogenized in a sterile physiological solution and serial dilutions were made. Each dilution was inoculated (0.1 ml) on MEA. After 2 days of cultivation at 25°C, identification of the mold was performed. After 3 days of fostering, seemingly different mold colony grew on 12 plates. A small sample from the middle of the colony was dug and transferred to new plate to culture the different colonies. Strains that were previous activated had a small amount of sterile water with the pipettor added. The mold was then extracted from the solution and inoculated on a slant medium. This was repeated.
Then slightly digged a small piece of colony from the center of the same kind. Mold were transferred to a new plate one by one until separating into single and stable colonies plates. Take strains previously activated and kept in the AGAR slant tube, added some amount (bacteriostatic ring according to the experimental plate number) of sterile water with the pipettor on the clean work platform, then moved the pipettor under the liquid level, absorbed a little liquid and gently hit out, beat the mold on the inclined medium, let more mold into the suspension, make of the mold suspension.

The mold was observed through bio-microscopic observation. A small amount mold hyphae was dropped onto the water slide to make a suspension. At this point, the class slide was turned 45 degrees slowly to cover the mold to make observations of the bubbles. The slide was put on the staged and to observe the structure of the mold. Computer software was then used to characterize the mold.

Cultural and micromorphological characteristics of mold colonies observation and identification was performed through using identification ways.

The collection consisted of deteriorated animal specimens, including 12 species of animal specimens, 1) Porcupine, 2) wood, 3) bird, 4) turtle, 5) antelope, 6) deer 1, 7) deer 2, 8) wale, 9) Chimpanzee, 10) Raja porosa.

2.2.2. PCR Detection

The isolated single colonies and ITS primers were used to do PCR (S1000 Thermal Cycler BioRad, USA). Primers were ITS1 and ITS4. The mold DNA was amplified using the highly conserved fungal rRNA gene primers (ITS1F and ITS4) as White and Gardes previously described [20]-[22]. The sequence of ITS1F and ITS4 are

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\text{ITS1F} \quad 5'\text{-CTTGGTGCATTTAGAGGAAGTAA-3'}, \quad \text{ITS4} \quad 5'\text{-TCCTCCGCTTATTGATATGC-3'}.
\]

PCR reaction conditions was 94°C predenatured for 5 min, 98°C denatured for 30 s, 58°C annealed for 30s, 68°C extended for 2 - 3 min. After the total 27 cycles. They were detect by agarose gel electrophoresis.

2.2.3. Sequencing and Phylogenetic Tree Analysis

The isolated single colonies were sent to Shanghai Shenggong Company, Ltd to be sequenced. And the results were analyzed. Distance-matrix sequence alignment methods used Clustal W. Maximum Likelihood (ML) phylogenetic tree for the fungal Operational Taxonomic Units (OTUs) constructed based on fungal ITS gene sequences obtained using fungal specific primer set ITS1F/ITS4. These sequences resulted in 8 fungal Operational Taxonomic Units (OTUs) with ITS and 19 OTUs with 18S rDNA primer sets respectively by taking into account the 2% sequence divergence cut-off for species delineation. Topology was built using Mega v.5.03 from a ClustalW 1.83 alignment. Numbers below branches indicate bootstrap values (>50%) from 1000 replicates. New sequence types are marked with triangle. B: Proportional distribution of different fungal taxa in the clone library constructed with fungal specific primer set ITS1F/ITS4.

3. Results and Discussion

3.1. Mold Microscopic Observation Results

Identification of the mold was confirmed by culturing swabs from infected specimens and examining mold hyphae both light bio-microscopes. Mold microscopic observation of 12 mold infected species were identified.

The examined animal specimens in Shen Zhen museum showed clear signs of biodeterioration. Superficial colonies of mold were clearly visible and abundant. A total of 12 mold species were identified from all analyzed samples with 12 taxa identified (Figure 4).

The mold hyphae was detected used 40 fold light bio-microscopes. It need to improve the fold of bio-microscopes. It would have better observation results.
3.2. Single Colony-PCR Detection

Because the mold collected from the museum was mixed mold samples, they were still unable to determine the type and the single degree, so PCR was used for species identification. The isolated single colonies were detected by PCR primers ITS1 and ITS4. The results showed that the DNA bands were right (Figure 5).

Twelve animal species mold samples DNA were amplified through colony PCR amplification. Then it were done the species identification. The results proved that the colony PCR experiments could be successfully used for species identification.

Mold genetic materials were amplified in colony PCR expansion test and species identification was performed. The results proved that the colony PCR experiments could be successfully used for species identification.

Figure 5 showed the electrophoresis bands after the PCR amplification of 1 - 12 strains. It could be seen the bright and clear 12 bands from the figure, Colony PCR experiment successfully copied the genetic material of 12 strains.

Colony PCR constructed based on fungal ITS gene sequences obtained using fungal specific primer set ITS1F/ITS4. But the general primers sometimes had some limited. It would have better PCR results after screening advanced specific primers [23].

3.3. Sequenced and Analyzed, Phylogenetic Tree Analysis

The single isolated colony was sent to Shanghai Shenggong Company, Ltd to be sequenced. The results were analyzed. Distance-matrix sequence alignment methods used Clustal W (Figure 6). The results showed the mold from animal specimens mainly belong to Aspergillus and Neurospora. The results were benefit for different novel mold inhibitor screening according to the different types of mold that molecular species identification.

Phylogenetic tree for the fungal Operational Taxonomic Units (OTUs) constructed based on fungal ITS gene sequences obtained using fungal specific primer set ITS1F/ITS4. These sequences resulted just in fungal Operational Taxonomic Units (OTUs) with ITS and 19 OTUs with 18S rDNA primers that with different fungal taxa in the clone library constructed with fungal specific primer set ITS1F/ITS4. But fungal strains used for specificity test of new 28S rDNA primers would be more significant results [24]-[26].
Figure 5. Colony-PCR. ITS primer be used to identify of mold samples. 1-12 mold samples collected from 12 species. 1) Porcupine, 2) wood, 3) bird, 4) turtle, 5) antelope, 6) deer 1, 7) deer 2, 8) wall, 9) chimpanzee, 10) Raja porosa. M: Marker, Del2000. 1-12 was the mold samples PCR that was collected from animal specimens. This figure was the electrophoresis stripes after the PCR amplification of 1-12 strains, could see the bright and clear 12 stripes from the figure, colony PCR experiment successfully copied the genetic material of 12 strains.

Figure 6. Mold genetic sequenced and analysed, phylogenetic tree analysis. The results showed the prevailing mold species isolated from animal specimens was *Aspergillus niger* Tiegh (62.5%), followed by *Neurospora crassa* Shear & B. O. Dodge (25%) The signal colony that isolated sent to Shanghai Shenggong Company, Ltd. to be sequenced. And the results was analysed. Distance-matrix sequence alignment methods was used Clustal W.
4. Conclusion

All in all, after identifying the mold species from animal specimens, by PCR with ITS primer, bio-microscopic observation and sequencing and phylogenetic tree analysis, the results showed the mold from animal specimens mainly belonged to *Aspergillus* and *Neurospora*. This study established the foundations of controlling and restoring the mold that infected animal specimens and guided a new methodology of preparation and environmental friendly exhibition for animal specimens. It was benefit for screening different novel mold inhibitor according to the different types of mold.

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