Molecular characterization of Aspergillus infections in an Iranian educational hospital using RAPD-PCR method

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ABSTRACT
Objective(s): The nosocomial infections by Aspergillus species are associated with constructions and increased dust loads in hospital indoors. Our main object was to find the environmental sources of Aspergillus species causing hospital acquired infections.

Materials and Methods: The clinical and environmental samplings were performed during 18 months from spring 2010 to summer 2011 in Imam educational hospital, Urmia, Iran. A morphological diagnosis was performed including microscopic characterization of isolated aspergillus from cultured specimens and polymerase chain reaction - restriction fragment length polymorphism (PCR-RFLP) for the identification in the level of species. Random amplified polymorphic DNA - PCR RAPD-PCR using random primers for rDNA gene was performed to compare Aspergillus isolates of clinical cases with the relevant environmental sources.

Results: Use of RAPD method resulted various differential patterns, so that some Aspergillus isolates from the clinical and hospital indoor were completely matched (matched pairs) and some other Aspergillus isolates were not matched. In the case of matched pairs, Aspergillus niger and A. flavus isolated from broncoalveolar lavage and sinus discharge were relevant to those of air conditioner and walls surfaces, respectively.

Conclusion: The hospital sources for the Aspergillus clinical isolates included air condition and walls. RAPD-PCR analysis can play a trivial role to find the hospital sources of Aspergillus clinical isolates.

Introduction
The incidence of life threatening invasive aspergillosis has been developed following the increased immune compromised cases, bone or organ transplant receivers and patients with leukemia, lymphoma and other malignancies (1). Incidence rate of invasive Aspergillus infections is about 17 to 26%, 5 to 24% and 5 to 15% in lung transplant, acute leukemia and allogenic bone marrow transplant patients, respectively. Aspergillus fumigatus is the most common species isolated from invasive Aspergillus infections, although Aspergillus flavus, Aspergillus terreus, Aspergillus niger and other species also frequently cause invasive aspergillosis (1). Hospital acquired aspergillosis is known to be an airborne infection due to constructions and renovations which make the higher rates of airborne conidia. Detailed studies have found Aspergillus spores in many parts of hospital, including shower heads, dusty air conditioners, hospital plants, clothing and other sources (1). Small outbreaks of nosocomial aspergillosis have been reported, sometimes in association with environmental sources of conidia. Elucidation of the complex epidemiology in such cases requires detailed molecular typing studies. However, not all of the molecular procedures are applicable to fungal typing (2). Recently, a study of A. terreus infections in immunosuppressed patients showed that environmental and some patient isolates had identical band patterns by RAPD analysis (3).

A recent study on the immune suppressed patients at the nephrology and transplant ward of Imam University Hospital, Urmia, Iran showed 36 (66.6%) and 17 (31.4%) frequencies of Candida and Aspergillus species (respectively) isolated from clinical specimens. Moreover, 110 fungal isolates of Candida, Saccharomyces, Aspergillus, Penicillium, Alternaria and Cladosporium species were obtained from hospital indoor and outdoor. Among all, 35 (31.5%) Candida and 48 (43.2%) Aspergillus species were detected. Regarding high frequently of

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Aspergillus spp. isolated from clinical and environmental sites in the nephrology ward, the authors tried to perform a molecular epidemiologic study to find accurate and exact environmental sources for Aspergillus infections and colonization in the large, general hospitals. Other studies have used molecular methods for the identification and epidemiological aims on pathogenic bacteria and viruses including Mycobacteria (4, 5), Escherichia coli (6) and Rotavirus (7). Present study uses the molecular method of random amplification of polymorphic DNA (RAPD) to study the environmental sources of Aspergillus species which recently isolated from all cases with hospital acquired infections (HAIs) or colonization.

Materials and Methods

Study subjects
Clinical and environmental samples were obtained during 18 months from spring 2010 to summer 2011 at the large, training hospital, Imam Khomeini. The clinical specimens including sputum, bronco alveolar lavage (BAL), urine, sinus discharge, skin and nail touching on the surfaces and wall splits. Air samples were collected from various places close to patient’s bed carpets and meadows (8). The diagnosis was then confirmed by molecular identification using restriction fragment length polymorphism (RFLP) for the epidemiological linkage among Aspergillus species. The method was performed for both clinical and environmental isolates of species. For the restriction enzyme digestion, 13 µl of each PCR product was digested by 5 U (0.5 µl) of the restriction enzyme MwoI at 37°C enabled us to differentiate most of Aspergillus isolates in level of species. The method was performed for both clinical and environmental isolates. For the restriction enzyme digestion, 13 µl of each PCR product was digested by 5 U (0.5 µl) of the restriction enzyme MwoI, 1.5 µl of the enzyme buffer, and then incubated at 37°C for 180 min (10). Digested PCR products were subjected to a 2% agarose gel electrophoresis and visualized in trans illuminator (Gel Doc system). Aspergillus isolates were identified comparing the electrophoretic DNA patterns with standard measures (Table 1).

Culture and Identification
All specimens were transported to Medical Mycology Center, Urmia Medical Sciences University (UMSU). A morphological diagnosis was performed using culture media, Czapekdox agar (CZA) and Sabouraud glucose agar (SGA) for the primary identification of isolated molds (8). The diagnosis was then confirmed by molecular identification using restriction fragment length polymorphism (RFLP) followed by random amplified polymorphic DNA (RAPD) for the epidemiological linkage among clinical and environmental sources.

DNA extraction
Aspergillus mycelial mass was harvested from the 12 to 24 hr fungal liquid cultures, filtered and purified. The genomic DNA was extracted by glass beads and phenol-chloroform method,(a solution of 1mM EDTA, 1% SDS, 100mM NaCl, 10 mM Tris-HCl and 2% Triton X-100, in distilled water, pH 8.0 was used as LysisBuffer). The extracted DNA was checked by using 1.5% agarose gel electrophoresis (3).

PCR for identification
The PCR assay was performed using 5 µl of the DNA template in a total reaction volume of 50 µl (consisting of PCR buffer [20 mMTris- HCl at pH 8.0], 50 mM KCl, 0.1 mM each of forward [ITS: 5'-TCC GTA GGT GAA CCT GCG G-3'] and reverse [ITS-4: 5'- TCC TCG GCT TAT TGA TAT GC-3'] primers for ITS regions of rDNA [purchased from Mirhendi Molecular Biology Lab, TUMS], and 1.5 U of Taq DNA polymerase). We used universal primers for the amplification of Aspergillus ITS regions (forward primer: 5'- TCC GTA GGT GAA CCT GCG G - 3', reverse primer: 5'- TCC TCG GCT TAT TGA TAT GC - 3') (9).The reactions were performed in a thermo cycler (XP Cycler, BIOER, China). Thermal program included an initial DNA denaturation at 95°C for 5 min that followed by 30 cycles, consisting of the stages; denaturation at 95°C for 30 sec, annealing at 55°C for 30 sec, and extension at 72°C for 1 min, with a final extension at 72°C for 5 min following the last cycle. The DNA fragments were length separated by electrophoresis through 1.5% agarose gels in Tris Borate EDTA (TBE) buffer and 0.50 mg of ethidium bromide per ml. Results were documented using a UV trans illuminator (SynGene SYD4/680X, UK) (10).

Digestion of PCR products using RFLP method
Digestion of amplified ITS fragments by using a novel restriction enzyme, MwoI at 37°C enabled us to differentiate most of Aspergillus isolates in level of species. The method was performed for both clinical and environmental isolates. For the restriction enzyme digestion, 13 µl of each PCR product was directly digested by 5 U (0.5 µl) of the restriction enzyme MwoI, 1.5 µl of the enzyme buffer, and then incubated at 37°C for 180 min (10). Digested PCR products were subjected to a 2% agarose gel electrophoresis and visualized in trans illuminator (Gel Doc system). Aspergillus isolates were identified comparing the electrophoretic DNA patterns with standard measures (Table 1).

Random amplification of polymorphic DNA (RAPD)
For the RAPD-PCR, six single primers which successfully tested before were used as random primers on Aspergillus genomic DNA. The primers: P1 (Rp4): 5'-CAGATGCTTC -3', P2 (Rp1): 5'-TAGGATCGGA -3', P3 (SOY): 5'-AGGTACCTGA -3', P4 (RP2): 5'-AAGGATACGA -3', P5 (R108): 5'-GATTGGCCCT -3', P6 (UBC90): 5'-GGGGTTAGG-3' were randomly selected from many available primers (Purchased from Molecular Biology Lab, TUMS). The primers were run into a PCR master mix containing 3 mM MgCl2, 200 pmol of each primer, and 5 ng of DNA, in final volume of 100 µl. The thermal protocol used in thermo cycle system included 5 min at 95°C, 45 cycles of 94°C for 45 sec,
Molecular epidemiology of Aspergillus using RAPD-PCR

35°C for 1 min, 72°C for 1.5 min and a final 72°C for 5 min (11). Agarose gel electrophoresis of the PCR products followed by a ethidium bromide staining showed various patterns making DNA bands which analyzed by using Image Master software (Gene Snap Tool, SynGene, version: 4.01.0, UK). Reproducibility of DNA patterns was demonstrated by the analysis of two to three Aspergillus subcultures. The RAPD-PCR patterns were compared between the clinical and environmental Aspergillus isolates for each case and the similarity of RAPD patterns was analyzed.

Results

Totally 454 samples including 198 clinical and 256 environmental specimens were collected. The experimental studies on the clinical specimens resulted total of 93(47%) fungal and bacterial isolates. Among all, 54 (58%) were fungal isolates which obtained from cases with symptomatic infection started after hospitalization. Fungi included Candida spp. 36 (66.6%), and Aspergillus spp. 17 (31.4%). Aspergillus isolates included A. flavus (47%), A. fumigatus (29.4%) and A. niger (23.6%) as the most frequent species. Our findings of Aspergillus morphologic identifications were confirmed by PCR-RFLP method. Also, 256 specimens were collected from the relevant environmental sites including beds, carpets, walls, trolleys, sinks, medical devices, air-condition systems and air samples. Morphologic examinations resulted to 110 fungal isolates including Candida, 35(31.5%) and Aspergillus, 48(43.2%) species and other fungi (saprophytic molds), 20(20.3%) such as Alternaria, Saccharomyces, Mucorals, Penicillium, Cladosporidium and Pheohyphomycetes. Aspergillus isolates included A. niger (46.7%), A. flavus (41.8%), and A. fumigatus (14.7%) (Table 2). It means that five A. fumigatus isolates, eight A. flavus and four A. niger were obtained from the clinical specimens of cases with invasive or non-invasiveaspergillosis mostly in the intensive care units, nephrology and surgery wards. Also 7, 20 and 21 isolates of A. fumigatus, A. flavus, and A. niger (respectively) were detected in samples and other environmental specimens.

From 65 clinical and environmental isolates of Aspergillus species, 28 isolates were common between cases and their environments in the level of species. For making an exact correlation among the clinical and environmental isolates, the findings of RAPD-PCR were analyzed. The random primers, P1-P6 made different electrophoretic DNA patterns for most cases. For example, application of P1 resulted in different DNA patterns for the pairs (clinical and environmental isolates) 16, 36 and 37 from 65 cases of Aspergillosis. Just in one case (pair 31) same DNA patterns were observed (Figure 1c). Use of primer P2 resulted in identical patterns for three pairs: 16, 31 and 37 as shown in Figure 1d. At the same way, the other primers created common patterns not more than one. Comparison of clinical and environmental pairs with RAPD-PCR using six random primers revealed similar electrophoretic DNA patterns for clinical and environmental Aspergillus isolates (Figures 1a, b) just in two of 28 pairs including 32 and 45. A. niger and A. flavus were included in the pairs 32 and 45 respectively. The Aspergillus isolate of bronco alveolar lavage of case (32) was completely similar to that of relevant air conditioner used in the private room. Also, DNA pattern of A. flavus isolated from sinus discharge of case (45) was similar to the isolate of wall swabs. Other Aspergillus pairs showed no similarity (Table 3).

Table 1. Frequency of Aspergillus and other opportunistic fungi isolated from hospital indoor and outdoor

<table>
<thead>
<tr>
<th>Environmental specimen</th>
<th>Case</th>
<th>HCWs</th>
<th>Visitor</th>
<th>Carpet</th>
<th>Walls Bed &amp; blanket</th>
<th>Sink</th>
<th>Trolleys</th>
<th>Medical devices</th>
<th>Air</th>
<th>Air conditioner</th>
<th>Outdoor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Contaminants</td>
<td>Cae</td>
<td>HCW</td>
<td>Visitor</td>
<td>Carpet</td>
<td>Walls Bed &amp; blanket</td>
<td>Sink</td>
<td>Trolleys</td>
<td>Medical devices</td>
<td>Air</td>
<td>Air conditioner</td>
<td>Outdoor</td>
</tr>
<tr>
<td>Candida</td>
<td>1</td>
<td>5</td>
<td>1</td>
<td>2</td>
<td>4</td>
<td>3</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Aspergillus</td>
<td>3</td>
<td>1</td>
<td>0</td>
<td>9</td>
<td>7</td>
<td>7</td>
<td>2</td>
<td>5</td>
<td>1</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Other</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>4</td>
<td>6</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>13</td>
<td>6</td>
<td>1</td>
<td>14</td>
<td>16</td>
<td>15</td>
<td>6</td>
<td>10</td>
<td>7</td>
<td>9</td>
<td>4</td>
</tr>
</tbody>
</table>

HCWs: health care workers

Table 2. List of Aspergillus spp. Commonly isolated from cases and hospital indoor

<table>
<thead>
<tr>
<th>Aspergillus clinical isolates</th>
<th>Species</th>
<th>Clinical</th>
<th>Environmental</th>
</tr>
</thead>
<tbody>
<tr>
<td>KDH3</td>
<td>A. niger</td>
<td>BAL</td>
<td>carpet</td>
</tr>
<tr>
<td>KDH16</td>
<td>A. flavus</td>
<td>nail scraping</td>
<td>walls, carpet</td>
</tr>
<tr>
<td>KDH24</td>
<td>A. niger</td>
<td>sputum</td>
<td>carpet, walls</td>
</tr>
<tr>
<td>KDH27</td>
<td>A. flavus</td>
<td>sputum</td>
<td>air conditioner</td>
</tr>
<tr>
<td>KDH28</td>
<td>A. fumigatus</td>
<td>BAL</td>
<td>trolleys</td>
</tr>
<tr>
<td>KDH30</td>
<td>A. flavus</td>
<td>air</td>
<td>air</td>
</tr>
<tr>
<td>KDH31</td>
<td>A. fumigatus</td>
<td>BAL</td>
<td>carpet</td>
</tr>
<tr>
<td>KDH32</td>
<td>A. niger</td>
<td>BAL</td>
<td>air conditioner</td>
</tr>
<tr>
<td>KDH33</td>
<td>A. niger</td>
<td>BAL</td>
<td>carpet</td>
</tr>
<tr>
<td>KDH34</td>
<td>A. flavus</td>
<td>BAL</td>
<td>walls</td>
</tr>
<tr>
<td>KDH36</td>
<td>A. fumigatus</td>
<td>BAL</td>
<td>carpets</td>
</tr>
<tr>
<td>KDH38</td>
<td>A. flavus</td>
<td>BAL</td>
<td>beds</td>
</tr>
<tr>
<td>KDH45</td>
<td>A. flavus</td>
<td>sinus discharge</td>
<td>walls</td>
</tr>
<tr>
<td>KDH47</td>
<td>A. niger</td>
<td>sputum</td>
<td>trolleys</td>
</tr>
</tbody>
</table>

Table 3. Frequency of RAPD based correlation between the clinical and environmental Aspergillus isolates

<table>
<thead>
<tr>
<th>Species</th>
<th>Total C&amp;E by RFLP</th>
<th>Common species in C&amp;E</th>
<th>Same C&amp;E isolates by RAPD</th>
<th>Different C&amp;E by RAPD</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. fumigatus</td>
<td>12</td>
<td>4</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>A. flavus</td>
<td>28</td>
<td>6</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>A. niger</td>
<td>25</td>
<td>5</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>Total</td>
<td>65</td>
<td>28</td>
<td>2</td>
<td>18</td>
</tr>
</tbody>
</table>

C: clinical aspergillus isolates; E: environmental aspergillus isolates
**Discussion**

There is plenty of evidence supporting the role of opportunistic fungi as important agents of hospital acquired infections. During 1980 to 1990, Aspergillus spp. emerged as causing agents of life-threatening infections in immune compromised patients (12). Nosocomial aspergillosis still poses a significant clinical problem (1). Although PCR based methods for detection of fungi have been described recently (13, 8, 2), there is a problem of finding the real source of opportunistic fungi such as Aspergillus spp. causing hospital acquired infections. In the present study, the PCR based technique, RAPD-PCR was employed to find a correlation between the environmental and cases isolated Aspergillus. Before using RAPD method for epidemiologic surveys, it has to be evaluated the extent of the differences in DNA patterns that can be generated. The previous data verify that RAPD analysis is useful for fingerprinting *A. fumigatus* (9, 15), *A. flavus* (3, 17), *A. niger* (18) and *A. terreus* (17, 19).

Use of RAPD in the present study was based on the previous studies on discriminatory power of this method using the random primers among medically important Aspergillus species (17). Comparing the findings of RAPD on the environmental isolates with those which obtained from cases shows that some Aspergillus isolates match completely in DNA patterns. In fact, we considered a 100% similarity of RAPD patterns between two relevant Aspergillus isolates. Moreover, among all Aspergillus isolates of this study, only two cases were correlated with environmental sources. In these cases, *A. niger* and *A. flavus* were included as the Aspergillus species causing nosocomial aspergillosis which obtained from hospital indoor. *A. niger* was isolated from both specimens of bronco alveolar lavage and air conditioner in the patient room and *A. flavus* from sinus discharge and walls.

Likewise our findings showed that of more than 27% of Aspergillus species have no correlations with the clinical and environmental Aspergillus isolates. From all tested environmental samples, the carpet, walls and beds were the most contaminated surfaces by the above fungi. Isolation of aspergillus spore from air samples was not considerable in our study, although 5 aspergillus cases were obtained from the air of wards. The main route of Aspergillus nosocomial infection is inhalation of airborne spores by immune compromised patients and pneumonia is the result of lung tissue invasion in the patients (1). Because of their size, aspergillus spores remain suspended in air for a long time. Moreover, it is supposed that Aspergillus contamination of other sources was due to air contamination of hospital indoors. Building works or constructions are capable of liberating large numbers of airborne fungal spores into the environment (20).

Our results of Aspergillus species identification in hospital specimens is confirmed by other studies, so that in a similar study, *A. fumigatus* and *A. flavus* were the most frequently isolated Aspergillus spp. in

**Figure 1.** RAPD patterns of clinical and environmental for the strains 32 and 45 are similar using each tested primers (a, c). Application of P1 in RAPD resulted different patterns for the pairs of 16, 36 and 37 (b). P2 resulted same patterns for Aspergillus groups 16, 31 and 37 (d).
the patients with proven aspergillosis (8). Some studies reported that Aspergillus airborne spores have also been isolated from unfiltered air, ventilation system, and dusts displayed during hospital renovations and constructions (21). Although our findings of RAPD analysis showed a molecular correlation between some clinical Aspergillus isolates and their environmental sources as mentioned before, no such sources were detected upon detailed screening of the other wards; no same strains were detected in carpets, trolleys and air samples, and no aspergilli were grown from the samples taken from medical sets.

Conclusion
As a conclusion, RAPD-PCR can be applied as a simple, rapid, and useful method, but it plays a trivial role in finding the hospital sources of Aspergillus clinical isolates. It should be noted, however, that selecting the random primers is an important point to find the highest level of molecular similarity.

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