MOLECULAR CHARACTERIZATION OF MYCOBACTERIAL ISOLATED FROM HUMAN AND ANIMALS

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ABSTRACT

Sputum from clinically confirmed patient and lung tissues from tuberculin positive cattle were taken from el Menia governorate. The samples were subjected to mycobacterial isolation on Lowenstein Jensen media. Human samples gave 8 out of 20 positive culture with isolation rate of 40% where animal samples gave 11 out 15 positive culture with isolation rate of 73.3%. Biochemical identification revealed that all the human isolates belonging to M. tuberculosis where animal isolates where exclusively M. bovis. Molecular characterization using PCR utilizing primers set amplifying IS6110 gene revealed that all human and animal isolated were positive giving amplicon with relative molecular size of 254bp. When using primer set for detecting of the RD8 region, only human isolates gave a clear amplicon migrating about 673 bp which confirms that these isolates were belonging to M. tuberculosis.

Keywords: M. bovis; M. tuberculosis; RD8; IS6110; mycobacteria isolation

INTRODUCTION

Mycobacteria can infect and harbor a wide range of living hosts including both warm and cold blooded livings. Due to the close contact between animals and human especially in Upper Egypt, we have investigated the distribution of both M. bovis and M. tuberculosis public health. Bovine tuberculosis is caused by intracellular infection with the acid-fast bacterium, Mycobacterium bovis. In cattle, exposure to this organism can result in a chronic disease that jeopardises animal welfare and productivity, and in some countries leads to significant economic losses (Pollock and Neill, 2002). When disease develops, the associated granulomatous pathological changes are seen mainly in the lower and upper respiratory tract (Neill et al., 1994) and, because of this pattern, it is considered that infection most often follows aerosol exposure to M. bovis (Neill et al., 2001). Infected cattle are an important source of infection for other cattle and even human especially if they under stress or immune-compromised (Phillips et al., 2003). M. bovis is a member of the tuberculosis complex of organisms and represents a serious zoonotic risk to public health concerns, there is an urgent need for improved methods to combat bovine tuberculosis. Knowledge of bovine immune responses to M. bovis can be utilised to develop new methods for disease diagnosis. Bovine tuberculosis infection in cattle is usually diagnosed in the live animal on the basis of delayed hypersensitivity reactions. Delayed hypersensitivity test is the standard method for detection of bovine tuberculosis. The comparative intradermal tuberculin test with bovine and avian tuberculin is used mainly to differentiate between animals infected with M. bovis and those sensitized to tuberculin due to exposure to other mycobacteria (environmental) or related genera. This sensitization can be attributed to the antigenic cross-reactivity among mycobacterial species and related genera. This study was undertaken for isolation, identification and molecular characterization of Mycobacteria from human and animal cases in Menia governorate.

MATERIAL and METHODS

Mycobacteria isolation and identification:

Animal samples:
Lungs showing P/M lesions from tuberculin positive cross breed dairy cattle \((n=15)\) were collected from Menia Governorate. The samples were spliced into small sections 3 mm thick and transferred to a sterile mortar containing washed sterile sand. Two ml of sterile distilled water were added to the crushed tissue, homogenized and ground till suspension was obtained. Two ml of 4% \(\text{H}_2\text{SO}_4\) acid were added to the mixture, and then incubated at 37 \(^\circ\)C for 30 minutes. The mixture was diluted with 16 ml of sterile distilled water and centrifuged at 3000rpm for 20 minutes. The supernatant fluid was poured off into disinfectant (5% phenol solution) and the obtained sediment was inoculated into two pyruvated Lowenstein – Jensen slants. Then incubated at 37 \(^\circ\)C in inclined position for overnight, then vertically for at least 6-8 weeks and examined daily over a week period then once a week. The obtained growths were observed for morphological character and for pigment production. Those gave acid fast bacilli were further cultured in Middlebrook 7H9 liquid medium supplemented with albumin dextrose-catalase enrichment (Difco,) at 37\(^\circ\)C for 4-6 weeks.

**Biochemical identification:**

The growing animal and human isolates were subjected to morphological and biochemical identification using nitrate reduction test, niacin production test, Pyrazinamidase test, and growth on L.J. slants with either sodium Pyruvate or glycerol according to Kubica, 1973; and WHO, 1998.

**Polymerase chain reaction:**

**Extraction of genomic DNA from field isolates:**

According to Sambrook et al. (1989), and Soliman et al., 2004

One hundred mg (wet weight) of the cell pellet of each isolate was resuspended in 0.5ml TE buffer, allowed for 2 cycles freezing and thawing. The cells were then homogenized in glass homogenizer, and then incubated 4 hours at 37\(^\circ\)C with 5ul lysozymes (final concentration 100\(\mu\)g/ml). Proteinase-K was added 25\(\mu\)l/0.5 ml (final concentration 100\(\mu\)g/ml) and incubated for further 3 hours at 56\(^\circ\)C with shaking. DNA was then extracted using Trizol reagent according to the manufacture instruction. Briefly, one milliliter Trizol was added and after 5 minutes of incubation at room temperature (RT), 0.4 ml of chloroform was added, vortex for 15 seconds, kept at RT for 3 minutes. Then centrifuged at 14,000 rpm/10 min at 4\(^\circ\)C. The upper aqueous phase containing RNA was completely discarded and the DNA in the interphase was precipitated with 0.6 ml absolute ethanol and kept at RT for 3 minutes. Then centrifuged at 14,000 rpm/10 min at 4\(^\circ\)C. The upper aqueous phase containing RNA was completely discarded and the DNA in the interphase was precipitated with 0.6 ml absolute ethanol and kept at RT for 3 minutes before centrifugation at 4000 rpm/5 minutes at 4\(^\circ\)C. The supernatant was completely discarded and the DNA pellet was washed twice with 0.1 sodium citrate in 10% ethanol. At each time, the DNA pellet was kept in the washing solution for 30 minutes at RT with periodical mixing and centrifuged at 4000 rpm/5
minutes at 4°C. Following the 2 washes, the DNA was resuspended in 2ml of 75% ethanol, kept at RT for 20 minutes with periodical mixing and then centrifuged. The DNA pellet was finally dried briefly for 5 minutes under vacuum and redissolved in 50ul of 8 mM NaOH. The pH was then adjusted at 8 by adding 115 µl/ml of 0.1 M HEPES. Ten µl of RNAase were then added and incubated at 37°C for 1 hour then stored at -20°C till used. The concentration and purity was measured using Nanodrop ND-8000 Spectrophotometer (NanoDrop Technologies).

**PCR amplification of IS6110 and RD8 gene:**

Both human and animal isolated were identified using primers that target both IS6110 that identify any typical mycobacterial isolates and the RD8 gene that identify only the human isolated (Mycobacterium tuberculosis). The genomic DNA (50ng) was mixed with 10 µl of dream tag master mix, 10 Q-solution and 100 nM of either primer A or B (table 1). The mixture was placed in the thermal cycler (T3000 professional, Biometra Inc, Germany), and adjusted at 95°C/3 min for Initial denaturing then 40 cycles of denaturing at 95°C/1 min, annealing at 56°C/45 seconds and extension at 72°C/1 min. A final extension step at 72°C/10 minutes was included. The product was then visualized by agarose gel electrophoresis. The product size was analysed using syngene software with 100bp DNA ladder as a marker (Sambrook et al. 1989 and Soliman et al., 2004).

<table>
<thead>
<tr>
<th>gen</th>
<th>Primer name</th>
<th>sequence</th>
<th>Product length</th>
<th>Strain to be amplified</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>IS6110</td>
<td>INS-6110-F</td>
<td>CGTGAGGGGATCGAGGTGGC</td>
<td>254 bp</td>
<td>M. tuberculosis and M. bovis</td>
<td>Mehdikhani and Rokni 2012</td>
</tr>
<tr>
<td></td>
<td>INS-6110-R</td>
<td>GCGTAGGCCTCGGTGACAAA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RD8</td>
<td>RD8-F</td>
<td>GGTGTGATTTGTTGAGACGATG</td>
<td>678 bp</td>
<td>M. tuberculosis</td>
<td>Stephen et al., 1999</td>
</tr>
<tr>
<td></td>
<td>RD8-R</td>
<td>AGTTCCCTCTGACTAATCCAGG</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table (1) the primer sequence and the expected product size for the IS6110 and RD8 genes used in the current study.

**RESULTS**

Isolation and identification of animal isolates: As seen in Fig (1), out of 15 specimens taken from the slaughtered tuberculin positive cattle, 11 (73.3%) mycobacterium isolates were obtained on Lowenstein-Jensen media the biochemical identification revealed that all isolates were M. bovis. (Table 2)

![Fig (1) Lung show suspected tuberculous lesion (left) and L.J. media showing the growth of M. bovis after 2 months of incubation (right).](image)
Isolation and identification of human isolates.

Samples from patients confirmed to be infected with tuberculosis through X-ray examination of the chest were subjected to isolation of *M. tuberculosis*. Out of 20 samples used in this study, 8 samples (40%) yield positive colonies on L.J. media by the week 8; these colonies were very few and tiny in appearance. After about 12 weeks nearly all the 8 samples gave clear colonies which appear as crumbly, “bread-crumb rough”, waxy, non-pigmented (buff colored). All isolates confirmed to be *M. tuberculosis* using biochemical identification (Table 2)

Results of PCR:

Two sets of PCR primers were used to detect and differentiate between *M. bovis* and *M. tuberculosis*. All the 11 animal and 8 human isolates were tested with *IS 6110* that detect all Mycobacteria and then tested with RD8 primers sets that detect only *M. tuberculosis*. As seen in photo 3 and 4, all isolates were positive on using the IS 6110 primers, indicating that all the isolates are Mycobacteria in nature. On testing the isolates with the primer sets that detect the RD8 region however, only the human isolates gave a positive amplicons migrating about 650bp which corresponding to *M. tuberculosis*
Table (2): Results of physical and biochemical identification of the isolated acid fast bacilli from suspected human and animal samples.

<table>
<thead>
<tr>
<th>Test or property</th>
<th>M. bovis</th>
<th>M. tuberculosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of isolates</td>
<td>11</td>
<td>8</td>
</tr>
<tr>
<td>Ziehl – Neelsen stain</td>
<td>Acid fast bacilli</td>
<td>Acid fast bacilli</td>
</tr>
<tr>
<td>Growth at 28°C</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Growth at 37°C</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Growth at 42°C</td>
<td>±</td>
<td>±</td>
</tr>
<tr>
<td>Colony morphology</td>
<td>Smooth</td>
<td>Rough</td>
</tr>
<tr>
<td>Dark pigmentation</td>
<td>Non pigmented</td>
<td>Non pigmented</td>
</tr>
<tr>
<td>Photo pigmentation</td>
<td>Non pigmented</td>
<td>Non pigmented</td>
</tr>
<tr>
<td>Niacin production</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Nitrate reduction</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>Tween 80 hydrolysis 10days</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Catalase semiquantitative &lt;45</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Catalase PH 7.0-68c0</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Arylsulfatase (3 days)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Arylsulfatase (2 weeks)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Urease</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Iron uptake</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Growth on TCH (2 μg/ml)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Growth on NaCl 5%</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Photo (3) PCR amplification of the IS6110 of the animal isolates. Note that all the samples gave a positive amplicon migrating at about 250 bp except sample #9 which serves as NTC (animal samples).

Photo (4) PCR amplification of a region in the RD8 which is absent in M. bovis relative to M. tuberculosis (human samples)

DISCUSSION

Tuberculosis infection in man and animals remains one of the most devastating disease with high morbidity and mortalities with subsequent severe economical impacts especially in the developing countries like Egypt, which requires maintained focusing on the behavior of the disease within the subjects and establishment a strict control measures.

Isolation of Mycobacterium tuberculosis from biological samples is essential in molecular study, drug resistance survey and initiating treatment for cases suspected to developed drug resistance (Philip et al., 2012).

The total isolation rates of Mycobacteria from carcasses of reactors (tuberculin positive cattle) with lesions were summarized up in table (2). From a total of 15 cases 11, positive cultures were recovered with an isolation rate of 73.3 %. Other investigators reported, however, lower rates (Gallo et al., 1983, 29.1%,
Payeur and Marquardt, 1988, 8.9%). On the other hand, Chul Soon Choi (1981) in Korea reported a much higher isolation rate amounting to 92.1%. These results depend mainly on the actual disease status present in the tested herd and to some extend on the experience of the investigators as well as the technique used for decontamination of tissue specimens.

In human cases, the isolation rate was much lower (8 out of 20 with a rate of 40%) this lowered rate may attributed to the drug treatment in patients under investigation. one of the most critical factors that cases failure in Mycobacteria isolation from sputum, is the lower volume of the sample and the presence of large amount of commensals and other pathogens in the immune-compromised patients (Anargyros et al., 1990).

The resurgence of tuberculosis in man and animals around the world especially in developing countries has renewed interest in understanding the epidemiology and pathogenesis of this disease. One important advance of molecular techniques is that it allows the identification and tracking of individual strains of Mycobacteria with very clear molecular probes. This new discipline, the molecular epidemiology, emerged with the identification of IS6110, a novel mycobacterial insertion sequence which formed the basis of a reproducible genotyping technique for M. tuberculosis complex. Although other typing methods, appear to offer certain advantages in terms of reproducibility, economic cost, ease of execution and general applicability to clinical standards, IS6110-based typing remains the internationally accepted standard (Burgos and Pym, 2002) and continues to provide new insights into the epidemiology of M. tuberculosis. In the current study all the isolated strains both from human and animal hosts gave positive bands at the expected molecular size indication that these isolates were belonging to the M tuberculosis complex.

In order to differentiate between the human (M. tuberculosis) and animal isolates (M. bovis) a primer set was used that detect a conserved region in RD8 of M. tuberculosis that is completely absent in M. bovis. RD8 encompasses a region of 5895 bp on genome of the M. tuberculosis. The deletion contains six ORFs with a seventh, lpqQ, which codes for a truncated antigen termed lipoprotein, (truncated at the 5’ end by deletion)

Among these six ORFs, Rv3619c and Rv3620c code for members of the ESAT-6 and QILSS families (Harboe et al., 1996 and Cole et al., 1998), and two encode PE and PPE proteins antigens that is used by Mycobacteria for energy transfer. The last two ORFs, ephA and Rv3618, code for a putative epoxide hydrolase and a possible mono-oxygenase protein antigens respectively.

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