EARLY DIAGNOSIS OF TUBERCULOUS PERITONITIS BY NESTED PCR AND AUTOMATED CULTURE TECHNIQUE

Apurba Sankar Sastry, Sandhya Bhat K, Anand Sankar Sastry, Kumudavathi M S,

1. Assistant Professor, MD, DNB, Department of Microbiology, Meenakshi Medical College, Kanchipuram, Tamilnadu.
2. Assistant Professor, MD, DNB, Department of Microbiology, Meenakshi Medical College, Kanchipuram, Tamilnadu
3. Senior Resident, MD, Department of Medicine, Karpaga Vinayaga Medical College, Kanchipuram, Tamilnadu.
4. Tutor, D (Bact), Department of Microbiology, Meenakshi Medical College, Kanchipuram, Tamilnadu.

CORRESPONDING AUTHOR
Dr. Apurba Sankar Sastry,
Department of Microbiology,
Meenakshi Medical College and Research Institute,
Kanchipuram, Near Chennai, India
E-mail: drapurbasasstry@gmail.com
Ph: 0091 9444327314.

ABSTRACT: INTRODUCTION: Early laboratory diagnosis of tuberculous peritonitis (TBP) is crucial to start antitubercular chemotherapy and to prevent its complications. However conventional methods are either less sensitive or time consuming. Hence the diagnostic potential of BacT/ALERT and polymerase chain reaction (PCR) was evaluated in this study.

MATERIAL AND METHOD: The study group comprised of 21 cases and nine controls. The cases were divided into confirmed subgroup (seven cases) - smear or culture or histopathologically proven and probable subgroup (fourteen cases) - clinically suspected cases. Ziehl Neelsen (ZN), Auramine Phenol (AP) staining, Lowenstein Jensen (LJ) culture, BacT/ALERT and nested Polymerase chain reaction (PCR) targeting IS6110 were carried out on all the patients.

RESULTS: Sensitivity of ZN, AP staining and LJ culture were found to be 23.8%, 28.5% and 57.1% respectively. Whereas BacT/ALERT and nested PCR showed a sensitivity of 76.1% and 90.4% respectively. The mean detection time of growth by LJ culture was 32.10 days whereas that of BacT/ALERT was 21.28 days. The contamination rates in LJ culture and BacT/ALERT were 5.0% and 10.0% respectively.

CONCLUSION: Nested PCR and BacT/ALERT found to be more sensitive compared to LJ culture and smear microscopy. BacT/ALERT detects mycobacterial growth at a faster rate with less contamination rate compared to LJ culture. As both false negative and false positive results are reported on nested PCR, so alone it should not be used as a criterion for initiating or terminating the therapy but should be supported by clinical, radiological, cytological and other microbiological finding.

KEY WORDS – Tuberculous peritonitis, nested PCR, BacT/ALERT

INTRODUCTION: Tuberculous peritonitis (TBP) accounts for 10-12 % of all cases of extrapulmonary tuberculosis. Although it is uncommon in western world, it remains a serious problem in Asia especially in HIV positive patients. Laboratory methods play a crucial role in establishing the diagnosis, early starting and monitoring the chemotherapy, preventing the transmission, identifying the changing pattern of epidemiology and detection of resistance to drug. Diagnosis of TBP is often difficult due to paucibacillary nature of the disease. Though ZN staining is rapid and cheap but it is neither sensitive nor specific requiring a minimum bacillary load of $10^4$/ml. Culture is more sensitive than ZN staining with detection limit of 10-100
bacilli/ml but requires prolonged incubation time of 6-8 weeks because of the long generation time of tubercle bacilli (3). Even histology is also at times not conclusive.

Many newer diagnostic modalities like automated microbial detecting systems and molecular methods have come up with increase isolation rate and early detection of the organism. BacT/ALERT Microbial Detecting System is based upon colorimetric detection of pH change which occurs due to CO₂ production by Mycobacteria (3). Not much studies are there evaluating diagnostic potential of BacT/ALERT for tuberculosis (TB) especially for TBP. Polymerase chain reaction (PCR) is another alternate for the early diagnosis of TBP. IS6110 is a transposone present as multiple repetitive elements in the genome of M. tuberculosis complex with variable copy numbers ranging from 0 to 26 (3). Due its repetitive nature, tests using IS6110 as primer yield higher sensitivity. Nested polymerase chain reaction (PCR) using IS6110 though has been widely evaluated in the diagnosis of TB but fewer studies are only available for the diagnosis of TBP. Hence we have evaluated the potential of BacT/ALERT and nested PCR for the diagnosis of TBP.

MATERIAL AND METHODS: The present study was carried out in the department of Microbiology of a tertiary care centre and was approved by the institute ethical committee. CASES: Samples were collected from 21 cases of TBP admitted to the hospital, during July 2010 to September 2012 and divided to the following subgroups.

a) Confirmed TBP subgroup: comprised of seven case confirmed by smear, culture or histopathology.

b) Probable TBP subgroup: comprised of fourteen clinically suspected cases having features like pyrexia of unknown origin, abdominal tenderness, weight loss, night sweat and/or positive Mantoux test and/or elevated ESR and/or positive peritoneal fluid cytology like pleocytosis, elevated protein level and reduced sugar level and/or evidence of pulmonary or any other organ tuberculosis and patient improving on antitubercular drug (ATT) (Table 1).

CONTROLS: Nine peritoneal fluids were collected from known cases of bacterial peritonitis. Approximately 2ml of sample was collected aseptically. One ml of the sample was used for both smear and culture, 0.5 ml was used each for BacT/ALERT & PCR. Smears were made from the samples and were stained by both Ziehl–Neelsen technique and auramine phenol method as per the standard methods (3). Cultures were carried out on Lowenstein – Jensen (L-J) egg based medium as per standard method (4).

BACT/ALERT: The processing of peritoneal fluid was done as per as the manufacturer’s instruction (5). Inoculated BacT/ALERT MP bottles were loaded in MB/BacT instrument for the incubation and the growth was monitored. BacT/ALERT MP bottle with mycobacterial growth produced a color change of sensor from dark green to yellow, changing the screen color to yellow. The positive BacT/ALERT MP bottles were unloaded, vortexed heavily to make the large clumps to break and suspend uniformly. The mycobacterial growth was confirmed by performing ZN staining from it.

POLYMERASE CHAIN REACTION: The PCR was carried out as per the method described elsewhere (6). The peritoneal fluid was centrifuged at 3000 rpm for 15 minutes, and then the supernatant was discarded. To the sediment, 50 µl lysozyme was added followed by incubation at 37°C in water bath overnight. 70 µl 14% SDS and 6 µl proteinase K were added and the
mixture was incubated at 65°C for 15 minutes. Subsequently, 10 µl of 5M NaCl and 80 µl of 10% CTAB (activated at 55°C) were added and the final mixture was incubated at 65°C for 10 minutes. Then, 800 µl of phenol, chloroform and iso amyl alcohol was added in ratio of 25: 24: 1. The mixture was spun at 10000 RPM for 10 minutes at 4°C. Supernatant was collected, 600 µl of ice cold isopropanol was added and was incubated at -20°C overnight. Next day it was spun for 10 minutes at 12000 RPM at 4°C. After the fluid was drained out, it was kept in the incubator for drying. Finally, 20 µl Tris buffer was added and it was stored at -20°C.

Primers were validated by blasting the primer sequence used for detection of M.tuberculosis in the genome database of all the organisms on the website (http://www.ncbi.nlm.nih.gov/blast/) and were found to be specific for the organism. The sequence of the TB PCR Primers were: first set conventional PCR round- forward primer (FL): 5’ CTC AAG TGA AGG AGG CAA CC – 3’ and reverse primer (FR): 5’ TGG GCT AGG GTG TTG ATC TC – 3’ where as for the nested PCR: forward primer (NFL): 5’ CGT CTG GAG CGT GAC CTA CT – 3’ and reverse primer (NFR): 5’ GAC ATC TCG ACG GTC AGT CA – 3’ respectively.

PCR mix consisted of 4µl of extracted DNA, 10 µl of 2X ready made master mix, 2µl of 10µM Primer (FL and FR for 1st round PCR and NFL and NFR for nested PCR) and 10µl of milliQ water. The settings of the thermocycler programmed were similar for both first round and nested PCR except for the primers used. Total 30 cycles were carried out in each round, each cycle comprised of initial denaturation at 94°C for 5 minutes, denaturation occurred at 94°C for 60s, annealing occurred at 56 °C for 60s, amplification occurred at 72 °C for 60s followed by final extension at 72 °C for 7 min. Agar gel electrophoresis was carried out and bands are visualized under UV rays. The MTB specific nested PCR product size is 219 bp length. The bands were separated in the agarose gel according to the molecular weight which then checked by comparing the bands with standard molecular weight marker.

RESULT: The microbiological findings of various tests were depicted in table 1. ZN staining and AP staining have shown 23.8% (5 out of 21) and 28.5% (6 out of 21) sensitivity respectively whereas LJ culture was found to be 57.1% sensitive (12 out of 21 cases). BacT/ALERT microbial detecting system detected sixteen out of 21 cases (76.1%). There was one case (in probable group) which was detected by BacT/ALERT but was negative by nested PCR. The contamination rate of BacT/ALERT and LJ culture in the present study was found to be 5.0% and 10.0% respectively. Mean detection time of BacT/ALERT in all the isolates was 21.28 days compared to 32.10 days taken by LJ culture. Gel image after nested PCR targeting IS6110 of M.tuberculosis was shown in figure 1. Nested PCR could detect nineteen out of 21 (90.4%) cases of TBP where as one out of nine (11.1%) in control group was also detected positive by nested PCR. There were four cases detected by nested PCR but was negative by BacT/ALERT. Both the cases and controls were age and gender matched.

DISCUSSION: Tuberculous peritonitis (TBP) is the sixth most frequent site of extrapulmonary involvement (7). Although an infrequent disease, TBP with its non-specific symptoms and sinister clinical course can be easily confused with other intraabdominal diseases. Both the incidence and severity of TBP are expected to increase with increasing incidence of HIV infection. Hence, early and prompt diagnosis of the condition contributes to early start of anti tubercular therapy, thus preventing thereby the complications due to the condition. However, accurate laboratory diagnosis of TBP still continues to be a challenge.
In the present study, acid fast microscopy by ZN and AP staining showed low sensitivity which can be explained because of the paucibacillary nature of peritoneal fluid. A high bacterial load (\(>10^4 - 10^5\) bacilli / ml) is needed in the specimen to render an AFB microscopy result positive\(^{(2)}\). Various studies documented similar sensitivities of smear microscopy for TBP of < 20 % \(^{(6)}\). AP staining is found to be more sensitive than ZN staining as the smears can be examined by AP staining at 250x or 450x, covering larger area in the same time \(^{(12)}\). Culture by LJ media showed an overall sensitivity of 57.1%. Similar sensitivities had been documented in different studies ranging from 20-75% \(^{(6)}\). Low sensitivity of LJ culture could be due paucibacillary nature of the peritoneal fluid as minimum detection threshold being 100-1000 bacilli per ml \(^{(3)}\). Other reasons could be due to presence of dead bacilli, intake of other broad spectrum antibiotics inhibiting Mycobacteria and prior starting of ATT.

The BacT/ALERT microbial detecting system showed an overall sensitivity higher than the LJ culture and specificity of 100%. In confirmed & probable sub group the sensitivity was 100% and 64.2% respectively. Out of sixteen mycobacterial isolates were recovered in our study, all (100%) were recovered by BacT/ALERT compared to 75.0 % (twelve isolates) recovery rate of LJ culture. Other studies also have shown recovery rate of 91% to 94% by BacT/ALERT \(^{(7,8)}\). The contamination rate of BacT/ALERT and LJ culture in our study was found to be 5.0% and 10.0% respectively. Similar reports were documented in different studies\(^{(7)}\). Mean detection time of BacT/ALERT in all seven isolates was 21.28 days compared to 32.10 days taken by LJ. The mean detection time of BacT/ALERT in smear positive specimen was 17days where as that in smear negative specimens were 22.8 days. Various studies documented mean detection time of 15-16 days by BacT/ALERT \(^{(7,8)}\). Early detection of the bacilli in these studies could be due to inclusion of both pulmonary and extrapulmonary specimens in these studies.

In the present study, nested PCR showed an overall sensitivity of 90.4 % which was higher than culture but differed in the confirmed (100%) and probable (85.7%) subgroup. In an European study by T.zoanopoulos et al, all the three suspected cases of TBP were found positive by PCR targeting IS6110 \(^{(9)}\). As the sample size was very less and response to ATT was considered as final inclusion criterion for TBP, so the sensitivity was found higher in this study which might be misleading.

In our study, there were two cases which could not be detected by nested PCR (false negative results). The reasons of lower sensitivity might be due to many reasons: First, presence of PCR inhibitors which are found to be more associated with extrapulmonary specimen compared to pulmonary specimen \(^{(10-11)}\). A better extraction procedure like immunomagnetic separation technique should be used which could capture all M.tuberculosis DNA, but not inhibitors \(^{(10)}\). Various resin matrixes like ‘Gene releaser preparations’ which absorb inhibitors without entailing further loss of DNA may also be used \(^{(10)}\). Second, poor lysis of Mycobacteria in the extraction procedure due to the complexity of the cell wall \(^{(9,12)}\). Third, some strains of M.tuberculosis in Asia lack the IS6110 sequence \(^{(13-15)}\). Hence few cases might have got undetected if Mycobacteria present in the samples lack the IS6110 sequence Fourth, low number of bacilli present in the peritoneal fluid.

The overall specificity of nested PCR in our study was 88.8%. There was one out of nine controls which was detected positive (false positive) by our study where actually Escherichia coli was isolated and patient responded to the antibiotics. The reasons of false positive results could be due to cross contamination during initial handling or due to amplicon carry over...
contamination which can be overcome by using a single tube nested PCR\cite{10}. It is also found that use of dUTP/Uracil-N-glycosylase could decrease the amplicon contamination\cite{10}.

There were seven cases which were detected by nested PCR but were rendered negative by LJ culture. The gain in the sensitivity (40%) over the so called "gold standard" can be explained by low detection limit of nested PCR of as few as 10 bacilli per ml. The sensitivity gain in the etiological diagnosis by nested PCR substantiates the diagnostic utility of this method. Moreover nested PCR offers the advantages like speed in obtaining the result, option of referring the sample rather than the patient to a specialized centre.

There was one case (in probable group) which was detected by BacT/ALERT but was negative by nested PCR whereas, there were four cases which was detected by nested PCR but was negative by BacT/ALERT. So the combined sensitivity of BacT/AERT and nested PCR in the probable group (92.8%, 13 out of 14 cases) is more than individual sensitivity of BacT/AERT (64.2%) and nested PCR(85.7%).

In conclusion, diagnosis of TBP is often difficult due to the atypical clinical presentation and paucibacillary nature of the sample. Both Nested PCR and BacT/ALERT found to be more sensitive compared to conventional LJ culture and smear microscopy. BacT/ALERT takes lesser time to detect growth and associated with less contamination rate compared to conventional LJ culture. We suggest that nested PCR should deserve a place in laboratory diagnosis of TBP but careful adherence to the test protocol is mandatory. As both false negative and false positive results are reported on PCR, hence PCR alone should not be used as a criterion for initiating or terminating the therapy. This should be supported by clinical, radiological, cytological and other microbiological finding (smear microscopy, culture by conventional and automated system) to guide the clinician in decision making for appropriate therapy whenever possible.

The limitations of the present study are as follows. 1) It was based on clinical criteria which though had been validated well before but cannot be considered as gold standard. 2) Internal controls were not used for nested PCR. We believe that use of internal control could have helped in eliminating the possibility of PCR inhibitors. 3) We have not evaluated the diagnostic potential of these tests in HIV positive patients, where the sensitivity would have been more. 4) Only IS6110 is targeted in our study which might be absent in some Indian isolates of M.tuberculosis. 5) The sample size is small. Since TBP is relatively rare, it is difficult to get more patients. If the sample size can be increased then it will provide a clear picture about the parameters.

REFERENCES:


Table- 1 : Various clinical, radiological, cytological and histopathological findings among the cases

<table>
<thead>
<tr>
<th>Findings</th>
<th>% of patients</th>
</tr>
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<tbody>
<tr>
<td>Fever</td>
<td>81.5</td>
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<tr>
<td>Abdominal tenderness</td>
<td>67.1</td>
</tr>
<tr>
<td>Weight loss</td>
<td>74.6</td>
</tr>
<tr>
<td>Night sweat</td>
<td>65.8</td>
</tr>
<tr>
<td>Abnormal peritoneal fluid cytology</td>
<td>57.14</td>
</tr>
<tr>
<td>Raised ESR</td>
<td>53.1</td>
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Table – 2: Microbiological findings among various subgroups of tuberculous peritonitis (TBP)

<table>
<thead>
<tr>
<th>Sub groups</th>
<th>No of cases</th>
<th>Number of cases positive for M. tuberculosis by</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Zn staining</td>
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<tr>
<td>Total cases</td>
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<td>05</td>
</tr>
<tr>
<td>Confirmed TBP</td>
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<td>03</td>
</tr>
<tr>
<td>Probable TBP</td>
<td>14</td>
<td>02</td>
</tr>
<tr>
<td>Control (Non TBP)</td>
<td>09</td>
<td>0</td>
</tr>
</tbody>
</table>

Figure-1: Gel image after nested PCR targeting IS6110 of M.tuberculosis

Lane M : 100bp DNA ladder,   Lane 2 : 219bp PCR product of positive control,

Lane2&3: 219bp PCR product of clinical control,

Lane 4 to 6 : Clinical samples negative for IS6110 of M.tuberculosis