Comparative Evaluation of Several Gene Targets for Designing a Multiplex-PCR for an Early Diagnosis of Extrapulmonary Tuberculosis

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Purpose: Diagnosis of extrapulmonary tuberculosis (EPTB) poses serious challenges. A careful selection of appropriate gene targets is essential for designing a multiplexer-polymerase chain reaction (M-PCR) assay.

Materials and Methods: We compared several gene targets of Mycobacterium tuberculosis, including IS6110, devR, and genes encoding MPB-64 (mpb64), 38kDa (pszt1), 65kDa (hsp65), 30kDa (fbpB), ESAT-6 (esat6), and CFP-10 (cfp10) proteins, using PCR assays on 105 EPTB specimens. From these data, we chose the two best gene targets to design an M-PCR.

Results: Among all gene targets tested, mpb64 showed the highest sensitivity (84% in confirmed cases and 77.5% in clinically suspected cases), followed by IS6110, hsp65, 38kDa, 30kDa, esat6, cfp10, and devR. We used mpb64+IS6110 for designing an M-PCR assay. Our M-PCR assay demonstrated a high sensitivity of 96% in confirmed EPTB cases and 88.75% in clinically suspected EPTB cases with a high specificity of 100%, taking clinical diagnosis as the gold standard.

Conclusion: These M-PCR results along with the clinical findings may facilitate an early diagnosis of EPTB patients and clinical management of disease.

Key Words: Mycobacterium tuberculosis, extrapulmonary tuberculosis, PCR, multiplex-PCR, diagnosis
co-infection, and EPTB exhibits serious challenges. Diagnosis of EPTB can be elusive due to paucibacillary load in the biological specimens, lack of adequate clinical sample volume, and localization of disease at sites that are difficult to access. Smear microscopy is the most extensively used method for the diagnosis of EPTB as it is inexpensive and rapid; however, the method has drawbacks due to low and variable sensitivity values (0–40%). Culture identification, though considered as a gold standard, has drawbacks including low and variable sensitivities (0–80%) in various clinical forms of EPTB, a slow turnaround time, and a labor-intensive process. The BACTEC system, histopathological/cytological examination, and interferon-γ release assays are also employed for the diagnosis of EPTB; however, these assays have their own limitations. Detection of adenine deaminase is utilized in diagnosing EPTB specimens, although there is no established cut-off value, and this test has been shown to be positive in other diseases such as lymphomas and collagen vascular diseases. Recently, nucleic acid amplification (NAA) tests such as PCR have emerged as potentially important tools for the diagnosis of EPTB specimens. Various mycobacterial gene targets such as insertion sequence (IS)6110; IS1081; 16S rRNA DNA; devR (transcriptional regulator, Rv3133c); genes encoding 65kDa (heat shock protein 65, hsp65; Rv0440), MP-64/MPT-64 (mycobacterial protein from species tuberculosis, mpb64; Rv1980c), 38kDa (phosphate-binding lipoprotein, pstS1; Rv0934), and MTP-40 (membrane associated phospholipase C1; Rv2351c) proteins; TRCA (conserved repetitive element); GCRS (guanine-cytosine-rich repetitive sequence); flb, encoding fibronectin-binding protein B (30kDa, Ag85B protein; Rv1886c); hupB, encoding histone-like DNA-binding protein (Rv2986c); and dnaJ, encoding chaperone protein (Rv0352), have been exploited for such tests. However, limited information is available for the utilization of the M. tuberculosis-specific region of difference (RD) 1 esat6, encoding early secretary antigenic target-6 (ESAT-6, Rv3875), and cfp10, encoding culture filtrate protein-10 (CFP-10, Rv3874), gene targets. Moreover, there is a high variation in these tests, owing to different gene targets employed as well as the different gold standards adopted in various laboratories. Recently, the introduction of the MTB/RIF GeneXpert (Cepheid, Sunnyvale, CA, USA) assay, which targets conserved rpoB gene encoding RNA β polymerase subunit, has been a major breakthrough in the diagnosis of EPTB; however, its use is limited in resource-poor settings due to its high cost. Therefore, we planned a study to compare several gene targets (IS6110, devR, mpb64, hsp65, 30kDa, 38kDa, esat6, and cfp10) for PCR assays on EPTB specimens to identify the two best gene targets exhibiting high sensitivities, using clinical diagnosis as the gold standard, and to design a simple, reliable, and cost-effective multiplex-polymerase chain reaction (M-PCR) assay with the two resulting targets for an early diagnosis of EPTB.

MATERIALS AND METHODS

Clinical specimens

The clinical specimens were collected from the patients visiting the clinical departments of Rajan Babu Institute for Pulmonary Medicine & Tuberculosis, Delhi; Vallabhbhai Patel Chest Institute, University of Delhi; and the Postgraduate Institute of Medical Sciences, Rohtak during the period of June, 2013 to September, 2014. Written informed consent was obtained from all patients participating in this study. This work was part of a project approved by the Maharshi Dayanand University Institute Ethical Committee under the protocol number MDU/CBT/11/793. This study was double blinded, as the laboratory was not aware of the clinical data and the clinicians were not aware of the laboratory data until all of the analyses were complete.

The sample size (n=155) was determined to be statistically significant with a 95% level of confidence, taking into account the prevalence rate of the study population (Delhi/Rohtak). Samples were divided into two groups. Group 1, the EPTB group (n=105) was subdivided into 1) confirmed EPTB cases (n=25; smear positive (n=21) or culture positive (n=4)) and 2) clinically suspected EPTB cases (n=80) that were smear negative and culture negative yet suspected on the basis of imaging, clinical findings, histological/cytological observations, and response to anti-tubercular therapy (ATT). Group 2 was the control group (n=50), which comprised non-TB individuals.

Among 105 EPTB specimens, there were 64 pleural fluids, six ascitic fluids, 20 pus specimens from different origins, two lymph node aspirates, ten urine specimens, two pleural biopsies, and one synovial fluid specimen. Similarly, pleural fluids (n=23), pus (n=5), ascitic fluids (n=16), and urine (n=6) were collected from non-TB individuals with renal failure, trauma, or cancer as controls. All EPTB patients were in the age group of 15–60 years, and both sexes were represented. The inclusion criteria in the selection of TB pleuritis and TB lymphadenitis patients were 1) fever, cough, pleuritic pain, malaise, anorexia, and exudative pleural effusion with lymphocytic predominance on diagnostic tap and 2) untreated patients with peripheral (superficial) TB lymphadenitis of cervical or axillary regions. Similarly, the inclusion criteria for the selection of abdominal TB patients were fever and weight loss with one or more of the following: diarrhea persisting >1 month; ascites; abdominal lymphadenopathy or mesenteric masses on ultrasound; or hepatomegaly, splenomegaly, generalized pain, or tenderness persisting over 7 days. The exclusion criteria were 1) patients with a history of TB in the preceding 2 years, those already receiving ATT, those with other coexisting medical illnesses (e.g., HIV-positive individuals), pregnant women, and diabetes mellitus patients and 2) smear- and culture-negative suspected EPTB patients who did not respond to ATT even up to 8 weeks of treatment. All EPTB and non-TB specimens except urine were collected in the minor operation theater, fol-
owing all standard aseptic surgical precautions of the participating institutes. All specimens were stored at 4°C and processed within 24 h.

**Sample processing and DNA extraction**

All clinical specimens including pleural biopsies (after homogenization) were digested and decontaminated using 1% N-acetyl-L-cysteine combined with 4% NaOH (NALC-NaOH). Briefly, an equal volume of NALC-NaOH solution was added to a clinical sample in a 15-mL centrifuge tube, mixed by inversion and incubated at room temperature for 20 minutes. Later, 0.067 M phosphate buffer (pH 6.8) was added, and the reaction mixture was subjected to centrifugation at 3000 × g for 15 min. The supernatant was discarded, and the pellet was used for DNA extraction, smear microscopy, and the culture. The decontamination and centrifugation steps were performed in a biosafety hood. The mycobacterial genomic DNA was extracted according to the method of van Helden, et al. with few modifications using the cetyl-N,N,N-trimethyl ammonium bromide (CTAB)-phenol-chloroform method. Briefly, the pellet was suspended in 500 μL of Tris-EDTA buffer, heated at 100°C for 10 minutes, and incubated with 20 μL of lysozyme (10 mg/mL) at 37°C for 2 h. Then, 30 μL of 10% SDS and 10 μL of proteinase K (20 mg/mL) were added, briefly vortexed, and incubated at 37°C for 1 hour, followed by incubation with 5 M NaCl (100 μL)+10% CTAB (80 μL) at 65°C for 20 min. Later, an equal volume of phenol-chloroform-isooamyl alcohol (25:24:1) mixture was added, mixed by inversion and centrifuged at 14000× g for 10 minutes. This step was repeated twice for the removal of DNA contaminants. The supernatant was discarded, followed by the addition of 3 M sodium acetate and isopropanol for DNA precipitation. The precipitate was washed with 90% ethanol and air dried. The DNA pellet was suspended in 100 μL of Tris-EDTA and stored at -20°C until further analysis.

**PCR**

We designed primers for *IS6110, mpb64, esat6, cfp10, 38kDa, 30kDa, devR*, and *hsp65* (targets of *M. tuberculosis* H37Rv) using Primer 3 software for PCR and validated the designs with a primer-designing software tool (http://www.ncbi.nlm.nih.gov/tools/primer-blast/). All primers were obtained from Eurofins Genomic India Pvt. Ltd., Bengaluru. Table 1 shows the primer sequence as well as the annealing temperature used for different targets. For PCR amplification of individual gene targets, 1 μL of purified *M. tuberculosis* H37Rv DNA (1 ng/mL) was added to 25 μL of PCR master mix containing 2.5 μL 10× PCR buffer, 1.5 μL of 25 mM MgCl₂, 0.5 μL of 200 μM (each) of the four deoxyribonucleoside triphosphates, 10 pM of each forward and reverse primer, PCR grade water, and 0.625 U of *Taq* DNA polymerase (Bangalore Genei Pvt. Ltd., Bengaluru, India). The DNA amplification was performed in a thermal cycler (T100™ BIORAD). After an initial denaturation at 95°C for 3 min, each cycle was composed of three steps: denaturation at 95°C for 30 s followed by annealing of respective primers at an appropriate temperature (Table 1) for 30 s, extension at 72°C for 45 s for 30 cycles, and a final extension at 72°C for 5 min. The PCR-amplified products were resolved by gel electrophoresis using 1.5% agarose in 1X TAE buffer containing ethidium bromide (0.5 μg/mL) at 80 volts for about 40 min and viewed under the Gel Documentation System (MiniLumi, DNR Bio-Imaging Systems Ltd., Jerusalem, Israel) to determine the presence of a specific size band for each gene target. The PCR amplification for DNA isolated from the clinical specimens was performed in a similar manner. All samples were run in duplicate. In each experiment, a positive control (*M. tuberculosis* DNA at 1 ng/mL) and two negative controls (no template DNA, only PCR grade water) were included.

| Table 1. Primer Sequences Used for Different Gene Targets of *M. tuberculosis* H37Rv |
| Gene target | Primers sequence (5’→3’) | Annealing conditions (°C/30 s) | Expected amplicon size (bp) |
| IS6110 | Forward GAAGAATCCCGGAGATAAAC | 53 | 258 |
| mpb64 | Forward GACTCTCTGGTCGAGTGAAAC | 54 | 163 |
| 38kDa (pstS1) | Forward ACACCTCTGTCGACAGTA | 53 | 383 |
| 30kDa (tpb8) | Forward TGCTGATGGTCTCTGTC | 55 | 190 |
| hsp65 | Forward GGCTCTGTGCGGTA | 52 | 400 |
| devR | Forward GAGTGTCTTTACCCG | 49 | 241 |
| esat6 | Forward GTCTGCTCTTACCG | 53 | 220 |
| cfp10 | Forward CAGAGATGAAGACCGATG | 54 | 203 |
Table 2. Combination of Different Concentrations of Two Primer Pairs Used for Optimization of M-PCR

| Primer pairs | Concentration of primers (µM) |
|-------------|-------------------------------|
| mpb64      | 0.2 | 0.2 | 0.4 | 0.4 |
| IS6110     | 0.2 | 0.4 | 0.4 | 0.2 |

M-PCR, multiplex-polymerase chain reaction.

M-PCR
M-PCR (mpb64+IS6110) amplification was performed with the same concentrations of master mix components as for monoplex PCR, with the exception of the different primer pair concentrations. DNA amplification was performed for 30 cycles as described above, although the annealing step was performed at 53°C for 30 s. For the optimization of M-PCR, the combinations of different concentrations of two primer pairs were evaluated (Table 2) using 1 µL of purified M. tuberculosis DNA (1 ng/mL) to choose an appropriate ratio for mpb64:IS6110, which showed the clear amplicons of specific sizes for the two gene targets. PCR amplification for DNA isolated from the clinical specimens was performed in a similar manner and determined the presence or absence of a single band or two bands of specific size(s) under the Gel documentation system. Both positive and negative controls were run as explained above. The PCR/M-PCR results were compared using clinical diagnosis as the gold standard (comprising smear, culture, clinical findings, imaging, histological and cytological observations, response to ATT, and all together).

Statistical methods
The sensitivity, specificity, positive predictive value (PPV) and negative predictive value (NPV) were calculated as detailed by Srivastava, et al.14

RESULTS
Prior to taking up this study in EPTB samples, validation of the PCR tests using different gene targets was carried out with few sputum samples (n=6) from PTB patients, which were found to be smear-positive and culture-positive for M. tuberculosis.

Comparison of PCR results among EPTB specimens with different gene targets
Fig. 1 shows the amplicons of different size(s) using different gene targets (IS6110, mpb64, esat6, cfp10, 38kDa, 30kDa, devR, and hsp65) with the purified M. tuberculosis DNA via PCR. After optimization of PCR with the purified DNA, we performed PCR using the clinical EPTB as well as non-TB samples. Representatives of positive clinical samples using different gene targets have been depicted (Fig. 1). The order of sensitivity with different targets in 105 EPTB cases (both confirmed and suspected cases) in descending order was mpb64+IS6110>hsp65>38kDa>30kDa-esat6>cfp10>devR (Table 3). However, the specificity of the PCR assay was high (88–100%) with all gene targets employed. The PPV, NPV, and accuracy observed with the individual gene targets have been summarized as percentages in Table 3.

Optimizing the primer-pair concentrations for M-PCR
Considering that mpb64 and IS6110 showed the highest positivity among all eight targets examined, we chose these two targets for M-PCR. We optimized the primer concentrations for mpb64+IS6110 (Table 2) to develop an M-PCR. Among the different combinations of primer concentrations, a combination of 0.2 µM of mpb64 and 0.4 µM of IS6110 showed sharp amplicons with the purified M. tuberculosis DNA (Fig. 2). Therefore, the same combination of primers was incorporated in the PCR master mixture to evaluate the clinical EPTB samples.

M-PCR
The M-PCR targeting mpb64+IS6110 showed better % sensitivity (90.5%) than monoplex PCR (Table 3 and 4) with individual mpb64 (79%) and IS6110 (75%) with a high specificity (100%) in 105 EPTB and 50 non-TB specimens (controls). Among the 25 confirmed EPTB cases, M-PCR revealed 24 positive cases, thus leading to 96% positivity. Out of 24 positive cases, 17 were positive with both IS6110 and mpb64, three were positive with IS6110 only, and four were positive with mpb64 only. Out of 80 clinically suspected EPTB cases, M-PCR showed 71 positive cases, thus leading to 88.75% positivity. Out of 71 positive cases, 50 were positive with both IS6110 and mpb64, nine were positive with IS6110 only, and 12 were positive with mpb64 only (Fig. 3). The M-PCR test demonstrated 100% PPV, 83.33% NPV, and
Table 3. PCR Results Using Various Gene Targets

| Results (+/-) | mpb64 | IS6110 | 38kDa | hsp65 | 30kDa | esat6 | cfp10 | devR |
|--------------|-------|--------|-------|-------|-------|-------|-------|------|
| n=25         | +21   | -4     | +20   | -5    | +18   | -7    | +16   | -9   |
| Sensitivity (%) confirmed EPTB cases | 84    | 100    | 80    | 81.81 | 76    | 82.62 | 80    | 77.27 |
| PPV (%)      |       |        |       |       |       |       |       | 84.90 |
| NPV (%)      | 92.59 | 90.90  | 86.79 | 88.46 | 83.63 | 84.90 |       |       |
| ACC (%)      | 86.15 | 94.66  | 83.33 | 85.33 | 86.66 | 82.66 |       | 76.66 |
| Group I      |       |        |       |       |       |       |       |      |
| n=80         |       |        |       |       |       |       |       |      |
| Sensitivity (%) clinically suspected EPTB cases | 77.5  | 100    | 73.75 | 63.75 | 67.50 | 58.75 | 60    | 56.25 |
| PPV (%)      |       |        |       |       |       |       |       | 89.58 |
| NPV (%)      | 100   | 100    | 92.72 | 93.10 | 92.15 | 90     |       | 53.75 |
| ACC (%)      | 86.15 | 83.94  | 74.61 | 76.92 | 71.53 | 63.88 |       | 66.55 |
| Total EPTB cases | 62    | 18     | 59    | 21    | 51    | 29    | 54    | 26   |
| Sensitivity (%) | 79.04 | 100    | 75.23 | 65.71 | 65.92 | 60.0  | 60    | 59.04 |
| PPV (%)      |       |        |       |       |       |       |       | 57.14 |
| NPV (%)      | 100   | 100    | 94.52 | 93.83 | 92.53 | 92.22 |       | 50.0  |
| ACC (%)      | 85.16 | 83.22  | 74.19 | 74.19 | 70.32 | 69.23 |       | 65.16 |
| Group II     |       |        |       |       |       |       |       |      |
| Controls     |       |        |       |       |       |       |       |      |
| n=50         |       |        |       |       |       |       |       |      |
| Sensitivity (%) | 100   | 100    | 92.92 | 92.92 | 92.92 | 92.92 |       | 92.92 |

n, number of specimens; +, PCR positive cases; -, PCR negative cases; PPV, positive predictive value; NPV, negative predictive value; ACC, accuracy; EPTB, extrapulmonary tuberculosis; Total EPTB cases, confirmed clinically suspected cases.

The diagnosis of EPTB remains inconclusive in most cases. The sensitivity of conventional bacteriological methods, such as smear staining and culture (6,12), has been reported for the diagnosis of TB pleuritis (Table 5). Based on a meta-analysis of six studies (598 samples) on the Xpert assay, lower sensitivity and specificity (75%) for the diagnosis of TB pleuritis were observed with the Xpert assay (Table 5). A meta-analysis revealed high sensitivity (95.3%) and specificity (100%) for M-PCR using IS6110 and mpb64 from pleural fluids, pleural biopsy, and gastrointestinal TB; however, the sensitivity and specificity of Xpert assay were lower than the reported one using an imperfect gold standard. ART, etc. validation study, when diagnosing EPTB specimens, with varying sensitivities, which can lead to inaccurate PCR results. Several researchers have used clinical diagnosis and composite reference standard (CRS; combination of smear, culture, histology and cytology, clinical findings, response to ATT, etc.) to validate PCR results when diagnosing EPTB specimens. In-house PCR tests provide high specificities (95-100%) for the diagnosis of TB lymphadenitis, osteoarticular TB, and gastrointestinal TB; however, the sensitivity (100%) for each of these tests is considerably lower than the reported one using an imperfect gold standard. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. The sensitivity and specificity of each of these tests are different than the reported one using an imperfect gold standard. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. The sensitivity and specificity of each of these tests are different than the reported one using an imperfect gold standard. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests. Culture is the most widely used gold standard, though it is suboptimal for validating EPTB specimens when heterogeneous sensitivities and specificities are observed with commercial tests.
pleural fluids, using CRS as the reference standard. Therefore, the World Health Organization (WHO) has not recommended the use of the Xpert assay for the diagnosis of TB pleuritis, though the same assay has been recommended for the diagnosis of TB lymphadenitis and TB meningitis. Interestingly, our M-PCR test was positive in six of ten urine samples collected from the suspected TB pleuritis patients (Table 5). Among those six positive cases, two were missed by IS6110 yet were detected by mpb64. The detection of *M. tuberculosis* in urine samples by PCR can be a useful method for the diagnosis of TB pleuritis and other clinical EPTB forms in which sample collection is difficult and requires aggressive techniques, though it needs further validation in a large number of specimens. The PCR/M-PCR assay is also limited in that it does not discriminate between alive and dead or degraded bacilli. However, this is likely a theoretical limitation, and its relevance must be judged in light of the overall picture of EPTB cases.

Genotype studies of *M. tuberculosis* strains in the Indian population using spoligotyping and IS6110-restriction fragment length polymorphism reveal the East African Indian family to be the major clade in the south India, whereas the Central Asian family (CAS) predominates in the northern part of India, and the largest clade among the CAS belongs to the CAS1-Delhi lineage, which possesses variable IS6110 copy numbers. IS6110, which belongs to a family of IS of the IS3 category has been widely employed in PCR tests due to the presence of its multiple copies in the *M. tuberculosis* complex genome, which is believed to confer higher sensitivity. However, in recent years, several clinical investigations raised concerns over IS6110 specificity as well as sensitivity in the diagnosis of TB owing to false-positive (due to homology with other target DNA besides *M. tuberculosis*) or false-negative (due to a single copy or absence of copies of IS6110) results, particularly from Southeast Asian countries, thereby suggesting the use of other potential gene targets for monoplex or M-PCR assays. Our PCR results with individual IS6110 and mpb64 (Table 3) are similar to prev-

### Table 4. Sensitivity and Specificity of M-PCR

| Type        | Specimens Results | M-PCR (mpb64+IS6110) | Both IS6110 & mpb64 | Only IS6110 | Only mpb64 | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) | ACC (%) |
|-------------|-------------------|-----------------------|---------------------|-------------|------------|----------------|----------------|---------|---------|---------|
| **Group I** |                   |                       |                     |             |            |                |                |         |         |         |
| Confirmed   | EPTB cases n=25   | +                     | 24                  | 17          | 3          | 4              | 96             | 100     | 98.03   | 98.66   |
| Suspected   | EPTB cases n=80   | +                     | 71                  | 50          | 9          | 12             | 88.75          | 100     | 84.74   | 93.07   |
| Total       | EPTB cases n=105  | +                     | 95                  | 67          | 12         | 16             | 90.47          | 100     | 83.33   | 93.54   |
| **Group II**| Negative control  | n=50                  | 0                   | 0           | 0          | 0              | 100            |         |         |         |

n, number of specimens; M-PCR, multiplex-polymerase chain reaction; +, PCR positive cases; -, PCR negative cases; PPV, positive predictive value; NPV, negative predictive value; ACC, accuracy; EPTB, extrapulmonary tuberculosis; Total EPTB cases, confirmed+clinically suspected cases.

### Table 5. Sensitivity of M-PCR in Individual EPTB Specimens

| Groups          | Type of EPTB specimens Results | M-PCR (mpb64+IS6110) | Both IS6110 & mpb64 | Only IS6110 | Only mpb64 | Sensitivity (%) |
|-----------------|--------------------------------|-----------------------|---------------------|-------------|------------|----------------|
| **Confirmed**   | Pleural fluids (21) +          | 20                    | 13                  | 3           | 4          | 95.23          |
|                 | Lymph node aspirates (2) +     | 2                     | 2                   | -           | -          | 100            |
|                 | Pleural biopsies (2) +         | 2                     | 2                   | -           | -          | 100            |
| **Suspected**   | Pleural fluids (43) +          | 41                    | 25                  | 7           | 9          | 95.34          |
|                 | Ascitic fluids (6) +           | 5                     | 3                   | 1           | 1          | 83.33          |
|                 | Pus (20) +                     | 18                    | 15                  | 1           | 2          | 90             |
|                 | Urine (10) +                   | 6                     | 4                   | -           | 2          | 60             |
|                 | Synovial fluid (1) +           | 1                     | 1                   | -           | -          | 100            |

n, number of specimens; +, PCR positive cases; -, PCR negative cases; M-PCR, multiplex-polymerase chain reaction; EPTB, extrapulmonary tuberculosis.
mous studies\(^{10,16}\) yet differ from the other studies\(^{23,24}\) reporting a sensitivity of <50%. Balne, et al.\(^{25}\) recently reported a very low sensitivity of 9% only with \(IS6110\)-based PCR in ocular TB specimens, whereas 70.2% and 40% sensitivities were observed with individual \(mpb64\) and \(38kDa\), respectively. The reason for low sensitivity values with \(IS6110\) in many studies could be due to south Indian isolates, 40% of which are shown to possess zero or low copy numbers of \(IS6110\), while only 10–15% of North Indian isolates are shown to possess zero or low copy numbers of \(IS6110\).\(^{22}\) Other reasons for low sensitivity with \(IS6110\) or other targets could also relate to insufficient lysis of bacterial cells, loss of DNA during purification, or the presence of PCR inhibitors in the EPTB samples.

The \(fbpB\) gene (Rv1886c), encoding 30 kDa protein, was included in these assays, as this protein constitutes up to 41% of total mycobacterial proteins in log-phase culture supernatants. The mRNA coding for 30 kDa protein has been utilized as a target for reverse transcriptase-PCR\(^{26}\) in order to diagnose EPTB, and it can also detect viable mycobacteria; however, it is cumbersome to work with mRNA routinely. We performed PCR targeting the 30kDa gene (Table 3), which revealed a slightly lower sensitivity (70%) than earlier observations (87.5% sensitivity) made by Kidane, et al.\(^{27}\) when diagnosing TB lymphadenitis. The PCR results with the individual \(hsp65\) and \(38kDa\) gene targets (65.7–69.5% sensitivity) (Table 3) are almost similar to previous studies with EPTB specimens, such as pleural fluids, pus, and skin or abdomen biopsies.\(^{8,26,28}\) When performed individually, our combined \(IS6110\) PCR and \(devR\) PCR tests showed a sensitivity of 80.9% and a specificity of 88% in 105 EPTB specimens (Table 3), which is comparable to such combined PCR tests reported by Chakravorty, et al.\(^{8}\) in pleural fluids and lymph node biopsies using the universal sample processing method. However, M-PCR in a single tube has advantages, as it reduces errors of cross contamination as well as cost and increases the sensitivity of the test.

The simultaneous identification and differentiation of \(M.\) \(tuberculosis\) and \(M.\) \(avium\) complexes and the non-tuberculous mycobacteria directly from PTB and EPTB specimens have been documented for an M-PCR using genus-specific primers targeting \(hsp65\), \(esat6/cfp10\), and \(M.\) \(avium\) complex-specific primer pairs targeting 16S–23S rDNA internal transcribed spacer-1 sequences.\(^{29,30}\) Similarly, Kim, et al.\(^{31}\) devised an M-PCR for the identification of \(M.\) \(tuberculosis\) complex to the species level in 37 strains and 178 clinical isolates targeting \(rpoB\) as well as RD1 and RD8 sequences. While performing PCR with the individual RD1 \(esat-6\) and \(cfp10\) gene targets for the diagnosis of EPTB specimens, lesser sensitivities (57–59%) (Table 3) were observed as compared to \(mpb64\) or \(IS6110\), likely due to the genetic variability of RD1 genes or low copies of ESAT-6 gene clusters in the genomes of certain \(M.\) \(tuberculosis\) isolates.\(^{32,33}\) We recently detected RD1 and RD2 proteins (\(ESAT-6\), \(MPB-64\), etc.) from both PTB and EPTB specimens using a novel ultrasensitive immuno-PCR assay (PCR amplified immunoassay) with a good diagnostic efficacy,\(^{34}\) though it needs to be better optimized for routine use.

Similar to our studies, Dubey, et al.\(^{35}\) used \(IS6110+mpb64\) in designing an M-nested PCR for the detection of \(M.\) \(tuberculosis\) directly from the blood samples of PTB as well as EPTB patients, using a Blacklight Card Disc with high sensitivity (95.7%) and specificity (100%). However, Elnifro, et al.\(^{36}\) earlier suggested avoiding the use of nested primers for the M-PCR assay as this could lead to false positive results. Other researchers developed M-PCR using an \(IS6110+38kDa\), \(IS6110+devR\) or \(IS6110+15S\) like element \(B9\), for the diagnosis of PTB and EPTB.\(^{37-39}\) The combination of three gene targets (\(IS6110+dnaA\)+\(hsp65\) or \(IS6110+16S\) rRNA+\(devR\)) for designing an M-PCR has also been document-
ed as a diagnosis tool for EPTB. However, we were unable to further raise the sensitivity of M-PCR using three gene targets, such as IS6110+mpb64+38kDa or IS6110+mpb64+htsp65 (data not shown), as our M-PCR exhibited substantially high sensitivity and specificity with two gene targets.

In conclusion, this study demonstrates the superiority of mpb64, followed by IS6110, among the several gene targets tested and further indicates the effectiveness of mpb64+IS6110 in the design of an efficient M-PCR test that facilitates an early diagnosis of smear-negative and culture-negative paucibacillary EPTB specimens, which are difficult to diagnose with the available standard methods. To the best of our knowledge, this is the first report to evaluate M-PCR with mpb64+IS6110 for the efficient diagnosis of TB pleuritis. Presently, we are validating these results in a large number of samples from varied clinical EPTB types in order for this simple and cost-effective test to be included in the diagnostic panel on a routine basis, which may prove to be a better alternative than the rpoB-based Xpert assay in resource-poor settings. This test might work well in varied clinical EPTB types from various countries that harbor M. tuberculosis isolates with high copy numbers of IS6110 and also countries that harbor M. tuberculosis isolates with zero or low copy numbers of IS6110 (e.g., South Indian isolates), as most can be detected by mpb64.

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