Role of GeneXpert in Rapid Molecular Detection of Extrapulmonary Tuberculosis in Tertiary Care Hospital

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Abstract

Introduction: Tuberculosis, the most common infectious disease with prevalence of 9.6 million globally. Most prevalent (23%) in India. Extrapulmonary tuberculosis (EPTB) accounts for 20% of total burden of tuberculosis. Rapid detection of Mycobacterium Tuberculosis (MTB) is essential for effective disease management. CBNAAT (Cartridge Based Nucleic Acid Amplification Test) or GeneXpert MTB/RIF assay - novel diagnostic tool to detect MTB and RIF resistance simultaneously. WHO recommends its utility for non-respiratory samples also. Burden of EPTB and drug resistance vary from place to place. Objective: Study was conducted to gather information about burden of disease in our locality and to asses utility of CBNAAT in detecting MTB and rifampicin resistance in suspected EPTB cases. Methods: Retrospective analysis of 281 samples from suspected cases collected in falcon tubes and processed using CBNAAT. Result: Total of 281 extrapulmonary samples received, 67(23.8%) were positive and 214(76.1%) were negative for MTB. Of 67 positives, RIF resistance detected in 1(1.49%) case. Maximum number of MTB detected in the age group 21-30 years (n=23, 34.3%). Among 165 males and 116 females, MTB detected in 44(26.6%) and 23(19.8%) respectively. Out of 281 patients, 24(8.54%) were HIV positive. Of these 24, only 8(33.3%) found positive for MTB. Among 257 non-HIV patients, MTB detected in 59(22.9%). Among different samples received, maximum number were Pleural fluid n=115(40.9%) and Maximum MTB positives found in FNAC (of lymphnodes) samples [n=35(52.2%)]. Conclusion: CBNAAT is a rapid test to detect MTB and rifampicin resistance simultaneously in EPTB and it reduced the treatment abuse in suspected cases.

Key words: CBNAAT, Extrapulmonary TB, MTB, rifampicin resistance.

Introduction

Tuberculosis (TB) is the most common infectious disease and in developing countries like India, it is major health problem. Due to inadequate diagnostic assays it remains as a challenge to public health. According to the WHO Tuberculosis report in 2015, there are 9.6 million people infected with TB. India has the maximum number of cases, that is 23% of the total cases all over the world. Globally, an estimated 3.3% of new TB cases and 20% of previously treated cases have MDR-TB [1]. Extra Pulmonary TB (EPTB) can affect any organ in the body and it accounts for 20% of total burden of tuberculosis globally. In majority of cases it remains undetected for a longer time. A major hindrance to the diagnosis of EPTB is the atypical presentation, often simulating neoplasia and/or inflammatory disorders and clinical specimens of deep organs are difficult to obtain. Adding to this, the Mycobacterium Tuberculosis Bacilli (MTB) load is generally very low in non-respiratory samples, therefore strongly affecting the sensitivity of acid-fast microscopy [2]. Also quick and reliable laboratory diagnostic methods for detecting tubercle bacilli in EPTB specimens are not easily available. This adds to the increased rates of morbidity and mortality in EPTB patients [3].

As the conventional laboratory methods are slow and cumbersome, Foundation for Innovative New Diagnostics (FIND) introduced cartridge-based nucleic acid amplification assay (GeneXpert MTB/rifampicin [RIF]). It is a molecular test which is fully automated and detects MTB directly from clinical samples within
two hours as well as RIF resistance which is the surrogate marker of MDR-TB conferring mutations in 81 bp RIF resistance determining region (RRDR) of the rpoB gene, which codes for a beta subunit of RNA polymerase of MTB, is the genetic basis of RIF resistance [4]. In 2014, The GeneXpert MTB/RIF assay has been strongly recommended by WHO for testing non-respiratory specimens from patients suspected of having EPTB to diagnose MTB and multidrug-resistant TB over the conventional tests (including microscopy, culture or histopathology). The test is currently recommended as a “first line” fast diagnostic test in high TB burden countries like India [5,6]. Burden of EPTB and drug resistance vary from place to place, therefore this study was conducted to assess the burden of EPTB and drug resistance in our locality.

Materials and Methods

Type and place of study: Study was a retrospective observational record-based analysis conducted in the RNTCP lab attached to the Department of Microbiology in Bowring and lady Curzon hospital.

Sample collection: Total of 281 clinically suspected EPTB cases were received in RNTCP lab from various department of our hospital and other private hospitals. Few details of the patients like Name, Address, Age, Sex, HIV status, Treatment received and Name of the referring centers were noted down.

Study period: From May 2016 to April 2018.

Sampling method: Extra pulmonary samples (pus, aspirates, body fluids, lymph node [LN] tissue) were collected in special, plain, universal 30ml clear plastic container with cap (falcon tubes) under aseptic conditions. 5ml of samples were received to which buffer solution was added, then the mixture is loaded to cartridge which were processed by Xpert MTB/RIF assay (Cepheid-Sunnyvale-USA), as per the guidance document given by Central TB division, Government of India (RNTCP, 2013; RNTCP, 2012). The results can be read as MTB detected, MTB not detected, RIF resistance detected; RIF resistance not detected; RIF resistance indeterminate; or invalid/error with the help of positive beacons.

Inclusion criteria: Extra pulmonary samples (pus, aspirates, body fluids, LN tissue) from all the age group irrespective of gender.

Exclusion criteria: Blood samples, urine samples, sputum samples and any other EP sample contaminated with blood is not included in our study.

Results

Total of 281 extrapulmonary samples received during study period, 165(58.7%) were males and 116(41.2%) were females. Most of the cases were of age group 21 to 30 years. Out of 281 samples received, 73(25.97%) were of FNAC, 115(40.92%) Pleural fluid, 26(9.25%) Ascitic fluid, 36(12.8%) CSF, 22(7.82%) Pus and 9(3.2%) were aspirates (other body fluids like synovial fluid, vitreous humor etc) as shown in the table-1.

Among 281 samples, 67(23.8%) were positive and 214(76.1%) were negative for MTB as shown in figure-1. Out of 67 positives, 66(98.5%) were MTB-positive/RIF resistance-negative and 1(1.49%) was MTB- positive/RIF resistance-positive (in aspirate sample) as shown in the figure-2. Maximum number of MTB was detected in the age group of 20-30 years (n=23, 34.3%). Among 165 males and 116 females, MTB was detected in 44(26.6%) and 23(19.8%) respectively as shown in figure-3. Out of total 281 patients, 24(8.54%) were HIV positive. Of these 24, only 8(33.3%) were found positive for MTB in different samples as shown in the table-2. Among 257 non-HIV patients, MTB was detected in 59(22.9%) as shown in table-3. Among different samples received, Pleural fluid was in maximum number n=115(40.9%). Maximum MTB positives were found in FNAC samples n=35(52.2%) as shown in the figure-4.

Table-1: Different EPTB samples received and their percentages

<table>
<thead>
<tr>
<th>Specimen type</th>
<th>Frequency</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>FNAC</td>
<td>73</td>
<td>25.97%</td>
</tr>
<tr>
<td>Pleural fluid</td>
<td>115</td>
<td>40.92%</td>
</tr>
<tr>
<td>Ascitic fluid</td>
<td>26</td>
<td>09.25%</td>
</tr>
<tr>
<td>CSF</td>
<td>36</td>
<td>12.81%</td>
</tr>
<tr>
<td>Pus</td>
<td>22</td>
<td>07.82%</td>
</tr>
<tr>
<td>Aspirates</td>
<td>09</td>
<td>03.20%</td>
</tr>
</tbody>
</table>
Table-2: MTB detected in different samples received from HIV patients

<table>
<thead>
<tr>
<th>Samples</th>
<th>Total Positives</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>FNAC</td>
<td>05</td>
<td>62.5%</td>
</tr>
<tr>
<td>Pleural fluid</td>
<td>01</td>
<td>12.5%</td>
</tr>
<tr>
<td>Ascitic fluid</td>
<td>00</td>
<td>00.0%</td>
</tr>
<tr>
<td>CSF</td>
<td>00</td>
<td>00.0%</td>
</tr>
<tr>
<td>Pus</td>
<td>02</td>
<td>25.0%</td>
</tr>
<tr>
<td>Aspirates</td>
<td>00</td>
<td>00.0%</td>
</tr>
</tbody>
</table>

Table-3: MTB positives among HIV and Non-HIV patients.

<table>
<thead>
<tr>
<th>HIV status</th>
<th>Total</th>
<th>Mtb positive</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>24</td>
<td>8</td>
<td>33.3%</td>
</tr>
<tr>
<td>Negative</td>
<td>257</td>
<td>59</td>
<td>22.9%</td>
</tr>
</tbody>
</table>

Figure 2: CBNAAT Results

Figure-3: RIF Resistance
Extra pulmonary tuberculosis constitutes 20% of burden of TB globally. EPTB is a pauci-bacillary disease as number of bacteria are less. Conventional methods (histology and smear microscopy) are not diagnostic and diagnostic methods (culture methods) are time consuming. Therefore, a need for new and rapid diagnostic methods, nucleic acid amplification techniques like GeneXpert (CBNAAT) [7]. Numerous studies have assessed the yield of PCR techniques for the diagnosis of EPTB [8,9,10]. Sanjay G M et al., conducted a study on trends of EPTB and found higher detection of EPTB cases in younger age group [11]. Similar results were found in our study. Maximum cases were in the age group 21-30 years were MTB was detected in 23 (34.3%) cases. Our several findings are consistent with TB details of other studies like Balasubramanian et al., where males have higher MTB diagnostic rates compared to females [12]. A study conducted by Ullah I et al., on EPTB using CBNAAT detected MTB in 60 (35.7%) samples, similar to our study where MTB detected in 67(23.8%). CBNAAT showed, 86% of Sensitivity, 88.4% of specificity and 71.7% of positive predictive value while negative predictive value was high i.e 95.1% in their study [13]. Scott L E et al., conducted a study on diagnostic accuracy of CBNAAT for EPTB specimens taking culture as reference, CBNAAT’s overall sensitivity was low (59%) while specificity was high.
(92%) and Highest sensitivities was obtained for pus (91%) followed by lymph node aspirates (80%). Study also found that CBNAAT is not much affected by bacterial contamination reducing diagnostic delay compared to traditional methods like culture [14]. Sanjay G M et al found maximum cases of pleural effusion-133(49.81%) with MTB detected in 13 (9.77%). In our study maximum MTB cases were detected in FNAC samples-35 (47.9%) of total 73 cases. Other samples processed were 115 cases of pleural fluid in which MTB detected in 13(11.3%), 22 pus samples with MTB detected in 8(36.3%), 26 ascitic fluid in which MTB detected in 2(7.69%), 36 CSF samples with MTB detection in 4(11.1%) and 9 aspirates where 5 (55.5%) were detected positive for MTB.

In a study done by Armand et al, among individual extrapulmonary samples, the sensitivity of CBNAAT was highest among lymph nodes (94.7%). Inclusion of CBNAAT in the initial diagnosis of tubercular lymphadenopathy in addition to the FNAC would decrease the over diagnosis of tuberculosis and injudicious use of anti-tuberculosis treatment [15]. Similarly, in our study maximum MTB was detected in FNAC of LN samples.

Mittal M et al., on comparison of diagnostic yield of CBNAAT and ZN (Ziehl-Neelsen) staining from HIV and non-HIV patients with extra-pulmonary tuberculosis in their study showed of 81 extra pulmonary samples processed, 21(25.9%) from HIV and 60(74%) from non-HIV patients. Out of 21 HIV patients, 4(19.05%) and 9(42.85%) were positive for ZN stain and GeneXpert respectively. Among 60 non-HIV patients 7(11.66%) and 19 (31.66%) were positive for ZN stain and GeneXpert respectively. Studies found that CBNAAT is more sensitive than ZN staining in EPTB while both HIV and non-HIV patients have a same yield [16]. In our study MTB was detected in 8 out of 24(33.3%) HIV patients while in non-HIV patients, MTB detected in 59 out of 257(22.9%). Molecular techniques have subsequently changed the field of tuberculosis diagnosis in both pulmonary and extra-pulmonary cases and have been proven to yield rapid results [17].

Conclusion

CBNAAT is a rapid test to confirm presence of MTB with simultaneous detection of rifampicin resistance in EPTB. Before the advent of CBNAAT, diagnostic test for all clinically suspected EPTB cases use to receive empirical ATT which was a burden on both patient and health care system. Introduction of CBNAAT for EPTB, has an impact on early detection, treatment and outcome as most presumptive cases have a confirmed diagnosis. It has dramatically reduced the treatment abuse in suspected cases because of lack of confirmatory tests. This study also helped to know epidemiology of our locality.

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Reference


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