Multicentric validation of indigenous molecular test Truenat™ MTB for detection of Mycobacterium tuberculosis in sputum samples from presumptive pulmonary tuberculosis patients in comparison with reference standards

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Background & objectives: Early case detection is essential to interrupt transmission and to prevent further spread of tuberculosis (TB) in high endemic settings. Nucleic acid amplification tests (NAATs) with visual read-outs are ideal as point-of-care tests. Truenat™ MTB is an indigenous chip-based NAAT for detection of Mycobacterium tuberculosis, which involves extraction of DNA and real-time polymerase chain reaction (PCR) using portable, automated, battery-operated instruments. The current multicentric study was aimed to evaluate Truenat for detection of MTB in sputum samples obtained from patients with presumptive pulmonary TB with reference to culture as gold standard and Xpert as a comparator.

Methods: The study was conducted at four sites, namely ICMR-National Institute for Research in Tuberculosis, Chennai; All India Institute of Medical Sciences, New Delhi; ICMR-National JALMA Institute for Leprosy and Other Mycobacterial Diseases, Agra; and National Institute of TB and Respiratory Diseases, New Delhi. Patients suspected to have TB were screened for eligibility. Two sputum samples were collected from each patient. Tests included smear, Xpert and Truenat directly from the sputum sample and culture by Lowenstein-Jensen (L-J) medium and MGIT960 from decontaminated pellets. Sample used for Truenat assay was coded. Resolution of Truenat false positives was done using an in-house PCR with TRC4 primers.

Results: The study enrolled 2419 presumptive TB patients after screening 2465 patients, and 3541 sputum samples were collected from the enrolled patients. Results of 2623 samples were available for analysis.
Tuberculosis (TB) control in disease endemic high-burden countries poses a challenge and requires rapid, cost-effective, reliable and user-friendly diagnostics. Although TB is treatable, the disease burden in India remains worrisome and the mortality rate continues to be high. The National TB Control Programme under the Government of India has been pushing various agencies to focus their efforts in search of reliable and speedier diagnostic tools. At present, the primary diagnostic tool used in most peripheral settings is sputum smear. However, due to its low and variable sensitivity, almost 50 per cent of cases are missed. Thus, limitation in diagnosis becomes the source for new infections allowing false-negative cases to spread the disease.

Culture methods including the commercially available fully automated liquid mycobacterial growth indicator tube (MGIT) are complex, slow, require specialized laboratories and are still not widely available in high-endemic countries. In addition, the high contamination rate observed in liquid culture poses another challenge.

Affordable, high-sensitivity nucleic acid amplification (NAA)-based technologies using closed systems could be the answer to improve access to diagnosis in high-burden settings. In addition, technologies requiring minimal infrastructure, especially those that are portable, automated with visual readout of results, could bring testing as near to the patient as possible. Commercial NAA tests (NAATs) such as Xpert MTB/RIF assay (Xpert, Cepheid, USA) and line probe assay (LPA) (GenoType MTBDR plus, Hain Lifescience, Germany) have been widely studied, and the WHO endorsed these tests for detection of Mycobacterium tuberculosis complex (MTBC) and rifampicin resistance (RR). Xpert simultaneously detects both MTBC and RR within two hours while LPA detects MTBC, RR and resistance to isoniazid (INH) in 48 h. Both these technologies have been introduced and are being scaled up within the National Tuberculosis Elimination Programme (NTEP) of India. With the successful implementation of these technologies, many new technologies were developed along similar lines as fast followers that require stringent evaluation for use in India.

The Truenat™ MTB (Molbio Diagnostics, Goa) is a chip-based NAAT developed for detection of M. tuberculosis from sputum samples. The test involves sputum processing using Trueprep-MAG™ (a nanoparticle-based protocol run on a battery-operated device) and real-time polymerase chain reaction (PCR) performed on the Truelab-UNO™ analyzer (handheld battery-operated thermal cycler). The Truenat™ MTB has been evaluated in India in comparison with acid-fast bacillus (AFB) smear, culture and an in-house PCR as a comparator. The test was reported to have a sensitivity and specificity of 91.1% [confidence interval (CI): 89.1-94.7%] and 100% (CI: 90.0-100.0%), respectively, as compared to the in-house nested PCR with 90.58% (CI: 85.5-94.3%) sensitivity and 91.4% (CI: 76.9-98.2%) specificity.

The present multicentre study was aimed to evaluate Truenat™ MTB assay for rapid detection of MTB from sputum samples of presumptive TB patients in comparison with culture as a gold standard and Xpert as a comparator and to resolve discrepancies using a comprehensive reference standard.

**Material & Methods**

The blinded, cross-sectional, multicentre study was conducted at four sites in India [ICMR-National Institute for Research in Tuberculosis (NIRT), Chennai; All India Institute of Medical Sciences (AIIMS), New Delhi; National Institute for Tuberculosis and Respiratory Diseases (NITRD), New Delhi; and National JALMA Institute for Leprosy and Other Mycobacterial Diseases (NJIL), Agra]. The study protocol was approved by the individual Institutional Ethics Committees. All four sites were certified by the NTEP to perform culture by solid, liquid and molecular.
methods and participated in intra-laboratory quality assurance for all methods.

**Study design:** Sputum samples from presumptive TB patients were collected and subjected to smear, Xpert and Truenat from raw sample followed by culture on solid Lowenstein-Jensen (L-J) medium and MGIT. Performance of Truenat in comparison with culture was analyzed and discrepancy resolution of culture-negative - Truenat-positive samples was done using an established in-house PCR. In the current study, in-house PCR using TRC4 primers was used instead of the widely tested IS6110 primers considering its discriminating limitations with regard to south Indian MTB strain\(^1^9\). Following discrepancy resolution and identification of true positives defined by demonstration of MTB in any test other than culture (smear or Xpert or TRC4 PCR), further analysis to evaluate the ability of Truenat to identify true positives was done using a comprehensive reference standard including all TB diagnostic tests used in the study.

**Sample size:** Based on an earlier study\(^1^8\), the sample size was calculated to achieve critical performance targets among groups with (i) sensitivity of 99 per cent (CI of 2%) among smear-positive, culture-positive patients requiring 150 patients; (ii) sensitivity of 75 per cent among smear-negative, culture-positive patients and CI of 7.5 per cent requiring 180 patients; and (iii) specificity of 98 per cent among symptomatic non-TB patients and CI of two per cent requiring 250 patients. Hence, the minimum required group size was 700 after adjusting for either 20 per cent refusal or dropout rate. All values were calculated based on the estimation procedure. In order to enrol 700 patients, an estimated 2000 presumptive TB patients were needed to be screened and recruited across the four sites.

**Inclusion and exclusion criteria:** Adults (≥18 yr) with clinical suspicion of pulmonary TB and persistent productive cough for ≥two weeks were included. Patients who had received one or more doses of anti-TB medication in the 60 days before screening were excluded\(^3\).

**Sample population:** During the study period between December 2014 and April 2017, consecutive adult presumptive pulmonary TB patients were screened at the four centres for eligibility and written informed consent was taken from each participant before sample collection. Less than two per cent of patients refused consent. Two sputum samples of 4 ml each were collected from each patient - spot specimen on day 1 and morning specimen on day 2. Samples were transported to the laboratories and processed on the same day except on holidays when the samples were stored at 4-10°C in the laboratories. Whenever the patient failed to provide a second sample or if the second sample was found to be less than 4 ml, all tests were done with the first sample. A detailed study workflow is presented in Figure 1.

**Methods:** Sputum samples were homogenized using sterile glass beads. Five hundred microlitres of sputum from each sample from the same patient was pooled and subjected to Xpert testing\(^9\). Another 0.5 ml aliquot of the sample was used for Truenat testing. The aliquot for Truenat testing was coded and the technician performing Truenat was blinded to the results from other tests. Briefly, to 0.5 ml of homogenized sputum two drops

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**Fig. 1.** Workflow of the study samples. NaLC-NaOH, N-acetyl-L-cysteine-sodium hydroxide; L-J, Lowenstein-Jensen medium; MGIT, mycobacterial growth indicator tube; ICT, immunochromatography test. All positive cultures confirmed by ICT.
of liquefaction buffer provided with the Trueprep® Auto Sample Pre-treatment Pack (TSPP) was added, shaken gently and allowed to stand for five minutes with intermittent shaking. The liquefied mixture was transferred to 2.5 ml of lysis buffer from TSPP, shaken vigorously and allowed to stand for 3-5 minutes. The whole content was transferred to the sample chamber of the cartridge provided in the Trueprep® Auto Sputum Sample Prep Kit using a Pasteur pipette. The cartridge was placed in the Trueprep® Auto Sample Prep Device. The device automatically extracted DNA in 25 min which was collected in the DNA elution chamber covered with aluminium foil. The DNA was transferred to a screw-capped cryovial. Six microlitres of the DNA was placed on the Truenat™ MTB micro-PCR chip placed on the chip tray of the Truelab™ Uno (V1.5) analyzer. The test was completed in 35 min and the displayed results were recorded. The remaining DNA was stored at −20°C. Samples that yielded invalid or error results were repeated once again to obtain interpretable results and those which gave similar results on repeat testing were considered invalid/error and were excluded from the analysis.

The remaining sputum sample was subjected to smear by fluorescence microscopy and subsequent decontamination using the standard N-acetyl-L-cysteine-sodium hydroxide (NaLC-NaOH) method20. The resultant pellet was used for inoculating MGIT960 tube (Becton Dickinson, USA) and L-J slopes (in-house preparation) (Fig. 1) as per the standard protocols. Positive cultures were tested by the standard identification tests including AFB smear, contamination check by inoculation on brain-heart infusion (BHI) agar and confirmation of the presence of MTBC using the MPT64 antigen-based immunochromatographic lateral flow assay21. Cultures that showed growth on BHI agar (HiMedia, Mumbai) were considered as contaminated. Cultures that showed the presence of MTBC along with contamination were decontaminated, and a repeat culture was done in MGIT system22. Once flagged positive by MGIT instrument, all three identification tests were repeated on the culture and final results were based on the second culture. Results from all the tests were compared after excluding invalids/errors/ contaminations. Evaluation of the Truenat MTB test was performed in comparison with culture as the gold standard and Xpert as a comparator for detection of M. tuberculosis.

Performance characteristics such as sensitivity, specificity and kappa were calculated using the Statistical Package for the Social Sciences (SPSS) version 25.0 (IBM, Corp., Armonk, NY, USA). P=0.05 was considered statistically significant for all tests.

For resolution of false positives, DNA elution was subjected to an in-house PCR using TRC4 primers. The primers used were TRC4 primers 1 (5'-GACAACGACGTGCGCCTACT-3') and 2 (5'-GACCGAATTAGCGTAGCTCC-3'). The 18-mer primers amplify 173 bp long single target of the repetitive element of TRC4. The fragment is located in the open reading frame of RV0697 in the MTB genome23. The two-step PCR conditions for DNA amplification were an initial denaturation step at 95°C for one minute, followed by 45 cycles of 95°C for 10 seconds and 65°C for 34 seconds. The final product assessed by two per cent gel electrophoresis was a 173 bp fragment24 as shown in Figure 2.

Results

The study enrolled 2419 adult presumptive TB patients after screening 2465 patients, and 3541 sputum samples were collected (Table I). Among 3541 samples, valid results for all tests were available for comparison from 2623 samples after elimination of errors, invalids and contaminations and these were considered for analysis. The results for all samples tested and among the analyzable set are presented in Table II.

Among the target groups, the sensitivity of Truenat for identifying MTBC was 92.9 per cent among smear- and culture-positive samples and 75 per cent among smear-negative culture-positive samples in comparison with 86 and 60.6 per cent, respectively, by Xpert. The
### Table I. Site-wise distribution of patients enrolled and samples collected

| Site                                                                 | Patients | Samples |
|----------------------------------------------------------------------|----------|---------|
| All India Institute of Medical Sciences, New Delhi                  | 747      | 747     |
| National JALMA Institute for Leprosy and Other Mycobacterial Diseases, Agra | 550      | 550     |
| ICMR-National Institute for Research in Tuberculosis, Chennai        | 572      | 1144    |
| National Institute for Tuberculosis and Respiratory Diseases, New Delhi | 550      | 1100    |
| **Total**                                                           | **2419** | **3541**|

Four sites collectively enrolled 2419 patients including smear positives and smear negatives.

### Table II. Distribution of all test results in 2623 samples

| Test       | Overall samples (n=3541), n (%) | Samples taken for comparative analysis (n=2623), n (%) |
|------------|---------------------------------|------------------------------------------------------|
|            | Positive | Negative | Invalid/error/contamination | Positive | Negative | Invalid/error/contamination |
| Smear      | 999 (28.2) | 2542 (71.8) | 0 (0.0) | 777 (29.6) | 1846 (70.4) | 0 (0.0) |
| Culture    | 1113 (31.4) | 2317 (65.4) | 111 (3.1) | 941 (35.9) | 1682 (64.1) | 99 (2.8) |
| Xpert<sup>@</sup> | 1055 (29.8) | 1904 (53.8) | 32 (0.9) | 972 (37.1) | 1651 (62.9) | 32 (0.9) |
| Truenat    | 1590 (44.9) | 1707 (48.2) | 244 (6.9) | 1272 (48.5) | 1351 (51.5) | 237 (6.7) |

<sup>@</sup>550 samples could not be processed due to non-availability of Xpert cartridges at the time of study and were excluded from the analysis. Truenat yielded maximum positivity of 44.9% among all samples tested and 48.5% among those considered for comparative analysis while the same for Xpert was 29.8 and 37.1%, respectively.

### Table III. Performance of molecular tests among the target groups

| Target group  | Total | TN positive (%) | Xpert positive (%) | P     |
|---------------|-------|-----------------|--------------------|-------|
| Sm+ Cult+     | 697   | 648 (92.9)      | 602 (86)           | <0.001|
| Sm− Cult+     | 244   | 183 (75)        | 148 (60.6)         | <0.001|
| Sm+ Cult−     | 80    | 65 (81.2)       | 62 (77.5)          | 0.702 |
| Sm− Cult−     | 1602  | 376 (23.4)      | 160 (9.98)         | <0.001|

TN: Truenat; Sm+ Cult+, smear-positive/culture-positive samples; Sm− Cult+, smear-negative/culture-positive samples; Sm+ Cult−, smear-positive/culture-negative samples; Sm− Cult−, smear-negative/culture-negative samples. MTBC, *Mycobacterium tuberculosis* complex

Differences in sensitivities between the two tests in these target groups were significant (Table III). Among the 80 smear-positive culture-negative samples, Truenat detected MTBC in 65 (81.2%) while Xpert was positive in 62 (77.5%), however, the difference was not significant. Truenat and Xpert were also positive in 376 (23.4%) and 160 (9.98%) samples, respectively, among 1602 smear- and culture-negative samples (Table III). Direct comparison with Xpert showed that among 972 samples that were positive by Xpert, Truenat was positive for 859 (88.3%), and among 1651 samples that were Xpert negative, 1238 (75%) were negative by Truenat.

Concordance analysis of Truenat and Xpert with culture yielded an overall sensitivity of 88.3 (95% CI: 86.1-90%) and 79.7 per cent (95% CI: 77-82.2%) for Truenat and Xpert, respectively (Table IV). The overall specificities of Truenat and Xpert were 73.8 per cent (95% CI: 71.6-75.9%) and 86.8 per cent (95% CI: 85.1-88.4%). Analysis of smear-positive and smear-negative samples independently yielded similar patterns of higher sensitivity for Truenat and higher specificity for Xpert in both types of samples (Tables V and VI).

Among all 441 samples that were culture-negative Truenat positive, 311 samples showed evidence of TB by other tests [smear +ve, 18; Xpert +ve, 160; smear and Xpert +ve, 62 and PCR (TRC<sub>4</sub> primer) +ve, 71] and were considered true MTB positives. The ability of Truenat to identify true positives in comparison with a comprehensive reference standard (CRS)
Table IV. Performances of Truenat and Xpert in comparison with culture

|          | Truenat | Xpert |       |       |       |       |       |       |       |       |       |
|----------|---------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
|          | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total |       |
|          | 831     | 441    | 1272   | 750    | 222    | 972   |
| Positive | 110     | 1241   | 1351   | 191    | 1460   | 1651  |
| Total    | 941     | 1682   | 2623   | 941    | 1682   | 2623  |

Sensitivity: 88.3% (86.1-90.3%)   Specificity: 73.8% (71.6-75.9%)   Sensitivity: 79.7% (77.0-82.2%)   Specificity: 86.8% (85.1-88.4%)

Among all samples tested, Truenat demonstrated higher sensitivity than Xpert and difference was significant \(P<0.001\). Xpert showed significantly higher specificity \(P<0.001\) than Truenat.

Table V. Performances of Truenat and Xpert in comparison with culture among smear-positive samples

|          | Truenat | Xpert |       |       |       |       |       |       |       |       |       |
|----------|---------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
|          | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total |       |
|          | 648     | 65     | 713    | 602    | 62     | 664   |
| Positive | 49      | 15     | 64     | 95     | 18     | 113   |
| Total    | 697     | 80     | 777    | 697    | 80     | 777   |

Sensitivity: 93% (90.8-94.8%)   Specificity: 18.8% (10.9-29%)   Sensitivity: 86.4% (83.6-88.8%)   Specificity: 22.5% (13.9-33.2%)

Sensitivities of Truenat and Xpert for detection of MTB among smear-positive samples in comparison with culture were 93 and 86.4%, respectively. The difference in sensitivity was significant \(P<0.001\) while specificities were 18.8 and 22.5%, and the difference was not significant.

Table VI. Performances of Truenat and Xpert in comparison with culture among smear-negative samples

|          | Truenat | Xpert |       |       |       |       |       |       |       |       |       |
|----------|---------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
|          | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total |       |
|          | 183     | 376    | 559    | 148    | 160    | 308   |
| Positive | 61      | 1226   | 1287   | 96     | 1442   | 1538  |
| Total    | 244     | 1602   | 1846   | 244    | 1602   | 1846  |

Sensitivity: 75% (69.1-80.3%)   Specificity: 76.5% (74.4-78.6%)   Sensitivity: 60.7% (54.2-66.8%)   Specificity: 90.0% (88.4-91.4%)

Sensitivity of Truenat and Xpert in detecting MTB among smear-negative samples in comparison with culture was 75 and 60.7%, while specificities were 76.5 and 90.0%, respectively. The differences in sensitivity and specificity between Truenat and Xpert were significant \(P<0.001\).

Table VII. Performances of Truenat in comparison with comprehensive reference standard

|          | Truenat | CRS    | Total |       |       |       |       |       |       |       |       |
|----------|---------|--------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
|          | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total | Positive| Negative| Total |       |
|          | 1142    | 130    | 1272   | 1122   | 130    | 1252  |
| Positive | 110     | 1241   | 1351   | 1241   | 130    | 2552  |
| Total    | 1252    | 1371   | 2623   | 1252   | 1371   | 2623  |

Sensitivities of Truenat and Xpert for detection of MTB among smear-negative samples in comparison with culture were 93 and 86.4%, respectively. The difference in sensitivity was significant \(P<0.001\) while specificities were 18.8 and 22.5%, and the difference was not significant.

Discussion

Truenat was evaluated earlier in India by Hinduja Hospitals, Mumbai. Sputum samples from 226 presumptive TB patients were tested by smear, culture, Truenat and an in-house PCR. A CRS including smear, culture, clinical findings and response to treatment was used to evaluate Truenat with the in-house PCR as a comparator. Truenat demonstrated a sensitivity including smear, culture, Xpert and TRC\(_4\) PCR showed an increase in sensitivity and specificity to 91.2 and 90.5 per cent, respectively (Table VII).
of 91.1 per cent (CI: 86.1-94.7%) and a specificity of 100 per cent (CI: 90.0-100%) in comparison with the CRS\textsuperscript{18}. The current evaluation on a larger sample size showed similar findings with Truenat yielding a sensitivity of 91.2 per cent and a specificity of 90.5 per cent in comparison with a CRS including a battery of diagnostic tests for TB. Direct comparison of Truenat with the comparator Xpert in the current study yielded a specificity of 75 per cent similar to the earlier study\textsuperscript{18} where, in comparison with the in-house PCR, Truenat yielded a specificity of 72 per cent.

Truenat demonstrated the highest overall positivity of 44.9 per cent among all the tests performed. The overall invalid rate of Truenat was 6.9 per cent in comparison with 0.9 per cent with Xpert. This finding was similar to the early error rates reported for Xpert\textsuperscript{25} following implementation.

In the present study, the higher sensitivity of Truenat in comparison with culture among the target groups demonstrated the assay’s capability to detect higher number of true positives. In comparison with culture, the assay achieved a sensitivity of 92.9 per cent as against the expected 99 per cent among smear-positive culture-positive samples, while among smear-negative culture-positive samples, the expected sensitivity of 75 per cent was achieved. These values were significantly higher than those of Xpert. However, Xpert showed a higher specificity of 90 per cent among the target group smear-negative culture-negative than Truenat (76.5%) while the overall specificities of Xpert and Truenat were 87 and 74 per cent, respectively.

The ideal way to eliminate false positivity by a TB diagnostic test is to demonstrate tubercle bacilli or its component in the same sample by other standard tests or by looking for clinical evidence for TB in the patient including response to treatment. Following discrepancy resolution, performance of Truenat was compared to a comprehensive reference standard including all available TB diagnostic tests such as smear, culture, Xpert and in-house TRC, PCR as done in an earlier work by Nikam et al\textsuperscript{18}. The sensitivity and specificity of Truenat increased to 91.2 and 90.5 per cent, respectively. Truenat failed to identify 11.6 per cent samples that were positive by culture, but it was less than those missed by Xpert (20.2%).

One key limitation of the study was the use of an in-house PCR for discrepancy resolution instead of sequencing. This was unavoidable as the only sample available for any further investigation was the DNA from the Truenat assay in the case of Truenat-positive/culture-negative samples.

The present multicentre study helped identify the operational advantages of Truenat. The assay required minimal training as technicians with minimal or nil exposure to molecular tests could perform the assay following a short training session. Truenat instruments were battery operated and temperature stable thereby demonstrating the robustness of the instruments while Xpert required continuous power supply and air-conditioning\textsuperscript{26}. No instrument failures were reported from any of the sites during the study period. Functional remote monitoring of the devices was another notable feature. Another study\textsuperscript{27} has shown Truenat to be the most cost-effective test in a point-of-care setting.

In conclusion, the study demonstrated that the precision of detecting MTB was higher with Truenat as compared with Xpert. The study also identified other parameters of the assay that may prove more advantageous over Xpert. Demonstration of these features through feasibility studies at peripheral field settings under routine NTEP conditions will be required for inclusion of the assay in the diagnostic algorithm of the national programme.

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**Conflicts of Interest:** None.

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