Evaluation of four molecular methods for the diagnosis of tuberculosis in pulmonary and blood samples from immunocompromised patients

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The present study analysed the concordance among four different molecular diagnostic methods for tuberculosis (TB) in pulmonary and blood samples from immunocompromised patients. A total of 165 blood and 194 sputum samples were collected from 181 human immunodeficiency virus (HIV)-infected patients with upper respiratory complaints, regardless of suspicious for TB. The samples were submitted for smear microscopy, culture and molecular tests: a laboratory-developed conventional polymerase chain reaction (PCR) and real-time quantitative PCR (qPCR) and the Gen-Probe and Detect-TB Ampligenix kits. The samples were handled blindly by all the technicians involved, from sample processing to results analysis. For sputum, the sensitivity and specificity were 100% and 96.7% for qPCR, 81.8% and 94.5% for Gen-Probe and 100% and 66.3% for Detect-TB, respectively. qPCR presented the best concordance with sputum culture [kappa (k) = 0.864], followed by Gen-Probe [k = 0.682]. For blood samples, qPCR showed 100% sensitivity and 92.3% specificity, with a substantial correlation with sputum culture (k = 0.754) and with the qPCR results obtained from sputum of the corresponding patient (k = 0.630). Conventional PCR demonstrated the worst results for sputum and blood, with a sensitivity of 100% vs. 88.9% and a specificity of 46.3% vs. 32%, respectively. Commercial or laboratory-developed molecular assays can overcome the difficulties in the diagnosis of TB in paucibacillary patients using conventional methods available in most laboratories.

Key words: tuberculosis - paucibacillary - Gen-Probe - Detect-TB - real-time PCR

The diagnosis of tuberculosis (TB) remains based on direct examination and solid cultures in the majority of countries with high TB prevalence (Smith et al. 1996, Chan et al. 2000, Silva et al. 2012, Nakiyingi et al. 2013). Frequently, among paucibacillary patients such as children, human immunodeficiency virus (HIV) co-infected patients and patients with the extrapulmonary form of TB (regardless of age), smear-based microscopy is negative and the sputum-culture sensitivity is low (Bollela et al. 1999, Burroughs et al. 1999, Pai et al. 2004). The use of molecular biology techniques has increased the ability to diagnose TB in paucibacillary TB (Almeda et al. 2000, Portillo-Gómez et al. 2000, Al Zahrani et al. 2001, Boehme et al. 2011, Lawn et al. 2013) and has also shortened the time of diagnosis in relation to culturing in solid media, from four-six weeks to 24-48 h (Richeldi et al. 1995, Kivihiya-Ndugga et al. 2004, Hida et al. 2012). The World Health Organization (WHO 2008) conducted a comparative analysis of 19 different commercial kits for the molecular diagnosis of TB, using the standard protocol for each test and considering endemic and non-endemic countries. The results demonstrated a mean sensitivity of 27% and specificity greater than 90% in only 30% of the tests. In that study, TB-HIV co-infection reduced test performance, regardless of whether TB was endemic to the country. The Gen-Probe Amplified-MTD Direct Test kit was the first test for detecting DNA of the Mycobacterium tuberculosis complex to gain approval from the Food and Drug Administration (FDA) (Vlaspolder et al. 1995).

Xpert MTB/RIF (Cepheid, USA) is the first fully automated assay based on hemi-nested real-time polymerase chain reaction (PCR) for the detection of TB and rifampicin resistance performed directly on untreated sputum, with results obtained in less than 2 h. The assay was approved by the WHO in 2010 (Lawn et al. 2013, Nakiyingi et al. 2013).

COBAS® Amplicor is a commercial test based on amplification, hybridisation and detection of the mycobacteria multicopy IS6110 insertion element that also has been approved by the FDA (McAdam et al. 1990, Thierry et al. 1990, Almeda et al. 2000, Huyen et al. 2013). The Detect-TB Ampligenix Biotech Company kit is a test in the pre-clinical stage of development, which associates the amplification of the mycobacteria IS6110 sequence with the reversed hybridisation of the amplification product with probes complementary to internal

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Acid-fast bacilli were detected by means of Ziehl-Neelsen staining on sputa smears. Sputa were decontaminated in accordance with the Petroff method and aliquots of 0.1 mL were placed on Lowenstein-Jansen solid medium. Blood samples did not undergo microbiological testing.

Microbiological analysis - Acid-fast bacilli were detected by means of Ziehl-Neelsen staining on sputa smears. Sputa were decontaminated in accordance with the Petroff method and aliquots of 0.1 mL were placed on Lowenstein-Jansen solid medium. Blood samples did not undergo microbiological testing.

Genomic DNA extraction - The patients’ blood samples underwent Ficoll-Paque Plus (GE Healthcare Biosciences AB, Sweden) fractioning for leucocyte isolation. An aliquot of 10⁶ leucocytes and the first of the three sputum aliquots underwent genomic DNA extraction using proteinase K protocol. Briefly, the material was suspended in a 400 μL lysozyme solution (1.25 mg/mL) and was incubated at 37°C for 16 h. Then, 75 μL of proteinase K (20 mg/mL) and 10% sodium dodecyl sulfate solution were added and the mixture was incubated at 65°C. Next, 100 μL of 5 M NaCl and 100 μL of CTAB/NaCl were added and the samples were incubated at 65°C. After 10 min of incubation, the DNA was extracted with phenol/chloroform solution and then concentrated by means of precipitation with isopropanol. The DNA precipitate was obtained through centrifugation at 16,000 g in a microcentrifuge, washed with 70% ethanol and dissolved in 50 μL of sterile water.

Conventional PCR - First, the genomic DNA underwent PCR for amplification of a target sequence on human DNA, acting as a positive control for the DNA extraction, thereby ensuring the absence of PCR inhibitors in the reaction. Next, PCR was performed with specific primers for amplification of the mycobacteria IS6110 genetic element. The total reaction volume was 50 μL and included 1.5 mM of MgCl₂, 25 pmol of each primer (sense: 5’cagctgcgcgatggcgaac and antisense: 5’taggtctgggttcgccgaaag, synthesised by Invitrogen, USA), 200 μM of each dNTP and 1.25 units of DNA polymerase (Invitrogen). The reaction conditions were one cycle of 96°C for 3 min, 30 cycles of 96°C for 30 s for denaturation, 65°C for 30 s for annealing, 72°C for 30 s for extension and, finally, a cycle of 7 min at 72°C using a thermocycler (Gradient Eppendorf, Germany). The PCR product was visualised on 2.5% agarose gel under ultraviolet light. The reaction was considered positive when the ethidium bromide-stained band corresponded to 245 bp.

Real-time PCR - Quantitative PCR (qPCR) was performed using the same set of primers for IS6110 amplification as used in conventional PCR and an internal probe (Applied Biosystems, USA) in the presence of the TaqMan® Universal PCR Master Mix and the DNA extracted from the patients’ sputum or blood. The conditions for amplification of the IS6110 element were as follows: denaturation for 15 min at 95°C followed by 45 cycles of 15 s at 94°C and 60 s at 60°C. In all PCR runs, standard curves were obtained using plasmid DNA encompassing the mycobacteria IS6110 sequence as the reaction positive control and all samples were tested in duplicate. Reaction with no DNA added was used as the negative control, also in duplicate. The reaction was run in an ABI 7500 machine with setup for absolute quantification, using a standard curve prepared with serial 10X dilution from 10 ng (cycle threshold (Ct) = 14) to 100 fg (Ct = 37) of plasmid control DNA in triplicate. The efficiency of the amplification reaction was 1.8, with a threshold set at 0.02 based on data from 26 different experiments. The slope was -3.84 and R² = 0.999.

Detect-TB - The second sputum aliquot was tested for the presence of mycobacteria genome using the Detect-TB Ampligenix Biotech Company kit (Belo Horiz...
All DNA extractions were performed at the Molecular Biology Laboratory of IMIP, inside a biosafety cabinet located in a room with restricted use to clinic sample handling. The PCR reactions were prepared in another room specifically designed for this purpose, free of biological samples and also inside a biosafety cabinet. The hybridisation procedures and the analysis of PCR products in gel were performed in the main area of the molecular biology laboratory.

For every conventional or real-time PCR run, two negative controls were run: one reaction was performed without the DNA template and another included genomic DNA of a healthy individual to identify possible non-specific DNA amplification.

To avoid interpretation bias, the research teams at the hospitals, LACEN-PE and the Molecular Biology Laboratory at IMIP, were different and the samples received a number at the hospital, a different number at LACEN-PE and yet another at the Molecular Biology laboratory. All patient identification numbers were noted in the patient database at the Molecular Biology laboratory; however, the codes were only broken for analysis at the end of the cohort study.

**Statistical analysis** - The sensitivity, specificity and the positive and negative predictive values (NPV) for each of the molecular tests evaluated for the diagnosis of TB in HIV-infected patients were calculated considering sputum culture as the gold standard, using a 2 x 2 table. The concordance between tests was evaluated using the kappa (k) test (Epi Info v.6.04 software). The receiver operating characteristic (ROC) curves were performed using MedCalc software, Ostend, Belgium.

**Ethics** - The Aggeu Magalhães Research Centre Institutional Review Board approved this study under the registration CAAE: 0007.0.095.000-07. The authors declare no conflict of interest and do not have a direct financial relation with the commercial identity mentioned in this manuscript.

**RESULTS**

**Sputum samples** - Among 194 sputum samples from 181 different HIV-infected patients, 19 were culture-positive for *M. tuberculosis*, of which only eight were smear-positive.

Comparing the ROC curves of the different tests with sputum culture as the reference, the best results were associated with qPCR methodology (Fig. 2A). qPCR presented 100% sensitivity, 96.7% specificity, 78.6% positive predictive value (PPV), 100% NPV and 97% accuracy (Table I). qPCR results demonstrated excellent concordance with culture (k = 0.864) and moderate concordance with Gen-Probe results (k = 0.586) (Table II). The Ct average of culture-positive, smear-positive sputa (32.5; range, 24.4-36.4) was similar to culture-positive, smear-negative sputa (33.4; range 30.3-35.4). Three samples presented culture-negative and positive-qPCR results, with Ct of 18, 19 and 21.

Gen-Probe presented the second-best results with 81.8% sensitivity, 94.5% specificity, 64.3% PPV, 97.7% NPV and 93% accuracy. A substantial concordance between Gen-Probe and culture results was observed (k = 0.682)
Detect-TB demonstrated a sensitivity of 100%, but a specificity of 66.3%, corresponding to a PPV of 24.4%. ROC curves constructed with different cut-off values for testing Detect-TB showed that the best results for sensitivity and specificity occurred with a cut-off of 300 (100% sensitivity, 67.4% specificity) and 275 (100% sensitivity, 66.3% specificity). In this study, the cut-off used was 275, which was indicated by the manufacturer (Fig. 2B). However, higher cut-off values might improve specificity with some compromised of sensitivity.

The lowest specificity was associated with conventional PCR performed from sputum (46.3%) and blood (32%) samples (Table I).

**Blood samples** - Concerning the molecular tests performed on the patients’ blood samples, we only took into consideration samples from patients for whom sputum also had been collected for definition of the TB cases based on sputum culture.

Regarding blood samples, qPCR showed 100% sensitivity, 92.3% specificity, 61.1% PPV and 100% NPV in comparison to the sputum culture, whereas the in-house PCR showed sensitivity and specificity values of 88.9% and 32%, respectively (Fig. 2C). A substantial concordance was observed between qPCR results obtained from blood and positive culture on sputum (k = 0.754) and also between qPCR results from blood and sputa (k = 0.630) (Table II).

**DISCUSSION**

Many studies on the accuracy of molecular assays have been published on the diagnosis of pulmonary and non-pulmonary TB, both associated with smear-positive or smear-negative sputum samples and in HIV-infected or non-infected patients (Peter et al. 2012, Lawn et al. 2013, Theron et al. 2013, Walusimbi et al. 2013, Raizada et al. 2014).

**TABLE I**

| Reference x tests (sample) | n   | PPV (%) | NPV (%) | Sensitivity (%) | Specificity (%) |
|---------------------------|-----|---------|---------|-----------------|-----------------|
| Culture x PCR (sputum)    | 194 | 16.8 (9.4-29.5) | 100 (88.8-100) | 100 (67.9-100) | 46.3 (32.7-53.7) |
| Culture x qPCR (sputum)   | 102 | 78.60 (48.8-94.3) | 100 (94.8-100) | 100 (67.9-100) | 96.70 (90-99.1) |
| Culture x Detect-TB (sputum) | 194 | 24.36 (12.2-37) | 100 (91.6-100) | 100 (67.9-100) | 66.30 (47.4-68.3) |
| Culture x Gen-Probe (sputum) | 102 | 64.30 (35.6-86) | 97.70 (91.3-99.6) | 81.80 (47.8-96.8) | 94.50 (87.1-98) |
| Culture x PCR (blood)     | 165 | 13.8 (8.3-21.7) | 95.9 (89.9-99.3) | 88.90 (63.9-98.3) | 32 (24.7-40.2) |
| Culture x qPCR (blood)    | 102 | 61.1 (36.1-81.7) | 100 (94.6-100) | 100 (67.9-100) | 92.30 (84.3-96.6) |

Detect-TB is a test in developing and Gen-Probe a commercial molecular test for TB diagnosis. Blood and sputum samples were collected on the same day. PCR: in house polymerase chain reaction; qPCR: quantitative real-time PCR.
Smear-based microscopy is the most commonly used approach for pulmonary TB detection, demonstrating 70% sensitivity compared with culture associated with a clinical definition of the disease as the gold standard; however, in HIV-infected patients the smear-based sensitivity might be lower than 40% (Boehme et al. 2011, Lawn et al. 2011, Scherer et al. 2011). In our study, the smear-based sensitivity was 42%. This was mainly related to the low bacillary load in sputum from HIV-infected patients (O’Grady et al. 2012).

We observed a great variation in the accuracies of the four tests evaluated considering that all were PCR-based assays and all used samples obtained from the same sputum. qPCR demonstrated superior performance with 97% accuracy, followed by Gen-Probe (93%), Detect-TB (70%) and conventional PCR (52%) for the detection of pulmonary TB infection from sputum. Recent publications regarding the molecular detection of pulmonary TB in HIV-infected patients also have shown variations in assay sensitivity (53-100%) between different or even the same assay (Table III).

For TB detection in sputum, our qPCR assay presented 100% sensitivity and 96.7% specificity, comparable to the sensitivity of 80-95% and the specificity of 95-100% reported by others who also developed the TaqMan assay for detection of the IS6110 element (Gomez et al. 2011, Lira et al. 2013, Albuquerque et al. 2014). qPCR was positive for 100% of the smear-negative, culture-positive patients, providing earlier TB diagnosis for more TB-suspected patients than smear-based microscopy (7% vs. 4%) in HIV-infected patients. Improvement in pulmonary TB detection (25%) by means of molecular assay was reported in Chile, a TB non-endemic area, confirming the power of molecular tools in TB diagnosis for TB-control programs (Balcells et al. 2012). Three patients with false-positive qPCR from sputum received empirical treatment for TB within six months of being tested and the clinical-symptom improvement confirmed their disease status. The withdrawal of a considerable (1.5 mL) amount of sputum for molecular analysis prior to sputum culturing might explain the absence of acid-fast bacillary growth on solid media, likely due to the loss of bacillary viability as a consequence of the sputum decontamination procedure (Michelon et al. 2011).

The sensitivity and specificity of the Gen-Probe assay as reported in the literature ranges from 92-100% and from 85-100%, respectively, but in HIV-positive patients the sensitivity might be lower than 42% (Syre et al. 2009, Davis et al. 2011, Papaventsis et al. 2012, Roberts et al. 2012). In our study, the performance of Gen-Probe was similar to qPCR (93% vs. 97% accuracy), with 81.8% sensitivity and 94.5% specificity for TB detection in sputum. Five false-positive cases were observed, two of whom received empirical TB treatment with improvement in clinical symptoms. In contrast to the qPCR assay, the Gen-Probe assay was associated with two false-negative cases, possibly due to the insufficient target DNA in the 0.5 mL sputum aliquot used, considering that the Gene-Probe protocol recommends the collection of 10 mL of sputum for decontamination and concentration procedures prior to use. The presence of PCR inhibitors in the sample may also have induced false-negative reactions with the Gen-Probe assay, which were not observed in the qPCR assay because, in this latter case, DNA was extracted using proteinase K digestion followed by a purification step performed with phenol-chloroform.

Cases identified as false-positives by the Detect-TB assay were responsible for the low specificity (66.3%) and PPV (24.4%) observed in our study. The causes for this performance are not clear, because a sensitivity of

### Table II

Concordance between microbiological and molecular tests for diagnosis of tuberculosis (TB)

| Tests (sample)                                | n  | Agreement | Kappa | Z       | p       |
|----------------------------------------------|----|-----------|-------|---------|---------|
| Culture X PCR (sputum)                       | 194| 0.490     | 0.139 | 2.76    | 0.0029  |
| Culture X qPCR (sputum)                      | 102| 0.971     | 0.864 | 8.80    | 0.0000  |
| Culture X Detect-TB (sputum)                 | 194| 0.628     | 0.231 | 3.65    | 0.0002  |
| Culture X Gen-Probe (sputum)                 | 102| 0.931     | 0.682 | 6.95    | 0.0000  |
| Culture X PCR (blood)                        | 165| 0.382     | 0.062 | 1.83    | 0.0337  |
| Culture X qPCR (blood)                       | 102| 0.931     | 0.754 | 7.59    | 0.0000  |
| PCR (sputum) X qPCR (sputum)                 | 102| 0.520     | 0.179 | 3.17    | 0.0007  |
| PCR (sputum) X Gen-Probe (sputum)            | 102| 0.647     | 0.301 | 3.15    | 0.0008  |
| qPCR (sputum) X Detect-TB (sputum)           | 102| 0.480     | 0.112 | 1.99    | 0.0236  |
| qPCR (sputum) X Gen-Probe (sputum)           | 102| 0.657     | 0.294 | 4.19    | 0.0000  |
| PCR (blood) X qPCR (blood)                   | 102| 0.902     | 0.586 | 5.92    | 0.0000  |
| qPCR (sputum) X qPCR (blood)                 | 102| 0.461     | 0.157 | 2.95    | 0.0016  |
| qPCR (sputum) X Gen-Probe (sputum)           | 102| 0.902     | 0.630 | 6.44    | 0.0000  |
| qPCR (sputum) X Detect-TB (sputum)           | 102| 0.520     | 0.179 | 3.17    | 0.0007  |

Detect-TB is a test in developing and Gen-Probe a commercial molecular test for TB diagnosis. Blood and sputum samples were collected on the same day. Culture was performed only in sputa. PCR: in house polymerase chain reaction; qPCR: quantitative real-time PCR.
| Reference | HIV-infected population | Biological sample | Sampling (n) | Methodology | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) | Target suspected cases | Gold stantard |
|-----------|-------------------------|-------------------|--------------|-------------|----------------|----------------|---------|---------|-----------------------|---------------|
| This paper | Recife, Brazil          | Blood             | 102          | qPCR        | 100            | 92.3           | 61.1    | 100     | In and outpatient     | Solid culture |
|           |                         | Sputa             | 102          | qPCR        | 100            | 96.7           | 78.6    | 100     |                       |               |
|           |                         |                   | 102          | Gene-Probe  | 81.8           | 94.5           | 64.3    | 97.7    |                       |               |
|           |                         |                   |              | Detect-TB   | 66.3           | 100            | 24.4    | 100     |                       |               |
| Albuquerque et al. (2014) | Recife, Brazil | Sputa             | 140          | qPCR        | 87.2           | 98.9           | 97.6    | 93.9    | In and outpatient     | Solid culture |
| Feasey et al. (2013) | Malawi, Africa         | Blood             | 104          | Xpert MTB/RIF | 21             | 100            | 100     | 64      | In and outpatient     | Sputum and solid culture |
| Abed Al-Darraji et al. (2013) | Malaysia            | Sputa             | 125          | Xpert MTB/RIF | 53.3           | 100            | 94      |         | Prisoners             | Liquid culture |
| Silva et al. (2012) | Santa Catarina, Brazil | Sputa             | 85           | PCR         | 59             | 33             | 87      | 10      | Inpatient             | Solid culture |
| O’Grady et al. (2012) | Lusaka, Zambia        | Sputa             | 408          | Xpert MTB/RIF | 88.2           | 95.1           | 92.2    | 92.5    | Inpatient             | Liquid culture |
| Peter et al. (2012) | Cape Town, South Africa | Urine             | 113          | Xpert MTB/RIF | 48             | 98             | -       | -       | Inpatient             | Liquid culture |
| Baleells et al. (2012) | Santiago, Chile       | Sputa             | 160          | Xpert MTB/RIF | 91.7           | 99.3           | 94.7    | 99.3    | Inpatient             | Liquid culture |
| Carriquiry et al. (2012) | Lima, Peru            | Sputa             | 131          | Xpert MTB/RIF | 97.8           | 97.7           | 95.7    | 98.8    | Inpatient             | Liquid culture |
| Scherer et al. (2011) | Porto Alegre, Brazil  | Sputa             | 74           | PCR dot blot | 72             | 85             | 85      | 72      | In and outpatient     | Solid culture combined with clinics |
| Lawn et al. (2011) | Cape Town, South Africa | Sputa             | 445          | Xpert MTB/RIF | 73.3           | 99.2           | 94.8    | 94.8    | Outpatient            | Liquid culture |
| Boehme et al. (2011) | Multi-site, Brazil    | Sputa             | 602          | Xpert MTB/RIF | 82.4           | 99.2           | 98.3    | 91.3    | Outpatient            | Liquid culture |
| Michelon et al. (2011) | Multi-site, Brazil    | Sputa             | 47           | Detect-TB   | 75             | 100            | -       | -       | Outpatient            | Culture       |
| Scott et al. (2011) | Johannesburg, South Africa | Sputa             | 124          | Xpert MTB/RIF | 84             | 96             | 92      | 92      | Outpatient            | Liquid culture |
| Davis et al. (2011) | Uganda, South Africa  | Sputa and bronchoalveolar lavage | 170 | MTB/RIF | 39             | 95             | -       | -       | Inpatient             | Solid culture |

HIV: human immunodeficiency virus; NPV: negative predictive value; PCR: in house polymerase chain reaction; PPV: positive predictive value; qPCR: quantitative real-time PCR.
75% and a specificity of 100% were reported using the same assay for TB detection in spontaneous sputum samples obtained from patients suspected of having TB (Michelon et al. 2011). Difficulties in handling the kit, reagents’ stability and delays in reading the reaction were considered. The handling was performed exclusively by laboratory personnel that had received training in executing the Detect-TB protocol. To read the samples, we placed the microplates on a cold surface to transport them to another laboratory in a connected building and thus some delay in reading might have contributed to these results, although we also should consider that some of these false-positive cases might indeed have been positive cases that had not been identified by culturing.

Our conventional PCR assay showed the poorest performance in sputum with 46.3% specificity. A published meta-analysis demonstrated a sensitivity variation of 32-92% with a mean of 72% and a specificity variation of 93-100% with a mean of 96% for the PCR simplex methodology for TB detection, in comparison with sputum culture results and clinical criteria (Sarmiento et al. 2003). In our study, we used only culture as the gold standard. Nevertheless, the fact that qPCR and conventional PCR results were so divergent surprises us because both assays used the same primers and sample processing. The main differences were (i) qPCR used a fluorescent internal probe for accuracy improvement, which per se justifies this finding, while the conventional PCR did not have the probe hybridisation step, (ii) samples were run in duplicate or triplicate in qPCR, whereas a single sample normally was run for conventional PCR and (iii) qPCR data were measurable and positive cases were defined based on a standard curve, whereas in conventional PCR, the results were based on the identification of a band in agarose gel corresponding in size to the amplified PCR product. For all these reasons, the conventional PCR test may not be reliable (Al Zahrani et al. 2001, Chakravorty et al. 2005).

The difference in sample processing is a limitation for comparative analysis of the accuracy of different molecular assays for TB diagnosis. A study on the performance of the Xpert MTB/RIF test with an IS6110-TaqMan qPCR assay for the detection of mycobacteria in respiratory specimens processed by the standard procedure of decontamination and centrifugation reported no differences in overall sensitivity (79% vs. 84%, respectively) of both tests, which presented 100% specificity for smear-positive sputum, with no differences observed in smear-negative sputum sensitivity (57% vs. 68%, respectively) (Armand et al. 2011). Alternatively, Park et al. (2013) compared the performance of the Xpert MTB/RIF test using 1 mL of non-processed respiratory specimens with the COBAS TaqMan MTB assay using decontaminated-concentrated respiratory specimens obtained from ~5 mL sputum. The performance of both assays using culture results as the gold standard showed that the overall sensitivities of the Xpert (67.9%) and COBAS (71.4%) assays were similar. The Xpert sensitivity did not significantly differ between smear-positive and smear-negative (67% vs. 69%) sputum; however, the reduction from the 87% sensitivity in smear-positive sputum to 54% sensitivity in smear-negative sputum observed with the COBAS assay suggested that the sputum-decontamination processing included in the COBAS protocol contributed to reducing the bacillary viability in culture, even though the proportional sputum volume used was larger than that used in the Xpert test (Park et al. 2013).

Comparing the performance of qPCR from sputum and blood samples of corresponding patients, the qPCR test exhibited an accuracy of 93% for the detection of TB in blood, showing good concordance with culture results (75%) and the qPCR results from sputum (63%). One explanation for this higher concordance with sputum culture than with qPCR testing of sputum was the high sensitivity of the qPCR test in sputum in relation to culture. The three cases of negative-culture and positive-qPCR results obtained from sputum were also negative-qPCR for blood, suggesting that mycobacteriaemia might be associated with advanced disease. In this context, published data on the Xpert test for TB detection in blood revealed a 21% sensitivity and 100% specificity compared with sputum culture (56% concordance), but a positive association was reported between positive Xpert assay from blood samples and the mortality rate in HIV-infected patients (Feasey et al. 2013). The main difference between the Xpert study and our protocol was in sample processing: we used leukocytes purified from 5 mL peripheral blood, whereas the Xpert protocol requires the collection of 18 mL of peripheral blood for cellular lysates and concentration into a 1 mL sample for introduction into the Xpert cartridge for diagnosis. In this context, it is noteworthy that blood components such as haem, lactoferrin and immunoglobulin G have been associated with reduced amplification yield (Al-Soud et al. 2000, Al-Soud & Rådström 2001) and the efficient removal of PCR-inhibitors from blood-extracted DNA contributes to improved sensitivity of the PCR-based diagnostic assay.

In HIV-infected patients, confirmation of the diagnosis of pulmonary TB from sputum is more difficult due to the low bacillary load in sputum and this difficulty is also associated with external factors that influence the sample quality, such as inadequate sputum collection, sample transportation, time delays in sample processing and the methodology employed. We believe that qPCR is one of the molecular tests that should be considered as a tool for the detection of pulmonary TB in smear-negative patients.

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