Detection of *Mycobacterium tuberculosis* from Clinical Samples using Different Real-Time PCR System

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### Abstract

During the study period (July to December 2016) a total of 188 samples (sputum and blood from each person) have been tested in the National Reference Laboratory (NRL) at the Institute of Chest and Respiratory Diseases / Baghdad. These cases are considered as suspected pulmonary tuberculosis. This study for these 188 suspected specimens declare that, 103 (54.78%) cases were positive using direct assay (AFB smear microscopy) and 85 (45.21%) were considered to be negative cases by smear microscopy. Of the 103 TB patients, 88 (85.43%) were represented as new TB cases; the others 15 (14.57%) patients appeared to follow up assessment of the directly observed treatment (DOT) program.

Detection of *M. tuberculosis* by real time PCR GeneXpert showed that among total 188 specimens (Suspected Group), 117 (62.23%) of the specimens were positive in GeneXpert from sputum and 71 (37.76%) negative specimens by GeneXpert. This study, declares the percentage of sensitivity, specificity, PPV and NPV of GeneXpert which were 94.11 %, 92.75 %, 95.72 % and 90.14 % respectively.

Detection of *M. tuberculosis* by real time PCR using Realline kit showed that among the total 188 specimens (Suspected Group), 112 (59.57%) of the specimens were positive and 76 (40.4%) negative which detected by Real-Time PCR using Realline Kit. All these samples are sputum. This study showed the percentage of sensitivity, specificity, PPV and NPV of Realline kit were 92.43%, 97.1%, 98.21% and 88.15 % respectively.

Detection of *M. tuberculosis* by real time PCR using MTB Robogene kit showed that among the total 188 specimens (Suspected Group), 111 (59.04%) of the specimens were positive, and 77 (40.95%) negative specimens by using MTB Robogene kit, the samples are sputum. The sensitivity, specificity, PPV and NPV of our study was 87.3%, 97.1%, 96.6% and 80.51% respectively. Detection of *M. tuberculosis* by real time PCR using MTB DNA Blood showed that among total 188 specimens (Suspected Group), 59 (31.38%) were positive in PCR and 129 (68.61%) were negative, DNA was extracted directly from blood and then tested in MTB Robogene kit. The present showed the percentage of the sensitivity, specificity, PPV and NPV to be 47.89%, 97.1%, 96.6% and 51.93.

### Keywords

*M. tuberculosis*, AFB, GeneXpert
Introduction

TB is considered the deadliest bacterial infections in the world. Infection with this disease can be fatal and it has a large impact on the global health getting international attention with increasing number of cases worldwide where the developed and developing countries were critically included (WHO, 2015) MTB is the main causative agent of TB which is an aerobic pathogenic bacillus that establishes infection in the lungs. It is believed that two billion people are carrying non-eradicated intra-granulomatous Tuberculosis bacilli as LTBI (short for latent tuberculosis infection) and around 10% of those people will be infected with active tuberculosis during their lifetime (Gerald L. Mandell, John E. Bennett, Raphael 2010). The MTB complex consists of: M. tuberculosis, M. canetti, M. microti, M. pinnipedii, M. africanum, M. bovis and M. caprae most of infections in humans are caused by M. tuberculosis with a small number of infections caused by M. africanum and M. bovis (LoBue et al., 2010).

It is estimated that 1/3 of the population (around 2 billion people) are infected with TB bacilli however only 5-10% of people get infected with the active disease (WHO 2007) the remaining 90% will initially be experience no symptoms of the disease and will experience latent disease which reactivation may happen the world health organization estimate that in 2008 there were 9.4 million cases of TB lead to 1.3 million deaths (WHO2009). The incidence infections and disease varies greatly geography wise as the estimated number of cases in 2008 occurred in Asia (55%) and Africa (30%) (WHO 2009).

Iraq is one of the 7 countries of the Region with a high infection rate of TB and has 3% of total number of cases worldwide there are and estimated number of 20000 TB patients in Iraq and more than 4000 estimated deaths due to TB (WHO2014). Smear microscopy is the most widely used test for the diagnosis of tuberculosis (TB). The majority of laboratories use conventional light microscopy to examine Ziehl-Neelsen stained direct smears, documented to be highly specific in areas with a high prevalence of TB but with varying sensitivity (20-80%) (WHO, 2009). A definitive diagnosis of tuberculosis can be made by culturing Mycobacterium tuberculosis organisms from a patient specimen most often sputum, but may also include pus, CSF, biopsied tissue etc. (Kumar et al., 2007). Many types of cultures are available. Traditionally, cultures have used the Lowenstein-Jensen (LJ) and Middlebrook media (7H9, 7H10, and 7H11) (Drobniewski et al., 2013).

Mycobacterial culture remains the gold standard for detection and drug susceptibility testing (DST). Traditionally, culture on an egg-based solid medium, known as Lowenstein-Jensen medium, may take as long as 4-6 weeks. Liquid culture systems by using mycobacterial growth indicator tube (MGIT) offer a more sensitive and rapid alternative to conventional solid culture and may detect growth in 1-3 weeks (Cruciani et al., 2004; Dinnes et al., 2007). The emergence and spread of multidrug resistant (MDR) and extensively drug resistant (XDR) Mycobacterium tuberculosis complex (MTBC) strains poses significant challenges to disease control (WHO, 2010). In order to overcome conventional methods low sensitivity and diagnostic delays, nucleic acid amplification (NAA) tests have been introduced. The Xpert MTB/RIF is a cartridge based, automated diagnostic test that can identify Mycobacterium tuberculosis (MTB) and resistance to rifampicin (RIF). It was co-developed by Cepheid, Inc. and Foundation for Innovative New Diagnostics, with additional financial support from the US National Institutes of Health (NIH) and
technical support from the University of Medicine and Dentistry of New Jersey (UMDNJ). In December 2010, the World Health Organization (WHO) endorsed the Xpert MTB/RIF for use in TB endemic countries (WHO, 2010) and declared it a major milestone for global TB diagnosis. This followed 18 months of rigorous assessment of its field effectiveness in TB, MDR-TB and TB/HIV co-infection (Small and Pai, 2010). This test, and others that are likely to follow, have the potential to revolutionize the diagnosis of TB (Small and Pai, 2010; Van Rie et al., 2010). The recently introduced Xpert MTB/RMP assay detects the presence of MTB DNA and its susceptibility to rifampin (RMP) in a single reaction (Helb et al., 2010). Microbiological diagnostics is being revolutionized by real-time PCR because of the sensitivity and specificity of detection for determination of variants the sensitivity of real time PCR in clinical specimens ranges from 71.6%- 98.1% more than conventional staining (46.3%) and culture techniques (41.8%) and it has a 100% mycobacteria detection specificity (Parashar et al., 2006). It is also more cost efficient than traditional culture methods in determining the quantity of a given pathogen in a clinical specimen and it takes way less time than traditional methods which take from 1-14 days where as real time PCR takes around 30-50 minutes and those results are obtained in similar or better sensitivities. Unlike traditional culture methods which focus on measuring live pathogens real time PCR can measure both live and dead pathogens (Hein et al., 2001).

The aim of this project is the molecular characterization of M. tuberculosis isolated from patient (Baghdad). Identification of M. tuberculosis basically depend on microscopic characterization and then on PCR Technique, different system and have rapid diagnostic methods, with high sensitivity and specificity Lab method to diagnose diseased patients.

**Materials and Methods**

**Patients and sample collection**

The study was conducted at National Reference Lab (NRL) of tuberculosis/Baghdad, total of 188 patient suspected and 80 normal persons. Two samples have been gathered from every patient. first, one became taken from patient when he just reached the institute; second pattern accrued at early morning earlier than breakfast, the early morning collection represents the pulmonary secretions accrued in a single day, and consequently it typically has a better positivity. Gathered specimens were stored at –20°C until use (IUATLD, 2000; Sengooba et al., 2012). Ziehl-Neelsen stain sputum smears were tested for the presence of pulmonary acid fast bacilli. Also forty specimens from healthful men and women have been accumulated as negative control.

**DNA extraction using spin column (innuPREP Mycobacteria DNA Kit)**

Transfer 200 µl of sputum into 1.5 ml reaction tube (DNA_D1 which already supplied with the amplification kit MTB DNA Qualitative kit, containing Internal control DNA) and add 200 µl NAC buffer. Vortex shortly and incubate the sample at room temperature under continuous shaking for 20 minutes. Centrifuge the sample at 10000 rpm for 15 minutes. Remove the supernatant carefully, but completely. Add 200 µl TE buffer to the bacterial pellet and re-suspend the pellet completely. Add 15 µl Lysozym (stock solution 10 mg / ml in TE buffer). Mix by pulsed vortexing for 5 s. Incubate at 37°C for 30 minutes. After lysis with lysozyme add 200...
µl lysis solution TLS and incubate the tube at 95 °C for 20 minutes. After incubation at 95 °C place the tube on ice for 2 minutes. Open the tube and add 25 µl Proteinase K and incubate at 50°C for 30 minutes. Apply 600 µl of the sample to the Spin Filter (Blue) located in a 2 ml receiver tube. Close the cap and centrifuge 12000 rpm for 2 minutes. Discard the receiver tube with the µfiltrate. Place the spin filter into new 2 ml receiver tube. Open the spin filter and add 500 µl washing solution HS, close the cap and centrifuge 12000 rpm for 1 minutes. Discard the receiver tube with the filtrate.

Place the spin filter into new 2 ml receiver tube. Open the spin filter and add 650 µl washing solution MS, close the cap and centrifuge 12000 rpm for 1 minutes. Discard the 2 ml receiver tube. Place the Spin Filter into 1.5 ml Elution Tube. Carefully open the cap of the spin filter and add 50 µl pre-heated Elution Buffer. Incubate at room temperature for 2 minutes. Centrifuge at 8000 rpm for 1 minutes.

Fully automated Real-Time PCR System Xpert

Label each Xpert MTB/RIF cartridge with the sample ID. (Write on the sides of the cartridge or affix ID label.) Note: Do not put the label on the lid of the cartridge or obstruct the existing 2D barcode on the cartridge. Transfer at least 0.5 mL of the total resuspension pellet to a conical, screw-capped tube for the Xpert MTB/RIF using a sterile transfer pipette. Alternatively, the entire sample may be processed in the original tube. Store re-suspended sediments at 2–8 °C if they are not immediately processed for Xpert MTB/RIF. Do not store for more than 12 hours. Add 1.5 mL of Xpert MTB/RIF Sample Reagent (SR) to 0.5 mL of resuspended sediment sample using a sterile transfer pipette and shake vigorously 10 – 20 times. Note: One back-and-forth movement is a single shake. Incubate the specimen for 15 minutes at room temperature, between 5 and 10 minutes of the incubation, again shake the specimen vigorously 10 – 20 times. Samples should be liquefied with no visible clumps of sputum. Particulate matter may exist that is not part of the sample. Start the test within 30 minutes of adding the sample to the cartridge.

Open system Real-Time PCR qTOWER³

Centrifuge the MTB_D4 (RoboGene® MTB DNA Qualitative Kit) briefly at full speed to collect the lyophilized Reagent Mix on the bottom of the tube. Add 200 µl PCR grade water DNA to MTB_D4; close the tube, mix by brief vortexing followed by brief centrifugation at full speed.

Incubated at 37°C for 20 min using a thermal mixer (800 –1,000 rpm), mix by brief vortexing followed by brief centrifugation at full speed. Dissolved reagent mix can be stored at 2-8 °C and always protected from light up to 14 days (do not freeze!).

Add 20 µl 1x Master Mix to sample tubes and each tube with standards. Add 5 µl PCR grade water to tubes that serve as NTC and to all quantification standards containing the 1x Master Mix.

Do not exceed a final reaction volume of 25 µl. Add 5 µl of eluate from DNA isolation to the respective sample tubes containing the 1x Master Mix. Do not exceed a final reaction volume of 25 µl.

Program the applied real-time PCR platforms as indicated in Table 1 and start the program.
Results and Discussion

Population Studies

The data set in Table 2, exhibited the percentage of infected patients were male 73 constituted (70.87) to female 33 that constituted (32.03 %), with over all male to female ratio of 2.21 (73/33) with a highly significant difference (P≤0.01). These results were in accordance with previous results reported by Shaker (2013) and Sabah (2015). The attributable for higher men incidence are poorly understood, and require additional research to identify correlating hazard agents. Difference between male and female susceptibility to TB may result partly from biological differences (i.e. sex differences), cultural and the economic state of the society cause to gender differences having access to health care. Various studies introduced evidence for a possible role of the X chromosome and sex hormones (i.e. testosterone) in susceptibility to TB (Neyrolles and Quintana-Murci, 2009; Bayingana et al., 2014; Boum et al., 2014). Hormones related with sex may be a compelling factor for this difference, testosterone for example decreases the macrophage activation process as well as pro-inflammatory cytokines production, on other hand estrogens are proinflammatory mediator’s inducer. Recent study mentions that this difference may be due to more existence of men in the community and more an unexpected meeting with carriers and the disease’s risk factors such as cigarettes and narcotic materials (Bini EI et al., 2014; Babamahmoodi et al., 2015).

Extraction and concentration of DNA

268 samples which were suspected and healthy persons underwent to deoxyribonucleic acid (DNA) extraction. The range of DNA concentrations using, MTB Robogene kit was (1.34- 13.23) ng/μl. The mean concentration was 7.3ng/μl this value is somewhat similar to Sabah (2015) with range of DNA concentrations (1.24-12) ng/μl and mean concentration 3.7 ng/μl (Table 3). The concentration of DNA was enough to successful DNA amplification when detected by real time PCR. This result was accepted in a study done by AL-Noomani et al., (2010) (Figure 1 and Table 3).

Detection of M. tuberculosis by real time PCR GeneXpert

The results revealed among total 188 specimens (Suspected Group), 117(62.23%) of the specimens were positive in GeneXpert from sputum and 71 (37.76%) negative specimens by GeneXpert (Table 4). Figure 2 showed the positive case and Figure 3 showed the negative case

The diagnostic of Mycobacterium tuberculosis by culture method can be accomplished in two way, one of them is solid (Lowenstein Jensen media) and another one is liquid broth system (MGIT 320). The time required to get the result by using liquid culture assay is less than the time required to get the same result by using LJ medium.

In our study, results from Lowenstein Jensen media were taken. GeneXpert characterized with very small volume and so easy to use that mean friendly user and the requirement of training is available and need low level of knowledge for the end user. The time for getting the first result is so short period only 2 hours, in another hand the result period from culture method takes very long time reach to couple of weeks (WHO, 2011, Van Rie, 2010). A lot of studies have showed the performance and advantages of GeneXpert in diagnosis of PTB (Bowles, 2011; Miller, 2011). Our study, declare the percentage of sensitivity, specificity, PPV and NPV of GeneXpert which were 94.11 %, 92.75 %, 95.72 % and 90.14 % respectively that is comparable with other studies (Table 5) like
the study done by (Reechaipichitkul et al., 2017) with 84% sensitivity, 92% specificity, 91% PPV and 85% NPV. On account of its rapid and easy performance, the GeneXpert MTB/RIF assay has been recommended for initial testing or as an add-on to smear microscopy for TB diagnosis (Lawn et al., 2011; Van Rie et al., 2010; Opota et al., 2016). Moreover, this technique can detect resistant strains (Lin and Desmond, 2014).

Our study motivates and promotes the usage of GeneXpert in smear positive cases of pulmonary as recommended by WHO (WHO 2011). In special cases of patients with results of smear microscopy and GeneXpert which was not clear for diagnostic pulmonary on another hands all the clinical symptoms was occur such as HIV positive, in these cases it should begin to take the treatment after positive diagnostic by using culture method (International standard for tuberculosis care 2014) In despite of all better features of GeneXpert, this technique could not reduce the role of the traditional diagnostic assay like culture and smear and both advance and traditional works parallel as well as the anti-tubercular drug sensitivity should be done just to asset the development and response to the treatment and to identify resistance situation to Rifampicin for example (International standard for tuberculosis care, 2014)

Detection of *M. tuberculosis* by real time PCR using Realline kit

The results revealed among total 188 specimens (Suspected Group), 112 (59.57%) of the specimens were positive and 76 (40.4%) negative which detected by Real-Time PCR using Realline Kit, all these samples are sputum (Table 6 and Figure 4).

<table>
<thead>
<tr>
<th>Table.1 Cycling conditions for RoboGene® MTB DNA Qualitative Kit</th>
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<tbody>
<tr>
<td>Stages</td>
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<tr>
<td>Stage 1</td>
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<tr>
<td>Stage 2</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Table.2 Distribution of TB patients according to their gender</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Female</td>
</tr>
</tbody>
</table>
**Table 3** DNA extraction method related with concentration and purity

<table>
<thead>
<tr>
<th>DNA extraction method</th>
<th>Range of DNA concentration ng/μl</th>
<th>Mean of DNA concentration ng/μl</th>
<th>DNA Purity 260/280 mean</th>
<th>260/230 Mean</th>
</tr>
</thead>
<tbody>
<tr>
<td>MTB Robogene</td>
<td>1.34-13.23</td>
<td>7.3</td>
<td>1.84</td>
<td>1.12</td>
</tr>
</tbody>
</table>

**Table 4** Distribution of TB patients according to their gender using GeneXpert

<table>
<thead>
<tr>
<th>Gender</th>
<th>Healthy n (%)</th>
<th>Suspected n, +ve n (%)</th>
<th>Suspected n, -ve n (%)</th>
<th>Total n, +ve n (%) (GeneXpert)</th>
<th>Total n, -ve n (%) (GeneXpert)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>42(100%)</td>
<td>110, 84(76.36)</td>
<td>110,26, (23.36)</td>
<td>152.84(55.26)</td>
<td>152,68(44.73)</td>
</tr>
<tr>
<td>Female</td>
<td>38(100%)</td>
<td>78,33 (42.3)</td>
<td>78,45,(57.69)</td>
<td>116,33(28.44)</td>
<td>116,83(71.55)</td>
</tr>
</tbody>
</table>

**Table 5** The percentage of sensitivity, specificity, PPV and NPV of GeneXpert in comparing with sputum culture

<table>
<thead>
<tr>
<th>GeneXpert</th>
<th>L.J culture +ve</th>
<th>L.J culture -ve</th>
<th>total</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GeneXpert +</td>
<td>112</td>
<td>5</td>
<td>117</td>
<td>94.11</td>
<td>92.75</td>
<td>95.72</td>
<td>90.14</td>
</tr>
<tr>
<td>GeneXpert -</td>
<td>7</td>
<td>64</td>
<td>71</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>total</td>
<td>119</td>
<td>69</td>
<td>188</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

**Table 6** Distribution of TB patients according to their gender using Realline Kit in PCR

<table>
<thead>
<tr>
<th>Gender</th>
<th>Healthy n (%)</th>
<th>Suspected n, +ve n (%)</th>
<th>Suspected n, -ve n (%)</th>
<th>Total n, +ve n (%) (Realline Kit)</th>
<th>Total n, -ve n (%) (Realline Kit)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>42 (100%)</td>
<td>110, 75(68.18)</td>
<td>110,35,(31.81)</td>
<td>152,75(49.34)</td>
<td>152,77(50.65)</td>
</tr>
<tr>
<td>Female</td>
<td>38 (100%)</td>
<td>78,37 (47.43)</td>
<td>78,41, (52.56)</td>
<td>116,37(31.89)</td>
<td>116,79(68.1)</td>
</tr>
</tbody>
</table>
Table.7 The percentage of sensitivity, specificity, PPV and NPV of RealLine *Mycobacterium tuberculosis* in comparing with sputum culture

<table>
<thead>
<tr>
<th>RealLine TB kit</th>
<th>L.J culture +ve</th>
<th>L.J culture –ve</th>
<th>total</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>PPV (%)</th>
<th>NPV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RealLine TB kit +</td>
<td>110</td>
<td>2</td>
<td>112</td>
<td>92.43</td>
<td>97.1</td>
<td>98.21</td>
<td>88.15</td>
</tr>
<tr>
<td>RealLine TB kit -</td>
<td>9</td>
<td>67</td>
<td>76</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>total</td>
<td>119</td>
<td>69</td>
<td>188</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

Figure.1 Genomic DNA electrophoresis extracted by MTB Kit

Figure.2 Positive case detected by GeneXpert

Figure.3 Negative case detected by GeneXpert
Our study showed the percentage of sensitivity, specificity, PPV and NPV of Realline kit were 92.43%, 97.1%, 98.21% and 88.15 % respectively that if we compare to other studies (Table 7) is somewhat similar like the study done by (Pinhata et al., 2015) that found sensitivity of 90.3%. Specificity values was 98.6% and PPV, NPV were 86.2%, 99.1% respectively

In this study, RT-PCR detected a higher number of confirmed TB cases in comparison with microscopy, as was also found by Nimesh et al., (2013). Moreover, RT-PCR showed high specificity, as found by Broccolo et al., (2003) and Armand et al., (2011).

It is well documented that molecular methods show good sensitivity in smear-positive samples, but are less sensitive in paucibacillary samples (Greco et al., 2006; Armand et al., 2011).

False-negative results in molecular tests can be explained by a low load of mycobacteria, the presence of polymerase inhibitors and the uneven distribution of bacilli in paucibacillary samples (Reischl et al., 1998). There were nine false-negative samples by RT-PCR in our study. It is possible that mycobacterial DNA was lost during sputum aliquoting or pre-treatment for DNA extraction.

According to Garcı’a-Quintanilla et al., (2002), PCR sensitivity loss may be due to the detection of a single-copy gene (Lee et al., 2010).

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2368