Rapid Detection of Mycobacteria using automated Detection/ Incubation System in Exudative Pleural Effusion in Alexandria Main University Hospital

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ABSTRACT

Studies have been targeting to evaluate newer automated culture systems techniques for different types of samples to provide a more rapid diagnosis relying on sensitive and specific principle with the ability to discriminate between viable and dead mycobacterial bacilli. This study aimed at assessment of bactalert, biomerieux ® culture system as a rapid diagnostic tool for tuberculous pleural effusion fluid in comparison with Lowenstein Jensen conventional culture and Zeil Neelsen stained smears. Pleural aspirate collected from 60 patients presented with exudative pleural effusion was centrifuges and the deposit divided to be inoculated on bactalert bottles and specific principle with the ability to discriminate between viable and dead bacilli. Bact Alert system has been considered a reliable, easy, rapid screening test for tuberculous pleurisy that needs confirmation via other laboratory assays.

Introduction

Tuberculosis has been long considered as the leading curable infectious disease causing death (Dye et al., 2008; Kumar, 2015). Studies in populations with a high prevalence of tuberculosis reported that 30% of patients with tuberculosis are associated with tuberculous pleurisy as one of the most common forms of extrapolmonary tuberculosis (TB) (Aljohaney et al., 2012; WHO, 2013). The pathogenesis of primary TB effusion was described to be mainly due to delayed hypersensitivity response to mycobacterial antigens and activation of pleural cells for cytokine production. Many studies have investigated the usefulness of measuring different parameters in pleural fluid for an early diagnosis of tuberculous pleurisy. The definitive diagnosis of TB pleural effusions depends on the demonstration of acid-fast bacilli in the sputum, pleural fluid, or pleural biopsy specimens (Lima et al., 2003). The diagnosis can be established in a majority of patients from the clinical features, pleural fluid examination, including cytology, biochemistry, and bacteriology, and pleural biopsy. Culture on Lowenstein Jensen media is considered as the gold standard for definite diagnosis still it needs as long as 42 days to reveal a result. Detection of Acid fast bacilli via Zeil Neelsen staining of smears of pleural
fluid have been considered to be a rapid diagnostic tool but with a low sensitivity as well as inability to discriminate between actively replicating and dead bacilli (Steingart et al., 2007; Uddin et al., 2013). Similarly, rapid molecular techniques as PCR for detection of TB nucleic acid lacked the ability to discriminate between active and inactive TB cases positive for mycobacterial DNA. On the other hand, cytologic examination inspite specific had a reported low sensitivity while biochemical assays including adenosine deaminase and interferon gamma detection in the pleural fluid lacked specificity inspite of their very high sensitivity (Kaisemann et al., 2004; Tay et al., 2013).

Studies have been targeting to evaluate newer culture systems techniques to provide a more rapid diagnosis relying on sensitive and specific principle with the ability to discriminate between viable and dead mycobacterial bacilli. We intended in this study to assess bactalert culture system as a rapid diagnostic tool for tuberculous pleural effusion in comparison with Lowenstein Jensen and Zeil Neelsen stained smears.

**Patients and Methods**

60 patients presented to the department of chest medicine Alexandria main University hospital diagnosed as exudative pleural effusion according to light's criteria (Light et al., 1973; Light, 2002), have been counseled and consented for the requirements of the study. In case of approval, Data concerning age, previous medical history, signs and symptoms have been collected from the patients.

Pleural effusion fluid has been collected from all patients via thoracocentesis under strict aseptic technique in heparinized syringe to avoid clotting. Centrifugation was performed in sterile tubes for 10 minutes at 5000Xrpm, supernatant have been discarded less the lowermost 1 ml. The deposit has been suspended in this 1 ml and divided into three parts:

500ml was inoculated on bactalert MB enrichment (fluid culture media with supplement) vials kit (bioMérieux, USA) following the manufacturer instructions and inserted into Bactalert incubation/detection system (bioMérieux, USA) results were considered positive when a positive alert was revealed from the detection system and a growth curve was demonstrated via software (Metchock et al., 1999).

300 ml were used to make a direct smear subjected to Zeil Neelsen staining and examination by oil immersion lens under the light microscopy results were considered positive when acid fast bacilli were revealed (Master, 1992).

200ml have been inoculated on Lowenstein Jensen media (sigma-Aldrich, USA). Results were considered positive in case of growth of colonies that were identified for confirmation to be a mycobacterium tuberculosis growth (Master, 1992; Lowenstein, 1931).

**Results and Discussion**

Among the patients enrolled in the study 28(46.6%) were males and 32 (54.4%) were females, those results agree with similar reports noting a higher female to male ration (Fader et al., 2010; Noertjojo et al., 2002; Forssbohm et al., 2008) with an age range 25-80 years with the mean + SD 49.23+12.6, these results were confirming those of previous studies reporting approximately similar means of ages for patients presenting with pleural effusion (Hasanean et al., 2003). Highest rate of positivity among pleural effusion samples was revealed by the BACT/
ALERT incubation/detection system, and the lowest positivity detection was by direct ZN smear, this might be justified by small number of acid fast bacilli within different samples as it depends on whether or not the caseating lesions have got access to open in the pleural cavity or not. Similarly, reports from numerous researches have revealed different sensitivities for Zeil Neelsen direct smear (Lai et al., 2012; Maurya et al., 2011; Gill et al., 2013; Uddin et al., 2013).

Considering direct Zeil Neelsen smear as gold standard for comparison, it was revealed that

The sensitivity of the results of bact/alert incubation / detection system (100%), specificity (76.5%), Positive predictive value (42.8%), Negative predictive value (76.4%), Accuracy (71%).

Considering LJ culture as gold standard for comparison, it was revealed that the sensitivity of the results of bact/alert incubation/detection system (100%), specificity (95.1%), positive predictive value (90.47%), negative predictive value (100%), accuracy (88%) (Table 1).

Table 1: Comparison of the results of the performed laboratory assays

|                  | Bact/alert n=60 | Lowenstein Jensen media n=60 | Direct Zeil Neelsen smear n=60 |
|------------------|----------------|-------------------------------|-------------------------------|
| Positive         | 21 (35%)       | 19 (31.67%)                  | 9 (15%)                      |
| negative         | 39 (65%)       | 41 (68.33%)                  | 51 (85%)                     |

Table 2: Comparison of bact/alert results versus direct Zeil Neelsen smear

|                  | Bact/alert | Direct Zeil Neelsen smear |
|------------------|------------|---------------------------|
|                  | n=21       | n=51                      |
| Positive (n=21)  | 9 (100%)   | 12 (23.5%)                |
| Negative (n=39)  | 0 (0%)     | 39 (76.5%)                |

Table 3: Comparison of bact/alert results versus Lowenstein Jensen culture results

|                  | Lowenstein Jensen culture |
|------------------|---------------------------|
|                  | Positive (n=19)            | Negative (n=41)            | Total (n=60) |
| Positive (n=21)  | 19 (100%)                 | 2 (4.9%)                  | 21 (35%)     |
| Negative (n=39)  | 0 (0%)                    | 39 (95.1%)                | 39 (65%)     |

Both data in table 2 and 3 show a very high sensitivity (100%) of BACT/ALERT incubation/detection system for detection of mycobacterium TB in exudative pleural effusion still with a lower specificity (76.5% and 95.1%) compared to Zeil-Neeleen direct smear and Lowenstein Jensen culture respectively. This agrees with data released from many researchers reporting a high false positive rate of results with many developed automated incubation detection systems (Jorgensen, 1997; Weinstein, 1996; Morgan, 1983; Ichiyama, 1993). This finding have been justified by many authors who audited automated systems results as (Alfa, 1995; Martinez and Lakshmi, 2001 and Tortoli et al., 1998) that have reported false positive signals from automated detection systems for mycobacterial growth associated with presence
of high counts or malignant lymphocytes and polymorphs in the inoculated samples due to release of carbon dioxide from respiration of these white blood cells that might activate sensors of the detection system to reveal false positive signals.

The present study results confirm those of previous researchers that reported very high sensitivity of BACT/ART to detect mycobacterial growth with pleural effusion fluid, still its low specificity might mandate confirmation of the results using other methods with higher specificity. Due to its high sensitivity (100%), low labour, simple methodology, and rapid results BACT/Alert might be recommended as a reliable screening method for tuberculous pleurisy.

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