COMPARISON OF TUBERCULIN SKIN TEST AND LATERAL FLOW RAPID TEST FOR DETECTION OF BOVINE TUBERCULOSIS IN DAIRY CATTLE

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ABSTRACT

Tuberculosis is the most important zoonotic bacterial disease that is hazardous to both man and animals. A huge economic loss which could be direct or indirect is associated with the disease, so rapid diagnostic tests for tuberculosis are needed to facilitate early detection and prevention of disease transmission. The aim of this work is the detection of bovine tuberculosis by application of different serological tests. Tuberculin skin test applied on 1900 cattle, only 50 (2.6%) showed positive results, and then slaughtered. Forty five (90%) of slaughtered animals showed visible lesions on post mortem examination, while the other five (10%) showed non visible lesions. The bacteriological examination of the 50 samples revealed Mycobacterium bovis form 40 processed samples (80%). Results of Anigen Rapid Bovine TB Ab test and ELISA test had detected 42% and 48% of tuberculin positive cattle respectively. It was concluded that the Anigen Rapid Bovine TB Ab kit test is rapid, safe, simple and easy to perform and provide yes or no results within 15 to 20 minutes but it is not efficient for detection of bovine tuberculosis in cattle and could be useful as a complementary for tuberculin test.

Keywords: Antigen, bovine tuberculosis, ELISA, M. bovis, Rapid TB Kit.

INTRODUCTION

Bovine tuberculosis is a worldwide disease that causes a great harm on dairy farms and poses health risks to the population that consumes products of animal origin. It is still a problem with public health and economic importance in large areas of the world (Ritacco et al., 1987). The economic losses caused by the disease are not only a reduction of 10-20% in milk production and weight, but also due to infertility and condemnation of meat. The loss is estimated to be 10-25% of the reproductive efficiency, excluding the losses from mortality (Lilenbaum et al., 2001).

The disease has been difficult to control in livestock because of the lack of an effective vaccine and the presence of wildlife reservoirs. Currently, the primary methods used for the detection of TB in humans and ruminants include the measurement of a delayed type hypersensitivity (skin test) to purified protein derivative (PPD) and an indirect in vitro assay that measures the concentration of gamma interferon (IFN-γ) produced in response to stimulation with PPD (Monaghan et al., 1994, Wood et al., 1992, Wood and Jones 2001).

Some infected animals may have antibody response in absence of cell – mediated response, particularly when the bacterial load is high. A number of Enzyme Linked Immunosorbent Assays (ELISA) have been described based on complex M. bovis antigens, such as Purified Protein Derivatives (PPD) and phosphate antigens. All of these assays were successful in detecting circulating antibodies to mycobacteria but have been considered to lack specificity (Mcnair et al., 2001).

Serological assays are generally simple, rapid and inexpensive, but the development of improved serodiagnostic assays also require understanding the bovine tuberculosis humoral immune mechanisms which is characterized by heterogeneous antigen recognition (Lyashchenko et al., 1998). Advances in humoral based responses tests have led to the recently
development of two membrane-based antibody detection methods, Multi Antigen Print Immuno Assay (MAPIA) and a lateral-flow test. Ab Test Kit is a solid phase chromatographic immunoassay for the qualitative detection of *Mycobacterium bovis* antibody in serum or plasma (Greenwood et al., 2003). The purpose for conducting this study was to compare between the sensitivity of recent lateral flow rapid test and ELISA for diagnosis of bovine tuberculosis of tuberculin skin test reactor cattle.

**MATERIALS AND METHODS**

I. **Tuberculin skin test:** A total number of (1900) cross- breed dairy cattle from different farms were tested by Single Intradermal Cervical (SIC) tuberculin test as performed by OIE (2009).

II. **Serum samples:** From the positive tuberculin reactors cattle, blood samples were collected and serum samples were separated and stored at - 20°C till used in serological test.

III. **Post – mortem examination:** After slaughtering of tuberculin positive reactors, Post – mortem examination was done to detect the presence of any suspected tuberculous lesions such as caseation, calciﬁcation, or congestion that might be present in any lymph node (head, bronchial, hepatic, mesenteric, prescapular, popliteal and internal iliac lymph nodes). Moreover, specimens were collected from the lung, liver, kidney, diaphragm and peritoneum which showed congestion or suspected tuberculous lesions.

IV. **Tissue samples:** The internal organs (livers, spleens, and lungs lymph nodes) and lymph nodes showing tuberculous-like lesions were collected using aseptic techniques, placed in an ice pax and submitted as soon as possible to the laboratory where they were processed for isolation and identification of the organism.

V. **Bacteriological examination:** The organs, lymph nodes and/or tissues showing gross lesions were prepared for bacteriological examination. Prepared sections were stained with Ziehl – Neelsen’s stain. Samples were cultured on four tubes of Lowenstein-Jensen slants after being decontaminated with 4% H2SO4. Obtained isolates were identified by conventional methods (rate of growth, colonial morphology, pigmentation, and biochemical properties) according to Brasil (1994).

VI. **Enzyme Linked Immunosorbent Assay (ELISA):** The Enzyme Linked Immunosorbent Assay (ELISA) was applied in sera of tuberculin positive cattle according to Collee et al. (1996) using Bovine Purified Protein Derivatives (B – PPD). The optical density was measured at 405 nm using spectra III ELISA reader. Sample was considered positive if it yield a mean OD of each group equal to / or greater than the cut off value {Cut off value was calculated according to Nassau et al. (1976) which equal to the mean OD of negative serum plus 2 standard deviation}.

VII. **Lateral flow test Kit:** The testing of sera was carried out according to the manufacturer’s instructions using the Anigen Rapid Bovine TB Ab test kit as follows:

1. The foil pouch of test kit was removed and placed on a flat, dry surface.
2. Test units were labeled with samples names.
3. Four drops of serum were added slowly to sample well with the specimen dropper (if the migration has not appeared after one minute, one more drop of the specimen was added to the sample well).
4. A test result will be seen as a band in the result window of the kit.
5. The test results were interpreted within 20 minutes (no interpretation after 20 minutes).

**Interpretation of the test**

- Negative result: The presence of only one color band within the result window.
- Positive result: The presence of tow color bands (T band and C band) within the result window. (Even if the intensity of the band color is faint it should be consider as positive).
- Invalid: If the color band was not visible within the result window after performing the test, the result was considered invalid and the specimen was re-tested.

**RESULTS**

1. **Results of tuberculin skin test and post mortem finding of slaughtered tuberculin positive cattle**

   The results in table (1) illustrated the prevalence of tuberculin reactors in dairy cattle from different farms and PM findings of slaughtered tuberculin reactor cattle. From total 1900 tuberculin tested cattle, 50 were found to be reactors with a prevalence rate of 2.6%. On other hand, the number of non – visible lesion (NVL) reactors amounted to 5 animals (10%), while the number of visible lesions found to be 45 animals with an overall percentage of 90% as shown in the same table.

2. **Results of postmortem finding in slaughtered tuberculin reactor cattle according to the site of lesion.**

   Out of 50 tuberculin reactor animals, 45(90%) showed visible and 5(10%) had non-visible lesions, on the same time the visible lesions showing 6(12%) head, 24(48%) pulmonary, 10(20%) digestive and 5(10%) generalized as shown in Table (2).
3. Results of Anigen Rapid Bovine TB Ab test kit from tuberculin reactor cattle in comparison to the type of lesions

It is cleared from table 3, that 21(42%) of tuberculin reactor cattle were positive with Ani-gen Rapid Bovine TB kit. While, 29 (58%) were negative with Ani-gen Rapid Bovine TB kit.

4. Comparison between the results of bacteriological isolation, ELISA and Anigen Rapid Bovine TB Ab test kit on samples obtained from tuberculin positive animals

The obtained results in table (4) showed that, from 50 carcasses, 40 cultures positive for \textit{M. bovis} were recovered with an isolation rate of 80%. While, Anigen Rapid Bovine TB AB Test Kit has detected 42% of tuberculin positive cattle and the ELISA with B-PPD antigen has detected 48% of tuberculin positive reactor cattle as shown in the table.

Table 1: Results of tuberculin skin test and post mortem finding of slaughtered tuberculin positive cattle.

<table>
<thead>
<tr>
<th>No. of tested cattle</th>
<th>Reactor cattle</th>
<th>PM finding</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Visible Lesion</td>
</tr>
<tr>
<td>No.</td>
<td>%</td>
<td>No.</td>
</tr>
<tr>
<td>1900</td>
<td>50</td>
<td>2.6</td>
</tr>
</tbody>
</table>

PM: Postmortem. No.: Number.

Table 2: Results of postmortem finding in 50 slaughtered tuberculin reactor cattle according to the site of lesion.

<table>
<thead>
<tr>
<th>Reactor cattle</th>
<th>PM finding</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>VL (45) NVL(5)</td>
</tr>
<tr>
<td></td>
<td>Head Pulmonary Digestive Generalized</td>
</tr>
<tr>
<td>No.</td>
<td>%</td>
</tr>
<tr>
<td>50</td>
<td>2.6</td>
</tr>
</tbody>
</table>


Table 3: Results of Antigen Rapid Bovine TB Ab test kit from tuberculin reactor cattle in comparison to the type of lesions.

<table>
<thead>
<tr>
<th>PM finding</th>
<th>Number of positive tuberculin reactor</th>
<th>Anigen Rapid Bovine TB kit</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td></td>
<td>No.</td>
<td>%</td>
</tr>
<tr>
<td>Visible</td>
<td>45</td>
<td>20</td>
</tr>
<tr>
<td>Non visible</td>
<td>5</td>
<td>1</td>
</tr>
<tr>
<td>Total</td>
<td>50</td>
<td>21</td>
</tr>
</tbody>
</table>

PM: Postmortem. TB: Tuberculosis.
The objective of this study was to investigate the epidemiology of tuberculosis in beef cattle in Egypt. From the data collected, it was found that 24% of the tested cattle were reactors with a prevalence rate of 33.3%. The prevalence rate recorded in the present study is comparatively lower than that given by other investigators.

The results in table (1) illustrated the prevalence of tuberculosis in dairy cattle from different farms. The high prevalence rate of 50% was found to be reactors with a prevalence rate of 33.3%.

DISCUSSION

Bovine tuberculosis is an important zoonotic disease transmitted by direct contact, respiratory pathway, ingestion of unpasteurized milk and milk product, raw or uncooked meet. Tuberculosis can be difficult to diagnose based only on the clinical signs. Tuberculosis is usually diagnosed in the field with the tuberculin skin test, sputum and other body fluids may be collected for microbiological examination (Kaya et al., 2015). The tuberculin skin test was found to be the test of choice. Although the tuberculin test is a very reliable diagnostic method, yet as in all biological test, difficulties have been encountered, the greatest problem is the occurrence of so-called false negative responses, which sometimes see soon after infection, in the late stages of the disease (anergic cattle), in animals with poor immune responses and in those that have recently calved. To obviate this problem, an extensive effort has been under way to identify and characterize antigens unique to M. bovis that could be used in diagnostic assay (El-Mahrouk and El-Balawy, 2010). The objective of this study was to compare the sensitivity of two serological test for diagnosis of bovine tuberculosis in cattle.

The results in table (1) illustrated the prevalence of tuberculosis in dairy cattle from different farms and PM findings of slaughtered tuberculosis reactor cattle. From total 1900 tuberculin tested cattle, 50 were found to be reactors with a prevalence rate of 2.6%. The prevalence rate recorded in the present study is comparatively lower than that given by other investigators in Egypt (Lotfy et al., 1960, 6.9% ; Guindi et al., 1965, 26.5% ; El-Sabbani et al., 1992, 24% and El-Battawy 2008, 4.6% and in other countries of the world Oliveira et al. 1983, 3.2% in Brazil ; Ameni and Erkihun, 2007 in Ethiopia 11.6% ; Borna et al., 2009, 8% in Chad.

In the present study, the low incidence of infection could be attributed to many factors such as herd size, density of animals, breeding and management system, uncontrolled animal movements, unhygienic local habits and stress factors due to other diseases and mass vaccination against various diseases (Abuo – Eisha et al., 1995). At the same time, the prevalence recorded in the present study is comparatively higher than that given by other investigators (Shirma et al., 2003) and (Cleaveland et al., 2007) in Tanzania, as it was 1.3% and 0.9%, respectively. The high prevalence rate of tuberculosis reactors, in general, is a function of different factors such as the past history of tuberculosis in the dairy herd (Thornton and Gracey, 1976), method of breeding and housing besides the susceptibility and age of the animals (Guindi et al., 1980; Sharama et al., 1985).

On other hand, the number of non-visible lesion (NVL) reactors as shown in table (1) amounted to 5 animals (10%), a finding which may be attributed to the non-specific reaction to the tuberculin test which may be due to sensitization by other mycobacteria rather than M. bovis or even closely related microorganisms especially of the genus Nocardia or a combination of liver fluke infestation with saprophytic mycobacteria (Karlson, 1962, Cortina and Vera, 1986). Moreover, O’Reilly (1992) and Huitema (1994) ascribed the cause of non-specific reaction to the assumption that reactors may be slaughtered at stage of the disease where the tuberculous lesions are invisible or the lesions may be found in parts of carcass such as bone or skin.
As shown in table (1) the visible lesions found with an overall percentage of 90% which are higher than that reported by Oliviera et al. (1983), 75%; Zivkovic et al. (1984), 75.2% and Nasr (1997), 73.4%. On the other hand, other authors claimed a much higher percentage (Kilian, 1982, 96.3% in Germany and El-Sabban, 1992, 100% in Egypt).

Table (2) showed that the higher lesion severity was observed in the pulmonary lymph nodes (48%), this may be due to the intensive husbandry systems which make the respiratory excretion the main route by which animal – to – animal transmission occurs (Smyth et al., 2001).

The total isolation rates of M. bovis form carcasses of tuberculin reactors with and without lesions demonstrated in table (4). From a total of 50 carcasses, 40 positive cultures were recovered with an isolation rate of 80%. The obtained results were near to that mentioned by Tammemagi et al. (1973) (89.1%) and Naglaa (2008) (70.59%). Other investigators reported lower M. bovis recovery rate, by Adawy (1986) (17.5%) and Zschoc et al. (1990) (1.6%). These results depend mainly on the actual disease status present in the tested herd and to some extent on the experience of the investigators as well as the technique used for decontamination of tissue specimens. Additionally the harsh decontamination which was used to destroy contaminating bacteria other than mycobacteria in a sample may also have a harmful effect on the M. bovis causative organism (Victor et al. 1992 and Quinn et al., 1994).

It is cleared from table (3) that 21(42%) of tuberculin reactor cattle were positive with Anigen Rapid Bovine TB kit. While, 29 (58%) were negative with Anigen Rapid Bovine TB kit. The obtained results indicated the difference between the tuberculin test and Anigen Rapid Bovine TB kit for detection of tuberculosis in cattle. The Negative Anigen Rapid Bovine TB kit explained by the fact that low titer of antibodies to mycobacterial antigens which may be associated with heavy infection and that antigens may be released into the blood circulation and cause temporary suppression of antibody formation Krambovitis (1986) and that agree with Thorns and Morris (1983) who cleared the level of specific antibodies in many M. bovis infected cattle may be low or undetectable. Again this is supported with Amadori et al. (1998) who pointed that antibodies to mycobacterial antigens were investigated with various rates of success since the humeral immune response to M. bovis is late and irregular during the course of the disease.

Table (4) showed that the Anigen Rapid Bovine TB AB Test Kit has detected 42% of tuberculin positive cattle. While the ELISA with B-PPD antigen has detected 48% of tuberculin positive reactor cattle. The results agreed with those of Danbirni et al. (2013) who mentioned that the antigen rapid bovine TB Ab test alone is not efficient in diagnosis of TB and that the serological tests like ELISA must be used to validate results and disagreed with Danbirni et al. (2009) who found that (62%) of cows gave positive in antigen rapid bovine test TB Ab test. Moreover, Kalaf et al. (2014) showed that out of (28) cows examined with comparative tuberculin test, (21) cows showed positive tuberculin reactions (75%) and twenty two cow with a percentage of (78.57%) showed positive results for antigen rapid bovine TB.

The differences may be attributed to that the humoral immune test using the Enzyme-Linked Immunosorbent Assay (ELISA), immunochromatography (lateral flow) assay and other serologic based test may complement test of cellular immunity in anergic hosts (Awah-Ndukum, 2010).

CONCLUSION

It could be concluded that the Anigen Rapid Bovine TB Ab kit test is rapid, safe, simple and easy to perform and provide yes or no results within 15 to 20 minutes but alone it is not efficient for detection of bovine tuberculosis in cattle and may be useful as a complementary test for tuberculin test in some cases as in the late stages of the disease (anergic cattle).

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