LABORATORY DIAGNOSIS OF BRUCELLOSIS USING THE SLIDE AND STANDARD TUBE AGGLUTINATION METHODS*

BY

YAHYA FAYDI $^{\triangle}$ and SULEIMAN AL-KHALIL $^{\triangle}$.

ملخـــص

في هذا البحث قمن بفحص ١١٢ عينية دم لتشخيص مرض الحمى المالسطية بطريقية وتقد وجدنا ان ٥٦ ٪ من المرضى عندهم Rapid slide and standard tube agglutination method أقل من 20:1 بينما % 33 عندهم titer أعلى من 80:1 ٪

وقد وجدنا ان هناك تناسبا في استعمال الطريقتين حيث كانت نتائج كلا الطريقتين مشابهة لنتائج الطريقة الاخرى .

Abstract

Both the rapid slide and standard tube agglutination methods were used for the diagnosis of 112 blood specimens for Brucellosis. We found that 56% of the patients have a titer < 1:20 while 33% have a titer > 1:80.

We found that the rapid slide agglutination method correlates well with the tube agglutination method.

Introduction

Brucellosis is an epidemic disease in the West Bank of Jordan . It is a zoonotic disease in which infection is transmitted to humans from domestic animals through direct contact or mostly through consumption of unboiled milk and milk products taken from infected animals.

 \triangle Medical Laboratory Sciences Department , An-Najah National University – Nablus . * This paper was presented at the 30th scientific week held in Damascus , Syria , November 1990 . In the West Bank , people most commonly get the infection from eating raw white cheese prepared from the milk of infected goats , sheep or cows . The people who are involved in cheese production , do not boil or pasteurize milk before using it The disease starts to appear early in Spring and continues through Summer . This is

due to the fact that white cheese appears in the market in Spring.

Several serological tests are used for the diagnosis of the disease $^{(1,2,6,7)}$. The agglutination method is one of these methods. It is easy to do in private small laboratories. We undertook this study to determine whether to use the rapid slide agglutination method or the standard tube agglutination method.

Experimental

A total of 112 blood specimens withdrawn from patients showing the symptoms of Brucellosis . Sera of the patients blood were separated after clot formation . **Brucella abortus** antigen (Gamma diagnostics) , 0.85% saline , positive and negative control sera , serological pipettes , test tubes 13 x 100 mm and slides divided into 12 squares were used in the study .

Both the slide and tube methods were carried out according to the instructions of the manufacturer 3 .

Rapid Slide Method

Using a micropipette , the following serum volumes were delivered to the already divided slide : 0.08ml, 0.04ml, 0.02ml, 0.01ml and 0.005 ml. These serum volumes are approximately equivalent respectively to the following titers 1:20, 1:40, 1:80, 1:160 or 1:320. Then one drop of **B. abortus** antigen after shaking the vial was added to each dilution. The dropper of the vial was used to deliver the drop of antigen which is equal to 0.05ml. Using an applicator stick , the serum and antigen in each square were mixed and spread. The slide was rocked gently back and forth for no longer than 3 minutes. After that agglutination was observed both macroscopically and microscopically . The highest dilution which showed agglutination was considered the titer .

Both negative and positive control sera were treated as mentioned above.

Standard Tube Method

A series of $10(13 \times 100 \text{ mm})$ test tubes were placed in a rack. Then 0.9ml saline was delivered in the first test tube and 0.5ml in each of the remaining test tubes. After

that 0.1ml of the tested serum was added to the first test tube . After mixing , 0.5ml of the diluted serum was transfered to the second test tube . Then 0.5ml diluted serum was transfered from the second test tube to the third test tube , and so on until the contents of tube 10 were mixed , from which 0.5ml diluted serum was discarded . The resulting dilutions in the 10 test tubes ranged from 1:10 in tube no. 1 to 1:5120 in tube no 10 . As an antigen control another tube was added to the series containing 0.5ml saline . Then 0.5ml **B. abortus** antigen diluted 1:50 in saline , was added to each tube to make a final dilution varying from 1:20 to 1:10240 . After shaking the rack well , it was placed in a 37 °C water bath for 48 hours . The same procedure was repeated at the same time for positive and negative controls .

Results and Discussion

Up to date the agglutination method is used by most laboratories for the diagnosis of brucellosis. However, reports dealing with the usefulness of ELISA for diagnosing human brucellosis have described highly satisfactory results $^{(1,6)}$. Still the bacteriological examination is the most sensitive method $^{(7)}$.

In our study as shown in table 1,63(56%) of the patients have a titer of < 1:20 using the slide and tube methods. While 9 (8%) of the patient have 1:20 titer using the slide method and 1:40 using the tube method. Two patients (1.8%) have 1:40 titer using the slide method and 1:80 using the tube method. One patient (0.89%) has 1:80 titer in both methods. 2(1.8%) patients have 1:80 titer using slide method and 1:160 titer using tube method. Five patients (4.5%) have 1:320 titers using both slide and tube methods. One patient (0.89%) has 1:160 in slide method and 1:320 using tube method. Twenty eight (25.2%) patients have a titer > 1:320 using slide method and titers ranging from 1:640 to 1:5120 using tube-method. Patients having titers above 1:80 or showing a raising titer are considered to be infected with brucellosis⁽⁴⁾.

Our results show that the slide method correlates well with the tube method. This is in accordance with other studies⁽⁵⁾. Both methods have almost the same results with slight differences in the titer in few cases. Thus we suggest to use the rapid slide method as a screening test and to use the tube method to establish the titer in doubtful cases. We found that the tube method may give high false positive results in unexperienced hands due to false positive agglutination. In addition to this, glass should be completely clean and free of any contaminants.

Our recommendation is to do the rapid slide method, but doing the tube method just to find out the actual titer in some doubtful cases only.

Serum titers equal to 1:80 using the slide method should be preferably repeated using the standard tube method, as this is a borderline titer.

Number of Serum Specimens	Rapid Slide Method titer	Standard Tube Method titer	%
63	< 1:20	< 1:20	56.00
9	1:20	1:40	8.00
2	1:40	1:80	1.80
1	1:80	1:80	0.89
2	1:80	1:160	1.80
1	1:160	1:160	0.89
5	1:320	1:320	4.50
1	1:160	1:320	0.89
13	>1:320	1:640	11.60
7	1:320	1:1280	6.30
5	1:320	1:2560	4.50
3	1:320	1:5120	2.70
Total 112			99.87

Table 1: A comparison between rapid slide and tube agglutination methods .

Acknowledgement :

We would like to thank the participation of our finalist students for their help in carrying out this research work .

References

- Araj, G.F. LULU, A.R., Mustafa, M.Y., and Khateeb, M.I.: "Evaluation of ELISA in the diagnosis of acute and chronic brucellosis in human beings " J. Hyg. (Lond), 97 (1986), 457.
- 2- Bettelheim, K.A., Maskill, W.J., and Pearce, J. :" Comparison of standard tube and microagglutimation techniques for determining Brucella antibodies " J. Hyg. Camb., 90 (1983), 33.
- 3 Gamma Biologicals, Inc. Houston, TX 7709z.
- Jawetz, E., Melnick, J.L. and Adelberg, E.A. : Review of Medical Microbiology, Lange Medical book, Norwalk, Connecticut / Los Altos, California, (1987), 17 th ed., p. 266.

- 5- Lennette, E.H., Balows, A., Hausler, W.J., and Shadomy, H.J. : Manual of Clinical Microbiology, American Society of Microbiology, Washington, D.C., 1991, 5th, p. 461.
- 6- Marmonier, A., Stahl, J.P., Metz, = . and Micaud, M. "Interet et Valeur de la technique immunoenzymatique ELISA appliqute on diagnostic serologique des brucellosis humanies "Medicine et Maladies infectieuses, 99 (1979), 664.
- 7- Pellicer, T., Ariza, J., Foz, A. Pallares, R., and Gudiol, F.: "Specific antibodies detected during relaps of human brucellosis." J. Inf. Dis, 157 (1988), 918.