PRELIMINARY AGGLUTINATION EXPERIMENTS WITH

TREPONE MA PALLIDUM*†

BY

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Nelson and Mayer (1949) demonstrated the action of immobilizing antibodies on mobile and virulent Treponema pallidum in vitro. These antibodies do not only immobilize but also destroy treponemes exposed to such antibodies in the presence of complement lose their motility and also their virulence and their ability to multiply. However, as the first obvious manifestation of the antigen-antibody reaction was the loss of mobility of the treponemes, the reaction by which they were demonstrated was described as the Treponema pallidum Immobilization (TPI) test.

It was natural to try to demonstrate other specific antibodies against virulent Treponema pallidum. Studies of specific immobilization in vitro will reveal that many specific immobilized treponemes are not only immobile, but also appear to have changed morphologically as if they had been “chafed” or “gna wed”. Frequently, there is also a visible diminution in the number of organisms. The adherence disappearance reaction developed by Nelson (1952) may be due, at least in part, to a specific lysis.

Living, mobile treponemes are known to have a tendency to form agglutinates in certain circumstances. In our experiments we have also seen that the agglutination of live treponemes may occur in the absence of antibodies in the serum. Further, we know that physical factors, such as shaking, or merely sedimentation, will agglutinate treponemes. However, Cain (1953), McLeod and Magnuson (1953), Tani and Asano (1951), and Hardy and Hollander (1953) succeeded in suspending dead treponemes in such a way that they did not agglutinate spontaneously, even after having been left for weeks at -4° C., or after shaking. This made it possible to repeat experiments with the same antigen and to make adequate tests of reproducibility. This and the independence of complement are the two great advantages of this method. However, the fact that no complement is necessary for the agglutination reaction is also a drawback of agglutination tests if the results are intended for diagnostic use, because there is no satisfactory means of controlling the potential presence of non-specific agglutinating factors in the serum. If, however, a diagnosis of syphilis has been established, the consistent demonstration of agglutinins also opens up new possibilities in syphilis research.

The following investigations were inspired by such theoretical considerations and by the papers published by McLeod and Magnuson, by Cain, and by Hardy and Hollander on the Treponema pallidum Agglutination reaction (TPA) test. We wanted to compare the reproducibility, the sensitivity, and the specificity of the TPA test with those of TPI tests and of the old tests with lipoidal antigens by which the so-called reagins are demonstrated (reagin reactions).

Methods and Material

At the Statens Seruminstitut in Copenhagen we examined 194 sera, mostly human, for the presence of three groups of antibodies: immobile, agglutinins, and reagins.

The TPI test (H. Aa. Nielsen) was carried out in its latest form as described by Nelson and Mayer (1949), the only modifications being a four-fold increase of the sodium thioglycollate content of the medium and the taking of readings after 18 and 42 hrs’ incubation. In the latter case complement was added twice, i.e., immediately and after 24 hrs’ incubation, the final contents of complement being the same as that employed in the 18-hr experiments.

Reagins were demonstrated by Meinicke’s clarification test, Kahn’s Standard test, and Morch’s complement-fixation test with cardiolipin antigen (MR, KR, and C-WR-M).

For TPA tests (G. Ehrmann) as well as for TPI tests we used the pathogenic Nichols strain inoculated intratesticularly in rabbits. In order to avoid in vivo prestimulation of the treponemes, the rabbits had been irradiated on the same day or the day before inoculation. Rabbits weighing about 3 kg. were given a single universal radiation of about 1,000 r. Data: Focal distance 40 cm. Filter: 0.5 Cu, 1.0 Al, 215 kV 10 mA. The thinly sliced testes with early orchitis were shaken in a 0.85 per cent. saline solution. Each testis was shaken...
in 20 ml. saline three times for 30 min. at +4°C. The suspensions obtained were poured together after completion of shaking. A considerable amount of tissue debris, erythrocytes, and sperm was removed by 10 minutes' slow centrifugation. The clear suspension was then centrifuged for 60 min. at 3,000 rotations per minute in a centrifuge of 20 cm. radius. The sediment was washed once with saline, diluted to 10 to 15 million organisms per ml. of saline, and killed by heating in a water bath at 56°C for 30 min. Suspensions prepared in this way showed spontaneous agglutination in three batches out of twelve. The satisfactory suspensions were kept at +4°C ready for use during the whole period of the experiment (4 weeks). During our attempts to prepare a reliable suspension we made the following observations.

The testes were carefully prepared and were freed from fat, because fat contents will give rise to agglutinates. Such agglutinates, which may be distinguished from specific ones, may cause a loss of treponemes and impair the readings. This applies also if the preparation is shaken too fast, and if the work is performed at higher temperatures than about 4°C. Removal of tissue particles by filtration through coarse glass filters and through filter paper was not successful.

Apparently, morphologically damaged treponemes will adhere to the glass; this may frequently be observed when placing treponemes on slides, and will often result in a very undesirable loss of organisms—sometimes almost 100 per cent. Equally unsatisfactory results were obtained in attempts to kill the treponemes with penicillin. Thus agglutination with treponemes prepared in this manner is completely non-specific, even if the organisms retain their form better than when they are killed by heating.

It was found that the addition of merthiolate, as recommended by Hardy and Hollander (1953) was not necessary for the preservation of the treponemes. Finally, it was observed that several treponemal suspensions did not show any in vivo sensitization in the TPI test in spite of the fact that they did agglutinate spontaneously.

Technique.—Normal agglutinins were removed from human sera by means of sheep cells. Inactivated serum 1 ml. and 2 ml. 50 per cent. sheep cell suspension in saline were thoroughly mixed and left in a water bath at 37°C. for 90 min. and then placed in an icebox for 18 hrs at about +4°C. The sheep cells were subsequently removed by centrifugation.

Absorbed serum 0.025 ml. + antigen suspension 0.1 ml. were shaken for 2 hrs in a covered Kahn shaker, causing a rise in temperature of 28 to 30°C. The mixture was left in a water bath at 45°C. for 8 to 12 hrs. Antigen suspensions alone, without serum, were always used as controls.

Readings.—Without previous shaking, 0.01 ml. of the sediment was removed with a special pipette and examined in dark field. The agglutinates were first identified in low power, and then, in a higher magnification (× 512), identified as "genuine" agglutinates. The treponemes had by then settled lengthwise end to end, forming pointed, spiral-shaped bundles of variable thickness which, by strong agglutination, again assembled into larger groups. The characteristic spiral-shaped serrations and striped appearance of the agglutinates distinguish them from the pseudo-agglutinates; in the latter agglutinates the treponemes form loose balls, mostly with an amorphous centre.

Counts were made of 25 fields; the agglutinates thus identified were recorded in such a manner that the number of treponemes per agglutinate could be evaluated and averages calculated. At the same time, the number of free treponemes in the fields were counted. Three values were thus obtained for each serum:

1. Number of agglutinates in 25 fields;
2. Average number of treponemes per agglutinate;
3. Number of free treponemes in 25 fields.

A positive value was defined as having at least one agglutinate with ten or more treponemes observed in each of the 25 fields. The third figure served as a control reading, the result necessarily depending upon the first two figures. The more numerous the agglutinates and the higher the average of the agglutinated treponemes, the fewer were the free treponemes and vice versa.

In this manner 154 human and forty rabbit sera were examined; the results were compared with those obtained by the reagin reactions and TPI tests.

Of these, fifty (three rabbit and 47 human sera) were taken from the stock of lyophilized sera kept at the Statens Serum Institut in its capacity of WHO Reference Laboratory; the remaining 107 human sera were taken from the TPI routine, and the remaining 37 rabbit sera originated from various experiments set up for other purposes.

Results

(A) Human Sera.—The sera were divided into four clinical groups.

Group 1.—43 sera from 37 patients with definite anamnestic and clinical syphilis; all the patients of this group had been treated, several of them for many years.

Group 2.—Fourteen sera from fourteen patients with doubtful syphilis. Most of the patients from this group had been treated. As the results for Groups 1 and 2 were very much alike, they were compiled together (Table I).

<table>
<thead>
<tr>
<th>TABLE I</th>
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<tbody>
<tr>
<td>GROUPS 1 AND 2: DEFINITE AND DOUBTFUL SYPHILITICS (57 SERA FROM 51 PATIENTS)</td>
</tr>
<tr>
<td>Combined Result of Reagin Reactions</td>
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<tr>
<td>+ and ±</td>
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Group 3.—35 sera from 34 patients in whom the possibility of syphilis could be excluded with almost
**AGGLUTINATION EXPERIMENTS WITH T. PALLIDUM**

TABLE II

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Combined Result of Reagin Reactions</th>
<th>TPI₁₈</th>
<th>TPI₂₄</th>
<th>Clinical Findings</th>
<th>TPA</th>
<th>Time of Diagnosis</th>
<th>Whether Treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>Dark field positive, primary syphilis</td>
<td>+</td>
<td>1948</td>
<td>Yes</td>
</tr>
<tr>
<td>2</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>Dark field positive, primary and secondary syphilis</td>
<td>−</td>
<td>1951</td>
<td>Yes</td>
</tr>
<tr>
<td>3</td>
<td>±</td>
<td>–</td>
<td>−</td>
<td>Dark field positive, primary syphilis</td>
<td>+</td>
<td>1 month ago</td>
<td>Yes</td>
</tr>
<tr>
<td>4</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>Dark field positive, secondary syphilis</td>
<td>+</td>
<td>6 months ago</td>
<td>Yes</td>
</tr>
<tr>
<td>5</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>Aortitis</td>
<td>−</td>
<td>1939</td>
<td>Yes</td>
</tr>
<tr>
<td>6</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>No information on stage of disease</td>
<td>+</td>
<td>About 1924</td>
<td>Yes</td>
</tr>
<tr>
<td>7</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>Doubtful congenital syphilis</td>
<td>+</td>
<td>1918</td>
<td>Yes</td>
</tr>
<tr>
<td>8</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>Doubtful congenital syphilis</td>
<td>+</td>
<td>1947</td>
<td>Yes</td>
</tr>
<tr>
<td>9</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>Doubtful congenital syphilis</td>
<td>+</td>
<td>1952</td>
<td>Yes</td>
</tr>
<tr>
<td>10</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>Doubtful congenital syphilis</td>
<td>−</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>11</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>Doubtful secondary case</td>
<td>−</td>
<td>1949</td>
<td>Yes</td>
</tr>
<tr>
<td>12</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>&quot;Herpes genitalis.&quot; Reagin tests strongly positive at diagnosis</td>
<td>−</td>
<td>1950</td>
<td>No</td>
</tr>
</tbody>
</table>

Results of the five positive cases in TPA, which were not positive in TPI, are listed in Table IV. No significant difference could be demonstrated between the TPI and TPA tests.

In the controls (Group 4) the TPI and TPA reactions were negative in all sera, whereas three sera reacted in one or more of the reagin tests.

**Table IV**

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Symptoms</th>
<th>TPA Positive, TPI Negative Sera</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Retinopathy, hypertension</td>
<td>C-WR-M*</td>
</tr>
<tr>
<td>2</td>
<td>Lupus erythematosus Simple ulcer of the labium pudendi</td>
<td>11</td>
</tr>
<tr>
<td>3</td>
<td>Lesion healed up in 2 days after local treatment with mercurochrome. No glandular swelling. No induration</td>
<td>—</td>
</tr>
<tr>
<td>4</td>
<td>Rheumatoid arthritis Fertility, arthritis, pulmonary infiltration</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>—</td>
<td>1</td>
</tr>
</tbody>
</table>

The results of C-WR-M and KR are given in degrees of strength (Schmidt, 1951). The MR results are listed in the usual way: strongly positive (+ +), weakly positive (+), doubtful (±), and negative (−)
against various bacterial diseases were also included in the experiments.

(1) Three pools of serum from apparently healthy rabbits with weakly positive reagin reactions were negative in both TPI and TPA tests.

One pool from thirty rabbits infected 4 months previously (WHO TPI Control No. 2) was positive in all three types of reactions.

(2) Serum was tested in eight rabbits from the TPI routine before (Serum No. 1) and after inoculation with Treponema pallida (Serum No. 2). The rabbits were not treated. The results of the three types of reaction, time from inoculation, and information on x-ray treatment are given in Table V.

All No. 1 sera (except three with doubtful reagin tests) were negative in all three types of reaction. TPI18 was positive in one No. 2 serum only, and TPI42 was positive in the same serum and doubtful in another No. 2 serum. TPA was positive in six out of the eight No. 2 sera, and the reagin tests were positive or doubtful in seven out of the eight No. 2 sera.

(3) In an experiment set up for other purposes fifteen rabbits were infected intratesticularly with syphilis on May 5, 1952. Six weeks after inoculation all except one were treated with either penicillin or trepropal. Altogether twenty sera from these fifteen rabbits were tested (Table VI). It should be noted that all the TPA and all the TPI42 results originate from one experimental day; the TPI18 results are taken from different days. This fact accounts for the discrepancies between the results obtained for 18 and for 42 hrs, for which the day-by-day variations of the TPI test are responsible.

(4) Nine sera from nine rabbits immunized against typhoid fever, and two sera from rabbits immunized against leptospirosis were all negative in both TPI and TPA.

**Discussion**

Most of the strongly positive and clearly negative sera were examined once more in the TPA test. As was to be expected, the results showed no change. This good reproducibility was also attained in the probable, non-specific, positive cases of rheumatic fever and lupus erythematosus which were examined a few times and always showed the same weak but clearly positive results.

With regard to the sensitivity of the TPA test in comparison with that of the TPI test and the reagin reactions, altogether twelve discrepancies were found in 57 human syphilitic sera (Tables I and II). In syphilitic rabbit sera a relatively large number of positive agglutination reactions were found (Tables V and VI), especially when compared with the TPI test.

The introductory remarks about the specificity of the TPA test were confirmed by our experiments (Table III). The fact that one-third of the sera can agglutinate treponemes by non-specific normal agglutinin (Turner, 1953) shows that the test may take a non-specific course. These probably non-specific results from sera of patients suffering from rheumatoid arthritis and lupus erythematosus are due to an agglutinating factor which is already being
used for serological diagnosis of rheumatoid arthritis (Ehrmann, Ferstl, Neumayer, and Schmidt, 1952). For this reason, and because of the difficulty of obtaining definite controls for non-specific agglutinins, we shall probably have to forgo the agglutination test in its present shape as a diagnostic aid, especially as the TPI test yields good specific reactions that so far are unsurpassed. On the other hand, the TPA test may be helpful, as mentioned above, in the further research of syphilis pathology and consequently also in the prognosis and evaluation of therapy.

**Summary**

In 194 sera (154 human and forty rabbit) the results of a *Treponema pallidum* Agglutination (TPA) test, using heat-killed pathogenic treponemes, were compared with the results obtained in the *Treponema pallidum* Immobilization (TPI) test and in three different reagent reactions.

The material comprised definite and doubtful syphilitic sera, presumably biologically false positive sera and normal sera. This limited material did not allow any definite conclusions as to the sensitivity and specificity of the TPA test.

In presumably biologically false positive sera both TPA and TPI tests gave significantly fewer positive results than the reagent reactions. However, there was no significant difference between the TPA and the TPI tests in these sera.

In syphilitic rabbits, agglutinins appeared almost simultaneously with reagins, the TPI test being preponderantly negative. On the other hand, the TPA test remained positive longer than the TPI, the reagins lying somewhere between the agglutinins and immobilins.

Non-specific normal agglutinins may also produce agglutination which could not always be removed entirely by absorption with sheep cells. Presumably non-specific, positive TPA was obtained in rheumatoid arthritis and lupus erythematosus in spite of absorption.

Our thanks are due to Dr. Jeppe Ørskov and Dr. Alice Reyn of the Statens Serum Institut, Copenhagen, for their kindness and understanding in our work on the agglutination reaction.

**REFERENCES**


