A stable haemagglutinating antigen for detecting toxoplasma antibodies

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SYNOPSIS The production of a toxoplasma-sensitized cell preparation stable for at least one year at 5°C and its performance in the diagnosis of toxoplasmosis is described.

Although the decade 1960-1970 saw considerable and important developments in the biological and immunological aspects of toxoplasmosis, the diagnosis of the condition still presents considerable difficulties. When it is not possible to isolate the parasite, diagnosis is dependent largely on serological tests, and the dye test (Sabin and Feldman, 1948), or one of its modifications, is still the one which commands most confidence. However, the disadvantages associated with this test need no elaboration, especially the continuing need for live parasites and for accessory factor in addition to the heavy demands it makes on laboratory time. These factors have been instrumental in favouring the centralization of tests in a few specialized laboratories which perhaps to some extent acts as a limiting factor on the numbers of tests performed. Of all the possible substitutes for the dye test, haemagglutination (Jacobs and Lunde, 1957), direct agglutination (Fulton and Turk, 1959), fluorescent antibody methods (Goldman, 1957), and complement fixation (Fulton and Fulton, 1965), the haemagglutination test has many attractions, not only because of its sensitivity but because this kind of test can now be carried out in plastic plates in micro quantities enabling many sera to be tested readily.

In 1960 Mitchell and Green stated: 'If stable antigens can be prepared and distributed the (haemagglutination) test should lend itself to greater standardization than is possible with the dye test and warrants further investigation'. Some progress towards this goal was made by Maloney and Kaufman (1960) and Park (1961) by the use of formalized cells and by Jennis (1966) using pyruvic aldehyde-treated cells. However, all of these cell preparations had a relatively short life at 5°C.

The sensitized cell preparation we describe here is stable for at least one year at 5°C and in our hands has proved a useful and reliable tool in the diagnosis of toxoplasmosis.

Material and Methods

CELLS Sheep erythrocytes, obtained at the abattoir, were collected in Alsever's solution, stored at 5°C and used within a fortnight of collection. The treatment of the cells with pyruvic aldehyde1 followed essentially the method described by Jennis and these treated cells were stored at 5°C in Sorenson's phosphate-buffered saline, pH 7.2, containing 0.1% sodium azide. Cells prepared in this fashion may be stored without deterioration and are available for sensitization as and when required.

ANTIGEN Cotton rats, 12-15 weeks old, were inoculated intraperitoneally with the RH strain of toxoplasma and 66 to 68 hours later the peritoneal exudate was harvested in saline. Such exudates were rich in parasites with very few white cells; any very rare blood-tinged or cellular exudates were discarded. The exudate was washed once in saline and 1 part of the deposit was mixed with 9 parts of distilled water and frozen and thawed three times. After minimal centrifuging to remove gross particles, the supernatant constituted the sensitizing antigen which was stored at −20°C.

CELL SENSITIZATION Pyruvic aldehyde-treated cells were thrice-washed in saline and resuspended in Sorenson's buffered

1Pyruvic aldehyde (40% aqueous solution) obtained from Messrs Kodak, Ltd, Kirby, Liverpool.
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Saline pH 6.0. The packed cell volume was obtained from a haematocrit tube estimation and the cell concentration adjusted to 1-4% with the pH 6.0 buffer. Antigen was also diluted in this buffer to the required concentration as determined by previous chessboard titration against the WHO Toxoplasma Reference Serum. The antigen standard employed was the highest dilution, usually 1:200 to 1:500, producing a haemagglutinating antibody titre of 1:1000 with the reference serum.

The diluted antigen was mixed with an equal volume of the 1-4% cell suspension and sensitization proceeded for one hour in a water bath at 56°C. (It was found to be important in producing batches of cells comparable in sensitivity to that of the initial titration to sensitize cells in small volumes. Thus if each tube of an antigen titration was of a 10 ml cell/antigen volume then larger volumes were sensitized in approximately 10 ml amounts.) On removal from the water bath, the coated cells were cooled under running tap water, washed once in saline and resuspended to an 0·7% concentration in Sorenson's phosphate-buffered saline, pH 7.2, containing 0·1% sodium azide. Storage was at 5°C.

Testing of Serum Specimens for Antibody

Haemagglutination (HA) tests were all carried out using the Microtiter* apparatus in permanent lucite V-welled plates with a standard volume of 0·05 ml. Doubling dilutions of the test serum were made in 2% serum saline (diluent serum was either foetal bovine serum or human toxoplasma antibody-free serum). Of the well mixed sensitized cells, 0·05 ml was added to each well, the plate tapped to mix, and left undisturbed on the bench at room temperature. Cell and antibody controls were included in each test. Results were read after leaving overnight at room temperature but in fact a two-hour reading was equally satisfactory.

Dye tests were performed in this laboratory by the method used in the PHLS Toxoplasma Reference Laboratory, Leeds, but with parasites harvested from the peritoneal exudate of infected cotton rats instead of mice.

Ultracentrifugation of sera was carried out on sucrose density gradients formed from 10 to 50% sucrose concentrations in an MSE Superspeed 50 at 40 000 rpm for 18 hours.

Titres obtained using sensitized cells were compared with dye test titres on the sera of patients grouped as follows:

(1) Twenty-two sera received from Dr G. B. Ludlam of the Toxoplasma Reference Laboratory, Leeds, where a diagnosis of acquired toxoplasmosis was based on clinical findings substantiated by significant dye test titres.

(2) Eighty-four sera from patients attending ophthalmic clinics where choroiditis was detected and where the ophthalmologist considered the diagnosis of toxoplasmosis likely or possible on clinical grounds.

(3) A group of 161 sera from cases of lymphadenopathy including patients with the glandular fever syndrome, rheumatoid arthritis, and others in a miscellaneous group where the diagnosis of toxoplasmosis was considered a possibility by the clinician concerned and who submitted the serum for toxoplasma antibody examination.

(4) One hundred and twenty-four embryopathy sera from babies in the Western Region of Scotland including two cases of congenital toxoplasmosis diagnosed on clinical and laboratory evidence (Ross et al, 1972).

(5) A group of 459 sera from pregnant women from the north of Scotland who were attending routine surveillance clinics.

Results

In group 1 (Fig. 1), in general a high titre group, 20 of the 22 sera gave readings which were either the same or within a single dilution reading between the

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Fig. 2 Haemagglutination and dye test titres of 84 ophthalmic clinic sera (group 2).

The two sera shown in Fig. 4, where both HA and dye test titres were in excess of 1:256, gave final titres of 1:1024 and 1:4096. The respective dye test titres were 1:512 and 1:8192. Thirty sera, however, had comparatively high HA titres (≥ 1:32) in the absence of correspondingly high dye test titres.

Four hundred and twenty-one of the 459 sera in this group (group 5) agreed within four-fold limits. Of the 38 sera outside those limits the preponderance was entirely towards higher HA readings. The behaviour of the HA and dye test antigens was followed through in serial serum samples from one antenatal patient where rising dye test titres suggested active infection. The samples from mother and baby were fractionated on sucrose density gradients and the results are shown in Figure 6.

Discussion

Using freshly sensitized cells, Jacobs and Lunde originally found that 93% of their sera tested by both HA and dye test had titres which agreed within a four-fold difference while 4% showed differences greater than 16-fold. The stable HA

two tests. In the two sera where the difference was greater, the dye test showed respectively an eight-fold and a 16-fold increase over the HA titre of 1:512.

In group 2, the agreement between the dye and HA titres was very close, only one serum of the 84 submitted showing a difference more than four-fold, as shown in Figure 2.

Group 3 is a heterogeneous group of sera and includes serum from cases with the glandular fever syndrome as well as cases of rheumatoid arthritis. In this group of 161 sera, of which 16 were positive by the Paul-Bunnell test and 17 by the Rose-Waaler test, 127 showed agreement between the toxoplasma dye and HA tests within fourfold limits. Positive Paul-Bunnell sera were easily identified by their agglutination of pyruvic aldehyde-treated but non-sensitized sheep cells. Since sheep cell agglutinins are not readily completely removed from high titre Paul-Bunnell-positive sera it is easier to identify them thus than to attempt absorption in routine practice.

One hundred and twenty-four embryopathy sera, all gathered in the neonatal period and earlier tested against a variety of viral antigens, made up group 4.
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Fig. 3 Haemagglutination and dye test titres of 161 sera including cases of lymphadenopathy, glandular fever and rheumatoid arthritis (group 3). X = Positive Paul-Bunnell test sera.

Fig. 4 Haemagglutination and dye test titres of 124 embryopathy sera (group 4).
Haemagglutination and dye test titres of 459 antenatal clinic sera (group 5).

Fig. 6  Haemagglutination and dye test results of testing sucrose density gradient ultracentrifugation fractions of serial samples of serum and cord blood from antenatal patient of group 5. (Fractions 1 to 3 are IgM; fractions 5 to 10 are IgG.)

antigen, standardized against the WHO reference serum and tested on the 850 sera comprising groups 1-5, was found to agree, within four-fold limits, with the dye test titre in 745 (88%) of the sera; 105 sera (12%) had greater than four-fold differences between the two tests. However, the correlation of titres varied widely according to the particular group being examined.
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Ninety-one per cent of the sera in group 1 were either the same, or within a single dilution difference in both tests, and in the ocular cases in group 2 agreement within a four-fold limit was as high as 99%.

Group 3 included positive Paul-Bunnell and also positive Rose-Waaler test sera. Although the difficulty of Paul-Bunnell sera could have been overcome by the use of sensitized human, rather than sensitized sheep, cells the best correlation between HA and dye test titres has been found by those workers employing sheep cells (Chordi et al., 1964). The three highest Rose-Waaler titre (1:256) sera had neither toxoplasma HA nor dye test antibodies; conversely one of the lowest Rose-Waaler (1:32) sera had a toxoplasma HA and dye test titre of 1:256.

Within the four-fold criterion the agreement with HA and dye titres in group 3 was 79% but if the 16 Paul-Bunnell-positive sera are omitted the level of agreement is increased to 88%.

The 124 embryopathy sera in group 4 showed the poorest correlation of HA and dye test titres, 24% of the sera tested had positive HA titres at least four-fold higher than the respective dye titres. However, the two highesttitre sera were from cases diagnosed on clinical grounds as congenital toxoplasmosis.

Group 5 showed a 92% agreement of HA and dye test titres within the four-fold limit. When a second specimen obtained from one of the patients in this group showed a raised dye titre a further specimen of blood and also cord blood was obtained at delivery. The HA and dye results of fractions separated in sucrose density gradients by ultracentrifugation were of value in demonstrating the sensitivities of the two tests.

Whilst there is evidence that the dye test and HA test measure different antibodies (Fleck, 1961) we have not found a positive dye test in the absence of HA antibody in any of the acute human sera nor in sera from many experimental animals which we have infected. As in the acute form of the acquired disease and in the congenital form the early antibody is of IgM nature (Remington, 1968), the HA test is particularly useful in these cases because of the relatively greater haemagglutinating efficiency of IgM antibody as opposed to antibody of the IgG class which appears to be more sensitively measured by the dye test (Fig. 6). If this is so the correlation of HA and dye results may rest to some extent on the relative concentrations of the immunoglobulin class of toxoplasma antibody present in the serum.

In the entire series of 850 sera, no serum was found with an HA titre of 1:32 which had a dye test titre greater than this.

We believe that the employment of this reagent would enable laboratories to undertake a wider screening of sera than would be practical if all sera had to be submitted to the dye test.

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References


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