A simple, rapid ELISA method for the detection of DNA antibodies

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SUMMARY Employing an enzyme-linked immunosorbent assay (ELISA) technique the serum antibodies against native (double stranded) and denatured (single stranded) deoxyribonucleic acid (DNA) have been measured in various disease groups and a group of blood donor sera.

The ELISA method has been compared with a radioimmunoassay method using native (double stranded) DNA as substrate antigen and a latex-fixation technique using particles coated with soluble deoxyribonucleoprotein (SNP). It is concluded that ELISA offers an economic and reliable alternative to isotope techniques for the assessment of antibody content in systemic lupus erythematosus (SLE) and related disease states for the clinical laboratory.

Deoxyribonucleoprotein is the major component of the nucleus of which the nucleic acid moiety forms the genetic material of the chromosome. Antibodies directed against soluble nucleoprotein and its constituent products, purines, pyrimidines and polynucleotides have been detected in patients with rheumatoid arthritis and associated diseases but antibodies directed against native (double stranded) DNA are a hallmark of patients with systemic lupus erythematosus (SLE).1,2 Antigens to native DNA have been particularly associated with clinical disease activity.2-4 Furthermore, it is generally believed that anti-DNA antibodies are the principal antibody component of circulating immune complexes in SLE and that the pathogenesis of their glomerulonephritis is analogous to experimental serum sickness.5-6 Clinical laboratories are called upon to perform reliable and reproducible tests for the detection of antibodies to DNA and clinicians expect that the results will aid their diagnosis.7-8 Quantitative assays which are sufficiently sensitive for the detection of antibodies specific for DNA have hitherto relied upon a radioimmunoassay method using isotope-labelled DNA which when complexed with antibody is subsequently precipitated with ammonium sulphate or a coprecipitating antibody,9-12 or a fluorescein-labelled second antibody in an immunofluorescent staining procedure involving as substrate the kinetoplast of the protozoan Crithidiculae luciae.13-16

The possibility of enzyme-linked immunosorbent assays has been investigated by several workers using denatured DNA,17-19 and more recently native DNA coated to polystyrene tubes with the aid of protamine sulphate.20 During the past twelve months Cordis Laboratories, Florida, have developed and marketed two commercial kits using as substrate rigid plastic discs coated with native DNA (Cordia N) or soluble nucleoprotein (Cordia NP). The present paper describes a reliable microtitre plate method suitable for routine screening of test sera for antibody activity directed against both native and denatured DNA.

Material and methods

Material

Sera
Blood donor sera (100 samples) were kindly supplied by the Regional Blood Transfusion Service, Birmingham. All other sera were specimens, sent for analysis to this department, which were stored at −20°C containing 1/10000 sodium azide as preservative. All dilutions of sera were made in a buffer containing: NaCl 40 g; Na2HPO4 5.75 g; KH2PO4 1 g; KCl 1 g; NaN3 1 g; Triton X-100 50 ml; made up to 5 l with distilled water. Immediately before use bovine

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serum albumin was added to a final concentration of 1 g/100 ml.

Latex agglutination
Fisher diagnostic kits for DNP antibodies in SLE were purchased from AR Horwell Ltd. Kits were used according to manufacturer’s recommendations.

DNA for radioimmunoassay
Native DNA, from Escherichia coli intrinsically labelled with 14C was purchased from Amersham International as 50 µg amounts in 5 ml lots. In order to avoid any denaturation each batch was pipetted into 10 x 0.3 ml and 10 x 0.2 ml aliquots immediately upon receipt and stored at -70°C. Aliquots were diluted 1/10 in phosphate-buffered saline (PBS) immediately before the test.

DNA for ELISA
Highly polymerised DNA extracted from calf thymus was used for all tests shown in the text. DNA from alternative sources were compared in the initial stages of this work. These included DNA from rat liver nuclei, E coli and herring sperm. All were purchased from Sigma Chemical Co Ltd.

Residual protein from calf thymus DNA was removed by overnight digestion at 37°C of a 5 mg/ml solution of DNA with 200 µg/ml trypsin. The final incubation mixture contained 50 mM trisodium EDTA and 1 g per 100 ml sodium lauryl sulphate in 0.2 M TRIS-HCl buffer at pH 7.4. The solution was extracted twice with an isoamyl alcohol: chloroform mixture (1/24 vol/vol). The DNA was recovered from the aqueous phase by carefully rotating a glass rod in the solution after addition of two volumes absolute ethyl alcohol. The “spoiled” DNA was washed in 70% ethyl alcohol, dried and dissolved in PBS.

Denatured DNA was prepared by plunging a tube of native DNA solution into a boiling water bath for 10 min then rapidly transferring tube and contents to an ice bath. Single stranded DNA was prepared as required from aliquots of native DNA stored at -20°C.

Antisera
The antihuman IgG used as second antibody in the radioimmunoassay technique was raised in this laboratory by injecting rabbits intramuscularly with human IgG. The immunogen was eluted from a DE 52 cellulose column in 0.01 M phosphate buffer (pH 8) and emulsified in Freund’s complete adjuvant.

Enzyme-labelled antihuman IgG for ELISA was purchased from Miles Laboratories Ltd, and used at a dilution of 1/1000 in Tween-PBS buffer.

TWEEN-PBS (“Wash”) buffer for ELISA
NaCl 40 g; Na2HPO4 5.75 g; KH2PO4 1 g; KCl 1 g; NaN3 1 g; Tween 20-2.5 ml; made up to 5 l with distilled water.

Substrate buffer for ELISA
Diethanolamine 97 ml; NaN3 0.2 g; MgCl2 6H2O 0.1 g; distilled water 800 ml; adjust to pH 9.8 using 1 M HCl and make up volume to 1 l. Pellets (5 mg) of p-nitrophenyl phosphate substrate were purchased from Sigma Chemical Co (one tablet per 5 ml substrate buffer for working solution).

Scintillation fluid for radioimmunoassay
2:5 diphenyloxazole (PPO) 6-0 g; 1:4 di-2,5-phenyloxazoyl benzene (POPOP) 0-12 g; Triton X-100 333 ml: xylene (scintillation grade) 666 ml.

METHODS

Radioimmunoassay for antibodies to DNA
To duplicate 0.1 ml aliquots (≈100 ng) of 14C DNA in 50 mm x 6 mm polystyrene precipitin tubes (Sterlin RT20) was added 0.1 ml of a 1/10 dilution of serum in PBS pH 7.4. After thorough mixing the tubes were incubated at 37°C for 1 h and then placed at +4°C overnight to allow completion of complex formation. The IgG in each tube was precipitated with the second antibody (rabbit antihuman IgG) allowing incubation at 37°C for one hour and +4°C for two hours.

The precipitate was removed by centrifugation and washed by resuspending in PBS and recentrifuging. The washed precipitate was dissolved in two drops of 1 M NaOH at 56°C for 30 min neutralised with HCl in methanol (1% vol/vol) and quantitatively transferred to a counting vial using scintillation fluid. Counts were made over 10-minute periods and the tests calculated as a percentage binding of the total tube. Appropriate positive and negative controls were included in each batch of tests.

ELISA method for antibodies to native and denatured DNA
To each well of a 96-well microtitre plate was added 0.2 ml of a 1% aqueous solution of protamine sulphate (Sigma). After 30 min at room temperature the plate was emptied by simple inversion and washed five times with a jet of distilled water from a plastic wash bottle.

DNA solution (0.2 ml) at a concentration of 10 µg/ml was added to the desired number of wells and allowed to stand at room temperature for 30 min. (For convenience native DNA could be applied
to one half of each plate and denatured DNA to the remaining half—in this way 48 sera including controls could be tested under identical conditions against both antigens.)

After DNA attachment the plate was washed five times in Tween-PBS buffer and 0.2 ml of each sample applied to the appropriate wells. After further incubation at room temperature for 30 min the plates were carefully washed five times in Tween-PBS buffer and shaken almost dry. An amount (0.2 ml) of 1/1000 dilution of alkaline phosphatase-labelled antihuman IgG was pipetted into each reaction well and the plate allowed to incubate at room temperature for a further 30 min.

After washing with Tween-PBS buffer five times the plate was again shaken dry and finally 0.2 ml of p-nitrophenyl phosphate solution added to each reaction well. Incubation was allowed to take place at 37°C for 20 min when the reaction was halted by the addition of 0.05 ml of 2 M NaOH. Hydrolysis of the substrate to p-nitrophenolylate was measured by determination of the absorbance at 410 nm on a Dynatech automatic plate reader. Appropriate positive and negative sera together with a "no serum" blank were included in each batch of tests.

Results

Method controls

In an attempt to demonstrate that positive tests observed in the microtitre wells were due to reaction of enzyme-linked anti-IgG reacting with antibodies directed against DNA antigen a series of controls, including inhibition assays, were set up.

Use of protamine sulphate

In our ELISA method, omission of the protamine sulphate incubation stage resulted in total negative tests. In a separate series of tests it was shown that in the absence of protamine sulphate at least 16 hours were required for adequate binding of denatured DNA; furthermore, for native DNA it took as much as four days to achieve comparable colour intensity and reproducibility from a positive control serum (see Fig. 1).

Pretreatment with protamine sulphate for 15 min or more facilitated maximum binding of DNA in 30 min.

Inhibition of reactions using soluble antigens

In inhibition experiments, sera that were native DNA-positive were inhibited by pretreatment with free double stranded antigen; furthermore, sera that were denatured DNA-positive were inhibited with free single stranded antigen. Not surprisingly, some cross reaction occurred, resulting in absorption of antibodies to single stranded DNA using free native DNA and absorption of antibodies to native DNA using free denatured DNA.

Enzyme degradation of coated antigens

DNA-coated plates treated at 37°C for 30 min with the enzyme deoxyribonuclease (Sigma) resulted in complete removal of antigen from the polystyrene surface. Conversely, treatment with nuclease S1 (Sigma), which is specific for single stranded DNA, digesting double stranded DNA only at damaged sites,21 removed all the antigen from only denatured DNA-treated plates making little significant difference to those plates coated with native DNA.

Colour intensity and antibody concentration

Eleven positive sera were tested for DNA antibody activity using doubling dilutions of serum; the results were plotted and the slopes of the graphs compared (Fig. 2a). From the data it is clear that colour intensity is proportional to the amount of antibody bound to the DNA antigen and that the slopes are very similar. To demonstrate more clearly the exact nature of the slope the individual results have been superimposed around an OD of 0.75. Colour intensity is linear between optical densities of 0.45 and 1.2 (Fig. 2b). One positive serum was tested on three consecutive days to demonstrate the reproducibility of the colour reaction. The results are shown in Fig. 2c.
Sera from Disease Groups

Figure 3 shows a comparison of antibody activity against native and denatured DNA between normal sera (100), sera from known patients with SLE (48), patients with liver disease (100), Crohn's disease (38) and coeliac disease (14), and rheumatoid arthritis (101). Figure 3 also shows the results of 100 consecutive sera, taken from a day's work load, and 100 other sera from patients with connective tissue and other diseases, found to have a strongly positive antinuclear antibody (titre > 1/100), although negative for double stranded DNA antibodies by the radioimmunoassay method.

All sera were diluted 1/50 in modified Tween-PBS (see Discussion). Patients from the rheumatoid group were serologically positive by standard Rose-Waaler technique. Many of these sera were also known to have a positive antinuclear antibody (ANA).

Sera tested in the "liver diseases" group were from patients with jaundice or cirrhosis, selected on the basis of positive smooth muscle and mitochondrial antibodies by fluorescent antibody test, but these excluded all sera positive for ANA.

From the results in Fig. 3 it can be seen that on the one hand not one serum in 100 normal sera was found to have any antibody activity to either native or denatured DNA and on the other not one serum in the 48 SLE sera tested using the same antigens (see under next heading) was found to be negative. Weak antibody activity in both the rheumatoid and the liver disease groups is not unexpected for it is well documented that low activities of antibodies to ANA, DNA and RNA are present in many autoimmune diseases, probably directed against the protein moiety of cellular constituents. Of the "random" specimens taken from the daily work load one which gave an optical density reading of 1-5 was found to be a patient with SLE.

Sera Run in Parallel with Current Radioimmunoassay

During a six-month period all specimens entering
the laboratory requesting DNA antibodies were tested by both RIA and ELISA and the results compared. It can be seen from Table 1 that agreement was reached on over 85% of all specimens tested, almost two-thirds being unequivocally negative by both tests and a further quarter, positive by both tests. Of the remaining 15% of tests only two specimens from a total of 400 were found to be consistently negative by ELISA for antibodies directed against native, double stranded DNA but positive by RIA. Both of these specimens had considerable activity against denatured DNA in the ELISA test and probably reflect the quiescence of the disease state referred to earlier.\textsuperscript{5–4} In the RIA test they were only weakly positive, at a level of antibody binding where reproducibility is poor.

Over the same six-month period 570 specimens were tested by the latex agglutination method using deoxyribonucleoprotein sensitised particles (Fisher). From Table 2 it can be seen that there is total agreement with almost 80% of sera tested and that the enzyme-linked method picks up more “positive” sera than does the latex test.

The results obtained by ELISA were compared to the results obtained by radioimmunoassay (Fig. 4). Though sera were found to be positive by both methods, the correlation of colour intensity by ELISA with percentage binding by radioimmunoassay was not always good. Some specimens exhibited a seemingly disproportionately high colour intensity

\begin{table}[h]
\centering
\begin{tabular}{|l|l|l|}
\hline
Test reaction & No of sera & %
\hline
ELISA & - & 98 & 24.5
\hline
RIA & - & 56 & 14.0
\hline
ELISA & - & 244 & 61.0
\hline
RIA & - & 2 & 0.5
\hline
Total & - & 400 & 100.0
\hline
\end{tabular}
\caption{Comparison of ELISA and RIA in 400 sera sent for DNA antibody determination}
\end{table}
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Table 2  Comparison of ELISA and the DNP Slide Latex test in 570 sera sent for DNA antibody determination

<table>
<thead>
<tr>
<th>Test reaction</th>
<th>No of sera</th>
<th>%</th>
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</thead>
<tbody>
<tr>
<td>ELISA +</td>
<td>104</td>
<td>18-3</td>
</tr>
<tr>
<td>Latex</td>
<td>2</td>
<td>0-3</td>
</tr>
<tr>
<td>ELISA +</td>
<td>117</td>
<td>20-5</td>
</tr>
<tr>
<td>Latex</td>
<td>347</td>
<td>60-9</td>
</tr>
<tr>
<td>ELISA +</td>
<td>2</td>
<td>0-3</td>
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<tr>
<td>Total</td>
<td>570</td>
<td>100-0</td>
</tr>
</tbody>
</table>

Table 2 Continued

<table>
<thead>
<tr>
<th>Test reaction</th>
<th>No of sera</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Latex</td>
<td>2</td>
<td>0-3</td>
</tr>
</tbody>
</table>

Fig. 4  Scattergram of results from serum samples estimated for DNA antibody activity estimated by ELISA and RIA to show the correlation between values by the two tests. Results below the solid line drawn at 5% binding (RIA) are considered to be negative by this test whilst those above the broken line at 10% are unequivocally positive. Results below 0-20 OD at 410 (to the left of the vertical line) are considered to be negative by ELISA in the method as described. The hatched area represents results from 244 sera which were too numerous to plot individually.

compared to the DNA binding by RIA. Conversely very high binding by RIA did not necessarily yield an intense colour at a high dilution that one would expect. This is not a unique finding but has been reported previously for ANA by fluorescence compared to ELISA. However, all sera reacting with one preparation of DNA may not react identically with another preparation of DNA from a different source. The overall statistical correlation coefficient of results of the 76 sera positive by radioimmunoassay (> 10% binding) tested in parallel with ELISA was 0-64 (p > 0-001). Analysis of all 98 sera positive by RIA (that is, > 5% binding) gave a correlation coefficient of 0-62 (p > 0-001). The correlation was less good when results from sera that were positive by ELISA but negative by RIA were included in the analysis.

Discussion

The potential for ELISA assays to detect antibodies to DNA has long been recognised but difficulties encountered in current methodology and quantitative expression have resulted in a reticence to introduce the technique into routine laboratory practice. The present paper describes a simple, sensitive, rapid and reliable method for the quantification of antibodies to both native and denatured DNA which overcomes many of these problems.

There have been conflicting reports on the significance of antibodies to native and denatured DNA. Early reports that only sera from SLE patients reacted with double stranded DNA were followed by reports of anti-double stranded DNA in drug-induced SLE, but these reactions may be due to antibodies directed not against the nucleic acid but against deoxyribonucleoprotein or against histone. Cohen and Steward claimed that the vast majority of SLE sera react preferentially or even exclusively with single stranded DNA rather than double stranded DNA and work by Winfield recorded high avidity binding to single stranded DNA but differing avidity to double stranded DNA.

A major difficulty in current techniques is the selection and preparation of the antigen to be used in the test procedure. The antigen in its pure native form is exceptionally difficult to prepare without traces of denatured, broken or nicked molecules and as little as 0-1% denaturation of double stranded DNA will provide sufficient antigen sites for significant reaction with antibodies to single stranded DNA. In a series of papers on the quantification of DNA antibodies by the Farr assay, the effects of antigen size and structure and the influence of reaction conditions on the level of DNA binding have been described.

It is clear that discrepancies do arise when comparing results obtained for anti-DNA antibodies by
ELISA to those by methods using antigen in free solution. These discrepancies may be due to antibodies directed against new antigenic determinants exposed during conformational changes imposed upon the molecule when adhering to the solid phase carrier. An alternative explanation for the quantitative anomalies may be the difference in affinity and avidity of the antibodies from various patients for the antigenic sites of the DNA molecule.

**REDUCTION IN BLANK READINGS**

During the initial stages of this work considerable difficulty was experienced with high blank readings when using previously published methods. This was overcome by increasing the Tween (or Triton X-100) content to 1% (vol/vol) and adding 1 g/100 ml bovine serum albumin to the solution used for all serum dilutions.

High blank readings were also found when plates coated with protamine sulphate were stored for subsequent use. As the method developed here added only one 30-minute incubation to a technique which could be completed in just two and a half hours all plates were routinely precoated immediately before DNA addition on the day of the test to circumvent the latter difficulty.

**CHOICE OF ANTIGEN**

DNA from various sources (rat liver, calf thymus, *E. coli* and herring or salmon sperm) have been used as substrate antigen by different authors. A comparison of these antigens in both native and denatured form was made using microtitre plates with and without protamine sulphate pretreatment. Though minor differences did arise there was no significant difference between the antigens employed in the described test. DNA from rat liver seemed to be more resistant to denaturation by boiling, giving consistently lower results for single stranded DNA. As calf thymus DNA is readily available commercially in an acceptably pure form this antigen was adopted for routine use.

**MICROTITRE PLATES AND USE OF PROTAMINE SULPHATE**

Day-to-day variation of optical density was found to be minimal using the ELISA technique on DNA bound to polystyrene through protamine sulphate when using doubling dilutions of any one of the positive sera on microtitre plates. Changing the manufacturer or batches from the same manufacturer, made no significant difference to the results. The test was not subject to edge effect nor indeed to any positional effect within one plate. Microtitre plates designed for use in haemagglutination techniques ("U" well or "V" well) were perfectly adequate for the test obviating the unnecessary expense of purchasing flat bottomed trays specifically designed for protein adherence in ELISA methodology.

**EXPRESSION OF RESULTS**

Direct interlaboratory comparison of quantitative assessment of antibody activity against DNA is absolute terms is at present difficult. Results are frequently expressed as percentage binding of a given quantity of labelled DNA or as a percentage of the activity of an in-house control serum which has no direct relation to any other control serum. Throughout the present paper statistical data has been calculated by comparing percentage binding of a control standard serum by RIA with colour intensity produced by ELISA using the same standard serum at a known series of dilutions. That dilution of serum yielding an optical density of 0.75 is then expressed in U/ml. The dilution may have to be mathematically derived from the linear portion of the curve. In this way any day-to-day variation in colour intensity was removed by direct comparison to the standard serum, the value of which was fully evaluated. A positive control serum is included in each DNA kit supplied by Cordis but though this serum is given a value (in IU) it is as yet related only to the WHO international ANA standard for immunofluorescent techniques.

The ELISA method offers several advantages over others currently available: (a) it is quantitative for the antibody, (b) it is cheap, (c) it does not use radionuclides, (d) it may readily be automated, (e) it is sensitive to low antibody activities, (f) it is a simple, rapid technique which may conveniently be applied to other antigens. The method overcomes the subjectivity of fluorescent methods using kinoplasts which, though truly specific, are difficult to interpret in the weak positive range even by experienced personnel. Finally, in attaching the DNA molecule to a solid phase carrier through protamine sulphate one reduces the electrostatic charge which causes non-specific protein-protein interreactions in free solution. The development of a cheap ELISA method seemed a natural and desirable progression to enable the Department to provide an improved service in this field. Though absolute correlation of colour intensity with percentage binding by RIA has not yet been adequately explained we feel that the method described affords a simple, practical way of screening large numbers of sera for DNA antibodies.
References


Requests for reprints to: Dr RA Thompson, Regional Immunology Department, East Birmingham Hospital, Birmingham B9 5T, England.