Laboratory diagnosis of rheumatoid arthritis: a solid phase radioassay for IgG and IgM antiglobulins

L. J. NINEHAM, F. C. HAY, AND I. M. ROITT

From the Department of Immunology, The Middlesex Hospital Medical School, London W1P 9PG

SYNOPSIS  A technique suitable for the routine estimation of IgM and IgG antiglobulins has been devised. The assay involves the binding of antiglobulins to plastic tubes coated with rabbit immunoglobulin: the amount of antiglobulin bound is then determined by adding radiolabelled antihuman IgG or IgM. The conditions for the assay have been examined and optimal incubation times and amounts of reagents established. Verification of the antibody nature of antiglobulin activity has been obtained. Both IgG and IgM antiglobulins were raised in virtually all seropositive rheumatoid arthritics, and most seronegative patients gave elevated values for either IgM or IgG rheumatoid factors. The use of an anti-light chain reagent as a screening test for total antiglobulins was investigated. These tests should prove valuable in diagnosis and permit quantitative evaluation of research studies.

In rheumatoid arthritis, antiglobulins or rheumatoid factors are present in both serum and synovial fluid. Although most laboratory diagnostic procedures depend on the measurement of IgM rheumatoid factors, it has become increasingly apparent that IgG antiglobulins play an important role in the pathogenesis of the synovial inflammatory response. Previously in this laboratory a procedure was developed for the detection of IgG antiglobulins involving elution from an immunoabsorbent, followed by quantification by single radial immunodiffusion (Torrigiani and Roitt, 1967). The immunoabsorbent was prepared from rabbit immunoglobulin cross-linked with benzidine. This technique, although reliable for estimating antiglobulins in each immunoglobulin class, was too time-consuming for routine use.

Therefore we have developed an assay which provides a rapid and simple means of routinely estimating antiglobulins of known class. Catt and Tregear (1967) have shown that plastic tubes may be used to bind proteins and we have used this principle for our radio assay. Antiglobulins are bound to rabbit immunoglobulin linked to the surface of plastic tubes; the amount of antiglobulin bound is then determined by adding radiolabelled antihuman IgG or IgM. In the present paper, we amplify an earlier preliminary report (Hay et al, 1975) and look at the possible use of anti-light chain as a screening reagent.

Material and methods

ANTISERA  Anti-human IgG, IgM, light chain, and ovalbumin were prepared by immunizing rabbits with purified FcY, IgM, light chains or ovalbumin (Hudson and Hay, 1976). The anti-IgM was rendered specific by adsorption with a cyanogen bromide-activated Sepharose-4B immunoabsorbent of cord serum. The antisera were checked for specificity by immunoelectrophoresis, Ouchterlony immunodiffusion, and radioimmunoassay. Purified antibody was prepared by adsorbing the antisera to cyanogen bromide-activated Sepharose-4B immunoabsorbents of human IgG, IgM, Fab or ovalbumin, followed by elution of the specific antibody with 0-1 m glycine-HCl buffer, pH 2-8. The purified antibodies were then radio-labelled with 125I (Na 125I, IMS 4, Radiochemical Centre, Amersham) by a modification of Hunter and Greenwood's (1962) method. Antibody 1 mg in 1 ml phosphate buffered saline (PBS), 0-15 m, pH 7.2, was labelled with 500 μCi 125I by adding 60 μg chloramine T. After two minutes' incubation at room temperature 120 μg sodium metabisulphite was added to stop the reaction. Free iodine was removed on a Sephadex G25 column. The labelled proteins were stored at -20°C.
COATING OF THE PLASTIC TUBES
One-millilitre volumes of rabbit immunoglobulin (Miles Laboratories, Stoke Poges) at concentrations of 0-1, 1-0, 10, 100, 1000, 5000, and 10 000 μg ml⁻¹ in PBS were incubated in polystyrene tubes (LP3, Luckham Ltd, Sussex) for one hour at 37°C and overnight at 4°C. After three washes with PBS, the tubes were incubated at room temperature for two hours with 2 ml of 1% bovine serum albumin (Armour Pharmaceutical Co Ltd, Eastbourne) in PBS (BSA/PBS) to block any remaining free sites. After three more washes with PBS the tubes were inverted and stored at 4°C.

BINDING OF SERUM ANTIGLOBULINS
The test sera were heated at 56°C for 30 minutes to inactivate complement. Duplicate 5, 25, 50, and 100 μl samples were placed in the coated plastic tubes together with BSA/PBS to a final volume of 0-5 ml. Coated tubes containing 0-5 ml BSA/PBS were used as background controls. The tubes were incubated at 37°C for one, three or five hours and then at 4°C for a half, one, two, four or six hours. Unbound proteins were removed by three washes with cold PBS.

DETECTION OF BOUND ANTIGLOBULINS
Human antibody bound to rabbit immunoglobulin coated tubes was detected by the addition of 0-1, 1-0 or 10-0 μg of purified radiolabelled anti-IgG, IgM or light chain in 1 ml BSA/PBS. Tubes were incubated at 37°C for one, three or five hours and then at 4°C for a half, one, two, four or six hours. Unbound labelled reagent was then removed by three washes with cold PBS. The tubes were counted in a gamma-ray spectrometer, the amount of radioactivity bound being a measure of the IgG, IgM or total antiglobulin in the patient's serum.

PAPAIN DIGEST OF HUMAN ANTIGLOBULINS
Immunoglobulins were prepared from pooled rheumatoid arthritis sera by precipitation with 45% saturated ammonium sulphate. After dialysis against PBS containing 0-01 m cysteine and 0-002 m EDTA, the immunoglobulins were digested with papain (2 mg/100 mg) for four hours at 37°C and undigested immunoglobulin was removed by gel filtration on Sephadex G 100.

PATIENTS
Sera were obtained from 82 patients (25 men and 57 women) with rheumatoid arthritis whose sera reacted positively in the sheep cell agglutination test (SCAT) (Roitt and Doniach, 1969) and from 22 SCAT-negative rheumatoid arthritis patients (3 men and 19 women). All these patients conformed with the American Rheumatism Association criteria for probable, definite or classical rheumatoid arthritis.

Samples were also obtained from 12 patients with osteoarthritis (4 men and 8 women), six with traumatic injury (2 men and 4 women), and 13 normal controls working in the immunology laboratory (7 men and 6 women). All the sera were frozen on the day of collection and kept at −20°C until tested.

Results

VARIATION OF ASSAY CONDITIONS

Tube coating
Using a fixed 50 μl amount of patient's serum, there is maximum binding of IgG antiglobulins at a coating concentration of rabbit immunoglobulin between 1 and 10 μg ml⁻¹ (fig 1). At higher coating concentra-

![Graph of antibody binding variation](http://example.com/fig1.png)

**Fig 1** Effect of variation in concentration of rabbit Ig used for coating. Tubes were coated with 1 ml volumes of solutions containing different amounts of rabbit immunoglobulin. Patient and control sera were then assayed by the standard procedure.

Amount of patient's serum added
Varying amounts of patients' sera, from 5 to 100 μl,
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were added to the rabbit immunoglobulin coated tubes. Increasing antibody was detected up to 50 µl but thereafter a plateau was reached (fig 2). Therefore 50 µl was used in all subsequent assays.

**Fig 2**  Dose response curves obtained with increasing amounts of patients' sera and an albumin control.

**Time of incubation of patient's serum**
Varying the incubation time of patients' sera at 37°C from one to five hours did not alter the amount of binding whereas a small increase in binding occurred on incubation at 4°C from half an hour up to two hours. Prolonged incubation to six hours did not give greater binding. Therefore one hour at 37°C and half an hour at 4°C was used routinely.

**Amount of anti-IgG added**
Human antiglobulins bound to the coated tubes were detected by incubating the tubes with 0-1 to 10-0 µg ml⁻¹ of purified anti-IgG. Figure 3 shows a linear relationship between the amount of anti-IgG added and the amount bound over this range. In all further assays 1 µg was used.

**Time of incubation with labelled reagent**
Although significantly greater specific binding was obtained by incubating the anti-IgG reagent for five hours at 37°C, good discrimination was already observed after one hour. Similarly, at 4°C an increase was seen up to six hours but discrimination was just as good at half an hour as at six hours. Therefore one hour at 37°C and half an hour at 4°C were used routinely.

**ROUTINE ASSAY CONDITIONS**
For routine use we selected the minimum incubation times and amounts of serum and reagents which

**Fig 3** Dose response curves with increasing amounts of radiolabelled antihuman IgG tested against patient and control sera.

**Fig 4** Assay reproducibility. Sera from patients and controls were tested on different days. The samples were refrozen between each assay.
gave good discrimination between patients and controls. The tubes were coated with 10 μg rabbit immunoglobulin in 1 ml for one hour at 37°C and overnight at 4°C. Fifty microlitres of patients’ serum was incubated for one hour at 37°C and for half an hour at 4°C, followed by 1 μg of labelled antibody for one hour at 37°C and half an hour at 4°C. The total incubation time was only three hours.

REPRODUCIBILITY OF THE ASSAY
Sera from three patients and two normal healthy controls were assayed on seven to nine different days. Between assays the samples were refrozen at −20°C. Figure 4 shows the assay to be highly reproducible, the mean coefficient of variance for the three patients being only 0·11.

SITE OF ANTIGLOBULIN ACTIVITY
The possibility existed that the binding of IgG reflected a property of a particular type of Fc region rather than conventional antibody activity associated with the Fab region. Table I shows that with whole

<table>
<thead>
<tr>
<th>Antiglobulin preparation</th>
<th>Radiolabelled second antibody</th>
<th>Anti-light chain</th>
<th>Anti-Fc</th>
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<tr>
<td>Whole IgG</td>
<td></td>
<td>108</td>
<td>125</td>
</tr>
<tr>
<td>Papain digest</td>
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<td>59</td>
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Table I Localization of the binding site for rabbit IgG in human IgG antiglobulin assayed by solid phase radioassay

mg antibody bound per tube

IgG, isolated from a pool of patients’ sera, antiglobulins were demonstrable by radioassay with both anti-light chain and anti-Fc antisera. However, with the IgG papain digest, activity was detected only by the anti-light chain reagent, demonstrating that the binding of antiglobulins was dependent solely on the Fab portion of the molecule.

IgM ANTIGLOBULINS IN RHEUMATOID ARTHRITIS PATIENTS
Raised levels of IgM antiglobulins were obtained in all rheumatoid arthritis patients found to be SCAT-positive and also in some SCAT-negative rheumatoid arthritis patients. Table II shows that the means of both groups were significantly different from the mean for the laboratory staff. Patients with traumatic injury or osteoarthritis gave even lower values than the laboratory staff control group. There was a significant correlation between the SCAT titre and the amount of IgM antiglobulin (r = 0·43; P < 0·001).

<table>
<thead>
<tr>
<th>SCAT-positive RA</th>
<th>Mean ± standard error</th>
<th>P (v staff)</th>
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<tbody>
<tr>
<td>1·98 ± 0·06</td>
<td></td>
<td>&lt;0·01</td>
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<tr>
<td>SCAT-negative RA</td>
<td>0·87 ± 0·08</td>
<td>&lt;0·01</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>0·37 ± 0·06</td>
<td>&lt;0·1</td>
</tr>
<tr>
<td>Staff</td>
<td>0·60 ± 0·06</td>
<td></td>
</tr>
<tr>
<td>Traumatic injury</td>
<td>0·46 ± 0·10</td>
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</table>

Table II Serum IgM antiglobulins in patients with rheumatoid arthritis and controls

NSD = not significantly different

mg anti-IgM bound 1−1 serum

IgG ANTIGLOBULINS
IgG antiglobulins were also detected in both SCAT-positive and SCAT-negative rheumatoid arthritis groups (table III), the means being significantly

<table>
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<th>SCAT-positive RA</th>
<th>Mean ± standard error</th>
<th>P (v staff)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0·48 ± 0·03</td>
<td></td>
<td>&lt;0·01</td>
</tr>
<tr>
<td>SCAT-negative RA</td>
<td>0·29 ± 0·05</td>
<td>&lt;0·01</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>0·13 ± 0·01</td>
<td></td>
</tr>
<tr>
<td>Staff</td>
<td>0·12 ± 0·01</td>
<td></td>
</tr>
<tr>
<td>Traumatic injury</td>
<td>0·10 ± 0·01</td>
<td></td>
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</tbody>
</table>

Table III Serum IgG antiglobulins in patients with rheumatoid arthritis and controls

NSD = not significantly different

raised compared to the mean for control groups. Not all rheumatoid arthritis patients had raised levels of both IgG and IgM antiglobulins, but 96% of the samples examined gave raised values (over 2 standard deviations above the combined mean of the laboratory staff and traumatic injury patients) for either IgG or IgM antiglobulins (fig 5).

In seropositive rheumatoid arthritis there was a good correlation between the amounts of IgG and IgM antiglobulins (r = 0·64; P < 0·001). However, there was little correlation in seronegative rheumatoid arthritis (r = 0·12; 0·6 > P > 0·5). The SCAT titre in the seropositive patients correlated just as well with the level of IgG antiglobulin (r = 0·44; P < 0·001) as it had with the IgM antiglobulins.

TOTAL ANTIGLOBULINS
As the light chains are common to all immunoglobulin classes, we have tested some of the sera with anti-light chain as the radiolabelled reagent to detect antiglobulins in all classes. Figure 6 shows raised levels of antiglobulins in all SCAT-positive and 50% of SCAT-negative rheumatoid arthritis patients compared with the low levels in laboratory staff and osteoarthritis and traumatic injury patients.

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CONTROL FOR BINDING OF NORMAL RABBIT IgG BY IgM RHEUMATOID FACTOR

It was possible that the IgM rheumatoid factors might attach to the rabbit immunoglobulins on the tube through only some of its Fab regions, leaving the others free. These could then combine with the Fc region of the radiolabelled rabbit anti-IgG used for detecting IgG antiglobulins. To test this hypothesis, radiolabelled purified rabbit anti-ovalbumin was added to the tubes in parallel with anti-IgG and anti-IgM. Table IV shows that only very low levels of binding were obtained with anti-ovalbumin even in the presence of large amounts of rheumatoid factor.

Discussion

In the laboratory the diagnosis of rheumatoid arthritis traditionally depends on the detection of rheumatoid factors in the classical latex and sheep cell agglutination tests. Unfortunately, positive results with these techniques are obtained only with 19S IgM rheumatoid factors. However, there is now increasing evidence that IgG antiglobulins may be more important in the pathogenesis of rheumatoid arthritis. Winchester et al (1970) have shown that complexes in the synovial fluid of rheumatoid arthritis patients are composed almost entirely of IgG, and Winchester (1975) has since shown that at least 50% of this IgG has antiglobulin activity. The importance of these antiglobulin complexes has been demonstrated by their ability to activate complement and to cause a reduction in total haemolytic complement within the synovial fluid. Furthermore, Pope et al (1974) have shown that the intermediate complexes found in the serum of three rheumatoid patients were composed entirely of self-associated IgG antiglobulins, while Theofilopoulos et al (1974) have evidence which strongly suggests that IgG antiglobulins play a part in the development of rheumatoid vasculitis.

To simplify the detection of IgG antiglobulins,
several workers have modified the immunoabsorbent procedures (Bianco et al., 1971; Bianco et al., 1974; Florin-Christensen et al., 1974). However, all of these techniques are still time-consuming, and some of the immunoabsorbents, particularly activated agarose (Bianco et al., 1974), give uneconomically high binding of IgG with normal controls. In the present work we have found that binding to antigen-coated plastic tubes provides a convenient and rapid assay which is free from these problems. Repeated testing of samples on up to nine different occasions has shown the assay to be highly reproducible.

In determining the parameters for the assay, we have found that the concentration of rabbit immunoglobulin used for coating the tubes to be important, since less antiglobulin is detected at coating strengths both above and below the optimum. In order to obtain low backgrounds with controls, it is essential that only purified antibody should be used as the radiolabelled reagent. Most other variables can be considerably altered while still giving good discrimination, and thus for our routine assay we have chosen those conditions most economical with respect to incubation times and amounts of reagents. However, the results show that for increased sensitivity, prolonged incubation is required and a saturating amount of radiolabelled antibody should be added. Preliminary studies have shown that the use of an enzyme-linked anti-Ig in place of the iodine labelled antibody is perfectly feasible.

Our technique has enabled us to investigate the site of antiglobulin activity within the immunoglobulin molecule. After papain digestion of immunoglobulin from rheumatoid arthritis patients, only the Fab regions bound to the rabbit immunoglobulin coated tubes. Thus it seems likely that we are detecting a true antigen-antibody reaction in our assay rather than non-specific adherence through the Fc region.

Virtually all the SCAT-positive sera gave raised IgM and IgG antiglobulin values. Most SCAT-negative sera also showed raised antiglobulins though the mean was lower than that for the SCAT-positive group. When radiolabelled anti-light chain was used as a screening reagent for total antiglobulins, all the SCAT-positive and many of the SCAT-negative sera had raised levels. However, as our anti-light chain results showed a greater correlation with IgM than with IgG antiglobulins, we have found that with SCAT-negative sera, it is also necessary to look for IgG antiglobulins using an anti-IgG reagent.

Since the test allows the rapid estimation of both IgM and IgG antiglobulins it should be of value routinely in the diagnosis of adult rheumatoid arthritis and Still's disease.

We thank Mrs Noemi Ron for assistance with the anti-light chain assays. We are grateful to Miss Hilary Fischler for the preparation of the manuscript. This work was supported by the Medical Research Council of Great Britain.

References


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doi: 10.1136/jcp.29.12.1121

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