Measurement of rheumatoid factors by an enzyme-linked immunosorbent assay (ELISA) and comparison with other methods

J L M Bampton, T E Cawston, M V Kyle, and B L Hazleman

From the Rheumatology Research Unit, Addenbrooke’s Hospital, Hills Road, Cambridge CB2 2QQ

SUMMARY IgM rheumatoid factor (RF) was measured in the sera of 48 rheumatoid patients and of 48 age and sex-matched normal controls by the Rose-Waaler and latex agglutination tests, a rate nephelometer, and an enzyme-linked immunosorbent assay (ELISA). Good correlation was obtained between all assays. The rate nephelometer assay was the easiest and quickest to perform and gave results in international units/ml. The Rose-Waaler was the least sensitive assay and the most difficult to perform and interpret. Both the latex agglutination and the ELISA were sensitive, though some overlap of patient and control sera was seen with all the assays. In addition to IgM RF the ELISA was used to measure IgG RF and IgA RF in both rheumatoid and control sera. Although some normal sera had detectable amounts of IgG and IgA RF, the levels of both were significantly raised in the rheumatoid sera. IgG RF levels were lower after pepsin digestion of the sera, suggesting that IgM RF interfered with the assay for IgG RF unless this treatment was included.

Key words: arthritis rheumatoid, Rose-Waaler test, latex fixation test, nephelometry.

Rheumatoid factors (RF) are anti-IgG autoantibodies with specificity directed against antigenic determinants in the Fc region of IgG.1 They are present in the large majority of patients with rheumatoid arthritis and have long been found to be associated with the more severe forms of rheumatoid disease. The measurement of RFs is important in the diagnosis of rheumatoid arthritis and in determining prognosis, especially in high titres, as these patients tend to develop extra-articular complications. In addition it has been proposed that RFs are important in the pathogenesis of the disease.2 There is considerable evidence that RF contributes extensively to the immunopathology of RA, as it is the predominant antibody known to initiate immune complex formation and complement activation in the peripheral circulation and in tissue sites.2 3

The majority of routine laboratory tests for the detection of rheumatoid factor measure IgM RF by its ability to agglutinate sheep red blood cells, latex, or similar particles coated with IgG.4-6 However, recent publications have suggested that both IgG RF7 and IgA RF are also raised in seropositive RA, and IgG RF may be associated with the more severe extra-articular complications of the disease such as vasculitis.3 8

The majority of hospital laboratories still routinely measure IgM RF by either the Rose-Waaler or latex agglutination test. These methods are both difficult to quantitate, and we have found that there is a large variation in titres for the same sera between different laboratories. In addition it is not possible to measure either IgG RF or IgA RF by these methods. Although procedures have been described for the measurement of different classes of RF, specialised reagents and equipment are generally required. More recently suitable ELISA methods have been reported which can be routinely used in clinical laboratories.3 9-10

In this study we have compared four methods for the estimation of IgM RF and developed an ELISA method that can also measure IgG RF and IgA RF.

Materials and methods

All reagents were of analytical grade and were
obtained from Sigma Ltd, Poole, UK, unless otherwise indicated.

Patients and controls. Sera were obtained from 48 rheumatoid patients with active disease attending routine rheumatology outpatient clinics and from 48 age and sex matched normal healthy blood donors. All sera were stored at −70°C, thawed before use and heated at 56°C for 30 min before use to inactivate complement.

Development of ELISA. Rabbit IgG (R IgG) and human Fc were both tested as antigen in an ELISA. No significant difference was found between them, and R IgG was used in all further tests.

Evaluation of microtitre plates. Microtitre plates were obtained from four manufacturers: Nunc (Gibco Ltd, Paisley, Scotland), Falcon (Becton Dickinson UK Ltd, Cowley, Oxford), Linbro (Flow Laboratories Ltd, Irvine, Ayrshire, Scotland), and Dynatech (Dynatech Laboratories, Billingham, Surrey) and tested by measuring the amount of 125I-labelled R IgG bound to each well. Although Falcon plates bound slightly greater amounts of IgG than Linbro or Dynatech, no significant difference in final sensitivity was found, and the lower-cost Linbro plates were subsequently used. No significant difference was found between the amount of R IgG bound by wells at the centre and wells at the edge of any plate.

Binding of R IgG to microtitre plates. The amount of 125I-labelled R IgG bound to microtitre plates was estimated under different conditions of time (2–24 h), temperature (4°C, 22°C, 37°C), pH (2-0, 7-2, 9-6), and concentration (10–500 μg/ml). The optimum conditions chosen for binding of R IgG were an overnight treatment at 37°C of 50 μl of a 100 μg/ml solution in 50 mM sodium carbonate buffer pH 9-6. Blocking of non-specific antibody adsorption after IgG coating. No significant difference in titre was found between wells treated with bovine serum albumin (100 μg/ml) in either phosphate buffered saline (PBS) or sodium carbonate buffer (pH 9-6) for 1 hour at 37°C after R IgG adsorption and wells when this treatment was omitted.

Investigation of temperature and time of incubation. Various incubation times and temperatures were investigated for each stage of the ELISA assay. The optimum conditions chosen for a one-day assay were incubation of the plate with serum for 3 h at 4°C; incubation with enzyme conjugate for 2 h at 4°C, and incubation with enzyme substrate for 30 min at 37°C.

Investigation of enzyme substrate. The substrates o-phenylenediamine (OPD) and 2.2'-azino-di(3-ethyl-benzthiazoline-6-sulphonic acid) for peroxidase and p-nitrophenyl phosphate for alkaline phosphatase were investigated. The most sensitive substrate was OPD, and so horseradish peroxidase-antihuman immunoglobulin conjugates were subsequently used.

Final ELISA protocol. R IgG (50 μl) at a concentration of 100 μg/ml in 50 mM sodium carbonate buffer pH 9-6 was added to Linbro ELISA microtitre plates. Wells containing sodium carbonate buffer (50 μl) only were prepared as controls. The plates were sealed with Nescofilm (Jencons Scientific Ltd, Leighton Buzzard, UK), incubated overnight at 37°C, then washed three times with PBS + 0.05% Tween 20 (PBST), soaking for 3 min between each wash. Serum was diluted 1:10 and in a further seven 1:4 dilutions across the plate, so that each serum was tested in a dilution range of 1:10 to 1:138 240. Aliquots (50 μl) of each dilution were added to test and control wells. The plates were incubated for 3 hours at 4°C and then washed three times as previously described. The peroxidase-conjugated second antibody was diluted 1:1000 in PBST. Aliquots (50 μl) were added to each well and the plates incubated at 4°C for 2 h. OPD (20 mg) was prepared immediately before use in citric-phosphate buffer pH 5-0 (20 ml) and then H2O2 (8 μl of 30% solution) was added. Aliquots (50 μl) of OPD in buffer were added to each well and the plates incubated in the dark at 37°C for 30 min. The reaction was stopped by the addition of 2 M H2SO4 (50 μl) to all wells and the absorbance of each well read at 492 nm with a Titertek multiscan (Flow Laboratories, Irvine, UK). The results were expressed as log10 titre at an absorbance value of 0-4.

Measurement of RF immunoglobulin class. IgM, IgG, and IgA RF were detected with peroxidase-conjugated rabbit antihuman μ chain and antihuman α chain immunoglobulins (Dako, Mercia Brocades Ltd, Weybridge, UK). IgG RF was detected by means of a rabbit antihuman IgG/Fab antibody (Nordic Immunological Labs Ltd, Maidenhead, Berks), which was conjugated to horseradish peroxidase by the method of Wilson and Nakane 1978.

Treatment of sera with dithiothreitol and pepsin. Sera were reduced with an equal volume of 10 mM dithiothreitol in PBS at 37°C for 1 h12 and then assayed by the procedure outlined above.

Sera were pepsin digested by a modification of the method of Faith et al.10 The concentration of pepsin chosen was that which destroyed all IgM RF as detected by our ELISA. Sera were diluted 1:20, incubated with pepsin-acetate buffer 150 μg/ml pepsin in 100 mM acetate buffer (pH 4-4) incubated for 20 h at 37°C and then neutralised with an equal volume of 0-56 M disodium hydrogen phosphate in PBST. Control sera were incubated in 100 mM acetate buffer pH 4-4 for 20 h at 37°C and then neutralised as above. The sera were then assayed by the ELISA with the
lowest dilution being 1:40. IgG RF and IgM RF levels were measured in sera before and after reduction with DTT and before and after pepsin digestion.

**Other RF assays**

*Rose-Waaler test.* The sera were adsorbed by adding 0·1 ml of concentrated washed sheep red blood cells (SRBC) to 0·5 ml of serum and incubated at 4°C overnight. Sensitised SRBC were prepared by adding a 2% (v/v) solution of washed SRBC in saline to an equal volume of rabbit antiserum (Wellcome Reagents Ltd, Beckenham, England) at its sensitising dilution. The inactivated adsorbed sera were titrated against the sensitised cells in microtitre plates: 40 μl of saline was added to all wells. 40 μl of inactivated adsorbed serum was added to the first well, mixed, and 40 μl transferred to the next well. In this way the serum was diluted across the plate. 40 μl of sensitised cells was then added to each well, and the plates incubated at 37°C for 1·5 h. Cells which had settled into a completely negative button were reported as negative, and those sera with a titre of <1:16 were considered negative.

*Latex agglutination.* The test was carried out with the Mercia Broacades latex kit (Mercia Broacades, Weybridge, UK). Titres of <1:20 were considered negative.

*Rate nephelometry.* A rate nephelometric assay for RF was performed with the Beckman Immunochemistry System (Beckman R11C, High Wycombe, UK). The nephelometer measures the changes in light scatter when serum containing IgM RF is added to partially aggregated human IgG. The rate of formation of the complexes is estimated by reference to a precalibrated standard curve, and the value is displayed as international units/ml. Levels of <60 IU/ml were considered negative.

The Beckman ICS was also used to measure IgG, IgA, and IgM levels in all sera and results were expressed as mg/100 ml.

**Statistical analysis.** The results from the four assays were analysed by the Mann-Whitney U test and the effect of pepsin and DTT on IgG RF levels analysed by the Wilcoxon rank sum test. Correlations were determined by calculating Spearman’s rank correlation coefficient.

**Results**

**Comparison of assays for IgM RF**

The IgM RF levels of both normal and pathological sera as measured by each assay are shown in Fig. 1a. The results are shown as log₁₀ titre for the Rose-Waaler, latex agglutination, and ELISA assays and log₁₀ IU/ml for the nephelometer. The levels of IgM RF were significantly raised (p<0.0001) in RA patients’ sera compared with normal sera in all assays. IgM RF levels correlated well between all assays as shown in Table 1. The strongest correlation was found between the latex agglutination assay and the ELISA, though all correlations were ≥0.7000 and were significant at p<0.0001.

Results were considered negative if the titre was <1:16 (Rose-Waaler test) or <1:20 (latex agglutination test). All normal samples were negative for these two tests, while 60% of RA patients’ sera had a titre of ≥1:16 in the Rose-Waaler test and 90% of RA patients had a titre of ≥1:20 in the latex agglutination test. A value of 60 IU/ml was considered to be negative in the nephelometer method, and 98% of the normal sera were below this level; 69% of the RA patients had values above this figure. A value of 1·75 (log₁₀ titre) was chosen for the IgM RF ELISA, which corresponded to the mean +2SD of the normal sera. With this figure 92% of RA patients had elevated levels of RF while 94% of the normal persons were negative.

**Measurement of IgA RF and IgG RF**

Different immunoglobulin classes of RF were detected with the ELISA protocol, and second antibody-enzyme conjugates specific for IgG and IgA were used.

The levels of IgA RF in patients’ serum was measured by a rabbit antihuman α chain-peroxidase conjugate. The levels of IgA RF in RA patients’ serum were significantly raised (p<0.0001) relative to those in normal sera (Fig. 1b) and 88% were above the mean + 2 standard deviations of normal sera (1·211 log₁₀ titre). The majority (94%) of normal sera were below this level. The levels of IgG RF measured in RA patients and control sera with a rabbit antihuman IgG/Fab antibody-peroxidase conjugate are also shown in Fig. 1b. The levels of IgG RF were significantly raised (p<0.0001) in RA patients relative to normal sera. However, although the majority of RA patients sera (85%) were above the mean + 2 standard deviations of normal sera (1·903), and 96% of normal sera had IgG RF levels below this value, owing to the large range of normal values there was considerable overlap with the lower pathological sera values. For a more accurate estimate of IgG RF both normal and RA patient sera were treated with pepsin. After pepsin digestion IgG RF activity in all normal sera was reduced to below a titre of 1:40 and was significantly reduced (p<0.0001) in RA patients sera (Fig. 1c). (The method for pepsin digestion did not allow IgG RF titres to be measured below a value of 1:40.) All values for normal sera were <1:40 after pepsin digestion.
Fig. 1  Estimation of rheumatoid factor in rheumatoid control sera—comparison of assays. (a) The levels of IgM RF in 48 rheumatoid and 48 age and sex matched controls were measured by the Rose-Waaler, latex agglutination, rate nephelometric assays and ELISA as described in the 'Methods' section. (b) The levels of IgA RF and IgG RF were measured in rheumatoid and control sera by the ELISA. (c) The level of IgG RF was measured in rheumatoid and control sera after pepsin digestion. This treatment destroys IgM RF, which can interfere with the measurement of IgG RF. The horizontal bars indicate the mean values for each group.

Table 1  Correlation coefficients of the levels of IgM RF between assays

<table>
<thead>
<tr>
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<th>Latex agglutination</th>
<th>Nephelometry</th>
<th>ELISA</th>
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<tr>
<td>Rose-Waaler</td>
<td>0.8178</td>
<td>0.6998</td>
<td>0.8029</td>
</tr>
<tr>
<td>Latex agglutination</td>
<td>—</td>
<td>0.7635</td>
<td>0.8733</td>
</tr>
<tr>
<td>Nephelometry</td>
<td>—</td>
<td>—</td>
<td>0.8121</td>
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The correlation coefficients of the levels of IgM RF between the four assays were measured by calculating Spearman's rank correlation coefficient. All correlations were significant at p<0.0001.

digestion, while 46% of the RA patients sera were above this value. There was a high degree of correlation between all classes of RF measured by ELISA in RA patients sera (Table 2).

The total immunoglobulin levels for both normal and RA patients' sera were measured and are shown in Fig. 2. The levels of all immunoglobulins were significantly raised in RA patients' sera when compared with the levels in normal sera (IgG and IgM at p<0.0001 and IgA at p<0.0002). Table
Measurement of rheumatoid factors by an enzyme-linked immunosorbent assay

Table 2  Correlation between RFs of different immunoglobulin class in rheumatoid patients' sera

<table>
<thead>
<tr>
<th></th>
<th>IgA RF</th>
<th>IgM RF</th>
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<tr>
<td>IgG RF</td>
<td>0.7701</td>
<td>0.9184</td>
</tr>
<tr>
<td>Pepsin IgG RF</td>
<td>0.7403</td>
<td>0.7224</td>
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<tr>
<td>IgA RF</td>
<td>—</td>
<td>0.7329</td>
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</table>

The correlation coefficients between the levels of IgG, IgA, and IgM RFs in rheumatoid patients' sera were measured by calculating the Spearman's rank correlation coefficient. All correlations were significant at p<0.0001.

Illustrates the correlations obtained between the different classes of RF with the total immunoglobulin content of RA patients' sera. Good correlation was found between the levels of IgG and IgA in RA patients' sera, but correlation between other immunoglobulin classes in RA and normal sera were low.

Table 3  Correlation between immunoglobulin levels in rheumatoid and normal sera

<table>
<thead>
<tr>
<th></th>
<th>IgA</th>
<th>IgM</th>
</tr>
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<tbody>
<tr>
<td>RA patients:</td>
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<tr>
<td>IgG</td>
<td>0.6799</td>
<td>0.4111</td>
</tr>
<tr>
<td>IgA</td>
<td>—</td>
<td>0.3442</td>
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<tr>
<td>Normals:</td>
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<tr>
<td>IgG</td>
<td>0.1880</td>
<td>0.2798</td>
</tr>
<tr>
<td>IgA</td>
<td>—</td>
<td>0.0515</td>
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</table>

The correlation coefficients between the levels of IgG, IgA, and IgM in rheumatoid and normal sera were calculated.

Discussion

At present there is considerable interest in ELISA assays in the clinical laboratory. These assays represent a quick and cost-effective way of processing large numbers of samples while using simple
and relatively inexpensive equipment. A number of ELISA methods have been published for measuring RF, but they have had a restricted routine usage, and the more common Rose-Waaler and latex agglutination assays are still performed in the majority of hospital service laboratories. If the ELISA assay is to gain general acceptance, it must be carefully compared with other methods. In the present study we have compared the two traditional methods with a rate nephelometric assay and an ELISA for rheumatoid factor. There was good correlation for IgM RF values between all assays. Although the Rose-Waaler and latex agglutination tests are relatively easy to perform, the end points of both assays are difficult to determine without experience, and both assays are prone to observer variability. Consistent results can be obtained with care in one laboratory if the same personnel repeat the test from week to week, but interlaboratory variation for the same sera is high. The rate nephelometric assay was the easiest and quickest test to perform and gave quantitative results in a very short time. It is useful for following patients serially over a period of time, and consistent results are obtained from serum samples when assayed at different times. Of the four assays tested the rate nephelometric assay gave the most consistent results. Other assays had to be run with a standard and the titres adjusted up or down for all samples according to the result obtained for the standard. The ELISA was quick and easy to perform and would suit a routine laboratory. It was the most sensitive assay and recognised 92% of rheumatoid patients’ sera as positive. The corresponding figures for the other assays were latex agglutination test 90%, rate nephelometric assay 69%, and Rose-Waaler 60%.

The ELISA was also able to measure IgG and IgA RF in addition to IgM RF. The results were quoted in titres for the ELISA, but it can be adapted by incorporating a standard curve to give quantitative results in µg IgM RF/ml or in international units. We are currently evaluating quantitation of the assay using these methods and interfacing the ELISA reader to a computer to automate the estimations and allow the results to be calculated by reference to a standard curve.

The classical tests for RF usually measure only IgM RF, but in recent years increasing attention has focused on the role of both IgG and IgA RF. IgG RF was first measured quantitatively by Torrigiani and Roitt. Since that time many different assay protocols for the measurement of IgG RF and IgA RF have been published, and the most widely used are radioimmunoassay, immunofluorescent techniques, or ELISA. March et al. have published a simple microtitre plate assay using immunoglobulin coupled to red blood cells.

The significance of the raised levels of IgG RF in rheumatoid disease is not clear. Early reports suggested that IgG RF was significantly raised in both seropositive and seronegative arthropathies. However, it is now thought that these early results were incorrect, and subsequent studies using more sensitive and specific methods of detection have shown that significant levels of IgG RF are not associated with seronegative diseases. RF can interfere with the measurement of IgG RF in solid phase assays, as it can bind to the immobilised target IgG and to IgG in the serum. The bound serum IgG is then detected with the antibody conjugate as IgG RF. Two methods have been used to destroy IgM in the test serum and so prevent these interactions. These are either pepsin digestion or treatment with a reducing agent. In this study we used pepsin digestion and adapted the method of Faith et al. to adjust the level of pepsin so that all serum IgM was destroyed. We found that the results obtained after reduction and alkylation of sera were variable and that higher levels of IgG RF were present in control sera after this treatment. Similar findings have been reported by Carson et al. and Wernick et al. and so pepsin digestion was considered to be the method of choice.

High titres of IgG RF are associated with the more severe forms of rheumatoid disease, and patients with vasculitis or subcutaneous nodules have high levels of serum IgG RF. Immune complexes containing IgG RF have been detected in the serum, synovial fluid, and synovium of patients with rheumatoid arthritis, and the levels of IgG RF are higher in synovial fluid than in the corresponding serum sample. This suggests that locally produced IgG RF may be important in the pathogenesis of the disease maintaining the rheumatoid synovitis through its ability to activate complement by self-associating or in combination with IgM RF. The vasculitic lesions probably result from the deposition of pathogenic immune complexes in the microvasculature of the skin and other target organs, and IgG RF appears to be present in many of these complexes. In addition it is possible that IgG RF may aggregate platelets in vivo and so promote vascular damage.

Relatively little interest has been shown in the finding that the levels of IgA RF are raised in rheumatoid patients, though this finding had been previously reported. Dunne et al. demonstrated that IgA RF was present in not only the serum but also the saliva of rheumatoid patients and patients with Sjögren’s syndrome. This
IgA RF showed a pattern of reaction with human IgG subclasses similar to that of IgM RF from the same patients. Koopman et al. detected significantly raised levels of IgA RF in rheumatoid sera and synovial fluid compared with serum from patients with seronegative disease or normal control serum. In the present study, we obtained similar results, finding significantly raised levels of IgA RF in rheumatoid sera, but their significance in the immunopathogenesis of the disease is unknown.

Rheumatoid factor measurement by the conventional tests of Rose-Waaler and latex agglutination are difficult to quantify. We found that the nephelometric assay is rapid, reproducible, and allows results to be presented as international units/ml. The ELISA for rheumatoid factor is simple to perform and interpret and uses equipment which is present in the majority of routine hospital laboratories. We aim to develop a screening test to distinguish quickly between positive and negative sera and also use a standard curve in each assay so that results are quoted in either international units/ml or μg RF/ml.

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References