INTERACTIONS OF RHEUMATOID FACTOR WITH IMMUNE PRECIPITATE CONTAINING ANTIBODY OF HUMAN ORIGIN

BY

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The term “rheumatoid factor” is applied to the macroglobulin component of serum which is responsible for a group of serological reactions used as diagnostic procedures in rheumatoid arthritis. The rheumatoid factor is able to react with antigen-antibody complexes of diverse origins. It agglutinates sheep red cells sensitized with rabbit amboceptor (Waaler, 1940; Rose, Ragan, Pearce, and Lipman, 1948), Rh-positive red cells coated with selected incomplete anti-D antibodies of human origin (Foz and Batalla, 1956; Grubb, 1956; Waller and Vaughan, 1956), and Brucella abortus sensitized with a strong incomplete anti-Brucella antibody of human origin (Foz and Batalla, 1956).

Rheumatoid factor is adsorbed onto to immune precipitates consisting of various antigens and the corresponding rabbit antibodies (Vaughan, 1956; Edelman, Kunkel, and Franklin, 1958; Vaughan, Ellis, and Marshall, 1958; Corcos, 1960; Mellors, Nowoslawski, Korngold, and Sengson, 1961). Vaughan (1956) observed that absorption with an immune precipitate consisting of diphtheria toxin and human antitoxin did not reduce the activity of one rheumatoid serum in the Waaler-Rose test. Under the experimental conditions, a considerable solubility of the precipitate made the quantitative data uninterpretable. Additional data were not available in the literature concerning reactions of rheumatoid sera with immune precipitates containing antibody of human origin.

Recently, we have had the opportunity of studying a strong human precipitin and its reactions with rheumatoid sera. The present paper describes the results of these experiments. After completion of the experiments, we learned of the independent investigations of Aho, Kirpilä, Wager, and Virkkunen (1961) which confirm and extend our findings.

Materials and Methods

Precipitating Antibody.—The antibody was found by Dr. S. Blix during studies on fibrinolysis. Its properties are described in another paper (Blix, 1961), and only a few data will be given here.

The patient R.K., a 58-year-old male, had a 5-year history of peripheral arterial insufficiency in both legs. Bilateral femoral-popliteal by-pass operations had been performed and were followed by secondary thrombosis with purulent ulcerations. These were treated locally with Varidase* eight times during the first 6 months of 1960, and the antibody was demonstrated in November, 1960.

The present experiments were performed with serum samples obtained on different occasions in December, 1960, and January, 1961, and the antibody activity decreased slowly during this period. After chromatography on DEAE-cellulose (kindly performed by Dr. T. Reinskou), it was demonstrated by Dr. Blix that antibody activity was present in the fractions containing 7S γ-globulin. The patient was of type Gm(a+b+x+). The serum titre was less than 1:5 in the Waaler-Rose test and the F.II latex particle test was negative.

Tests for Rheumatoid Factor Activities

Waaler-Rose Test.—This was made with human Group O Rh-negative red cells (Podliachouk, Eyquem, and Jacqueline, 1958), using the corresponding amboceptor from rabbits (commercial preparation, Institut Pasteur, Paris, France). The sensitizing amboceptor was used in one-quarter of the minimum agglutinating dose.

F.II Latex Particle Test.—This was performed according to Winblad (1960, and personal communication) with the following slight modifications: the suspension was stabilized with human albumin, and 0·15 M

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* Varidase was the commercial preparation of Lederle, New York, N.Y., Lots No. 2201-108A and 2200-982A.
† Akryl plast particles (Latex) were kindly provided by AB Bofors, Nobelkrut, Bofors, Sweden.
SIGMA buffer pH 8.2 was used. To 0.5 ml serum dilution was added 0.5 ml suspension of γ-globulin coated particles; the results were read macroscopically without centrifugation after sedimentation overnight at 37°C.

Agglutination Tests with Red Cells Coated with Incomplete Anti-D.—These were made on slides as previously described (Harboe, 1959, 1960a). Rheumatoid sera containing anti-Gm(a) and anti-Gm(x) (Grubb, 1961a) were selected from the panel of sera used for Gm-typing in this laboratory. Anti-D R.A. was used to coat red cells for determination of Gm(a) and Gm(x) types. Titres of anti-Gm(a) and anti-Gm(x) are given as the titres of sera known to contain these substances, when investigated with red cells coated with anti-D R.A. Anti-D S.V. was used for Gm(b) typing.

Red cells coated with anti-D Mu. and Ri. are known to be agglutinated by nearly all rheumatoid sera showing positive Waaler-Rose and F.II latex-particle tests. Since these anti-D sera may be used diagnostically to the demonstration of rheumatoid factor activity, they are referred to as “diagnostic anti-D” in this paper. The antibodies are described in more detail elsewhere (Harboe, 1960b).

Absorption Procedure.—One ml. serum from the patient R.K. was incubated with 1 ml. of a solution of Varidase containing 10,000 units/ml. for 1 hr at 37°C. and 20 hrs at 4°C. Preliminary experiments indicated that these concentrations corresponded to the equivalence point of the precipitation curve. The precipitate was isolated by centrifugation for 30 min. at 1,800 G, and washed three times in 10 ml. chilled saline. The washed precipitate (containing about 0.12 mg. nitrogen as determined by micro-Kjeldahl analysis) was used to absorb 2 ml. rheumatoid serum diluted 1:5 for 24 hrs at 4°C. The absorption was performed in sealed tubes during continuous slow movement to secure optimal contact between precipitate and serum. After absorption, the precipitate was removed by centrifugation and the supernatant transferred to a second tube for repeated absorption. The procedure was repeated for a different number of times, as indicated in the text. After the final absorption, the supernatant was tested for rheumatoid factor activities as indicated. For control, other portions of the rheumatoid sera diluted 1:5 were treated in exactly the same way, except that no precipitate was added. In most sera, a spontaneous precipitate formed within a few days and was removed during centrifugation. For additional control, a third tube of each diluted rheumatoid serum was frozen down (−25°C.) at the beginning of the absorption procedure. The tubes were thawed at the end of the absorption procedure and tested simultaneously with the other materials. Spontaneous precipitates are known to contain rheumatoid factor (Christian, 1959), but there was no significant difference in activity between the frozen samples and the samples which were kept at 4°C.

Quantitation of Precipitated Protein.—To 0.75 ml. serum R.K. was added 0.75 ml. Varidase containing 20,000 units/ml. This concentration of approximately two times equivalence was chosen on the basis of data from Edelman, Kunkel, and Franklin (1958). The mixture was incubated for 1 hr at 37°C. before the addition of 0.75 U.L. rheumatoid serum A.H. or normal sera showing negative Waaler-Rose and F.II latex-particle tests for control. After additional incubation for 20 hrs at 4°C., the precipitates were spun down by centrifugation for 60 min. at 1,800 G at 4°C. The supernatant was carefully removed and the precipitates washed twice in chilled saline. Finally, the precipitates were dissolved in 4 ml. 30 per cent. urea in 0.2 N NaOH, and the optical density was determined at 280 mμ in a Beckman spectrophotometer, model DU-G2400.

Experiments and Results

Serological Activities before and after Absorption.—The Figure (opposite) shows the results of absorption experiments on a rheumatoid serum (198/LW) containing anti-Gm(a). After one absorption, anti-Gm(a) activity could no longer be demonstrated, nor were cells coated with “diagnostic” anti-D Ri. agglutinated. The titres in the Waaler-Rose and F.II latex-particle tests were also reduced, but to a less degree.

The Figure also shows that another rheumatoid serum (270/AA), which also contained anti-Gm(a), gave similar results.

Serum S.V. contained anti-Gm(x) with a titre of 1:640. After one absorption with the immune precipitate, no anti-Gm(x) activity could be demonstrated.

Two rheumatoid sera with fairly strong serological reactions, which did not agglutinate red cells coated with anti-D R.A., were then investigated. The serum titre was determined after four times absorption with the immune precipitate in the Waaler-Rose test and F.II latex-particle test, and against red cells coated with “diagnostic” anti-D Mu. and Ri., and compared with the unabsorbed control.

Table I (opposite) shows that there was a marked reduction in all serological activities by the absorption procedure.

Quantitative Studies.—The results of one experiment are shown in Table II (opposite). It may be seen that the amount of precipitated protein was greater when
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Figure.—Absorption of rheumatoid serum 198/LW and 270/AA containing anti-Gm(a) with human specific precipitate.

TABLE I

<table>
<thead>
<tr>
<th>Test</th>
<th>Waaler-Rose</th>
<th>F.II-Latex Particle</th>
<th>&quot;Diagnostic&quot; Anti-D Sera</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum</td>
<td>Before</td>
<td>After</td>
<td>Before</td>
</tr>
<tr>
<td></td>
<td>Reciprocals</td>
<td></td>
<td>Reciprocals</td>
</tr>
<tr>
<td>A.H.</td>
<td>320</td>
<td>5</td>
<td>1,280</td>
</tr>
<tr>
<td>O.H.</td>
<td>320</td>
<td>20</td>
<td>2,560</td>
</tr>
</tbody>
</table>

TABLE II

<table>
<thead>
<tr>
<th>Reagents Incubated</th>
<th>Varidase</th>
<th>Varidase</th>
<th>Varidase</th>
<th>Varidase</th>
<th>Varidase</th>
<th>Serum R.K.</th>
<th>Saline</th>
<th>Normal Serum 1</th>
<th>Normal Serum 2</th>
<th>Rheumatoid Serum A.H.</th>
<th>Rheumatoid Serum A.H.</th>
<th>Optical Density (280 μm) of Dissolved Precipitate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td>Varidase</td>
<td>Saline</td>
<td>Saline</td>
<td>Saline</td>
<td>Saline</td>
<td>Saline</td>
<td>Serum R.K.</td>
<td>Saline</td>
<td>Saline</td>
<td>Rheumatoid Serum A.H.</td>
<td>Rheumatoid Serum A.H.</td>
<td>0.120</td>
</tr>
<tr>
<td></td>
<td>0.138</td>
<td>0.200</td>
<td>0.210</td>
<td>0.138</td>
<td>0.200</td>
<td>0.210</td>
<td>0.120</td>
<td>0.120</td>
<td>0.138</td>
<td>0.200</td>
<td>0.210</td>
<td>0.010</td>
</tr>
</tbody>
</table>

For technique of investigation, see text.
the immune precipitate was incubated with rheumatoid serum A.H. than after incubation with control sera showing negative tests for rheumatoid factor or saline. These findings were reproduced on three occasions. The experiment was made with only one rheumatoid serum (A.H.) because, as purified rheumatoid factor was not available, the tests had to be carried out with neat rheumatoid serum. Under these conditions, spontaneous precipitation (Christian, 1959) often makes interpretation difficult or impossible. Serum A.H. was selected because it was the only one, in a series of rheumatoid sera with fairly strong serological reactions, which gave only a trace of spontaneous precipitate after dilution 1:5 in saline and several days’ storage at 4°C. Relevant controls are included in the Table.

Controls

(1) Varidase is a purified preparation of the streptococcal enzymes streptokinase and streptodornase. Streptokinase activates the fibrinolytic system of human plasma and serum resulting in formation of plasmin (fibrinolysin) which is a proteolytic enzyme. In the organism, the activity of plasmin is directed primarily towards fibrin and, to some extent, fibrinogen. In addition, it may digest other proteins (Sherry, Fletcher, and Alkjaersig, 1959), possibly including the macroglobulins responsible for the activity in the present serological tests.

Four rheumatoid sera were incubated with Varidase under conditions optimal for plasmin formation (Blix, 1961). No reduction of serological activity was observed in the sera when compared with similar incubation with saline for control.

(2) Serum R.K. was of type Gm(a+). During the absorption procedure, Gm(a+) γ-globulin might therefore loosen from the immune precipitate in amounts sufficient to inhibit anti-Gm(a). After absorption of serum 198/LW (Figure), the following experiments were made to study this possibility:

To demonstrate the presence of Gm(a+) γ-globulin, we used anti-Gm(a) Kouba and red cells coated with anti-D R.A. In this system, Gm(a+) normal serum showed full inhibition of agglutination in dilutions up to 1:1,280, whereas Gm(a−) normal serum did not inhibit in dilution 1:5. After three times absorption of serum 198/LW with the immune precipitate, dilution 1:5 had a definite inhibiting ability towards serum Kouba. At dilution 1:10, no inhibition was observed. The unabsorbed control of serum 198/LW did not inhibit in dilution 1:5 or higher—the latter experiment was performed after heating of diluted serum 198/LW to abolish its agglutinating ability (Grubb and Laurell, 1956; Harboe, 1960a). The results of the experiments are summarized in Table III.

### Table III
**ATTEMPT TO DEMONSTRATE Gm(a+) GAMMA-GLOBULIN LOOSENED FROM SPECIFIC PRECIPITATE DURING ABSORPTION**

<table>
<thead>
<tr>
<th>Dilution of Test Material</th>
<th>Test Material</th>
</tr>
</thead>
<tbody>
<tr>
<td>198/LW</td>
<td>Gm(a−)</td>
</tr>
<tr>
<td></td>
<td>Before Absorption</td>
</tr>
<tr>
<td>1 : 5</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 10</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 20</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 40</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 80</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 160</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 320</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 640</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 1280</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 2560</td>
<td>+++++</td>
</tr>
<tr>
<td>1 : 5120</td>
<td>+++++</td>
</tr>
</tbody>
</table>

### Controls

(3) Absorption of rheumatoid sera with immune precipitates consisting of various antigens and the corresponding rabbit antibodies may abolish serological activity in the Waaler-Rose test leaving the latex-fixation titres virtually unchanged. Corcos (1960) found that this was the case when human albumin or purified 7S human γ-globulin was used as antigen in the immune precipitate. He also found that immune precipitates containing aggregated human γ-globulin removed the activity in both serological tests, as does absorption with aggregated human γ-globulin alone. An important source of error in the present experiments might therefore be that the antigen itself, Varidase, might react with the rheumatoid factor. Inhibition experiments were made to clarify this point:

Four rheumatoid sera were diluted until they contained ten agglutinating doses in the different tests, and serial two-fold dilutions of a Varidase solution originally containing 10,000 units/ml. were added in order to test for inhibiting capacity. In the Waaler-Rose test, no
significant inhibition was found. In the F.II latex-particle test, slight inhibition was observed in the first two tubes. Anti-Gm(a) Kouba and 198/LW were not inhibited by Varidase (these experiments were performed using red cells coated with anti-D G63, see below). Whether the agglutination of red cells coated with anti-D Ri was inhibited by Varidase, could not be tested because it was found that the preparation agglutinated such cells.

Rh-positive red cells were coated with ten different strong incomplete anti-D sera and tested for agglutination by serial dilutions of a Varidase solution originally containing 10,000 units/ml. Red cells coated with one anti-D serum (G63) were only weakly agglutinated in a narrow concentration range of Varidase, while cells coated with any of the other anti-D antibodies were strongly agglutinated. A “prozone phenomenon” was observed with six of the anti-D antibodies; no agglutination was observed by the highest concentrations of Varidase, whereas lower concentrations of the preparation showed strong agglutinating ability. The basic nature of this agglutination is unknown. It is probably not of enzymic nature, as the activity was somewhat stronger at 4° C than at 37° C. The agglutination was inhibited by low concentrations of both human albumin and γ-globulin.

It was concluded from these experiments that Varidase itself did not inhibit the rheumatoid factor to a significant degree and, accordingly, that such inhibition could not explain the findings of the absorption experiments.

Discussion

The rheumatoid factor is able to react with γ-globulin of diverse origins. Theoretically of great importance is whether it can react with human γ-globulin in vivo, and whether this reaction is of any pathophysiological consequence for the individual.

Experiments in this laboratory (Harboe, 1961) indicate that anti-Gm(a) is a separate component of the complex of closely related macroglobulins usually designated as rheumatoid factors. Anti-Gm(a) and the other specific agglutinating substances of the Gm system are inhibited by native human 7S γ-globulin (Grubb, 1961b). Similar reactions are directly observed in experiments using red cells coated with selected incomplete anti-D antibodies, where prozones often occur in agglutination tests. It has been demonstrated that these prozones are frequently caused by the simultaneous presence of an agglutinating substance and its specific inhibitor in individual rheumatoid sera (Swahn and Grubb, 1958; Harboe, 1960a). Additional evidence for a reaction between rheumatoid factor and human γ-globulin in vivo is the presence of the 22S complex in some rheumatoid sera. Rheumatoid factor is composed of 19S γ-globulins which often exist in the circulation bound to 7S γ-globulin, and this complex has a sedimentation coefficient of 22S (Franklin, Kunkel, Müller-Eberhard, and Holman, 1957).

In the beginning of the experiments described in this paper, the behaviour of rheumatoid factor with immune precipitates containing antibody of human origin was scarcely known. The only data available were those on rheumatoid serum antibody, which was absorbed by Vaughan (1956) with a precipitate consisting of diphtheria toxin and human antitoxin. There was no reduction in activity after absorption as judged by the Waaler-Rose test. The reasons for the discrepancy between Vaughan’s findings and ours are probably that we did repeated absorptions, whereas his serum appears to have been absorbed only once, and that the other tests (demonstration of anti-Gm(a) and agglutination of red cells coated with “diagnostic” anti-D) are more sensitive than the Waaler-Rose test (Figure). The present findings showed that different components of the rheumatoid factor were adsorbed on to the specific precipitate containing human antibody.

After completing the present investigations, we learned of the important paper of Aho, Kirpilä, Wager, and Virkkunen (1961). They immunized two patients with rheumatoid arthritis, who had strongly positive serological reactions, with diphtheria toxoid, and both patients developed strong precipitating antibodies. It was found that rheumatoid factor was adsorbed to, and could be eluted from, the precipitates, which consisted of diphtheria toxoid and the patients’ own antitoxins. Identical findings were made by three different tests for rheumatoid factor activity: the Waaler-Rose test, the F.II latex-particle test, and an agglutination test with red cells coated with anti-D Ri.

Studies on the Gm system have shown that components of the rheumatoid factor may react with the individual’s own native γ-globulin (Swahn and Grubb, 1958; Harboe, 1960a). The present experiments show that the rheumatoid factor is adsorbed on to immune precipitate containing antibody of human origin. The experiments of Aho, Kirpilä, Wager, and Virkkunen (1961) further demonstrate that immune complexes containing precipitating antibody from rheumatoid arthritis patients react with their own rheumatoid factor. It remains to be demonstrated whether these phenomena observed in vitro may indicate autoimmune mechanisms in rheumatoid arthritis.

Summary

The behaviour of rheumatoid factor was studied with an immune precipitate consisting of “Varidase” and a human 7S γ-globulin antibody. Different
components of rheumatoid factor [as defined by the Waaler-Rose test, F.II latex-particle test, agglutination tests with red cells coated with “diagnostic” anti-D, and demonstration of anti-Gm(a) and anti-Gm(x)] were all absorbed on to this immune precipitate.

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REFERENCES


Interactions du facteur rhumatismal avec un immun-précipité contenant des anticorps d’origine humaine

**RÉSUMÉ**

On étudia le comportement du facteur rhumatismal avec un immun-précipité consistant de “Varidase” et d’un anticorps humain, la globuline gamma 7S. De différents composants du facteur rhumatismal—définis par la réaction de Waaler-Rose, la réaction F.II d’agglutination de particules de latex, des réactions d’agglutination de globules rouges enduits d’anti-D “diagnostique” et mise en évidence d’anti-Gm(a) et anti-Gm(x)—furent tous absorbés dans ce immun-précipité.

**Interacciones del factor reumatoide con un inmun-préciptado conteniendo anticuerpos de origen humano**

**SUMARIO**

Se estudió el comportamiento del factor reumatoide con un inmun-préciptado conteniendo “Varidase” y un anticuerpo humano, la globulina gamma 7S. Diferentes compuestos del factor reumatoide—definidos por la reacción de Waaler-Rose, la reacción F.II de aglutinación de particulas de latex, reacciones de aglutinación de eritrocitos cubiertos de anti-D diagnóstico y comprobación de anti-Gm(a) y anti-Gm(x)—fueron todos absorbidos en este inmun-préciptado.
Interactions of Rheumatoid Factor with Immune Precipitate containing Antibody of Human Origin

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