RELATION OF HIGH MOLECULAR WEIGHT PROTEINS TO THE SEROLOGICAL REACTIONS IN RHEUMATOID ARTHRITIS*

BY

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The ability of sera from patients with rheumatoid arthritis to potentiate the agglutination of sensitized particulate bodies has been recognized since Cecil, Nicholls, and Stainsby (1931) noted the agglutination of certain strains of streptococci by these sera. More recently this property has been employed as the basis of a number of clinical tests for the diagnosis of rheumatoid arthritis. The tests most frequently used are the sensitized sheep cell agglutination, Fraction II agglutination, latex fixation, Rh agglutination, and gamma-globulin precipitation tests. The reaction common to all these tests seems to be the interaction of the rheumatoid factor or factors present in serum with gamma-globulin attached to a particulate body. The particulate body probably acts only as a carrier for the gamma-globulin since it has been demonstrated by Epstein, Johnson, and Ragan (1955) that Fraction II gamma-globulin can form a precipitate with rheumatoid sera directly. This is especially true when the gamma-globulin has previously been altered by heating or a number of other procedures (Franklin, Holman, Müller-Eberhard, and Kunkel, 1957).

Normal human gamma-globulin contains two major ultracentrifugal components. The main one has a sedimentation rate of 7S, while the minor one, which normally makes up from 5 to 10 per cent. of the total gamma-globulin, has a sedimentation coefficient of 19S (Müller-Eberhard, Kunkel, and Franklin, 1956). It has been demonstrated by numerous observers, particularly Svartz and Schlossman (1954) and by Ziff, Brown, Lospalluto, Badin, and McEwen (1956), that the rheumatoid factor is a gamma-globulin and that it can be concentrated in the euglobulin fraction of serum. Since the euglobulin fraction is rich in the minor high molecular weight fraction of gamma-globulin, it seemed likely that the rheumatoid factor might be associated with a protein of high molecular weight. An ultracentrifugal study of sera and serum fractions from patients with rheumatoid arthritis was therefore undertaken in an effort further to characterize the rheumatoid factors responsible for the positive serological tests. Each of these was found to be of high molecular weight. In addition, it was possible to detect in the sera of certain of patients with rheumatoid arthritis the presence of an unusual high molecular weight protein complex (Franklin and others, 1957).

Materials and Methods

These are similar to those previously described (Franklin and others, 1957) and will be outlined only briefly.

1) Ultracentrifugation.—Analytical examination of sera and protein fractions was carried out in a Spinco Model E ultracentrifuge. Preparative density gradient ultracentrifugation was performed in a Spinco Model L ultracentrifuge in a swinging bucket rotor through a sucrose gradient. The distribution of gamma-globulins of different sedimentation rates was determined by a quantitative immunologic assay, using antisera against the major 7S gamma-globulin and the minor 19S gamma-globulin fractions; it was further checked by analytical ultracentrifugation of the recovered material.

2) Electrophoresis.—Zone electrophoresis was carried out according to methods previously described, using starch as a supporting medium and barbital buffer pH 8.6, T° 0.1.

3) Serologic Tests.—The gamma-globulin precipitation test was carried out with gamma-globulin heated at 63° for 10 minutes as previously described. The sensitized sheep cell agglutination test was performed according to the method of Heller, Kolodny, Lepow, Jacobson, Rivera, and Marks (1955), the latex fixation test according to Singer and Plotz (1956) and the Rh agglutination test according to Waller and Vaughan (1956).
Results

(1) Presence of a High Molecular Weight Complex in the Serum of Certain Patients with Rheumatoid Arthritis.—Fresh sera from 31 patients with rheumatoid arthritis and 37 sera from normal control subjects and from patients with a variety of diseases many of which were associated with abnormalities in the serum proteins were examined in the analytical ultracentrifuge. In all the sera examined a peak with a sedimentation coefficient of 19S could be observed during the first hour of centrifugation at 52,640 r.p.m. This peak can be seen in normal serum (Fig. 1, opposite). In fourteen of the sera from patients with rheumatoid arthritis this peak divided into two parts. One of these sedimented with the normal 19S component. The other one sedimented more rapidly and has previously been called the 22S component (Franklin and others, 1957). In seven of these fourteen sera this component was present in concentrations between 75 and 350 mg. per cent., while in seven others it was present in amounts too small to be accurately measured and could be noted only as a distinct asymmetry preceding the 19S peak. The plasma was examined in two cases and was also shown to contain the component. Three sera from patients with rheumatoid arthritis with easily detectable amounts of this unusual complex are shown in Fig. 1 (opposite). None of the control sera contained measurable amounts of this high molecular weight material.

(2) Relationship of High Molecular Weight Complex to Serological Tests.—Table I demonstrates that, in general, there was some correlation between the amount of 22S material and the intensity of the precipitation reaction and sheep cell agglutination reaction. Subsequent studies on many additional sera have confirmed this observation and have shown a similar relationship of the high molecular weight fraction to the latex fixation and Rh agglutination reactions. In every case sera that were most highly active by the serological tests contained measurable amounts of 22S material. It appeared that the ultracentrifuge was relatively insensitive and that small amounts of 22S material might be present but remain undetected in sera that were less positive in the serological reactions.

Further direct evidence demonstrating that the biologic activities were associated with a high molecular weight protein was obtained by density gradient high speed centrifugation of whole sera. Table II shows the results obtained with fractions obtained from one patient with rheumatoid arthritis. None of the activity by any of the serological tests was present in the top fractions which contained the bulk of the 7S gamma-globulin. (The albumin coloured with bromphenol blue was localized in Fraction II.) All the activity by each of the four tests was found near the bottom of the tubes in those fractions shown to contain all of the 19S gamma-globulin as determined by a specific antiserum and less than 10 per cent. of the total gamma-globulin.

Table I

<table>
<thead>
<tr>
<th>22S Material (mg. per cent.)</th>
<th>Gamma-Globulin Precipitation</th>
<th>Sensitized Sheep Cell Agglutination (Dilution)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>1+ 2+ 3+ 4+</td>
<td>128 256 512 1,024</td>
</tr>
<tr>
<td>&lt;75</td>
<td>6 8 2</td>
<td>3 1 1 1</td>
</tr>
<tr>
<td>75–350</td>
<td>3 4</td>
<td>1 2 2 2</td>
</tr>
</tbody>
</table>

Table II

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Reaction with 7S As.</th>
<th>Reaction with 19S As.</th>
<th>Sensitized Sheep Cell Agglutination</th>
<th>Gamma-Globulin Precipitation Test</th>
<th>Anti-Rh Agglutination</th>
<th>Latex Fixation</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>_</td>
<td>_</td>
</tr>
<tr>
<td>II Albumin</td>
<td>2+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>_</td>
<td>_</td>
</tr>
<tr>
<td>III</td>
<td>4+</td>
<td>0</td>
<td>1/64</td>
<td>2+</td>
<td>1+</td>
<td>_</td>
</tr>
<tr>
<td>IV</td>
<td>1+</td>
<td>1+</td>
<td>1/164</td>
<td>4+</td>
<td>4+</td>
<td>1/1,024</td>
</tr>
<tr>
<td>V Bottom</td>
<td>1+</td>
<td>4+</td>
<td>1/512</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Numerous experiments of this type with both high and low titre sera always gave similar results with the serological activities in the bottom fractions. These experiments, while demonstrating an association of biologic activities with a high molecular weight protein, could not differentiate the 19S and 22S types.

The role of the 22S component in the gamma-globulin precipitation reaction in particular and in the other serologic tests as well was further demonstrated by repeated absorption of positive sera with altered gamma-globulin. After several absorptions there was a great decrease in 22S material in the supernatant with a concomitant loss of biologic activity.
(3) Characterization of 22S Complex.—Ultra-
centrifugal examination of various electro-
phoretically isolated protein fractions from sera of
patients with rheumatoid arthritis showed the high
molecular weight component to be associated only
with the gamma-globulin fraction. Fig. 2 shows it
to be distributed throughout the gamma-globulin
fraction with a mobility slightly greater than that of
the peak of the gamma-globulin. Similarly, the
mobility of the four tests corresponded quite closely
with that of the high molecular weight component.
None of the other protein fractions contained any
of the high molecular weight complex nor were they
highly active in any of the serological tests.

Although evidence of heterogeneity in the ultra-
centrifuge suggested the presence of more than one
molecular species in a number of instances, it was
possible to determine an approximate sedimentation
coefficient of the complex. The $s_{20,w}$ was calculated
to be 22·7S. Since it seemed possible that the 22S
component might represent a circulating complex
composed of 19S gamma-globulin and another
protein, attempts were made to dissociate it and to
study the constituent parts. The 22S material
proved to be readily dissociable by 4-6M urea, acid
at pH 3, and various other procedures. Although
the initial observations were carried out on electro-
phoretically isolated gamma-globulin fractions, it
soon became evident that, for purposes of accurate
quantitation of the composition of the 22S material,
the euglobulin fraction prepared by dilution of the
serum with 10 volumes of deionized distilled water
was superior. By this simple procedure it was
possible to achieve considerable purification of the
high molecular weight material.

Fig. 3 shows the appearance of the euglobulin
fraction prepared from equal aliquots of a highly
active rheumatoid serum and subsequently dissolved
in equal volumes of saline, 30 per cent. urea, and
glycine buffer pH 3. It can be seen that, whereas
the original euglobulin was composed of approxi-
ately equal amounts of 7S, 19S, and 22S material,
the urea and acid have resulted in complete dis-
appearance of the 22S peak with a concomitant
increase in material sedimenting with the 7S and 19S
fractions. Numerous observations of this type
to be reported separately indicated that the 22S
material was a complex composed of approximately
50 per cent. 7S and 50 per cent. 19S material.
Removal of the dissociating agent, especially if urea

![Image](http://ard.bmj.com/)

**Fig. 2.**—Electrophoretic distribution of the 22S component (above)
and the activity in the sensitized sheep cell agglutination and gamma-
globulin precipitation tests (below) compared with the protein
distribution in the gamma- and beta-globulin fraction of a serum
from a patient with rheumatoid arthritis.

**Fig. 3.**—Ultracentrifugal patterns of the euglobulin fractions prepared
from the serum of a patient with rheumatoid arthritis dissolved in
5 per cent. saline, pH 3, urea, and mercapto-ethanol. The
dissociation of the 22S peak in acid and urea is accompanied by an
increase in the amount of 7S and 19S material. The mercapto-
ethanol caused a loss of both 19S and 22S material.
was used, has yielded highly active 19S fractions but has not yet resulted in reaggregation of the 22S complex.

Dissociation of the 22S complex also sometimes occurred during preparative purification of the euglobulin fractions by density gradient centrifugation. The bottom fractions obtained by this procedure usually contained more than 90 per cent. 19S material and no detectable 7S and 22S molecules. These preparations were extremely active by the four serological tests used. Similarly it was possible to obtain active 19S material by gradient centrifugation of acid and urea-dissociated euglobulin fractions, particularly if urea was used. However, here too it was not possible to reform the 22S complex by the addition of gamma-globulin, even if the 7S material that was split off during dissociation was used.

**Dissociation with Sulphhydryl Compounds**

Recent observations (Ryle and Sanger, 1955; Deutsch and Morton, 1957) that many proteins consist of polypeptide chains linked by disulphide bridges suggested the possibility that the 22S component and its active 19S derivative might owe their large size to such a linkage of smaller units. Mercapto-ethanol at a concentration of 0·1 M and cysteine at a concentration of 0·1 M caused complete dissociation of both the 22S and the 19S components from euglobulin preparations. The result of one such experiment is shown in Fig. 3. All 19S and 22S material disappeared and there was an increase in smaller proteins mostly of the 7S class. With this dissociation there was a complete loss of serological activity both in the gamma-globulin precipitation test and in the sheep cell agglutination reaction.

**Analyses of the Fraction II - Rheumatoid Factor Precipitates**

Similar studies were carried out on the precipitates obtained by the addition of Fraction II gamma-globulin to rheumatoid sera. This precipitate, after washing, could be dissociated with urea and acid buffers to give soluble protein solutions. Examination in the ultracentrifuge showed that the precipitates were composed of 7S and 19S material in approximately equal amounts. Fig. 4 illustrates the ultracentrifuge pattern of two such dissolved precipitates. A similar composition was obtained when either urea or acid buffers were employed. Removal of the 7S material by density gradient centrifugation yielded 19S fractions soluble in isotonic saline which were highly active in each of the four serological tests. Of particular interest in these experiments was the finding that the material obtained by dissociation of the Fraction II - rheumatoid factor precipitate contained high agglutinating activity for sensitized sheep cells.

![30 per cent. urea](http://ard.bmj.com/)

Fig. 4.—Ultracentrifugal patterns of two Fraction II—rheumatoid factor precipitates, one dissolved in acid and the other in urea, showing the presence of approximately equal amounts of 7S and 19S material.

**Discussion**

It is well known that sera from many patients with rheumatoid arthritis contain a factor or factors which potentiate certain agglutination reactions. This property has been used as the basis of a number of serological tests characteristic of this disease. The present study has demonstrated the presence of an unusual high molecular weight protein complex in the sera of certain patients with rheumatoid arthritis who are strongly positive by the serological tests. This material is associated with the gamma-globulin fraction, has a sedimentation rate of approximately 22S, and appears to be an easily dissociated complex of 7S and 19S gamma-globulin fractions. Some evidence was obtained that the factors responsible for each of the serological tests were associated with the high molecular weight protein complex. This evidence was best for the gamma-globulin precipitation test. Occasional exceptions were encountered with the sheep cell reaction and further work is required to prove an absolute relationship.

The exact incidence of the 22S component in sera from patients with rheumatoid arthritis is difficult to determine because of the insensitivity of the ultracentrifuge. It can be seen by direct ultracentrifugal examination as a well-defined peak or as a broad inhomogeneity in about one-third of the patients. However, since it is detectable only in those sera which are strongly positive by the much more sensitive serological tests, it seems probable that this material may be present in amounts too small to be detected in a greater percentage of the cases. This
is in line with the known higher incidence of positive results by the serological tests.

The observation that the 22S material is readily dissociated into two components with sedimentation rates of 7S and 19S indicates that it is a complex of 7S and 19S material which exists in a soluble state in the sera of certain patients with rheumatoid arthritis. While the serological titres observed in whole serum seemed, in general, to parallel the amount of 22S material present, it has not been ruled out that some activity could reside in the 19S fraction of fresh serum. It has been readily possible to obtain highly active 19S material both by dissociation of the 22S complex and by centrifugal isolation from dissolved precipitates formed by the interaction of Fraction II gamma-globulin and rheumatoid factor. This would suggest that the true rheumatoid factor is a 19S gamma-globulin, but that it exists in vivo primarily as a circulating complex. Since 19S gamma-globulin is known to contain a number of antibodies (Franklin and Kunkel, 1957), it is possible that the 19S gamma-globulin active in the rheumatoid tests is also an antibody. The nature of the 7S portion of the complex remains to be determined. The possibility that the 22S material may be a circulating antigen-antibody complex appears worthy of further investigation.

Summary

An unusual high molecular weight protein complex was detected in the sera of certain patients with rheumatoid arthritis by direct ultracentrifugal examination. This material migrated as a gamma-globulin electrophoretically, had a sedimentation rate of approximately 22S and could be dissociated into two fractions with sedimentation coefficients of 7S and 19S.

Evidence was obtained of a relationship between the 22S complex and four of the commonly-used serological tests. In whole sera the titles generally paralleled the amount of the 22S component, although some exceptions were encountered. Density gradient centrifugation showed that each of these serological activities sedimented with fractions of high molecular weight. Dissociation of the 22S material followed by density gradient ultracentrifugation resulted in 19S material which was highly active in the serological tests.

Further dissociation could be accomplished with mercapto-ethanol and cysteine which split disulphide linkages. This resulted in a loss of serological activity.

Analyses of the precipitate formed on the addition of Fraction II gamma-globulin to rheumatoid sera indicated that it consisted of approximately 50 per cent. 19S material. The latter isolated by centrifugation was active in the sheep cell agglutination test as well as in the other rheumatoid tests.

The accumulated evidence suggests that the rheumatoid factors are proteins closely related to other 19S gamma-globulins and to the 19S antibodies, and that they exist in serum and plasma complexed to another type of protein.

REFERENCES


HIGH MOLECULAR WEIGHT PROTEINS IN RHEUMATOID ARTHRITIS

Relación de proteínas de peso molecular alto con las reacciones serológicas en la artritis reumatoide

SUMARIO
Un complejo de proteínas de peso molecular insólita-mente alto fue encontrado en los sueros de ciertos enfermos con artritis reumatoide por examen directo a la ultracentrifugadora. Esta substancia se movía electroforéticamente como una gama-globulina, tenía una velocidad de sedimentación de cerca de 22S y podía disociarse en dos fracciones con coeficientes de sedi-
mentación de 7S y de 19S.

Los datos obtenidos muestran una relación entre el complejo 22S y cuatro de las reacciones serológicas comúnmente empleadas. En los sueros completos las cifras fueron generalmente paralelas a la cantidad del componente 22S, pero se notaron algunas excepciones. La centrifugación por escala de densidad mostró que cada una de estas actividades serológicas sedimentaba con fracciones de peso molecular alto. La disociación del producto 22S, seguida de ultracentri-
fugación por escala de densidad, resultaba en el producto 19S, que era muy activo en las reacciones serológicas.

Se podía obtener una disociación más pronunciada con la ayuda de mercaptoetanol y de cistina, que rompían el lazo disulfúrico. Esto resultaba en la pérdida de actividad serológica.

El análisis de precipitado formado al añadir la fracción II de la gama-globulina a los sueros reumáticos indicó que éste consistía aproximadamente de un 50 por ciento del producto 19S. Este último, aislado por centri-
fugación, fue activo en la aglutinación de los glóbulos de oveja así como en otros tests reumáticos.

Los datos recogidos sugieren que los factores reuma-
ticos son proteínas estrechamente relacionadas a otras gama-globulinas 19S y a anticuerpos 19S que existen en el suero y en el plasma en forma de complejo con otro tipo de proteína.
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