OBSERVATIONS ON THE ULTRACENTRIFUGAL PATTERN OF SERUM PROTEINS IN RHEUMATOID ARTHRITIS

BY

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Recent evidence has demonstrated that the serum protein alterations commonly present in rheumatoid arthritis, though following the general pattern of chronic infectious diseases (that is hypoalbuminaemia, hyperalpha- and hypergamma-globulinaemia), show some distinctive features that may be specific for this disease. The most evident among these features is the presence of globulins with a high sedimentation constant (S_{w, 20} \approx 22) (Franklin, Holman, Müller-Eberhard, and Kunkel, 1957; Kuhn, 1959); it is believed that the 22S fraction derives from the association of a globulin of the normal 7S class with a protein macromolecule (18S) that is responsible for the serological reactions termed collectively the rheumatoid factor (Franklin and Kunkel, 1958; Kunkel, Franklin, and Müller-Eberhard, 1959; Vaughan, 1959). A close relation between the level of this component and the severity of the clinical picture has been observed (Franklin, Kunkel, and Ward, 1958; Kuhn, 1959). The exact significance of this fraction in the pathology of rheumatoid arthritis is still unknown; it has been suggested that it may play a role in determining vascular and perivascular lesions that are an essential feature of the disease (Christian, 1958; Epstein and Engleman, 1959); furthermore, its precipitation in tissues, as a consequence of an immunological reaction, may have a pathogenic role (Franklin, Kunkel, and Ward, 1958; Kunkel, 1959; Vaughan and Orbison, 1959). However, these hypotheses are far from being substantiated by direct evidence and a more detailed knowledge of the protein pattern in this disease seems to be necessary before an interpretation is attempted.

Data obtained in our Department (Salteri, Cirla, and Fasoli, 1960) in patients with acute exacerbations of rheumatoid arthritis corresponding to the picture of the “aggravation syndrome” that often develops after prolonged steroid treatment (Villa, Ballabio, and Sala, 1953; Ballabio and Grampa, 1960), may contribute to the characterization of the protein pattern of this disease.

Material and Methods

Fifteen patients with the clinical picture of rheumatoid arthritis according to current criteria have been studied: eleven showed the usual signs of the disease in its fully developed form and four presented the symptoms of the “aggravation syndrome”. Serum samples, obtained in the post-absorptive state, were diluted with normal saline to a protein content of 1-3 g. per cent. and analysed in the Spinco Model E Ultracentrifuge, at room temperature (about 20° C.), at 52,640 r.p.m. Photographs were taken at 16-min. intervals after the rotor reached full speed. The pictures were enlarged and the areas of the various peaks determined with a planimeter. The S values were calculated according to the usual formula, by extrapolating to zero concentration the values observed for the same samples at various dilutions.
Results

(A) Typical Rheumatoid Arthritis (Fig. 1)

The ultracentrifugal analysis of serum proteins does not show in most cases specific abnormalities: there is a moderate increase of the 7S fraction, and an inconstant tendency towards a diminution of the 4·5S fraction, while the 19S component is well within the normal range of variation. This picture corresponds, as a whole, to that of other chronic inflammatory diseases.

However, in two cases in this group another peak was present, close to the descending branch of the 7S peak, that corresponded to an S value of 9-12 and to a concentration of 0·3-0·5 g. per cent. (Fig. 2, opposite). This component is not evident with this technique (or is present only in trivial amounts) in the rest of the sera of the group, and in the conditions involving hyperglobulinaemia that have been studied so far, with the exception of some cases of plasmocytoma.

(B) Rheumatoid Arthritis with "Aggravation Syndrome" (Fig. 1)

The four cases in this group show a lower level of the 4·5S fraction than the preceding series, and no clear-cut differences exist with regard to the 7S and 19S components. All show relatively large amounts of the 9-12S fraction (0·2-0·5 g. per cent.), and three also have a 22S component of 0·1-0·15 g. per cent. (Fig. 3, opposite).

In three cases the analysis was repeated 1 to 4 months later, after a period of treatment with high doses of ACTH and with prednisone at a lower dose level than before admission. Two patients experienced a marked clinical improvement and, at the same time, showed an amelioration of the protein pattern—an increase of the 4·5S fraction and a decrease of the 7S and 9-12S fractions; the 22S component did not change to a significant degree. The third patient did not improve either clinically or with regard to the serum protein picture (Table).

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Protein Fractions (g. per cent.)</th>
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<tbody>
<tr>
<td></td>
<td>4·5S</td>
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<tr>
<td>1</td>
<td>Before</td>
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<tr>
<td></td>
<td>After</td>
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<tr>
<td>2</td>
<td>Before</td>
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Discussion

Our data correspond in general to the results obtained by other authors: however, they differ in showing the occasional presence of a 9-12S
fraction in patients with typical rheumatoid arthritis and in all cases of our series with the "aggravation syndrome". A fraction with similar sedimentation characteristics has been observed in trace amounts by Wallenius, Trautman, Kunkel, and Franklin (1957) in their ultracentrifugal studies of the electrophoretic components of normal human serum; it was seen that both the alpha₂ and beta globulins contained a small amount of material with an S value of 12. Christian (1959) has presented data suggesting that cold precipitates from rheumatoid arthritic sera may contain a component with an S value between 7 and 19.

Our data show that the level of the 9-12S component may be related to the severity of the rheumatoid arthritis symptoms, particularly in cases with the "aggravation syndrome".

In three of our cases a 22S fraction was present; our data are not sufficient to rule out a possible relation between the level of this fraction and that of the 9-12S component; however, they suggest that this is not the case, though both changes are often present at the same time.

It is noteworthy, on the whole, that rheumatoid arthritis is associated with serum protein changes that are rarely, if ever, present in other related or unrelated conditions; the presence in plasmocytoma or macroglobulinaemia of fractions having a sedimentation constant of the same order of magnitude as that of the two above-mentioned components observed in rheumatoid arthritis (9-12S and 22S) does not imply the chemical and/or biological identity of the fractions showing the same S values.

The tendency of rheumatoid arthritic patients to build up protein components that differ in sedimentation rate (and therefore in molecular weight) from the normal may indicate either a derangement of serum protein synthesis or the interaction of normal protein with other substances or with one another so that a larger particle is formed. Some data suggest that the latter mechanism is possibly the one responsible.

No data are available on the origin of the 9-12S component of rheumatoid arthritis; the possibility that it may derive from the interaction of normal components cannot be ruled out. The fact that a fraction with similar characteristics has been shown to be present, in minimal amount, in normal sera would suggest that it is related to this normal component. Obviously new evidence is necessary to clarify this question.
Summary

Ultracentrifugal analysis of serum proteins was performed in fifteen patients with rheumatoid arthritis. In eleven cases the disease had a typical course, and in four who had been treated with long-term steroid therapy the clinical picture of the so-called "aggravation syndrome" was present. In the former group the serum protein pattern showed a moderate rise of the 7S fraction as the most constant abnormality; in two cases a peak in the 9-12S region was observed. In the latter group (those with the "aggravation syndrome") a consistent diminution of the 4-5S fraction was recorded in addition to the rise of the 7S fraction. The 9-12S peak was evident in all cases, of the "aggravation syndrome" and in three of them a 22S component was also present.

The amelioration of the clinical picture was associated with a rise of the 4-5S fraction and a diminution of the 7S and 9-12S components.

The possible meaning of these findings for the pathological physiology of rheumatoid arthritis is tentatively discussed.

REFERENCES


Observations on l'image ultracentrifugue des proteines du serum dans l'arthrite rhumatismale

RÉSUMÉ

On analyse par centrifugation les protéines sériques de 15 malades atteints d'arthrite rhumatismale. Dans 11 cas la maladie présentait une évolution typique, mais chez 4 malades, traités pendant un temps prolongé par des stéroïdes, il y avait un tableau clinique nommé "syndrome d'aggravation". Dans le premier groupe on nota une élévation modérée de la fraction 7S comme la plus constante anomalie; dans deux cas on observa un sommet dans la région 9-12S. Dans le deuxième groupe (celui avec la "syndrome d'aggravation") il y eut une diminution régulière de la fraction 4-5S, en même temps qu'une augmentation de la fraction 7S. Le sommet 9-12S fut évident dans tous les cas de "syndrome d'aggravation" et dans trois d'entre eux il y eut aussi un composant 22S.

Une amélioration du tableau clinique fut associée à une élévation de la fraction 4-5S et une baisse des composants 7S et 9-12S.

On discute tentativement la possible importance de ces données dans la physiopathologie de l'arthrite rhumatismale.

Observaciones sobre el proteinograma sérico por ultracentrifugación en la artritis reumatoide

SUMARIO

Se realizó un análisis por ultracentrifugación de las proteínas séricas de 15 enfermos con artritis reumatoide. En once casos la enfermedad presentaba un curso típico, y en cuatro enfermos, tratados durante un largo tiempo con esteroides, estaba presente un cuadro clínico del llamado "síndrome de agravación". En el primer grupo se notó una moderada elevación de la fracción 7S como la anormalidad más constante; en dos casos se observó una cima en la zona 9-12S. En el segundo grupo (aquel con "síndrome de agravación") se presentó una coherente disminución de la fracción 4-5S, al mismo tiempo que la elevación de la fracción 7S. La cima 9-12S fue evidente en todos los casos con "síndrome de agravación" y en tres de ellos un componente 22S estuvo presente también.

La mejoría del cuadro clínico se vio asociada con una elevación de la fracción 4-5S y una disminución de los componentes 7S y 9-12S.

Se discute tentativamente el posible significado de estos hallazgos en la fisiopatología de la artritis reumatoide.
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