MIF production of lymphocytes from patients with rheumatoid arthritis with antigen–antibody complexes

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Eibl, M. M., and Sitko, C. (1975). Annals of the Rheumatic Diseases, 34, 117. MIF production of lymphocytes from patients with rheumatoid arthritis with antigen–antibody complexes. Immune complexes, human erythrocytes coated with human IgG antibody, inhibit the migration of lymphocytes from rheumatoid arthritis (RA) patients. No correlation could be observed between migration inhibition and Waaler-Rose titre (sheep cell agglutination titre) in individual patients. Production of migration inhibitory factor (MIF) could be detected in 16 of 20 cultures of RA lymphocytes incubated with antigen–antibody complexes, when the supernates were tested with guinea pig macrophages. Only two supernates of healthy persons showed migration inhibition activity under these conditions. The results suggest that migration inhibition by antigen antibody complexes in RA represents a true reaction of delayed hypersensitivity.

Lymphocytes of patients with rheumatoid arthritis (RA) react with aggregated homologous IgG as shown by us (Eibl, 1971) and others (Fröland and Gaarder, 1971; Brostoff, Howell, and Roitt, 1973), as well as with antigen–antibody complexes (Eibl, 1972). Isolated mononuclear cells show migration inhibition when tested with autologous or homologous gammaglobulin as antigen (Eibl, 1972). The same phenomenon has been shown with buffy coat cells by Fröland and Gaarder and was recently repeated by Brostoff et al. (1973). Migration inhibition could also be observed using blood group O D-positive erythrocytes and human IgG anti-D, as well as complexes consisting of measles antigen and human IgG antibody in slight antibody excess (Eibl, 1972). Additional indication for the reaction of RA lymphocytes with antigen–antibody complexes has been provided by Bach, Delrieu, and Delbarre (1970) who showed rosette formation of sheep erythrocytes coated with rabbit IgG antibody with lymphocytes of RA patients. We were able to show a similar reaction in a homologous system (Eibl, Sitko, and Thumb, 1973).

As for the migration inhibition observed by aggregated gammaglobulin and antigen–antibody complexes with cells of RA patients, the question arises whether this phenomenon is due to a T-cell effect or is caused by bridging, whereby the antigens react with an antiglobulin site on the surface of lymphocytes producing rheumatoid factors and would cause inhibition by cross-linking the cells. The experiments reported here were designed to analyse whether the lymphocytes of RA patients, showing migration inhibition by antigen–antibody complexes, can be stimulated in culture to produce migration inhibitory factor (MIF) by similar antigen–antibody complexes under experimental conditions where stimulation of cells derived from healthy persons do not occur.

Patients, materials, methods

Twenty patients with established RA were studied. Diagnosis was made according to the criteria of Ropes, Bennett, Cobb, Jacox, and Jesser (1959). Fourteen of the patients tested had recently been referred to hospital because of active disease. Six were outpatients with quiescent disease. The age of the patients, two males and eighteen females, varied between 40 and 71 yrs with an average age of 57. Steroid therapy did not exceed 10 mg prednisone per day. Twenty healthy blood donors, not matched for age and sex, served as controls. The age of the controls, ten males and ten females, ranged from 23 to 52 years with an average age of 44 years.

LYMPHOCYTES

Mononuclear cells were isolated from heparinized whole blood by a modification of the method described by Thorsby and Bratlie (1970) and used by Aiuti, Lacava, Garofalo, D’Amielto, and D’Asero (1973). 40 ml of venous blood were collected with phenol-free heparin. 4 ml per tube was layered gently over 2 ml of a solution composed of 4.78 g Ficoll (Pharmacia), 10 ml

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Ronpacon (Cilag-Chemie), and 65·2 ml aqua bidest. The tubes were centrifuged at 600 g for 5 min. The leucocyte-rich layer was aspirated, washed in TCM 199, containing 2% human albumin and adjusted to contain 5 × 10^6 cells per ml. The isolated cell suspension contained approximately 90-95% mononuclear cells, of which 6-10% were monocytes as judged by latex phagocytosis.

**ERYTHROCYTES**

Human D-positive erythrocytes of blood group O were used.

**ANTI-D ANTIBODIES**

An anti-D IgG preparation with an antiglobulin titre of 1:2000 was used in a dilution 1:100 to sensitize erythrocytes.

**ANTIGEN-ANTIBODY COMPLEXES**

*Sensitization of erythrocytes for MIF in the direct assay*

The IgG preparation with an anti-D titre of 1:2000 was diluted 1:100. To 2 ml of 1% suspension of D-positive erythrocytes 2 ml of the diluted antibody were added, and red cells were sensitized for 1 hour at 37°C. Sensitized cells were washed twice and resuspended in MEM to contain 5 × 10^8 cells per ml. To each tube of 5 × 10^6 lymphocytes 0·1 ml sensitized erythrocytes were added.

*Sensitization of erythrocytes used for stimulation of MIF production*

20 ml of a 5% suspension of human erythrocytes (D-positive) were incubated for one hour at 37°C with 4 ml of undiluted anti-D antibody. Erythrocytes were washed twice after sensitization and resuspended in 2 ml MEM. To each lymphocyte culture 0·2 ml of sensitized erythrocytes were added.

**MIGRATION INHIBITION OF MONONUCLEAR CELLS**

The method of Falk, Thorsby, Möller, and Möller (1970) was slightly modified. 5 × 10^6 mononuclear cells and 5 × 10^7 sensitized erythrocytes were incubated for 1½ hours at 37°C. Nonsensitized erythrocytes were added to the controls. After incubation, the cell mixture was centrifuged at 500 g for 5 min. The pellet was resuspended in the minimum possible amount of medium and the packed cells were filled in siliconized capillary tubes 1 mm in width, heat-sealed on one side. The capillaries were centrifuged at 500 g for 5 min, cut just below the cell fluid interphase, and fixed with sterile silicone grease on the bottom of a Syke-Moore tissue culture chamber. Chambers were filled with MEM containing Pen. Strep. and 10% fetal calf serum. The migration took place at 37°C for 24 h. Migration areas were magnified by standard technique, cut out of pergament, and weighed on a precision balance. Migration indices have been calculated in the following manner:

**Migration index:**

1. Migration area with sensitized erythrocytes
2. Migration area with nonsensitized erythrocytes.

**MIF PRODUCTION**

Lymphocytes isolated by the Ficoll-isopaque technique were used. A suspension of 2 × 10^7 lymphocytes/ml was prepared in MEM with Pen. Strep. and 5% human serum albumin added. 1·5 ml of lymphocytes were placed in a Leighton type tube (with bottom of 1·5 cm^2). 0·2 ml of sensitized erythrocytes were added to the experimental tubes while the same amount of nonsensitized erythrocytes was added to the controls. Final volume in every tube was 1·7 ml. The tubes were centrifuged for 24 hours at 37°C in a CO_2_ incubator. After 24 h. the tubes were centrifuged for 3 min at 2500 g. The supernates (1·2 ml each) were collected in individual tubes and tested on guinea pig macrophages for migration inhibitory activity. Controls were treated the same way, set up and tested on the same day. At least one patient and one control were tested with any one guinea pig macrophage suspension and also MIF was obtained in duplicate experiments and tested on two different macrophage preparations.

**MIF TESTING**

Peritoneal macrophages of guinea pigs were obtained 5 days after injection of 5 ml sterile paraffin oil into the peritoneal cavity of healthy outbred guinea pigs. The peritoneal cavity was rinsed with TCM 199 containing Pen. Strep. heparin and 2% human serum albumin. 5 × 10^6 macrophages were added to 0·5 ml MIF containing supernates and 0·1 ml fetal calf serum. After an incubation time of 90 min at 37°C the cells were centrifuged at 500 g for 5 min. The pellet was resuspended, put into siliconized capillary tubes, centrifuged at 500 g for 5 min, and cut at the cell fluid interphase. Two capillaries were fixed with silicone grease at the bottom of a Syke-Moore tissue culture chamber. The chamber was filled with MEM, supplemented by 20% fetal serum. 0·5 ml MIF containing supernates were added to every chamber. Incubation was carried out for 24 h. Migration areas were magnified and weighed. The effect of MIF was calculated as follows:

**Migration index:**

1. Migration areas of PGM with supernates of cultures with Ag–Ab–C
2. Migration area of PGM with supernates of cultures with Ag.

Waalser rose titres were measured as described by Steffen (1968).

**Results**

**MIGRATION OF MONONUCLEAR CELLS FROM RA PATIENTS AND HEALTHY PERSONS IN THE PRESENCE OF ANTIGEN-ANTIBODY COMPLEXES**

With a cell ratio of 5 × 10^7 sensitized erythrocytes to 5 × 10^6 lymphocytes significant migration inhibition could be shown in eighteen of the RA patients tested. Results are shown in the Table. Migration indices have been calculated as described above in Methods using the migration of lymphocytes in the presence of nonsensitized erythrocytes as controls. Migration indices below 0·8 were considered to represent significant inhibition. Mean value of migration indices was 0·45 with a standard deviation of 0·23, when lymphocytes of RA patients were...
investigated. Only two of the eighteen positive RA patients showed moderate inhibition of lymphocyte migration by antigen–antibody complexes with a migration index of between 0·6 and 0·8. In 16 cases there was a marked inhibition with migration indices between 0·1 and 0·6.

Lymphocytes of healthy persons did not show migration inhibition by antigen–antibody complexes in 19 of the 20 cases tested. Only one of the control lymphocytes showed moderate migration inhibition with a migration index of 0·72. Migration indices of the 19 control preparations varied between 0·81 and 1·2, with a mean value of 0·99 and a standard deviation of 0·127.

The differences of standard deviation in patients and controls gave significant results in comparison of variances by means of an F test (P < 0·01). For this reason the significance of the difference of migration indices between the RA and control groups had to be calculated by the test for different variances introduced by Welch with approximate T-value (Sachs, 1969). The difference of migration indices between controls and RA patients was highly significant (P < 0·001).

Results of Waaler-Rose tests and migration inhibition were correlated. No correlation could be observed between these two results (Fig. 1). Two patients showing no migration inhibition had sheep cell agglutinin titres of 1:512 and 1:128, respectively. Four of the patients tested had seronegative RA; migration indices in these patients were 0·79, 0·60, 0·42, and 0·20. On the other hand, four patients having Waaler-Rose titres of 1:512 and above showed migration indices of 0·98, 0·42, 0·42, and 0·37.

To test MIF production of lymphocytes in the presence of antigen–antibody complexes, we used a system whereby either nonsensitized erythrocytes or sensitized erythrocytes were added to lymphocytes, and supernates of these cultures were investigated. In experimental cultures, the erythrocytes added in the same quantity have been previously sensitized with anti-D antibody. Supernates of such cultures will be described as MIF-containing supernates.

Since we were interested in studying the effect of antigen–antibody complexes, we wanted to have as few factors as possible influencing the test system. For this reason, we chose lymphocytes incubated with erythrocytes alone as the most suitable controls. Supernates of these cultures are referred to as control supernates. Migration indices have been calculated

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<th>Migration index of RA lymphocytes</th>
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Mean migration index of the RA patients 0·45; standard deviation 0·23. Mean migration index of the controls 0·99; standard deviation 0·13. The difference between the RA and control experiments is highly significant (P < 0·001).

FIG. 1 Migration index of RA patients and controls with antibody-coated erythrocytes. No correlation could be found between Waaler-Rose titre and the extent of migration inhibition. ▲ controls, ● RA patients.
by comparing the migration of guinea pig macrophages in the presence of MIF-containing supernates to migration of the same cells with control supernates. Results are shown in Fig. 2. Each point represents means of duplicate experiments. Supernates of control and experimental cultures were always tested in parallel and at least one RA patient and one control were tested on any individual guinea pig macrophage preparation. Supernates of lymphocytes of RA patients showed migration inhibitory activity in 16 of the 20 cases tested. From the 4 supernates not showing such activity, 2 were from patients who also failed to be inhibited by antigen–antibody complexes in the direct migration assay.

The other 2 were supernates from patients whose cells did not show migration inhibition in the direct assay with indices of 0·6 and 0·52, respectively. The mean value of migration indices with RA supernates was 0·51 with a standard error of 0·245.

Two of the supernates of lymphocyte cultures from healthy persons caused in one case moderate, and in one case marked, migration inhibition with values of 0·77 and 0·44. Eighteen of the supernates from control persons did not lead to migration inhibition when tested on guinea pig macrophages. The mean value of migration indices with control supernates was 0·98 with a standard deviation of 0·268.

The significance of the results has been checked with the T-test for independent samples with equal variances. The difference between migration indices obtained by RA supernates and control supernates is highly significant (P < 0·001).

Discussion
As in our earlier work (Eibl, 1972), it could be shown in this study as well that antigen–antibody complexes inhibit the migration of lymphocytes from RA patients. Lymphocytes of eighteen of twenty patients tested showed significant migration inhibition with IgG-coated red cells. The migration index was below 0·8. Only one of twenty healthy persons tested showed migration inhibition under equal conditions. Brostoff and others (1973), who studied migration inhibition with aggregated gamma-globulin, could only show significant inhibition in 45% of the patients tested. Our previous work (Eibl, 1972) indicated that RA patients, whose lymphocytes react with antigen–antibody complexes, also react with aggregated globulin. For this reason we assume that the higher percentage of positive results in our studies is due to the more sensitive experimental system we used. Brostoff used buffy coat cells, a mixture of granulocytes, monocytes, and lymphocytes. We used isolated mononuclear cells consisting of lymphocytes and monocytes, and it is likely that granulocytes not reacting in this system dilute the effect.

To differentiate whether the migration inhibition observed is caused by the reaction of T-cells or by cells involved in the production of rheumatoid factors, we tried to correlate the presence and titres of rheumatoid factors in the patient's serum with migration inhibition of his lymphocytes. No correlation was shown, suggesting that early steps of rheumatoid factor production are not of major importance in the migration inhibition observed. Bach and others (1970), testing lymphocyte reactivity to antigen–antibody complexes by rosette technique, came to similar conclusions. In our previous studies using the rosetting system with human D-positive erythrocytes and human anti-D IgG antibody (Eibl and others, 1973), no correlation could be found with rheumatoid titres, either.

If migration inhibition has to be regarded as an in vitro correlate of delayed hypersensitivity, it has to be caused by soluble mediators produced by T-lymphocytes. It was, therefore, of interest to investigate whether such soluble mediators can be shown in the reaction between mononuclear cells of RA patients and antigen–antibody complexes. We preferred to use a particulate antigen, such as the red cell which can easily be separated from the MIF-containing supernate. To test the reactivity of RA lymphocytes with antigen–antibody complexes, we coated red cells with homologous IgG antibody. Lymphocytes incubated with erythrocytes...
alone served as controls. This type of control has been regarded as the most suitable one because the number of cells and metabolic conditions could be considered identical in control and experimental cultures. MIF production was shown in 16 of 20 lymphocyte preparations from RA patients, with guinea pig peritoneal macrophages as target cells. Only 2 of 20 lymphocyte preparations from healthy persons produced MIF after incubation with antigen antibody complexes. The difference between the patients and the controls was statistically highly significant. These data provide substantial evidence that mediators are produced by lymphocytes of RA patients on reaction with homologous antigen–antibody complexes and that the migration inhibition observed can be regarded in most of the patients as a true reaction of delayed hypersensitivity. We could not show the production of mediators in all of the patients tested and do not want to postulate that this is the only type of reaction between mononuclear cells of RA patients and antigen–antibody complexes.

The importance of sensitized lymphocytes in RA has been suggested by several previous studies and lymphocytic infiltration of the synovia in the early stage of the disease supports this. Delayed hypersensitivity to synovial tissue could be shown in RA patients by skin testing (Braunsteiner, Eibl, and Fellinger, 1961) and by migration inhibition (Rothenberger and Thiele, 1971). Kinsella (1973) showed that joint fluid of RA patients induced in vitro transformation of autologous peripheral blood lymphocytes. This response could either be due to the effect of soluble mediators or to antigen–antibody complexes present in the joint fluid of RA patients.

The results in this paper support the assumption that antigen–antibody complexes can trigger the production of mediators by T-lymphocytes in RA under conditions where cells of healthy persons do not react. We think the difference observed between RA patients and controls is quantitative and might represent amplification of a physiological mechanism in rheumatic disease.

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References


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