Antibodies to Cellular Antigens in Sjögren's Syndrome

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ABSTRACT An extract of human lymphocytes from continuous cell culture was used as the antigen source to detect antibodies in sera of patients with Sjögren's syndrome (SS). Using double diffusion in agarose, 85% of a selected group of patients had precipitating antibodies. Three precipitating antigen-antibody systems were detected and were shown to be different from those described previously in other systemic rheumatic diseases. The SS precipitating antibodies were temporarily classified as precipitins A, B, and C. SS patients with sicca syndrome but without clinical rheumatoid arthritis had precipitin systems A and/or B, and SS patients with associated rheumatoid arthritis had precipitin system C. Serum reactants were demonstrated by immunoelectrophoresis to migrate in the γ globulin region. The precipitating activity of the serum factors was not destroyed by treatment with 2-mercaptoethanol and was not removed by absorption of rheumatoid factor from the sera. The reactivity of the lymphocyte antigens was destroyed by treatment with trypsin but not by deoxyribonuclease or ribonuclease.

INTRODUCTION

Sjögren's syndrome (SS) is a chronic inflammatory autoimmune disease consisting of keratoconjunctivitis sicca and/or xerostomia in association with a connective tissue disease. It has been suggested that the clinical diagnosis of SS can be made when any two of the three features are found (1). Some of the more common connective tissue diseases found in association with SS are rheumatoid arthritis (RA), systemic lupus erythematosus (SLE), progressive systemic sclerosis (PSS), polymyositis, and polyarteritis nodosa (1, 2). Occasionally SS may be associated with plasma cell dyscrasia and lymphoma (3).

In the past, diagnosis of the sicca syndrome relied heavily on characteristic signs and physical tests such as the Schirmer test for keratoconjunctivitis sicca (4) and the Lashley cup test for xerostomia (5). A recent advance in the diagnosis of xerostomia is the evaluation of salivary glandular function by scintigraphy (6). Confirmation of SS is often made by lip biopsy (6). Although these techniques are adequate, more informative tests have been sought.

Various serological tests for SS have been evaluated. In studies where sera were tested for antinuclear antibodies, 48-68% were positive by immunofluorescence on tissue substrates (1, 2). Nuclear staining patterns of the homogeneous, speckled, and nucleolar types have been described. However, sera from other connective tissue diseases have also shown similar patterns of nuclear staining, and SS cannot be differentiated by this test alone (7). An immunofluorescent test measuring antiallivery duct antibodies has been used to test sera from SS patients. 70% of the sera from SS-RA patients reacted, but only 10% of the sera from patients with sicca syndrome alone were positive (6). Sera from patients with SS have been tested against various tissue extracts by the complement fixation test (1). The incidence of positive tests was high when sera from patients with sicca syndrome alone were tested (79%); however, the positives in the entire series were only 44%.

Immunodiffusion in gel has been used to detect serum precipitating antibodies to tissue antigens (8-12). In these studies, extracts of human thyroid, spleen, and calf thymus nuclei have been used. A higher incidence of precipitating antibodies were reported in the sera of patients with sicca syndrome without RA (29-81%),

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METHODS

Patient selection. 20 patients with SS were screened initially; however, only 18 sera were available in sufficient quantities for repeated studies. The clinical diagnosis of SS for each patient was established when at least two of the following criteria were present: keratoconjunctivitis sicca, xerostomia, or a connective tissue disease. Confirmation of the clinical diagnosis required the following laboratory evidence. The ocular tests used were either a Schirmer test (4) or a rose bengal staining test (1). The tests used for xerostomia were either salivary gland scintigraphy (5) or a lip biopsy (1) showing the abnormalities described in these reports.

The patients with SS were divided into two major categories, those with sicca complex without RA and those with sicca and associated clinical RA. The association with RA was made when symptoms and laboratory data met the criteria for classical and probable RA, as defined by the American Rheumatism Association (14). On the other hand, none of the patients with SS-sicca without RA had symptoms satisfying criteria for classical or probable RA. Other sera which were examined were obtained from 27 patients with SLE, 16 with PSS, 37 with classical and probable RA, 12 with mixed connective tissue disease (MCTD), 22 with discoid lupus erythematosus (DLE), and 22 from apparently healthy individuals. Patients with SLE met the preliminary criteria for this disease (14) and did not have features diagnostic of other connective tissue diseases. Patients were classified as having PSS on the basis of typical skin changes in the absence of polymyositis or diagnostic criteria for SLE. MCTD patients were diagnosed on the basis of classical symptoms and signs described previously (15) with the association of high titers of antibody to nuclear ribonucleoprotein (RNP). Patients with DLE had skin lesions which were considered to be compatible with this diagnosis, but, in addition, did not have other features meeting criteria for SLE (14).

Tissue extracts. A human cell line (Wil2) donated by Dr. R. Lerner, Scripps Clinic and Research Foundation, La Jolla, Calif.) was used as a source for tissue extraction. Wil2 is a continuously growing diploid lymphocyte line which was obtained from a patient with hereditary spherocytosis. The cells were seeded in Earle's minimal essential medium (MEM) supplemented with 2 mM each of glutamine, sodium pyruvate, and nonessential amino acids (Flow Laboratories, Inc., Rockville, Md.). The supplemented medium also contained 10% fetal calf serum and 10 ml/liter of antibiotic-antimycotic mixture (Grand Island Biological Co., Grand Island, N. Y.), and was adjusted to a final pH of 7.2 with 5.6% NaHCO3. Cells were allowed to grow on a gyratory shaker to a concentration of $1.5 \times 10^9$/mm$. After this, they were centrifuged at 350 g for 10 min, the packed cells resuspended in MEM without calf serum, and were centrifuged again at 900 g for 6 min. The washing process was repeated three times. After the final centrifugation an equal volume of 0.25 M sucrose with 0.004 M calcium chloride at pH 6.2 (0.01 M phosphate) was added. The cells were then sonicated with a model W185 sonicator (Ultrasonic Systems Inc., Farmingdale, N. Y.) at a maximum output for 30 s. The sonicate was then centrifuged at 105,000 g for 20 min. The supernate was saved and used as a source of antigen. The entire procedure was performed at 4°C.

Immunological techniques. For the demonstration of precipitating antibodies in human sera, a modification of the Ouchterlony double-diffusion method was used (16). Plastic plates (1 X 3 cm) containing 5 ml of 1% agarose were used. Reactions were allowed to proceed at room temperature, and precipitin lines were observed over a period of 72 h. Early in these studies, it was apparent that reactive sera were segregating into two groups. For each group of precipitating sera, a prototype serum was chosen and used as the reference serum at a dilution which gave optimum precipitin lines with lymphocyte extract. Sera were analyzed in double diffusion by placing them in wells adjacent to reference sera and by examination of precipitin lines for identity or nonidentity with these reference sera. In cases where precipitin lines were weak, manipulation of the system such as changing the dilution of sera or the concentration of antigens in extract produced stronger precipitin lines and allowed definite identification with reference sera.

As a result of previous studies, other reference sera were available which contained antibodies to DNA, solubilized deoxyribonucleoprotein (snP), streptomyein (Sm) antigen, RNP antigen, and nucleolar RNA (17). Antibodies to DNA, snP, and Sm antigens are present primarily in SLE, antibody to RNP in SLE and the MCTD, and antibody to nucleolar RNA in scleroderma and SS (17). With sera containing precipitating antibodies to Wil2 extract, it was possible to show that the antibodies in SS sera were not identical to those previously described for other connective tissue diseases.

Noble agar (1%) in 0.05 ionic strength barbital buffer, pH 8.5, was used for immunoelectrophoresis (IEP). Patients' sera were placed in the center well and electrophoresed for 1.5 h at 8 mA/slide. Lymphocyte extract was added to one trough and antihuman serum added to the second trough. Reaction was allowed to proceed at room temperature. Precipitin lines were observed over a period of 72 h. Appropriate control sera and antigens were treated in the same manner.

Enzymatic treatment of extracts. Ribonuclease (RNase), deoxyribonuclease (DNase), and trypsin were obtained from Worthington Biochemical Corp., Freehold, N. J. They were first tested in the appropriate buffers to determine if the enzymes were fully active in these buffers (18). These enzymes were employed in the treatment of lymphocyte extracts. The ratios by weight of enzyme to substrate used were as follows: RNase to substrate 1:1; 1:5, and 1:30,
DNase to substrate 1:2 and 1:50, and trypsin to substrate 1:20 and 1:100. All digestions were incubated at 37°C for 2 h. Trypsin digestions were performed in 0.04 M Tris buffer at pH 8.1 in the presence of 0.0115 M calcium ions. DNase digestions were performed in 0.01 M phosphate buffer, 0.15 M NaCl at pH 7.4 (PBS) in the presence of 0.006 M magnesium ions. The RNase digestions were performed also in PBS at pH 7.4. The treated extracts were then titered out against prototype sera in the Ouchterlony technique and compared with controls.

**Chemical treatment of patients' sera.** Patients' sera were treated with 2-mercaptoethanol (2-ME). Reactants were added to each other at volume of 1:1 leaving the final molarity of 2-ME at 0.1 M and the sera at a dilution of 1:2. After mixing, the reactants were left at room temperature for 15 h (19) before testing by the Ouchterlony technique. The treated sera were then titered out against the lymphocyte extract and compared with controls.

**Absorption of rheumatoid factor from patients' sera.** Human γ-globulins (fraction II) (Miles Laboratories Inc., Miles Research Div., Elkhart, Ind.) at a concentration of 25 mg/ml in PBS were heated at 63°C for 20 min. The aggregated globulins were then lyophilized and added to sera at 15 mg/ml and left overnight at 4°C. The samples were then centrifuged at 36,000 g for 20 min, and the supernate was tested for residual rheumatoid factor (RF) by the latex agglutination test.

**Chemical determinations.** Protein concentrations were determined by the Lowry procedure (20). DNA was determined by a modification of the diphenylamine reaction (21), and RNA was determined by the orcinol reaction (22).

**RESULTS**

**Characterization of crude extracts.** Some of the characteristics of the extracts have been determined by chemical methods. After the cells had been sonicated in a sucrose-calcium chloride solution, the protein concentration was 40 mg/ml. After centrifugation at 105,000 g, the supernate contained 25 mg/ml protein, 5.3 mg/ml RNA, and 0.018 mg/ml DNA. This supernate was used as a source of antigen for detecting antibodies in patients' sera by double diffusion in agarose.

**Preliminary screening of patients' sera by Ouchterlony technique.** The sera from 20 SS patients were tested against lymphocyte extracts containing 25 mg/ml protein in the Ouchterlony technique. 17 of these were positive (85%). When the SS sera which were previously positive were titered out against the undiluted extract, precipitin lines were observed with some sera at 1/512 dilution. When the extract was diluted out and tested against seven representative SS sera, the end points ranged between 5 mg/ml and 1.2 mg/ml of protein. Some of the extract was dialyzed against PBS at pH 7.4 and lyophilized. The lyophilized material was as reactive as unlyophilized extract and remained stable for at least 2 mo at 4°C.

Fig. 1 illustrates the three precipitin systems found in sera from patients with SS. Serum SS, which contained precipitins A and B, was from a patient with sicca syndrome. The serum labeled SS-RA was from a patient with sicca syndrome and RA which produced one precipitin system with the extract and showed complete nonidentity with the precipitins of the SS serum. The treated sera were then titered out against two different methods. The treated sera were then titered out against the lymphocyte extract and compared with controls.

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(closer to the Wil2 extract), and precipitin system B. Serum SS-RA produced one line with the extract (precipitin system C) that showed complete nonidentity with systems A or B. It was important to establish that the precipitin systems observed with SS sera were different from precipitating antibodies described in other connective tissue diseases. This was performed by using previously analyzed sera containing known antibodies, and such a study is shown in Fig. 2. Serum SS was from a patient with SS and reacted with Wil2 to give the two precipitin systems described in Fig. 1. Precipitin system A was much stronger than precipitin system B because the two tissue antigens were present in the extract in different concentrations, with A predominating over B. With this particular SS serum, antibody to the A antigen was present in higher titer than antibody to the B antigen. SLE1 was serum from a patient with SLE and gave two precipitin lines with Wil2, the sharper line closer to the antigen well has been identified in previous studies as the Sm system. The second line further from the antigen well has not been identified. It was clear that the Sm system of SLE was different from the systems in SS, MCTD1 and MCTD2 were two sera from two patients with MCTD. They also reacted with Wil2 extract, and this system has been identified as the RNP system of MCTD. The latter was also not identical immunologically with the SS systems. SLE2 was a serum with precipitating antibody to deoxyribonucleoprotein but did not react with Wil2, illustrating that the latter extract did not contain soluble deoxyribonucleoprotein. Studies not illustrated here showed that the SS-precipitating systems were not related to precipitating antibodies to native or denatured DNA. Similar studies were also conducted to analyze precipitin system C which showed that it was not related to precipitating systems involving Sm, RNP, sNP, and native and denatured DNA.

Of 17 sera which were positive in early screening studies, only 15 remained for identification of precipitin systems. Although only a limited number of SS sera were examined, the results appeared to show that precipitins A and B were present mainly in patients with sicca syndrome, and precipitin C was present in patients with SS-RA (see Table I). It was noted that the patient with sicca and juvenile RA had precipitin A, and the two patients with either SS-PSS or SS-Raynaud’s had precipitins A and C.

Sera from patients with diseases other than SS were also studied for precipitins A, B, or C. The only group with a significant incidence was the RA group in which 65% of the patients’ sera contained precipitin C, suggesting that the presence or absence of RA may influence which precipitins were detected (Table II).

**IEP of patients’ sera.** IEP was performed to determine if the serum factor(s) reacting with tissue antigens could be identified by this technique. Five sera (three from sicca patients and two from SS-RA patients) with the highest titers of serum precipitins in double diffusion also showed precipitins by IEP and in all five cases; serum precipitins were located in the γ-globulin region, and the precipitin arcs had the characteristic location and curvature of IgG. A representative example (serum GG) is shown in Fig. 3, where

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**Table I**

<table>
<thead>
<tr>
<th>Patient</th>
<th>Disease</th>
<th>Number of systems</th>
<th>Precipitins</th>
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<td>G. T.</td>
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<td>J. M.</td>
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<td>T. H.</td>
<td>ss</td>
<td>2</td>
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<tr>
<td>P. C.</td>
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<tr>
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<tr>
<td>M. C.</td>
<td>ss-PSS</td>
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<td>+</td>
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<tr>
<td>P. F.</td>
<td>ss-Raynauds</td>
<td>2</td>
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<td>M. M.</td>
<td>ss-RA</td>
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<td>J. C.</td>
<td>ss-RA</td>
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<td>H. O.</td>
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<td>A. D.</td>
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<td>M. G.</td>
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<td>E. R.</td>
<td>ss-RA</td>
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ss, sicca syndrome; JRA, juvenile rheumatoid arthritis.
serum was electrophoresed in the center well and developed with antivhole human serum in the top trough and with tissue extract in the bottom trough. Serum \( \gamma \)-globulin demonstrated two precipitin arcs with gamma mobility and in double diffusion; this serum had been one of the reference sera with the two precipitin systems A and B. The other two sera, containing precipitins A and B, produced two arcs in IEP, but the two sera from patients with SS-RA, containing precipitin C, produced one arc. Other sera with lower titers of precipitins were also tested by IEP, but precipitin arcs were not always detected, a feature which was thought to be related to the lower sensitivity of the IEP technique compared to the Ouchterlony technique.

2-ME treatment of patients' sera. Sera from five patients with SS were treated with 2-ME, tittered against extracts, and the end points compared with controls. All five sera tested were resistant to 2-ME. Three of the sera were from SS patients with sicca syndrome which contained precipitins A and B. The other two sera were from patients with SS-RA which contained precipitin C.

Effects of physical or biochemical treatment of extracts. Some of the physiochemical characteristics of individual antigens have been determined. The extract was treated with RNase, DNase, and trypsin and was tittered against various sera by the Ouchterlony technique. In these studies sera from two patients were used to represent systems A and B, and two sera from SS-RA patients were used to present system C in the Ouchterlony technique. All the antigens in the A, B, and C systems were rendered nonreactive by treatment with trypsin at both enzyme-substrate ratios tested (1:20 and 1:100), but were unaffected by treatment with nuclease enzymes at all ratios tested. The antigens were not affected by 2-ME treatment or by heating at 56°C for 1 h.

Absorption of sera with aggregated IgG. Sera from seven patients with sicca syndrome alone and sera from six patients with SS-RA were tested by the Ouchterlony technique against heat-aggregated IgG. Sera from patients with sicca syndrome were nonreactive with aggregated IgG. Five of the six sera from SS-RA patients reacted. All five had high-titer RF by the latex agglutination test. Sera from these patients with SS-RA were absorbed with aggregated IgG until RF was reduced from a titer of 2,560 to 20 or less. The absorbed sera were negative for precipitins to aggregated IgG, but were still positive against the lymphocyte extract, demonstrating that the precipitin reaction of the C system was not due to RF.

DISCUSSION

In the present studies, we have described three types of precipitating autoantibodies in the sera of patients with SS. With the limited number of SS sera examined, it appeared that two types of precipitating antibodies, temporarily identified as the A and B systems, were present predominantly in SS without RA, whereas the third type of precipitating antibody was present in SS with RA and not in SS without RA. This segregation of precipitating antibodies has been substantiated in a study of a larger number of SS sera and will be reported in a later communication. In these studies 56 patients with sicca syndrome and 33 patients with SS-RA were tested. 76% of the SS-RA patients had precipitin C and less than 9% had precipitins A and/or B. On the other hand, more than 79% of the patients with sicca syndrome had A and/or B, whereas less than 5% had precipitin C. It should be noted that in studies by previous investigators, primarily those of Anderson, Gray, Beck, and Kinnear (9), two types of precipitating antibodies, SiD and SiT, were described. It would appear to be more than coincidental that these workers also found that the two antibodies were present in SS without RA but not in other forms of SS. They observed that one of their antigens, SiT was destroyed by heating at 56°C for 1 h and by digestion with trypsin, but SiD was not affected by either treatment. In our studies, both the A and B antigens were stable to heat but both were destroyed by digestion with trypsin. At this time, it is difficult to explain the discrepancies especially if the two antigens were presumed to be identical. However, we hope to obtain answers to these questions by exchange of sera and reagents with these workers.

In contrast to previous workers who have found low incidence of precipitins to tissue extracts in SS with RA, we found precipitins to the C system in SS-RA in high incidence not only in the few selected sera reported here but also in a larger series. This difference may be due in part to the fact that our extracts were obtained from tissue culture cells, a procedure which permitted immediate extraction of tissue antigens without undergoing autolysis. Alternatively, the C antigen might be unique to lymphocytes or to certain tissue culture cell lines, possibilities which we have not investigated. In any case, an interesting observation that emerged was that serologically, SS-RA could be differentiated from the pure sicca syndrome or SS without RA, suggesting that these two forms of SS might not necessarily be closely related. This was also suggested by MacSween et al. (23) who proposed that a subclinical form of SS might be present in some patients with RA. This proposal was based on the finding that there was a high incidence of salivary duct antibody in SS-RA but a low incidence in SS without RA. Further work

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showing differences in the forms of SS were studies (24, 25) that peripheral lymphocytes from SS-RA responded less well to mitogenic stimulation with phytohemagglutinin, dinitrochlorobenzene, tetanus toxoid, and streptolysin, than lymphocytes from SS without RA.

The intracellular location of the tissue antigens which react with SS sera has not been clearly defined. In the study of Bloch, Buchanan, Wohl, and Bunim, the incidence of antinuclear antibodies (ANA) in a large series of SS patients was 68% (1). Studies in this laboratory have shown a 59% incidence of ANA in SS, but in these same patients, 86% had precipitins A, B, or C. This suggests that at least one or more of the precipitins may be antibodies against cytoplasmic or non-nuclear components. If normal human leukocytes were used as substrate for ANA, in addition to nuclear staining, there was significant cytoplasmic staining with sera which gave positive precipitin reactions, further suggesting that cytoplasmic antigens might be involved in some of the precipitins systems described here.

It is becoming clear from many recent reports that autoantibodies of certain specificities are segregated in their distribution. For example, antibodies to double-stranded DNA and to the Sm nuclear antigen were detected predominantly in SLE (17, 26), and antibodies to nuclear RNP were detected in the MCTD as well as SLE (15, 17, 27). It would thus be important to be able to define the antigenic or biochemical specificities of the many autoantibodies found in certain diseases, since there might be distinct profiles of autoantibody specificities for each disease. The usefulness of such an approach has already become apparent in SLE where tests for anti-DNA antibodies have helped in diagnosis, as well as in management, of SLE. Furthermore, these and other studies on the specificities of autoantibodies in SS might be anticipated to lead to investigations concerning the relationship of these autoantibodies to disease processes. At the present time, it is not known whether these antibodies participate in causing tissue injury or are epiphenomena of previous cellular damage from other causes. In any event it is clear that autoantibodies in SS react with cellular antigens which are immunochemically different from cellular antigens reacting with autoantibodies in other systemic rheumatic diseases. The processes which lead to the production of autoantibodies of such diverse immunochemical specificities and the reasons for the segregation of some types of autoantibodies in certain diseases are not understood at the present time.

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REFERENCES


