Gamma\(_2\), Gamma\(_{1A}\), and Gamma\(_{1M}\) Antinuclear Factors in Human Sera *

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The term “immunoglobulin” is used to refer to one or more of the three classes of serum protein, gamma\(_{1A}\), gamma\(_{1M}\), and gamma\(_{2}\), each of which is known to contain antibodies. The \(\gamma_{1A}\) and \(\gamma_{1M}\) immunoglobulins have also been referred to as beta\(_{2A}\) and beta\(_{2M}\), their electrophoretic mobilities being intermediate between the principal gamma and beta peaks in serum protein electrophoresis. \(\gamma_{1A}\)-globulin, like \(\gamma_{2}\)-globulin, is approximately 7 S in its ultracentrifugal characteristics, but differs significantly from \(\gamma_{2}\)-globulin by other chemical and immunochemical criteria. \(\gamma_{1M}\)-globulin is a macro-gamma globulin that has an ultracentrifugation sedimentation coefficient of approximately 19 S.

Determination of the immunochemical class of globulins responsible for the various autoreactivities found in various diseases may be important for our better understanding of the mechanisms of disease. The antinuclear factors (ANF), which are apparently autoantibodies to nuclear constituents, are found in the sera of patients with systemic lupus erythematosus (LE) and less frequently in other so-called connective tissue diseases, particularly rheumatoid arthritis (RA). Goodman, Fahey, and Malmgren (1) characterized ANF in two LE sera as globulins of both 7 S and 19 S type. Baum and Ziff (2) showed that ANF in LE sera were eluted from DEAE cellulose both

at pH 7.0, 0.01 M phosphate, and at pH 5, 0.3 M phosphate, corresponding to the distribution of 7 S and 19 S globulins. Their data suggest that LE patients tended to have ANF of \(\gamma_{2}\) (7 S) class, whereas RA patients tended to have ANF of \(\gamma_{1M}\) (19 S) class.

Our work, employing different techniques, was designed not only to compare LE and RA patients on the basis of their \(\gamma_{2}\) and \(\gamma_{1M}\) ANF but, in addition, to investigate the incidence of ANF in the more newly described \(\gamma_{1A}\) class of globulins. In vivo tests have suggested that human atopic reagins are \(\gamma_{1A}\)-immunoglobulins (3). Anti-insulin (4), anti-Brucella (5), antidiapheria (5), and antiragweed antibodies (6) have recently been described in all three immunoglobulin classes.

Methods

A modification of the two-layer indirect ANF test (7) was employed. Three separate air-dried slides of human peripheral blood were fixed in 95% ethanol and used as a source of nuclear material. The patient’s serum or a fraction of the patient’s serum was overlaid for 30 minutes at room temperature. Then the slides were washed with phosphate buffered saline, pH 7, for 30 minutes. In the two-layer indirect ANF test, a fluorescein-conjugated goat antiserum against human \(\gamma\)-globulins would be applied for 30 minutes, the slides then washed, mounted in glycerol-saline, and then examined with a fluorescent microscope. In the present test a rabbit serum against one of the three human immunoglobulins (i.e., anti-\(\gamma_{1}\), anti-\(\gamma_{2}\), or anti-\(\gamma_{1M}\)) was applied for 30 minutes as the second layer to each of the three slides. The slides were washed again and then goat antirabbit globulin, conjugated with fluorescein isothiocyanate, was applied as the third layer. After 30 minutes the slides were again washed and then mounted in 90% glycerol-saline with glass cover slips. Appropriate control slides were included in each test to rule out nonspecific adsorption of the rabbit or goat sera to human leukocytes. For quantitation of ANF, serial fourfold dilutions of the patient’s serum were used instead

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of the patient's undiluted serum in the first layer. The titer of ANF of each immunoglobulin class was the highest dilution of the patient's serum giving nuclear fluorescence with a constant amount of specific rabbit and goat sera. The slides were read with a Zeiss-fluorescent microscope, ultraviolet light source, H B O 200 watt (Osram), exciting filter, 420 mμ, BG 12, blocking filter, 500 mμ.

Antisera. Undiluted specific rabbit antisera against human γ2-, γ1A-, and γ1M-immunoglobulins were shown after appropriate absorption to react in immunodiffusion tests only with the homologous human serum globulins (8). In passive cutaneous anaphylaxis in guinea pigs and in hemagglutination tests, 1:10 dilution of the sera detected only the homologous globulins; lesser dilutions were not tried (8). In the current immunofluorescent studies these sera were used in 1:20 dilutions. Goat anti-rabbit serum was obtained from a single goat after multiple immunizations with Pentex Fraction II \(^1\) of rabbit globulins and whole rabbit serum in Freund's complete adjuvant. The goat antisera was fractionated by precipitation in half-saturated ammonium sulfate and conjugated with fluorescein isothiocyanate (7). The goat antirabbit globulin conjugate was absorbed with acetone-dried human spleen powder, whole fresh human serum, and human Fraction II, so that it gave no precipitate with human sera in double diffusion in agar gel and would not directly detect human ANF on leukocyte nuclei, i.e., in the absence of a rabbit antihuman serum-immunoglobulin as an intermediate layer.

Antigens. γ2-, γ1A-, and γ1M-immunoglobulins were prepared from human sera (8). On immunoelectrophoresis using an anti-whole human serum in the antisera wells, they gave only one line in the expected locations.

These purified rabbit antisera and purified human immunoglobulins were used in absorption and blocking experiments designed to demonstrate the specificity of the specific rabbit antisera used in the three-layer ANF test. In absorption studies, serial additions of specific rabbit antisera to a patient's serum were shown to remove only the corresponding class of immunoglobulin ANF. For instance, the addition of antisera specific for γ2- and γ1M-immunoglobulins resulted in the absorption of ANF of both γ2 and γ1M classes, but γ1A ANF was still present. In the blocking experiments, the addition of purified human γ2-globulin to the rabbit anti-γ2 sera blocked the detection of γ2 ANF, whereas the addition of γ1A- or γ1M-globulins did not. Analogous specificity was demonstrated in blocking experiments for the anti-γ1A and anti-γ1M rabbit sera.

Chromatography. Three LE sera were fractionated by DEAE cellulose column chromatography with starting buffer, 0.04 M Tris, 0.005 M PO₄, pH 8.6, and limit buffer 0.5 M Tris, 0.5 M PO₄, pH 5.5, using a Varigrad as described by Riesfeld, Bergenstal, and Hertz (9).

Ultracentrifugation. Sera from four LE and one RA patient were fractionated by ultracentrifugation in a sucrose gradient (10). One-half ml of a 1:4 dilution of the patient's serum in 10% sucrose was layered on top of the sucrose gradient solution. The tubes were spun for 16 to 20 hours at 25,000 rpm in the SW39L rotor head, Spinco model L centrifuge. The fractions were collected as previously described (11), six from above and three from below. The optical densities of each fraction read at 277 mμ were recorded. One tube in each centrifugal run contained Evans blue dye, an incomplete anti-D serum, and rheumatoid factor markers for albumin, 7 S and 19 S globulins, respectively. Each fraction was examined for the presence of γ2-, γ1A-, or γ1M-globulins in double diffusion using the specific rabbit antisera against the immunoglobulins.

Sulfhydryl sensitivity. Samples of five LE sera and one RA serum were diluted 1:8 in 0.1 M mercaptoethanol or in saline and kept at 20°C for 24 hours (12). After dialysis to remove residual mercaptoethanol the treated and control serum specimens were then diluted and tested for ANF in the three-layer test.

Enzyme treatment. Glass slides of peripheral blood smears were incubated for 3 minutes at 20°C with 0.1% solutions of DNase, RNase, or trypsin.\(^2\) The control slides were incubated in PO₄-CO₂ buffer (13). These enzyme-treated or control leukocytes were then used in the three-layer test to determine if the reactivity of the nuclear material was eliminated by enzyme treatment. Sera from 6 RA, 1 juvenile RA, and 3 LE patients were tested on these enzyme-treated slides to show the specificity of their ANF for enzyme-sensitive nuclear material.

LE cell preparations. Patients' bloods were defibrinated by shaking with glass beads (14). After 2 hours of incubation at 20°C, smears of the buffy coat were made and stained with Wright's stain. Each LE cell preparation was examined for 20 minutes at 100 X magnification. In all patients studied all LE cell preparations were obtained at the time of or within 1 week of obtaining blood specimens for study by the ANF test.

Diagnostic criteria for RA and LE. The patients with RA in the present study all had definite or classical RA, by the diagnostic criteria of the American Rheumatism Association (15) and Steinbrocker's system of functional classification and anatomical staging (16). All patients with a diagnosis of LE had positive LE cell preparations at some time during their illness, with one or more of the following signs: fever, rash, serositis, migratory polyarthritis, renal disease, thrombocytopenia, or anemia (15). None of the LE patients had joint deformities, rheumatoid nodules, or erosion of cartilage on radiologic examination.

Results

LE and RA sera were found to have ANF of one, two, or most often all three immunoglobulin classes. Nuclei that were either solidly fluorescent or rimmed (Figure 1) denoted a positive

\(^1\) Pentex Inc., Kankakee, Ill.

\(^2\) Worthington Biochemical Corp., Freehold, N. J.
ANF test, whereas only background staining of the leukocyte cytoplasm was seen when the test serum contained no detectable amounts of ANF.

**Chromatographic studies**

Three LE sera were fractionated by DEAE cellulose column chromatography. ANF of γ2, γ1A, and γ1M classes were detected in those eluates expected to contain each of these immunoglobulins (17–19). A single chromatogram is shown in Figure 2. The γ1A ANF was not eluted without γ1M ANF with the particular gradient used. Chromatographic separation of two other LE sera gave similar results.

**Sulfhydryl sensitivity and sedimentation characteristics**

1. **Gamma2 and gamma1M ANF.** The antibody activity of γ2-globulins is not destroyed by incubation with 0.1 M mercaptoethanol, whereas that of γ1M-globulins is (12). Figure 3 demonstrates the sensitivity to inactivation by mercaptoethanol of ANF of the three immunoglobulin classes in five LE and three RA sera. The titers of γ1M ANF were reduced at least 16-fold in all cases. The titers of γ2 were not reduced in any instances. ANF detected by mono specific anti-γ2 antisera consistently sedimented as 7 S globulins, whereas γ1M ANF sedimented as 19 S globulins (Table I).

2. **Gamma1A ANF.** Titters of γ1A ANF were reduced completely in two instances, partially in four, and none in two (Figure 3). In sera from Patients K and L the γ1A ANF sedimented as 7 S globulins, whereas in sera S1, D, and E the γ1A ANF appeared at least in part to sediment faster than did γ2 ANF (Table I). These data suggest that γ1A antibody may be quite heterogenous in regard to its sedimentation characteristics. Furthermore, the sedimentation characteristics of γ1A ANF in particular sera were found to cor-

![FIG. 1. RESULT OF TYPICAL THREE-LAYER ANTINUCLEAR FACTOR (ANF) TEST ON DILUTION OF LUPUS ERYTHEMATOSUS (LE) AND RHEUMATOID ARTHRITIS (RA) SERA. Fluorescent nuclei either solidly fluorescent or rimmed denote a positive test. The LE serum is shown to have ANF of γ2-γ1A- and γ1M-immunoglobulin classes in titer of 1:256 or greater. The RA sera diluted 1:64 contain γ1A and γ2M ANF in one case and only γ2M ANF in the other.](image-url)
relation with sensitivity to inactivation by 0.1 M mercaptoethanol (Figure 3). In sera K and L₁ where γ₁A ANF sedimented as 7 S globulins, the titers of γ₁A ANF were not affected by mercaptoethanol treatment. In sera S₁ and E where γ₁A ANF appeared to sediment, at least in part more rapidly than 7 S globulins, the titers of γ₁A ANF were reduced by mercaptoethanol treatment.

Relation of ANF to RF

Hall and associates (20) demonstrated that in sera from RA patients where both ANF and rheumatoid factors (RF) were present, ANF could be absorbed by histone without affecting the titer of RF. RF could be absorbed by latex particles coated with γ-globulin without affecting the titer of ANF. In our study, 16 of 17 LE sera had measurable amounts of γ₁M ANF. None of these sera had RF detectable at 1:20 serum dilution with the human Fr. II sensitized latex agglutination test. Since the RA sera had high titers of RF as well as ANF, however, it was possible that in these latter sera the titers of ANF, particularly in the γ₁M class, might be falsely elevated due to interaction of RF and ANF on the nuclei. If this were so, the titers of ANF might be reduced if the RF were removed by absorption. Three LE sera without RF and four RA sera with latex titers of 1:160 to 1:10, 512 for RF, were serially absorbed with normal human Fr. II, aggregated by bis-diazotized benzidine (21). The RF titers were reduced to less than 1:20, but the titers of ANF were unaffected in six out of the seven sera. In one serum, however, the titer of γ₁M ANF was reduced from 1:256 to 1:4 while the titers of γ₂ and γ₁A were unaffected. Unsuccessful attempts were made to elute ANF from the absorbing aggregated γ-globulin by acidification to pH 5.

This experiment suggested that the titer of γ₁M ANF in some RA sera might be factiously elevated due to RF binding to γ₂ ANF. To test this, sera containing only RF were added to dilutions of three other sera in which only γ₂ ANF could be detected. In one of the three sera (serum K, Table II) a γ₁M ANF test appeared. Serum K, which was from a patient with juvenile rheumatoid arthritis, gave similar results with addition of ANF-negative RA sera W, A, and H.
TABLE I
Sedimentation of \( \gamma_2, \gamma_{1A}, \gamma_{1M} \) antinuclear factors (ANF) by sucrose gradient ultracentrifugation

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* Patient with definite rheumatoid arthritis (RA), stage IV, class III, onset at age 12 with disease activity for 25 years. \( \gamma_2 \) ANF titer 1:4; \( \gamma_{1A} \), 1:64; \( \gamma_{1M} \), 1:4, not included in Figure 3. Patients K, L_1, S_1, and E are shown in Figure 4. 1+ 4+, more than 75% of the nuclei in a field brightly fluoresce; 3+, 50 to 75% of the nuclei fluoresce; 2+, 25 to 50% of the nuclei fluoresce; 1+, up to 25% of the nuclei fluoresce. Fractions giving 3+ or 4+ nuclear fluorescence give positive ANF when diluted 1:10. The titer of fractions giving 1+ or 2+ nuclear fluorescence was 1:10. Sera from Patients L_1 and S_1 were placed in adjacent cups of the SW39L rotor and centrifuged for the same period.

The two other sera used as sources of \( \gamma_2 \) ANF were from adult LE patients A and M_1. Serum M_1 (Figure 4) was diluted 1:16, and serum A was used undiluted. They exhibited no ANF by \( \gamma_{1M} \) antiserum after mixing with the same three RA sera that had been used above. Apparently only in selected instances will RF complex with \( \gamma_2 \) ANF to be detected as though there were \( \gamma_{1M} \) ANF. This may well depend on specificity of RF for particular \( \gamma \)-globulins (i.e., Gm specificity) (22) or on denaturation of \( \gamma_2 \) ANF globulins. Possibly, therefore, in some RA sera, three-layer ANF tests using specific anti-\( \gamma_{1M} \) antiserum detect not only \( \gamma_{1M} \) ANF but also \( \gamma_{1M} \) RF complexed to \( \gamma_2 \) ANF.

Specificity for different nuclear materials

Sera from five adult RA, one juvenile RA, and four LE patients were tested on cells treated with enzymes to assess the specificities of ANF of the different classes. DNAse treatment abolished the reactivity of the nuclei with all the ANF of all but two of the ten sera. In one of these two sera (Patient B, Table III) \( \gamma_{1M} \) activity persisted, but the fluorescent nuclei were fewer and not so bright as with control slides. In the other serum (Patient V), ANF activity persisted in all immunoglobulin classes, but the number of nuclei staining and the intensity of staining was reduced by DNase and possibly RNAse treatment. Thus, some of the ANF present in the sera of Patients B and V apparently bind to nuclear material not
entirely sensitive to the action of DNAse. Patient M's \( \gamma_1 \) and \( \gamma_1 \) ANF did not bind to trypsin-treated leukocyte nuclei or to DNAse-treated nuclei. Trypsin-treated leukocytes also gave lesser nuclear fluorescence with Patient B's \( \gamma_1 \) ANF than did untreated nuclei. Apparently, therefore, part of the ANF in serum from Patient B and all of that from M react with nuclear material sensitive to the action of trypsin. \( \gamma_2 \), \( \gamma_1 \), or \( \gamma_1 \) ANF in the remaining seven sera (not shown in Table III) gave negative results with DNAse-treated nuclei and positive nuclear fluorescence with control, RNAse, and trypsin-treated nuclei. Indeed, with these seven sera trypsin treatment resulted in brighter staining of more nuclei than was seen in the control slides. The possibility may be considered that trypsin treatment may make nuclear membranes more permeable to ANF directed primarily against nontrypsin-sensitive intranuclear materials.

**LE cell preparations**

LE cell preparations were positive in 14 of the 17 LE patients with ANF detected by the immunofluorescent tests. Of the 18 RA patients six had positive LE cell preparations; five of these had \( \gamma_2 \) ANF of four or greater. Only one of nine RA patients with \( \gamma_2 \) ANF of less than four had a positive LE cell preparation (p < 0.01).

**Three-layer ANF test in LE and RA**

High titers of ANF activities were frequent in all three immunoglobulin classes in sera from patients with LE. The \( \gamma_2 \) ANF titered 16 or better in 15 of the 17 LE sera (Figure 4). There was no absolute correlation between the titers in the various immunoglobulin classes. In our clinic 27% of the RA patients have been shown to have ANF as detected by a two-layer indirect immunofluorescent test using a rabbit antiserum directed primarily against human \( \gamma_2 \)-globulin (23). Eighteen RA sera selected on the basis of a positive two-layer ANF test were examined by the three-layer ANF test. Only four of the 18 RA sera had \( \gamma_2 \) ANF that titered to 16 or better (Figure 5). The mean \( \gamma_2 \) titer among the RA sera was significantly (p < 0.01) less than the mean \( \gamma_2 \) titer among the LE sera. The \( \gamma_1 \) ANF titers of the RA sera, however, were entirely comparable to those of the LE sera, whereas the \( \gamma_1 \) ANF, although lower in mean titer, were not significantly so at the p < 0.05 level.

The three-layer immunofluorescent test might well be expected to detect amounts of ANF below the limits of sensitivity of a two-layer ANF test (7). Sera from 32 patients attending the hematology clinic were negative for ANF by the two-layer test (Figure 6). When tested by the three-layer test, ANF of only \( \gamma_1 \) class was found in four of these 32 control sera. In none of the four was the titer of \( \gamma_1 \) greater than 1:4. Ten
sera from RA patients with no ANF by the two-layer test were examined in the more sensitive three-layer test. One exhibited \( \gamma_2 \) and \( \gamma_{1A} \) ANF. Five had detectable amounts of \( \gamma_{1M} \) ANF. The titers are shown in Figure 6.

Clinical correlations

Seven of the nine LE patients with disease less than 1 year had \( \gamma_2 \) ANF titers of 64 or greater, whereas only two of the eight patients with diseases more than 1 year had \( \gamma_2 \) ANF titers in this range (Figure 4). The eight LE sera with \( \gamma_{1A} \) ANF titers of 16 or greater were from patients with clinical disease of less than 1 year's duration. Of the remaining nine patients, five had measurable \( \gamma_{1A} \) ANF titers of 1:4 or less, and eight of those patients had been ill for more than 1 year \((p < 0.01)\). Thus, our LE patients with symptoms of disease for less than 1 year tended to have higher titers of both \( \gamma_2 \) and \( \gamma_{1A} \) ANF. Those patients with disease of more than 1 year tended to be less severely ill; they had been on steroid or chloroquine therapy, or both, for longer periods.

In the RA patients with positive three-layer tests for ANF there was no correlation between the duration of disease and the titers of ANF of any one immunoglobulin class (Figure 5). However, these 18 sera were from RA patients with severe deforming and debilitating disease of long duration (23). They are only 27% of the group
of RA tested, the rest being negative in the two-layer test and from patients generally with less severe disease and of shorter duration. Ten such RA sera with negative two-layer tests were tested in the three-layer ANF test (Figure 6). Five of these with disease of somewhat longer duration had detectable amounts of ANF.

Discussion

A three-layer indirect immunofluorescent technique was adopted to measure the amounts of $\gamma_2$, $\gamma_1A$, and $\gamma_1M$ ANF in human sera. The specificity of the method is dependent on the specificity of the specific rabbit antisera used in the test. Others have shown that such antisera give mono-specific lines of precipitation in gel with their homologous antigens (17–19). The antisera used here were also shown to be specific by this method and in addition by the more sensitive method of passive cutaneous anaphylaxis in guinea pig skin (8). Furthermore, the specificity of the test was corroborated by specific blocking and absorption experiments with the immunofluorescent technique. ANF were detected in all three immunoglobulin classes and were found to elute from DEAE cellulose columns as previously described for the various $\gamma$-globulins (17).
Since anti-insulin (4), anti-Brucella, antidiphtheria (5), and antiragweed antibodies (6) have recently been described in all three immunoglobulin classes, the detection of ANF in the same classes of globulin fortified the suggestion that ANF, too, is an antibody. The reactivity of ANF of γ2 class was not affected by incubation with mercaptoethanol, whereas the reactivity of γ1M ANF was greatly reduced or abolished as is typical of other antibodies of γ2 and γ1M immunoglobulin classes (12). In two sera where the γ1A ANF sedimented as 7 S globulins the reactivity of γ1A ANF was unaffected by mercaptoethanol. In five sera where γ1A ANF sedimented more rapidly than 7 S globulins, the γ1A ANF were, at least in part, inactivated.

The specificity of ANF for particular nuclear antigens may be of pathogenetic importance. Casals, Friou, and Teague (24) have suggested that the presence of antibody to DNA is particularly correlated with illness. In seven of the ten sera tested here the ANF of all three immunoglobulin classes reacted with nuclear material sensitive to DNAse but resistant to trypsin and RNase. In one case (Patient B, Table III) γ1M ANF reacted with nuclear material resistant, separately, to trypsin, DNAse, and RNase, whereas γ2 ANF in the same serum reacted with DNAse-sensitive material. Another patient, (M3, Table III) had a serum that contained both γ1A and γ1M ANF reactive with nuclear material sensitive to either trypsin or DNAse. Patient V with LE had a serum that contained γ2, γ1A, and γ1M ANF which were at least in part reactive with nuclear material sensitive to neither trypsin, DNAse, nor RNase. The speckled pattern obtained with γ1M ANF on DNAse-treated nuclei suggests that some of the ANF in this serum was directed against phosphate extractable nuclear antigen described by Holman, Deicher, and Kunkel (25) and frequently found in the sera from cases of Sjögren's syndrome (26).

RF and ANF may occur together in the same human serum. Other workers (20) have shown that they behave as distinctively separate antibodies with specificity for two unrelated antigens, γ2 globulin and nuclear material, respectively. Since the three-layer ANF test with an anti-γ1M
serum was positive in 17 LE sera where no RF was detected, it was concluded that indeed these sera did contain $\gamma_M$ molecules that behave as antibodies to the nuclei. Since a positive three-layer test for $\gamma_M$ resulted where, in one instance, RF was added to $\gamma_2$ ANF in an in vitro experiment, possibly the three-layer test may sometimes give spuriously high titers for $\gamma_M$ ANF in some RA sera, where high titer RF and ANF occur together.

The serum factor responsible for the LE cell test has been described as a 7 S $\gamma_2$ globulin (27). The relationship between ANF detected by immunofluorescence and the LE cell factors was not directly examined in this study, but positive LE cell tests in RA patients were associated with higher titers of $\gamma_2$ ANF. In no case, as yet, have we failed to detect $\gamma_2$ ANF in the serum of a patient with a positive LE cell test. This observation does not, however, exclude the presence of LE cell-inducing factors of $\gamma_{1A}$ or $\gamma_M$ class.

With the less sensitive two-layer test, others have had negative ANF tests in patients with positive LE preps (20). In our hands, the two-layer ANF test has routinely been positive whenever the LE cell test or three-layer ANF test has been positive. In one patient (C5 with RA, Figure 5) the two-layer ANF test was negative on the usual peripheral smears but was positive on peripheral smears made with blood shaken with glass beads so as to provide more damaged leukocyte nuclei, as in a case described by Svec and Kaplan (28). The negative ANF tests associated with positive LE cell tests reported by others are less likely due to this type of phenomenon, but, probably more usually, due to lesser sensitivity of the reagents used in their ANF test.

The titers of $\gamma_2$ ANF were found to be significantly lower in 18 selected RA patients than were found in 17 LE patients. The median titer of $\gamma_{1A}$ ANF was lower in the RA group, but the difference between the RA and LE groups for both $\gamma_{1A}$ and $\gamma_M$ ANF was not statistically significant. Of interest was the higher titer of $\gamma_2$ and $\gamma_{1A}$ ANF in those LE patients with acute disease of less than 1 year than in the patients with LE for more than 1 year who were in clinical remission and had been treated with steroids and chloroquin. The RA patients that exhibited positive ANF tests constituted only 27% of the clinic population. They had a higher incidence of severe disease, disease of longer duration, and a higher incidence of rheumatoid nodules (23). From the larger group of less severely ill RA patients (73%) with a negative ANF by the two-layer test, five with disease of longer duration of the ten selected were shown to have $\gamma_M$ ANF. The biological significance of having a particular antibody in one or another immunoglobulin class or in having specificity for one or another nuclear antigen, or both, is at present a matter for speculation only. Immunization of man and animals with some foreign antigens results first in the production of $\gamma_M$ antibodies and then $\gamma_2$ antibodies (29). By analogy the immunoglobulin class of ANF may signify a particular stage of immunization in the disease, and the correlation between presence of the various classes of ANF and the duration of RA is suggestive of this. A prospective study is underway to test whether $\gamma_M$ ANF appears as an earlier serologic entity in the course of RA than do $\gamma_2$ or $\gamma_{1A}$ ANF.

Summary

A modification of the indirect immunofluorescent technique was employed to detect antinuclear factors (ANF) of $\gamma_2$, $\gamma_{1A}$, or $\gamma_M$-immunoglobulin classes. $\gamma_{1A}$ ANF in two cases appeared to sediment as 7 S globulins and be resistant to sulfhydryl (SH) treatment, whereas in other cases, they appeared to sediment faster than 7 S globulins and be inactivated by the SH treatment. ANF and rheumatoid factors (RF), although frequently found together in the same serum, behaved as separate serological entities, but evidence was found that in some RA sera RF may interact with $\gamma_2$ ANF on nuclei to give the appearance of $\gamma_M$ ANF.

ANF of all three immunoglobulin classes was detected in sera from both rheumatoid arthritis (RA) and lupus erythematosus (LE) patients. $\gamma_2$ ANF was found in higher titer in LE than in RA sera. Among the LE sera $\gamma_2$ and $\gamma_{1A}$ ANF were found in higher titer, if the sera were from patients with disease for less than 1 year. LE patients with disease for more than 1 year, who generally were also in remission and on chloroquin or steroid therapy, had lower titers of $\gamma_2$ and $\gamma_{1A}$ ANF.

Among the RA sera only 27% had ANF by a
less sensitive immunofluorescent test. Among these ANF was found in those patients who had severe disease with a median duration of 9 years. RA sera lacking ANF by a less sensitive test were from patients with less severe disease and disease of shorter duration, but of ten such patients selected for study, five with disease of somewhat longer duration proved to have low titer γ1M ANF.

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References


