Antibodies to T Cells in Patients with Systemic Lupus Erythematosus Can Induce Antibody-dependent Cell-mediated Cytotoxicity against Human T Cells

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Abstract Patients with active systemic lupus erythematosus (SLE) often have circulating antibodies to T cells. These patients also often have leukopenia and diminished numbers of T lymphocytes. In addition, certain T lymphocyte functions are frequently impaired in patients with SLE. It has been previously considered that a complement-dependent cytotoxic mechanism was responsible for the above observations. We now demonstrate that antibody-dependent cell-mediated cytotoxicity (ADCC), a cytotoxic reaction mediated by antibody and effector cells in the absence of complement, can also kill T cells from normal individuals as well as from patients with SLE. Moreover, this ADCC could be observed using the plasma, effector cells, and target cells all obtained from the same individual with SLE.

Plasma of those patients with active SLE, and in whom anti-T cell antibodies could be demonstrated by the more classical complement-dependent cytotoxicity, was most often able to mediate such an ADCC reaction. The IgG fraction of the plasma was responsible for inducing ADCC, and aggregated IgG could block the reaction. The fact that the IgG fraction was often more effective than the unfractionated plasma suggested that immune complexes present in SLE plasma might partially block the expression of ADCC. Because a single SLE plasma could induce ADCC in T cells from several different unrelated individuals, it is unlikely that antibodies directed against particular human leukocyte antigens (HLA) or blood group antigens are involved.

Received for publication 12 March 1980 and, in revised form 6 October 1980.

INTRODUCTION

The sera or plasma from some patients with systemic lupus erythematosus (SLE) contain antibodies directed against normal leukocytes, especially when the disease is active. These antibodies are thought to be related to the leukopenia observed in many patients with active SLE. Their existence has been demonstrated by a variety of methods such as antiglobulin tests, complement-dependent cytotoxicity (CDC) assays (1), and immunofluorescence techniques (2, 3). The studies to date have indicated that some of these antibodies are reactive with T lymphocytes and that these antibodies are of the IgG, IgA, and the IgM classes (2–4). Anti-lymphocytic antibodies of the IgM class are especially efficient at mediating complement-dependent lymphocytotoxicity (5, 6). In general, IgG antibodies tend to be inefficient in assays employing CDC or immunofluorescence with live cells. However, IgG antibodies can be detected after elution from living cells, by their ability to inhibit mixed lymphocyte reactions (7), or by immunofluorescent techniques using fixed cells (2, 4, 8).

Recently there has been great interest in the regulation of the immune response, particularly with regard to functional activities of particular T cell subsets (9–15). Some investigators have proposed that anti-T cell antibodies could be responsible for decreased function of suppressor cells (16–18). New Zealand mice with a

1 Abbreviations used in this paper: ADCC, antibody-dependent cell-mediated cytotoxicity; CDC, complement-dependent cytotoxicity; E/T, effector-to-target cells; HBSS, Hanks’ balanced salt solution; SLE, systemic lupus erythematosus; SRBC, sheep erythrocytes.
lupus-like syndrome have antibodies to T cells and decreased T cell functions, and have also been reported to have a loss of T suppressor function (19–21). Sakane et al. (22) showed that IgM anti-lymphocyte antibodies from patients with active SLE, in the presence of complement, preferentially killed suppressor cell precursors; therefore, the loss of suppressor T cells in patients with SLE may (in part, or entirely) be the result of the effects of such antibody activity.

Although much information now exists regarding T cell damage by antibody and complement in patients with SLE, the role of a mechanism involving antibody in the absence of complement, antibody-dependent cell-mediated cytotoxicity (ADCC), has been less carefully explored. In this paper, we have therefore asked whether anti-T cell antibodies in the plasma of patients with SLE were capable of exerting such cytotoxic effects on normal T cells as well as on their own T cells. We found that most plasma of patients with active SLE had anti-T cell antibodies, which, in the absence of complement, could mediate T cell damage in the presence of mononuclear cells.

METHODS

Plasma of the patients with SLE. All patients met the preliminary criteria proposed by the American Rheumatism Association for the diagnosis of SLE. Clinical activity was assessed at the time of blood drawing by two physicians on the basis of signs and symptoms (active rash, serositis, arthritis, active central nervous system disease, and active renal disease). Patients lacking these symptoms or detectable signs of activity were categorized as inactive. The active patients in this study had at least two of the above criteria of activity. Moreover, all active patients had large quantities of anti-DNA antibodies.

At the time blood was obtained for these studies, patients with active disease had not yet received any treatment. In some instances blood was also drawn from these same patients at a later time when their disease had become inactive. Inactive patients had previously been treated with corticosteroids and occasionally with azathioprime, but were not receiving such treatment at the time of study. In addition, such patients had little detectable anti-DNA antibodies.

All the SLE plasma used had been frozen once and had not been previously thawed. Normal fresh frozen plasma was obtained from healthy adults. All plasma was heated to 56°C for 30 min to inactivate complement and then centrifuged at 105,000 g for 2 h at 4°C before use to remove aggregated materials.

Effector cells of cell-mediated cytotoxicity and target T cells. Peripheral blood from healthy human donors (obtained from the Blood Bank Department, Clinical Center, National Institutes of Health, Bethesda, Md.) or from patients with SLE was anticoagulated with heparinized saline (10 U/ml blood). The blood was then diluted with an equal volume of Hanks’ balanced salt solution (HBSS) and 40 ml of the diluted blood was layered over 10 ml of Ficoll-Hypaque solution (Ficoll-Hypaque, Pharmacia Fine Chemicals, Div. of Pharmacia Inc., Piscataway, N. J.) and centrifuged at 4°C for 30 min at 400 g. The mononuclear cells were then removed from the interface and washed three times in HBSS. These cells were resuspended in the complete medium, RPMI-1640 (Gibco Laboratories, Grand Island Biological Co., Grand Island, N. Y.) supplemented with 10% heat-inactivated fetal calf serum (Microbiological Associates, Walkersville, Md). The T cells and non-T cells were prepared as described (23). In brief, neuraminidase-treated sheep erythrocytes (SRBC) and mononuclear cells were mixed in a ratio of 40:1 in HBSS supplemented with 25% fetal calf serum. This mixture was incubated at 37°C for 15 min, spun at 200 g for 5 min, and incubated on ice for 1 h. The SRBC-lymphocyte suspension was then layered on Ficoll-Hypaque, the SRBC rosettes were pelleted at 400 g for 20 min at 4°C, and the rosetted lymphocytes in the pellet fraction were removed and further purified by rosette formation, which was accomplished by the addition of a second portion of neuraminidase-treated SRBC, and subsequent centrifugation. The rosetted lymphocytes in the pellet from the final Ficoll-Hypaque gradient were suspended in 0.83% ice-cold ammonium chloride-0.17 M Tris buffer, pH 7.2, to remove SRBC by hypotonic lysis, centrifuged, washed twice in HBSS, and then resuspended in the complete culture medium. We will refer to this preparation as T lymphocytes. The T cell fraction contained >98% rosette-positive cells and <1% Ig+ cells. The interface cells, obtained after the second cycle of the T cell purification procedure used above, were designated non-T cells. These contained less than 1% E rosetting cells, 50–60% surface immunoglobulin-positive cells, and 20–30% monocytes as judged by Giemsa staining. These non-T cells were then further separated into adherent and nonadherent populations by overnight incubation on plastic petri dishes at 37°C. The adherent cells were then obtained by removal with a rubber policeman. These adherent cells consisted of 95% monocytes as judged by Giemsa staining; 70% of the nonadherent cells stained for surface immunoglobulin.

T cells were used as the target cells for ADCC, unfractionated lymphocytes, T cells, and non-T cells (unseparated, adherent, and nonadherent) were all tested for their ability to act as effector cells for ADCC. The unfractionated lymphocytes gave the greatest ADCC (Fig. 1) and were therefore used as effector cells for the rest of the experiments described in this paper.

Preparation of IgG and IgM fractions of plasma by Sephadex G-200 column chromatography. Plasma was precipitated with 50% ammonium sulfate, dialyzed against buffer, and applied to a 1.5-m long Sephadex G-200 (Pharmacia Fine Chemicals Inc.) column. Individual fractions were collected, and the optical density at 280 nm of each fraction measured in a
spectrophotometer. Marker proteins were run to confirm the approximate size of molecules obtained from the resulting peaks. A good separation of IgM and IgG peaks was observed. The purity of each fraction was confirmed by radial immunodiffusion of 20-fold concentrated samples using specific antibodies to human IgG or human IgM. To check further their purity, the peaks were separately rechromatographed; no IgG was found in the IgM fraction, and no IgM was found in the IgG fraction. Thus, no IgM-IgG or IgG-IgM complexes were present in the IgM fraction because IgG was not detectable in this fraction. The IgM and IgG peaks were separately pooled and concentrated. Individual fractions were dialyzed overnight against phosphate-buffered saline, pH 7.2, and used in the ADCC assays (see below).

Detection of anti-T cell antibodies by CDC using the dye exclusion test. 10^7 T cells were suspended in 1 ml of the medium supplemented with 40% SLE plasma, fractionated plasma, or normal plasma and left at 4°C overnight. 200 ml of fresh AB serum was then added as a complement source, and the cells were cultured an additional 2 h at 37°C. The total number of cells and the percentage of viable cells (dye exclusion) were then determined using trypan blue at a final concentration of 0.05%.%

% cell killing

\[
\text{number of viable T cells after treatment} = 1 - \frac{\text{with plasma plus complement}}{\text{with viable T cells after treatment}} \times 100.
\]

ADCC assay. ADCC was performed by a modification of previously published methods (24–26). 10 \times 10^6 purified target T cells obtained from normal donors or patients with SLE were cultured at 4°C overnight in the presence of 40% normal plasma or SLE plasma. Effector and target T cells were always from the same donor except for some of the experiments shown in Fig. 8. Plasma was either from the same donor as the cell donor or from other individuals. In some cases the target cells were incubated as above with the IgG or IgM fraction of plasma. Here, the IgG and IgM fractions were diluted to be equivalent to the concentrations of IgG and IgM that would be present in 40% plasma. In the studies in which the IgG and IgM fractions were used to treat the target T cells, 10% heat-inactivated fetal calf serum was also added to the overnight cultures.

The plasma or IgG or IgM-treated T cells were washed one time and then incubated at 37°C for 1 h with 100 μCi/0.1 ml ^51Cr)sodium chromate (New England Nuclear, Boston, Mass.) and washed three times with HBSS. In some instances, the IgG-treated target cells were not radiolabeled and were used in cold target inhibition studies (Results). Autologous unfractonated mononuclear effector cells, 2.5 \times 10^6, were mixed with 1 \times 10^6 of the above ^51Cr-labeled T cells in wells of a Falcon V-bottom plate in a total volume of 200 μl in RPMI-1640 supplemented with 10% heat-inactivated fetal calf serum. The plate was incubated for 6 h at 37°C in a humidified atmosphere containing 5% CO2. Although the effector-to-target cell (E/T) ratio usually used was 25:1, in some experiments this ratio varied. This method is called the "pre-culture method."

In some experiments the T cells were first ^51Cr-labeled, and then the SLE or normal plasma was added together with the effector cells; the plasma was present continuously (at a concentration of 40%) during the 6-h cytotoxic assay. This is called the "co-culture method."

The percent cytotoxicity was calculated as follows: at the end of the incubation period, the plates were centrifuged at 300 g for 10 min, a 100-μl portion of the supernate was removed from each well, and its radioactivity was counted in a gamma spectrometer. Maximum possible ^51Cr release was determined by three freeze-thaw cycles. The spontaneous release from wells without effector cells was also determined:

\[
\text{% cytotoxicity} = \frac{\text{counts per minute test sample} - \text{spontaneous release} \times 100}{\text{maximum release} - \text{spontaneous release}}
\]

All experiments were done in triplicate. Statistical analysis was performed with Student's t test.

Detection of anti-T cell antibodies by CDC using ^51Cr release. To determine the magnitude of CDC on these same populations of plasma-treated T cells, portions of these same ^51Cr-labeled T cells in microtiter plates were cultured for 6 h at 37°C with 100 μl of 33% fresh normal AB sera (as complement source) instead of the effector cells. It should be noted that here the T cells were first incubated with the SLE plasma and then washed exactly as in the pre-culture method for the ADCC (see above). The percent cytotoxicity was calculated using the same formula that was used to calculate the percent cytotoxicity for ADCC.

Preparation of aggregated human IgG and its inhibition of ADCC. Cohn Fraction II of human IgG (Miles Laboratories, Inc., Elkhart, Ind.) was dissolved in phosphate-buffered saline at the concentration of 20 mg/ml and heated at 63°C for 15 min. The aggregates were homogenized in phosphate-buffered saline and centrifuged at 1,000 g for 30 min. The supernate was collected, and the protein concentration was measured by optical density (280 nm) using a spectrophotometer. These finer aggregates of IgG were then diluted in complete culture medium. The effector cells were either untreated or treated with various concentrations of the aggregates for 30 min and not washed. They were then added to the pre-culture ADCC assay as described above, and the percent cytotoxicity was measured.

RESULTS

Comparison of pre-culture and co-culture ADCC methods. We compared two methods of ADCC using highly purified T cells from normal individuals as targets and their own unfractonated lymphocytes as effector cells. In the pre-culture method the plasma from four patients with active SLE showed 17.1±2% (mean±SE) cytotoxicity at a 25:1 E/T ratio (Fig. 2). In contrast, if the target cells were labeled with ^51Cr)sodium chromate first and then incubated with the same SLE plasma and effector cells (co-culture), only 3.6±0.9% (mean±SE) cytotoxicity was observed (P < 0.01 vs. pre-culture method). These results demonstrated that when SLE plasma was continuously present during the entire culture period ADCC was inhibited.

To analyze why this result was observed, we incubated unfractonated effector cells overnight at 4°C with a concentration of purified IgG equal to that which would be present in 40% plasma (see above). These cells were then washed three times and used in the ADCC assay at various E/T ratios. As shown in Fig. 3, such treatment of the effector cells markedly diminished their capacity to mediate ADCC. One explanation for
this result is that “cold” T cells in the effector population become coated with anti-T cell antibodies and inhibit by a cold target mechanism. To determine directly whether this was the case, the following experiment was performed. Here various numbers of T cells, preincubated with IgG fraction of SLE plasma but which were not radiolabeled, were added to the standard ADCC system. T cells not treated with the IgG fraction of SLE plasma were used as a control. As shown in Fig. 4 the addition of IgG-treated cold T cells markedly inhibited the ADCC. For this reason all further experiments were performed using the preinculture method in which the T cells were first treated with SLE plasma and then washed.

Comparison of ADCC and CDC using SLE plasma. 14 normal plasma, 8 plasma from inactive SLE patients, and 9 plasma from active SLE patients were tested for ADCC activity. The percent cytotoxicity (mean±SE) for these three groups was 1.4±0.4, 2.6±0.5, and 11.6±2.0, respectively. The active SLE group was significantly different from the other two groups, P < 0.01. The upper limit (the mean ±2 SD) of normal subjects, 4.4%, was used as the upper limit of normal. By this criterion, eight of nine active SLE plasma showed significant ADCC (Fig. 5A). In contrast, all eight inactive SLE plasma tested for ADCC activity produced cytotoxicity levels that were within normal limits. These results demonstrate that normal T lymphocytes can be destroyed by autologous unfractonated mononuclear cells in the absence of complement when these T cells are pre-treated with the plasma of active SLE patients.

The same labeled target T cells used as targets for ADCC were also used for the simultaneous measurement of CDC (Cr release technique) using some of the above plasma (individually numbered 1–7). The normal and inactive SLE plasma showed no cytotoxicity. However, plasma from the five patients with active SLE studied showed a CDC of 11.2±0.8% cytotoxicity (P < 0.01 compared with the other two groups) (Fig. 5B). When CDC induced by seven of these plasma was measured by dye exclusion techniques, all the plasmas from active patients caused significant cytotoxicity, whereas only one plasma from an inactive SLE patient showed cytotoxicity (Fig. 5C). Thus, those sera numbered 1–6 that demonstrated anti-T cell antibodies by the classical technique (dye exclusion) were the same that were
able to mediate ADCC. However, their rank order of efficiency in the different assays varied somewhat. It should also be noted that plasma 7, which was able to mediate CDC by dye exclusion (Fig. 5C), was unable to mediate ADCC (Fig. 5A). The dye exclusion technique may give a somewhat greater degree of cytotoxicity than the $^{51}$Cr release technique because the procedure of $^{51}$Cr labeling and washing of T cells that had been treated with SLE plasma may have killed some of the more sensitive cells; therefore, they may not be present during the actual 6-h $^{51}$Cr release assay.

**Comparison of ADCC mediated by active plasma and inactive plasma from the same patient with SLE.** ADCC activity of three pairs of plasma from patients with active SLE was compared with the ADCC activity using plasma from the same patients when they were inactive (Fig. 6). A fourth pair of plasma came from identical twins discordant with regard to disease activity. These results revealed that plasma of SLE patients supported significant ADCC activity when obtained in the active phase but not in the inactive phase of disease.

**Reproducibility of ADCC with different T cell donors.** Three active SLE plasma were studied for their ADCC activity against T lymphocytes from three unrelated normal subjects. As shown in Table I, these

![Figure 5](image5.png)

**Figure 5** Using purified T cells as targets, the percent cytotoxicity on purified T cells by either ADCC (A), or complement-mediated cytotoxicity by $^{51}$Cr release (B), or complement-mediated cytotoxicity by dye exclusion test (C) is shown. The individually numbered values show the effects of the same plasma in the three different assays. Vertical bars show the mean±SE.

![Figure 6](image6.png)

**Figure 6** The percent cytotoxicity by ADCC using three plasmas from patients obtained during the inactive phase of disease and during the active phase of disease. The starred values (*) from plasma obtained simultaneously from a set of identical twins, one of whom was active and one of whom was inactive.

![Table I](table1.png)

**Table I**

Reproducibility of ADCC using Different Lymphocytes as Targets

<table>
<thead>
<tr>
<th>SLE plasma</th>
<th>Blood donors (normal)</th>
<th>Mean±SE</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A</td>
<td>B</td>
</tr>
<tr>
<td>3*</td>
<td>12.8±1</td>
<td>14.0</td>
</tr>
<tr>
<td>4</td>
<td>8.8</td>
<td>16.0</td>
</tr>
<tr>
<td>6</td>
<td>16.8</td>
<td>14.4</td>
</tr>
</tbody>
</table>

* Patient numbers correspond with the ones shown in Fig. 5. † Values are percent cytotoxicity. ND, not done. No normal plasma showed more than 4.3% of cytotoxicity against any target cells.

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three plasma mediated significantly higher ADCC against all three normal target cells than did plasma from normal subjects. These results suggest that active SLE plasma act on T cells from any subjects and are not restricted to T cells bearing any particular human leukocyte antigen (HLA) type.

**Specificity and antibody nature of the factor in the SLE plasma leading to cell-mediated lysis.** IgG has been implicated as the major antibody class involved in ADCC, though several investigators reported that IgM antibody could also induce ADCC (27, 28). To determine whether the effective fraction of SLE plasma that mediated ADCC was an antibody of the IgG or IgM class, the IgG and IgM fractions of three patients with SLE and one normal individual were obtained by gel filtration and used for ADCC. First, the plasma and plasma fractions were tested for anti-T cell antibody by dye exclusion (Table II). Patient I was active and had anti-T cell antibodies of the IgM class as measured by CDC. Patient II was mildly active, and patient III had inactive SLE. No antibodies were present in the plasma of these latter two patients or the normal individual by this method of CDC.

Fig. 7 shows the results of ADCC using unfractionated and fractionated plasma from these three patients. In patient I, higher ADCC activity was found in the IgG fraction than in unfractionated plasma; the IgM fraction showed minimal ADCC activity despite containing most of the antibody active in CDC. In patients II and III, ADCC activity was shown only with the IgG fraction, and not in unfractionated plasma. Neither the IgG nor the IgM fraction from plasma from a normal individual could mediate ADCC. These results suggest that IgG was the major antibody class effective in our cell-mediated assay of ADCC.

**The use of effector cells and target cells from patients with SLE.** To help establish the relevance of an ADCC mechanism to the lymphopenia and immune dysfunction in vivo patients with SLE, we thought it important to demonstrate that cells from SLE patients could also mediate this effect. In Fig. 8 we show that effector cells from patients with SLE could mediate ADCC against their own target T cells. A dose-dependent increase in cytotoxicity, characteristic of ADCC, was also observed. Moreover, in this experiment only plasma from patients with active SLE resulted in maximal ADCC.

**Inhibition of lymphocyte-mediated cytotoxicity with aggregated IgG.** Immune complexes and aggregated IgG are known to suppress the ADCC reaction (29, 30). To determine whether our cell-mediated cytotoxicity was also inhibited by aggregated IgG, several concentrations of aggregated IgG were added to our system. As shown in Fig. 9, ADCC activities observed in two active SLE plasma were inhibited in a dose-response fashion by aggregated IgG.

**TABLE II**

<table>
<thead>
<tr>
<th>Patients</th>
<th>Unfractionated plasma</th>
<th>IgG fraction</th>
<th>IgM fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>I* (active)</td>
<td>271</td>
<td>7</td>
<td>34</td>
</tr>
<tr>
<td>II (mildly active)</td>
<td>7</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td>III (inactive)</td>
<td>8</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Normal</td>
<td>2</td>
<td>3</td>
<td>0</td>
</tr>
</tbody>
</table>

* Patient numbers correspond with the ones shown in Fig. 7. Values are percent killing. The means of the underlined values are significantly greater than the percent killing observed with normal plasma (4.4±1%).

**Figure 7** The percent cytotoxicity by ADCC of unfractionated plasma from SLE patients and a normal individual and the IgG and IgM fractions of these plasma.
Although the reduction includes both B and T cells (4, 31, 32), B cell function remains hyperactive, whereas there is often an impairment of certain T cell functions. These T cell defects are most prominent in patients with active disease, and are minimal in patients with inactive disease (4). Moreover, it has been suggested that the excessive B cell activity may stem (at least in part) from reduced T cell suppressor function; a role for anti-T cell antibody in the T suppressor defect has been postulated (16–18).

Because autoantibodies to T cells were found in patients with SLE (2–6) as well as in New Zealand mice with a lupus-like syndrome (19–21), their participation in the reduction of T cell numbers and the impairment of T cell functions have been actively investigated (7, 14–18, 33, 34). The proposed mechanism for both of these effects has most often been considered to be complement-mediated cytotoxicity. Antibodies of the IgM class are thought to be principally involved in this cytotoxicity, and are capable of eliminating suppressor T cell function (14–18). However, IgG antibodies also occur in patients with SLE; these antibodies have also been reported to be able to impair certain lymphocyte functions, such as the mixed lymphocyte reaction and ADCC (7, 33, 34).

In this report we now demonstrate that T cell damage by plasma of patients with active SLE may also be mediated by a complement-independent mechanism, namely ADCC. ADCC was observed using both normal and SLE lymphocytes as target and effector cells. Furthermore, this ADCC was observed using the plasma, effector cells, and target cells from the same individual with SLE. Such an observation strongly suggests that such an ADCC mechanism may be one of the explanations for the lymphopenia and immune dysfunctions observed in patients with SLE. Thus, our results are similar to previous studies of various diseases or disease models in which specific naturally occurring antibodies injured target cells bearing specific antigen by cooperation with mononuclear cells (35–39). These examples include antibodies to tumor cells (35), erythrocytes (36), and viruses (37–39).

Moreover, anti-DNA antibodies in the serum of patients with SLE have been shown to act as specific antibodies using a model target cell, cell-mediated lysis of DNA-coated cells (40, 41). In addition, Glinski et al. (42) observed a reduction in a subpopulation of T cells in patients with SLE; an IgG fraction of SLE sera could also induce, in vitro, a reduction of this same fraction in normal lymphocyte populations. Because the reaction was inhibited by heat-aggregated human gamma globulin, they speculated that ADCC might have been the mechanism involved (42).

This speculation is now confirmed by our results. Indeed, the IgG fraction of sera from patients with
SLE, but not from a normal individual, were demonstrated to be capable of killing purified normal T lymphocytes by an ADCC mechanism. An additional point is that the specificity of this reaction and the role of IgG are shown by the fact that it could be blocked by cold T cells coated with the IgG fraction from SLE plasma (Fig. 4). It is also unlikely that this activity of plasma from patients with SLE is a result of the presence of specific isohemagglutins, or anti-HLA antibodies, because a single plasma could induce ADCC in populations of lymphocytes from several different unrelated individuals. Anti-T cell antibodies themselves, in the plasma of patients with SLE, are most likely to be the effector antibodies in the ADCC we have observed.

Although there have been some reports that antibodies of the IgM class are capable of mediating ADCC (27, 28), we observed that only the IgG fractions of plasma of patients with SLE were able to exert significant ADCC (Fig. 7). This is consistent with the finding of Longmire et al. (43), who found that an IgG fraction from supernates of cultured spleen lymphocytes from patients with Hodgkin’s disease could also mediate ADCC against autologous T lymphocytes. We assume that most patients with active SLE have both IgM and IgG anti-T cell antibodies. However, plasma from patient 7 in Fig. 5 demonstrated a high level of CDC and no ADCC activity. In this patient it is possible that only anti-T cell antibodies of the IgM class were present.

In the course of our studies we observed cases in which unfractionated plasma from patients with SLE could not mediate ADCC but in which the IgG fraction could. The presence of inhibiting immune complexes in the unfractionated plasma might explain these observations. The possibility is supported by the finding that aggregated IgG could inhibit ADCC by whole plasma. In addition, our observation that the ADCC assay performed in the continuous presence of SLE plasma (co-culture method) failed to show high levels of ADCC, whereas preincubation of the target cells with plasma followed by washing (preculture method) produced higher levels of ADCC, might also be explained by the presence of immune complexes in some of these plasma. In this regard, several other investigators have noted that SLE sera also suppressed ADCC activity of normal human peripheral blood mononuclear cells against Chang cells or chicken erythrocytes (34, 44–47). Another reason for the failure of the co-culture method is that unlabeled T cells present in the effector cell population could have interacted with the anti-T cell antibody in the SLE plasma; here “cold target inhibition” could have occurred, and, thus, could have reduced the degree of ADCC observed. Indeed the results of the experiments shown in Figs. 3 and 4 support these ideas. In addition, antibodies to the effector cells of ADCC may have also inhibited the reaction when plasma was co-cultured throughout the ADCC reaction.

Of interest was the fact that the IgG fraction of the plasma of patients II and III (Table II and Fig. 7) with inactive disease could mediate ADCC in the absence of demonstrable complement-mediated cytotoxicity. Thus, ADCC activity may be a very sensitive method to detect anti-T cell antibodies in patients with inactive disease and with low levels of anti-T cell antibodies.

We have not yet studied functional impairments that might be produced by a cytotoxic ADCC mechanism. However, a previous study has suggested that a subpopulation of T cells may be removed selectively by the ADCC mechanism (42). Moreover, Twomey et al. (16), using unfractionated plasma, and Sagawa and Abdou (18), using the IgG fraction of SLE sera, could, in the absence of complement, inhibit the development of complement-mediated ADCC in the absence of temperatures from 4°C to 15°C (14), ADCC appears to work effectively at higher temperatures. Thus, it is possible that at body temperature ADCC may be an important mechanism by which regulatory T cell subsets are deleted in patients with SLE.

ACKNOWLEDGMENTS

The authors are grateful to Doctors Paul V. Holland, Richard Davey, and Harvey Klein, and Ms. Jane E. Kendall, Blood Bank Department, Clinical Center, National Institutes of Health, for their help and cooperation in supplying the blood from normal humans in these studies. We also wish to thank Mr. Charles Hoes for his excellent technical assistance, and Mr. J. Patton Reeves for separating the SLE plasma into IgG and IgM fractions.

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