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Abstract

Human antiidiotypic antibodies to anti-DNA antibodies can be separated into at least two categories based on their binding to anti-DNA, antiidiotypic antibodies, and antigens. One type was found mainly in inactive stage of SLE. The antiidiotypic antibodies appear to be directed towards idiotypic (Id) determinants in the antigen-binding sites of anti-DNA antibodies. Antibody from patient T.K. acted like a mirror image of anti-single-stranded DNA antibodies, O-81, as determined by a competitive inhibition RIA. Antibodies from patient S.U. also seemed to be Ab 2γ and Ab 2γ to anti-double-stranded(ds) DNA antibodies, NE-1. Most of normal subjects, on the other hand, had antibodies that bound to the human monoclonal anti-ds DNA antibodies, NE-1, NE-13, 7F4, and O-81. The Id-anti-Id interaction was not inhibited by the addition of DNA. Thus, normal subjects had Ab2α activity that recognizes Id determinants in the framework region common among anti-DNA antibodies, whereas antiidiotypic antibodies in most SLE sera appear to show Ab 2γ and Ab 2γ activity. The results provide evidence that the Id network system regulates immunological tolerance to DNA in humans.

Introduction

Antiidiotypic antibodies were originally described by Oudin and Michel (1) and Kunkel et al. (2), and now are well known to play an essential regulatory role in immune network systems (3–5). Idiotype(Id) immunity is thought to be an important mechanism for the maintenance of immunological tolerance to self-antigens and autoimmunity might be attributed to the disturbance of down-regulation of autoreactive clones in an Id network (6–8). Experimentally induced anti-Id antibodies have been shown to bind to Id determinants of autoantibodies (9–12). Most recognize a unique Id of autoantibodies, such as anti-DNA antibodies, anticytotoxic T lymphocyte receptor antibodies, antithyroglobulin antibodies or rheumatoid factor. However, some appear to be directed to Id determinants common to different kinds of autoantibodies (13, 14). Clinical investigations have revealed the presence of anti-Id autoantibodies to self-reacting antibodies in human (15–17). It is interesting that the Id of anti-DNA antibodies was found in active stage of systemic lupus erythematosus (SLE), whereas the anti-Id antibodies occur spontaneously in the inactive stage (15). Anti-Id antibodies act in immune system by recognizing the complementary Id on the surface of immune competent cells (4, 5, 18). When administered in vivo, experimentally induced monoclonal anti-Id antibodies to anti-DNA antibodies could induce a transient suppression of anti-DNA antibody production in NZB/WFl mice (19), but failed in MRL/lpr mice (20). Thus, elucidating the properties of anti-Id antibodies could be important not only for understanding the pathophysiological role of Id-anti-Id interaction in autoimmunity, but also for manipulating autoantibody production in autoimmune diseases (21). However, it has not yet been established whether human anti-Id autoantibodies recognize a private Id or a cross-reactive Id of autoantibodies. We also do not know if human anti-id are directed against antigen-binding sites (paraprote) or the framework region of antibody molecules.

In order to shed light upon the above-mentioned problems, we have studied the interaction between Id and anti-Id using human monoclonal anti-DNA antibodies and murine monoclonal anti-Id antibodies (22, 23) and demonstrated that there are heterogeneous populations of anti-Id antibodies to anti-DNA antibodies in human (23a). In the present study, we demonstrate that anti-Id antibodies with different Id-recognizing properties occur in human SLE and healthy subjects, indicating that the Id network system regulates human self-tolerance.

Methods

Reagents. Calf thymus DNA was purchased from Worthington Biochemicals (Freehold, NJ). DNase I was from Sigma Chemical Co. (St. Louis, MO) and polynucleotides and S, nuclease from Miles Laboratories (St. Louis, MO). DNA obtained commercially was purified further as described previously (24) and used as double-stranded(ds)DNA or single-stranded(ss)DNA.

Sera and monoclonal antibodies. Sera were obtained from 40 female and 3 male patients with SLE, or 11 female and 4 male healthy subjects. Immunoglobulins (Ig) were precipitated twice with 33% saturated ammonium sulfate and isolated by DEAE-cellulose chromatography. Some were further prepared as F(ab’), fragments. Each sample was pretreated with DNase I (Sigma Chemical Co.) and absorbed with pooled human IgG- and IgM-coupled-Sepharose 4B before use. Some were then passed through DNA-coupled Sepharose 4B affinity column (25).

Human monoclonal IgM anti-ssDNA antibodies (O-81) or IgM anti-dsDNA antibodies (NE-1 and NE-13) were obtained from EBV-transformed B cell clones (26). Another IgM anti-dsDNA antibody (7F4) was from a human-human hybridoma using 6TG resistant B lymphoblastoid cell line G K-5 (a gift from Dr. Sato, National Institutes of health, Bethesda, MD) and lymphocytes from healthy subjects. Each anti-DNA antibody was purified using affinity chromatography of DNA-coupled Sepharose 4B (25). Murine IgG1 monoclonal anti-Id antibodies to O-81 or NE-1 were produced from hybridomas (21) and designated D1E2 or 1F5, respectively. The anti-Id antibodies were
purified twice using a DEAE-cellulose column and further prepared as F(ab')2 fragments. The characteristics of these monoclonal antibodies are shown in Table I.

**Measurement of anti-DNA antibodies and anti-Id activity to anti-DNA antibodies.** Anti-DNA antibody activity was measured mainly with a modified solid-phase RIA as described in other papers (27).

Anti-Id antibodies were measured by direct binding to anti-DNA antibodies using solid phase RIA. Each well was coated with 100 μl of affinity-purified anti-DNA antibodies (O-81, NE-1, NE-13, or 7F4) or with pooled human IgM (2 to 5 μg/ml) in 0.15 M PBS, pH 7.2, and was incubated for 2 h at room temperature. After being washed in PBS containing 0.2% BSA and 0.05% Tween 20 (washing buffer), the wells were covered with 1% BSA in PBS for 60 min at room temperature. Then, 1,500 diluted test samples were added to the wells and incubated at 37°C for 90 min. After being washed, 125I-labeled anti-human γ rabbit IgG F(ab')2 was added and incubated for an additional 2 h at room temperature. Bound radioactivity was determined using a γ-counter.

Anti-Id activity was also determined by showing that antibodies could block the binding of monoclonal anti-DNA antibodies to monoclonal anti-Id antibodies. First, 100 μl of 1:20 diluted sample, which was pretreated with DNase I followed by the absorption of pooled IgM-coated Sepharose before use, was mixed and settled with 100 μl (2 x 10^6 cpm) of 125I-labeled anti-DNA antibody (O-81 or NE-1) in a plastic tube at room temperature. After 90 min incubation, 100 μl of 1:500 diluted anti-Id antibodies (D1E2 or 1F5) was added and incubated for 90 min. 200 μl of anti-mouse γ rabbit IgG (F(ab')2) were added and tubes were incubated for another 90 min. After centrifugation, the radioactivity of the precipitates was measured in a γ-counter.

**Isoelectric focusing and immunoblotting.** Flat bed isoelectric focusing (IEF) was performed in an apparatus (Pharmacia Fine Chemicals, Piscataway, NJ) at 4°C. 5% polyacrylamide gels (10 x 20 x 0.1) were made using 0.1% Tween 20. 20-μl samples of IgG were applied to the gel surface using Whatman No. 1 filter paper wicks, and focused at constant power (30 W) to a maximum of 3,000 V at 4°C for 2 h. The pH gradient was measured using a flat membrane pH electrode or IEF calibration kit. The focused gel was carefully removed from the plate, placed in a gel transfer holder, and blotted electrophoretically on to a nitrocellulose membrane at (1.45 μm) in transfer buffer (0.7% acetic acid) for 30 min at 30 V and 30 min at 150 V. After blotting, free binding sites on the nitrocellulose membrane were blocked by incubating for 1 h at 37°C on a rocker platform in 100 ml of 1% BSA in PBS containing 0.1% Tween 20. The membranes were then rinsed three times with PBS containing 1% BSA and 0.1% Tween 20 (washing buffer) and then overlaid with 125I-human monoclonal anti-DNA antibodies, O-81 or NE-1, or IgM for 90 min at room temperature with rocking. Last, the membranes were rinsed overnight with several changes of washing buffer of the rocker at room temperature. The membranes were dried and exposed to Kodak X-Omat x-ray film at -70°C.

**Competitive inhibition RIA.** Competitive inhibition was determined by measuring inhibition of direct binding of anti-Id antibodies to monoclonal anti-DNA antibodies using a solid-phase RIA. Prior to inhibition, each sample was diluted to the middle point of the maximum binding to O-81, NE-1, or 7F4. A mixture of the test sample and 50 μl of inhibitors at varying dilutions was incubated for 90 min in wells coated with monoclonal anti-DNA antibodies at 37°C. Results were expressed as percentage inhibition of anti-DNA antibody-binding activity.

**Results**

Evidence for the direct binding ability of human anti-Id antibodies to anti-DNA antibodies. Anti-Id autoantibodies in human were first tested by solid phase RIA after sequential absorption of the sample on columns containing pooled human IgG and IgM. Fig. 1 shows the binding of sera from active and inactive SLE and healthy subjects. About 60% of sera from subjects with inactive SLE show specific binding to human monoclonal anti-DNA antibodies. 20–30% of active SLE and control sera also showed the binding ability to O-81 and NE-1. Binding of these antibodies is specific for Ig of monoclonal anti-DNA antibodies because the binding to O-81, NE-1, NE-13, or 7F4 is apparently higher than those to irrelevant antibodies IgM. The specificity of binding is also demonstrated in Fig. 2, where preincubation with DIE2 or 1F5

<table>
<thead>
<tr>
<th>Antibodies</th>
<th>Source</th>
<th>Ig class</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>O-81</td>
<td>EBV-transformed B cell clone</td>
<td>Human IgM</td>
<td>Anti-ssDNA, poly(dT)</td>
</tr>
<tr>
<td>NE-1</td>
<td>EBV-transformed B cell clone</td>
<td>Human IgM</td>
<td>Anti-ds/ssDNA</td>
</tr>
<tr>
<td>NE-13</td>
<td>EBV-transformed B cell clone</td>
<td>Human IgM</td>
<td>Anti-ds/ssDNA</td>
</tr>
<tr>
<td>7F4</td>
<td>Human-Human hybridoma</td>
<td>Human IgM</td>
<td>Anti-ds/ssDNA</td>
</tr>
<tr>
<td>D1E2</td>
<td>Murine hybridoma</td>
<td>Murine IgGl</td>
<td>Anti-idiotype Ab to O-81</td>
</tr>
<tr>
<td>1F5</td>
<td>Murine hybridoma</td>
<td>Murine IgGl</td>
<td>Anti-idiotype Ab to NE-1</td>
</tr>
</tbody>
</table>

*Figure 1. Direct binding of human anti-Id antibodies to human monoclonal anti-DNA antibodies, O-81 (left) and NE-1 (right). Each sample was incubated in wells coated with O-81, NE-1 or pooled IgM for 90 min at room temperature. After being washed with buffer, the ability to bind to 125I-labeled anti-human rabbit IgG F(ab')2 was determined as described in Methods. Data are expressed as total binding to O-81 or NE-1 minus that of IgM (the range of direct binding to IgM is ~200–1,000 cpm). The specific binding to Id determinants was regarded as positive by the inhibition of the samples' binding to Ig-coated wells by the coexistence of free Id(O-81 or NE).*

**Antidiotypic Antibody to Anti-DNA Antibody** 749
Figure 2. Direct binding of human anti-Id antibodies to human monoclonal anti-DNA antibodies. Wells coated with NE-1, NE-13 or O-81 were first incubated with 1:500 diluted DIE2 or IF5 (murine monoclonal anti-Id antibodies) at 37°C for 90 min, washed with the buffer several times and then incubated with 1:100 diluted human sera at room temperature for 90 min. The following procedure was the same as described in Fig. 1. M.M. and S.N. are inactive SLE. E.T. is a healthy male subject.

...tained with anti-O-81 antibodies, whose binding to 125I-O-81 was blocked only by unlabeled O-81 (data not shown).

Inhibition of antigen-antibody reactions by anti-Id antibodies. Next, we showed that small amounts of anti-Id autoantibodies blocked the binding of DNA by human monoclonal anti-DNA antibodies. Samples were pretreated with DNase 1 and IgM-coupled Sepharose and passed through DNA Sepharose to remove any antibodies in the sera. 10 of 14 sera that had shown binding to anti-DNA antibodies markedly inhibited the interaction of 125I-O-81 with ssDNA (Fig. 4) but not 125I-NE-1-binding to dsDNA (Table II). Some inhibited the binding of O-81 to ssDNA as well as that of NE-1 to dsDNA. Comparable amounts of some anti-Id positive sera did not show such inhibition when tested with each reagents (Fig. 4).

Comparison of the incidence of anti-Id antibodies determined by different assays. We expected that anti-Id autoantibodies would also block the binding of monoclonal anti-DNA antibodies to murine monoclonal anti-Id antibodies. Indeed, we found a correlation between inhibition of 125I-O-81 or 125I-NE-1 to murine monoclonal anti-Id, DIE2, or IF5 and direct binding in most of sera. The samples that strongly inhibited the reaction, had also high binding capacity (Fig. 5). Some, however, bound to O-81 or NE-1, but failed to block binding of O-81 to DIE2 or NE-1 to IF5.

The binding specificity of anti-Id autoantibodies. T.K. and T.N. both reacted with O-81 Id determinants as confirmed by direct binding assays. However, only T.K. blocked DNA-binding by O-81 (Table II). T.K. but not T.N. inhibited the interaction between 125I-O-81 and DIE2. Thus, the binding specificity of anti-Id autoantibodies differs. We used a competitive inhibition solid-phase RIA to study further the human autoantibody specificity. Inhibition produced by IgG and an F(ab')2 fragment that had bound directly, was measured by diluting each to a concentration that produced half-maximal

Figure 3. Analysis of anti-Id antibodies by isoelectric focusing and immunoblotting. Inactive SLE samples were focused, blotted to a nitrocellulose membrane, and finally incubated with 125I-NE-1 (A) and 125I-IgM (B) as described in Methods.

Saito, Muryoi, Takai, Tamate, Saito, and Yoshinaga
binding before being mixed with inhibitors. The results of the competitive inhibition studies were shown in Fig. 6 and the tested samples were from patients with inactive SLE, (T.K., N.S., S.S., N.O., N.S., N.T., and S.U.) and from healthy subjects (K.A., O.K., T.N., O.O., E.T., and J.K.). The binding of T.K. to O-81 was apparently inhibited by ssDNA, O-81, and D1E2 but not by, NE-1 and 1F5. N.S. or S.S.-binding to O-81 was also blocked by ssDNA or O-81, but not by anti-Id(D1E2), NE-1, or pooled IgM. However, comparable amounts of DNA or murine monoclonal anti-Id antibodies failed to suppress the interaction between humoral anti-DNA antibodies and serum anti-Id antibodies from most healthy subjects. We noted that the binding of O-81 to the wells was inhibited by the addition of NE-1, NE-13, 7F4, as well as O-81. The binding of all anti-DNA antibodies to wells was also inhibited by NE-1 but pooled IgM was not. To determine what sites were recognized by anti-Id antibodies in T.K. or S.U. sera, various types of polynucleotides were tested for their ability to inhibit the reaction between anti-DNA antibodies and anti-Id antibodies. This was compared with the results obtained using murine monoclonal anti-Id antibodies (Fig. 7). It was demonstrated that T.K. antibodies and D1E2 had similar specificities. The effects of polynucleotides on the interaction between NE-1 and S.U. were also analogous to those seen with 1F5.

**Discussion**

This paper described the binding properties of anti-Id autoantibodies to human anti-DNA antibodies. We used human monoclonal anti-DNA antibodies derived from SLE and healthy subjects because the Id-anti-Id has been difficult to study using polyclonal anti-DNA antibodies (15-17). O-81 reacts with ssDNA but not with dsDNA, while, NE-1, NE-13, and 7F4 bind not only to ssDNA, but also to dsDNA, RNA, or cardiolipin (27). The specificity of NE-1 differs somewhat from the antigen-binding specificity of NE-13 or 7F4. The murine monoclonal anti-Id antibodies D1E2 selectively bind to Id of O-81 and 1F5 recognizes Id determinants in paratope of NE-1 and NE-13 but not that in O-81. D1E2- or 1F5-binding to the corresponding Id was inhibited by the coexistence of DNA antigens (23). When D1E2 or 1F5 coupled with KLH were administered to mice, Ab3 with anti-DNA activity could be elicited in their sera (data not shown). Thus, both anti-Id

**Table II. Comparison of the Incidence of Anti-Id Antibodies Determined by Different Assays**

<table>
<thead>
<tr>
<th>Patient</th>
<th>Disease</th>
<th>Direct binding to*</th>
<th>Percent inhibition of6</th>
<th>Percent inhibition of7</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>O-81</td>
<td>NE-1</td>
<td>IgM</td>
</tr>
<tr>
<td>T.K.</td>
<td>Inactive SLE</td>
<td>2,803</td>
<td>443</td>
<td>625</td>
</tr>
<tr>
<td>T.N.</td>
<td>Healthy</td>
<td>2,514</td>
<td>1,754</td>
<td>576</td>
</tr>
<tr>
<td>N.S.</td>
<td>Inactive SLE</td>
<td>1,346</td>
<td>372</td>
<td>428</td>
</tr>
<tr>
<td>M.M.</td>
<td>Inactive SLE</td>
<td>3,855</td>
<td>2,392</td>
<td>772</td>
</tr>
<tr>
<td>T.S.</td>
<td>Active SLE</td>
<td>576</td>
<td>674</td>
<td>645</td>
</tr>
</tbody>
</table>

Each sample was measured for the anti-Id antibody activity by direct binding tests, inhibition tests to O-81-binding to ssDNA, NE-1-binding to dsDNA, and inhibition assays to O-81-binding to D1E2, NE-1-binding to 1F5 as described in Methods. * Total binding to each monoclonal anti-DNA antibody and pooled human IgM (counts per minute). 6 The percent inhibition was calculated using binding in the presence of 0.2% BSA-PBS as 100% as Figs. 5 and 6.
antibodies seem to behave almost as internal images of the antigens recognized by anti-DNA antibodies. The above mentioned data indicate that these monoclonal antibodies (O-81, NE-1, and NE-13) represent different kinds of Id markers of anti-DNA antibodies. Although we used human monoclonal anti-DNA antibodies, we have evidence that these Id occur in vivo because we have found anti-DNA antibodies with the same Id as O-81 or NE-1 in sera and on the surface of human lymphocytes (Takai, O., manuscript in preparation). Consequently, we feel that the system presented in these experiments represents the in vivo Id-anti-Id interaction.

The binding of serum anti-Id antibodies to human monoclonal anti-DNA antibodies is not due to antiallotypic activity as described in other papers (23a). Women's sera may contain a broad spectrum of antiallotype antibodies. The antibodies from male normal subjects, however, showed similar property of anti-Id antibodies with that of female controls. In addition, the repeated presorption of the sera by pooled IgG and IgM-coupled Sepharose did not change the level of total binding to O-81 or that to NE-1. This was also confirmed by Figs. 2 and 3.

The system used in this report allowed us to compare the Id determinants recognized by the anti-Id autoantibodies. It should be kept in mind that the assay was restricted to detect IgG type antibodies. The results suggest that there are at least two different specificities of anti-Id antibodies. One type was directed toward Id determinants in the antigen-binding sites of antibody molecules. An anti-Id, T.K. specifically bound to O-81, but never to other anti-DNA antibodies or pooled IgM. T.K. also blocked O-81-binding to DNA. Moreover, the binding of O-81 to T.K. could be blocked not only by ssDNA, but also by poly(dT), poly(I), poly(U), which preferentially bind to O-8 (23). T.K. has the same binding properties as D1E2, a murine monoclonal anti-Id antibody raised against O-81. Other anti-Id antibodies had similar reactivities to monoclonal anti-DNA antibodies (Figs. 5–7). These antibodies may belong to the categories of Ab 2β or Ab 2γ described by Bona (27). Another group also reacted with the anti-DNA antibodies. These sera, however, failed to block the binding of anti-DNA antibodies to DNA and O-81- or NE-1-binding to these antibodies could not be blocked by DNA. In addition, comparable amounts of D1E2 or IFS failed to block the Id-anti-Id interaction of these sera. Thus, the anti-Id activity in this group seems to be directed towards the framework regions of anti-DNA antibodies, showing Ab 2α activity (27). It is interesting that

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**Figure 6.** Inhibition of human anti-Id antibody-binding to O-81(A) or to NE-1(B) by DNA, human monoclonal anti-DNA antibodies or murine monoclonal anti-Id antibodies. Anti-Id positive samples were incubated with the indicated concentrations of free DNA (●), O-81 (●), NE-1 (●), 7F4 (●), pooled IgM (●), D1E2 (●) or 1FS (●), in wells coated with O-81(A) or NE-1(B). The binding ability of the samples was determined as described in Methods. Data are expressed as the percentage inhibition of uninhibited binding (1-3 x 10⁸ cpm). T.K., N.S., S.S., N.O., N.S. and N.T. were from SLE patients and K.A., O.K., T.N., O.O., E.T., and J.K. from normal subjects. S.S., J.N., N.T., and J.K. were used as F(ab')₂ fragments.

**Figure 7.** Comparison of murine and human anti-Id antibody-binding to anti-DNA antibodies. D1E2 or inactive SLE IgG(T.K.) was incubated with the indicated concentrations of free polynucleotides in wells coated with O-81. 1FS or other inactive SLE IgG(S.U.) was also incubated with free antigens in wells coated with NE-1. The binding ability of the anti-Id antibodies was determined as Fig. 6.

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Sasaki, Muruyo, Takai, Tamate, Saito, and Yoshinaga
O-81-binding to anti-Id antibodies in some sera was inhibited by NE-1, NE-13, 7F4, as well as O-81, but not by pooled IgM. Similar results were obtained with NE-1, NE-13, and 7F4-binding. These indicate that anti-Id antibodies may recognize the Id determinants in the framework common among these anti-DNA antibodies.

Recently, anti-DNA antibodies have been shown to cross-react with cardiolipin, vimentin, and bacterial polysaccharides (28-32) and to share Id determinants with monoclonal antibodies reactive to Klebsiella and Escherichia coli (33-35). Anti-DNA, anti-gp 70, and other antibodies may constitute a network of idiotypically related antibodies in mice (36). Schwartz and his co-workers have demonstrated the presence of a parallel set of anti-DNA antibodies, which shared Id in the framework of anti-DNA antibodies but failed to bind DNA (36). Thus, there are mounting data that anti-DNA and related Id might occur in the association with bacterial or viral infection (37, 38). In this respect, it is interesting that all dominant Id systems are related to antibacterial antibodies (39). If we agree with these interaction, it is reasonable to speculate that naturally occurring idiotypes may induce the anti-Id antibodies to cross-reactive to anti-DNA antibodies (7). Considering above mentioned viewpoints, it should be noted that anti-Id antibodies with Ab 2a activity to anti-DNA antibodies were detected in healthy subjects. Ab 2a has been demonstrated to be able to function in a manner similar to Ab 2b to suppress the specific immune response in vivo and in vitro (40-42). Diamond and co-workers speculated that Ab 2a to anti-DNA antibodies could regulate autoantibody production in vitro (43). Taken together, these results suggest that anti-Id antibodies (Ab 2a) evoked by the stimulation of the parallel sets might work to regulate anti-DNA production on idiotypic network. This might result in maintaining the homeostasis of the immune response to self-components under normal circumstances. The Id regulatory mechanism may be broken during an acute episode in SLE, although we do not know yet about the factors to contribute to the breakdown of self-tolerance. In this condition, large amounts of anti-DNA autoantibodies occur in the circulation. Consequently, the paratopes of anti-DNA antibodies might be antigenic and selectively activate the restricted clones that have affinity to the antigen-binding sites (paratope) of anti-DNA antibodies. Then, the regulatory mechanism might operate via feedback control by the resulting anti-Id antibodies. This might be the reason why anti-Id antibodies directed against the antigen-binding sites of anti-DNA antibodies occurred in remission. We are currently studying whether or not these anti-Id autoantibodies are able to regulate anti-DNA production in humans.

The finding just mentioned may also have clinical importance. Treatment with anti-Id alters the formation of anti-DNA antibodies, which clearly play an important role in the tissue damage seen in SLE (44, 45). The sequential administration of anti-Id antibodies could suppress the total amounts of circulating autoantibodies to acetylcholine receptor (46) or to DNA (19). In humans, anti-Id antibodies effectively regulate already established spontaneously occurring autoantibodies in vitro (42, 47). We do not know if anti-Id antibodies with sharp specificities would induce efficient immunosuppression of all autoantibodies produced in autoimmune states (20). Our results, using monoclonal antibodies, suggested that a majoring of pathogenic autoantibodies share a small number of Id. This finding may permit a new therapeutic strategy.

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