ANTI-IDIOTYPIC ANTIBODIES TO THE COOMBS ANTIBODY IN NZB F1 MICE*

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The network theory of the immune system proposes that idiotype-anti-idiotype interactions provide important immunoregulatory controls (1). In many experimental systems, the passive administration of heterologous anti-idiotype antibody has been shown to enhance (2) or suppress (3) immune responses. In addition, spontaneous auto-anti-idiotype antibody has been detected after antigenic challenge (4), generally appearing coincident with a decline in antibody titer (5). These findings have suggested that network controls might similarly regulate the formation of pathological autoantibodies (6). Two recent studies support this hypothesis. In one, rabbit anti-(anti-thyroglobulin) idiotype prevented spontaneous autoimmune thyroid disease in Buffalo rats (7). In the other, anti-(anti-DNA)-idiotype was found in the serum of systemic lupus erythematosus (SLE)1 patients when they were in remission (8).

In the current work, we have investigated the SLE mouse strain NZB for anti-(autoantibody)-idiotype antibodies that might play an immunoregulatory role. With age, these mice spontaneously produce IgG anti-erythrocyte autoantibodies which mediate a severe Coombs-positive hemolytic anemia (9). The F1 hybrids of NZB and several normal strains are known to develop a milder hemolytic disease with lower levels of Coombs antibodies (10). We have found that some of these hybrids spontaneously produce high titers of anti-(anti-erythrocyte)-idiotype antibodies that specifically recognize the NZB Coombs idiotypes. This finding suggests that network idiotype-anti-idiotype control might play a protective role in suppressing autoantibody formation in these animals.

Methods

Mice. NZB and CBA/J mice were obtained from The Jackson Laboratory, Bar Harbor, ME. Some NZB mice used in early experiments came from Southwestern Medical School, Dallas, TX. (NZB × CBA)F1 mice were bred in the animal facilities at the University of North Carolina.

Mice were bled from the retro-orbital plexus. To obtain erythrocyte (RBC) suspensions, blood was pipetted immediately into a large excess of Alsever's solution, centrifuged, and washed three times in saline before use. It was necessary to use fresh RBC for hemagglutination assays, as storage at 4°C overnight caused inconsistent results. Serum samples were obtained by

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1 Abbreviations used in this paper: BSA, bovine serum albumin; MRBC, mouse RBC; RBC, erythrocytes; SLE, systemic lupus erythematosus; SRBC, sheep RBC.
allowing blood to clot at room temperature and separating by centrifugation. Serum was stored in aliquots at −20°C before use.

Hemagglutination Assays. For direct Coombs assays, washed RBC from 9–10-mo-old NZB mice were resuspended to 1% in normal saline containing 1% bovine serum albumin (BSA) (Sigma Chemical Co., St. Louis, MO). For indirect hemagglutination assays, a 10% suspension of washed normal mouse RBC (MRBC) from CBA mice was incubated with an equal volume of a 1:10 dilution of NZB serum for 30 min at room temperature with rocking. In both assays, developing sera were diluted in BSA-normal saline in V-bottomed microtiter wells, and an equal volume of 1% RBC suspension was added. Hemagglutination was read after 1 h at room temperature. Titers represent the highest dilution giving ≥2+ agglutination on a 0–4+ scale.

The developing antiserum used for screening for Coombs positivity was rabbit anti-mouse IgG, which was raised by immunization with an IgG myeloma and rendered gamma-specific by absorption with insolubilized IgM and lambda and kappa light chains. Detection of anti-RBC autoantibodies of differing subclasses was performed by using serial dilutions of rabbit antiserum against mouse IgG1, IgG2a, IgG2b, and IgG3, all obtained from Litton Bionetics (Kensington, MD) and found to be monospecific by gel diffusion. (NZB × CBA)F1 hybrid sera were used as developing reagents in anti-idiotype assays.

Hemagglutination Inhibition Assays. For confirmation of specificity, NZB sera were serially diluted and incubated for 30 min at room temperature with selected (CBA × NZB)F1 sera at a concentration fourfold greater than the agglutination endpoint. Appropriate Coombs-positive RBC were then added. After 60 min at room temperature, the hemagglutination inhibition titer was read as the highest NZB serum dilution to completely inhibit agglutination.

Absorptions. Sepharose conjugates of monospecific goat anti-mouse IgG and of anti-trinitrophenylated ovalbumin were kind gifts of Dr. Ellen Vitetta, University of Texas Health Science Center, Dallas, TX. Mouse IgG conjugated to Sepharose was provided by Dr. Stella Robinson (University of Texas Health Science Center). MRBC were coated with mouse IgG (Miles Laboratories, Elkhart, IN), by the chromium chloride technique (11). IgG-coated sheep RBC (SRBC) were prepared by immunizing CBA mice three times intraperitoneally with 0.1 ml of 10% SRBC at 10-d intervals, and incubating the resulting immune serum (which contained high titer IgG anti-SRBC antibody) with SRBC for 30 min at room temperature, followed by extensive washing.

Preparation of Pepsin Fragments of Coombs and Anti-Idiotype Antibodies. 200 μl of each of four NZB indirect-Coombs-positive sera and four corresponding hybrid anti-idiotype sera were precipitated with 45% saturated ammonium sulfate. These partially-purified IgG fractions were digested with 2% pepsin at 37°C for 18 h in 0.5M acetate, pH 4.5. The resultant F(ab′)2 fragments were absorbed with goat anti-mouse IgG (pFc′-specific)-Sepharose 4B. Absence of intact IgG and pFc′ fragments was demonstrated in each case by double immunodiffusion analysis. Fab′ fragments were prepared from these F(ab′)2 preparations by reducing with 0.01 M dithiothreitol and alklylation with a 10% M excess of iodoacetamide.

CBA RBC were coated with Fab and F(ab′)2 fragments of NZB IgG by incubating 10% washed CBA RBC with equal volume of a 1:20 dilution of these preparations (equivalent to a 1:40 dilution of whole serum). Indirect hemagglutination assays were done as described above.

Results

F1 Sera Agglutinate NZB Erythrocytes. We initially observed that sera from certain (NZB × CBA)F1 mice agglutinated the RBC from some Coombs-positive NZB mice. RBC from Coombs-negative NZB mice were never agglutinated by any F1 sera. Sera from NZB mice that had free-circulating anti-RBC antibodies (indirect Coombs positive) could be used to coat normal MRBC so that they would be agglutinated by the same F1 sera that agglutinated the donor NZB RBC. The activity of the F1 sera was age related, as 0 of 47 6–8-wk-old (CBA × NZB)F1 sera were able to agglutinate, compared with 15 of 84 4–6-mo-old sera. No difference was seen between F1 offspring of either CBA or NZB mothers. Agglutinating activity was not seen in the NZB or CBA parents, nor was it found in the unrelated mouse strains BALB/c and C57BL.
TABLE I

Absorption of (CBA × NZB)F1 Serum 57 Agglutinating Activity

| Absorbant                          | Titer⁻¹ vs. NZB E1 RBC | Titer⁻¹ vs. NZB C2 RBC |
|------------------------------------|------------------------|------------------------|
| None                               | 12,800                 | 6,400                  |
| Sepharose-goat anti-mouse IgG      | <10                    | <10                    |
| Sepharose-goat anti-TNP-OVA*       | 6,400                  | 6,400                  |
| Sepharose-goat anti-KLH‡           | 12,800                 | 3,200                  |
| Mouse IgG-coated SRBC§             | 12,800                 | 1,600                  |
| Mouse IgG-coated SRBC∥             | 25,600                 | 6,400                  |

* Trinitrophenylated ovalbumin.
‡ Keyhole limpet hemocyanin.
§ Coated with mouse anti-SRBC.
∥ Coated using CrCl₃.

TABLE II

Reactivity of (NZB × CBA)F1 Sera with NZB RBC (Reciprocal Titers)

| NZB RBC | E1 | F1 | H1 | J1 | K1 | K2 |
|---------|----|----|----|----|----|----|
| Cl      | 42 | 51,200 | 3,200 | — | — | — |
| 45      | 1,280 | — | 1,280 | — | — | — |
| 47      | 12,800 | — | 51,200 | — | — | NT‡ |
| 54      | 102,400 | 2,560 | 51,200 | 2,560 | 1,280 | — |
| 56      | 51,200 | 12,800 | — | — | 10,280 | 2,560 | 12,800 | 3,200 |
| 57      | 12,800 | 51,200 | 12,800 | 200 | — | NT | NT | NT | NT |
| 63      | 640 | 320 | — | — | NT | NT | NT | NT | NT |
| 66      | 102,400 | 51,200 | 2,560 | 51,200 | — | NT | NT | NT | NT | NT |
| 96      | — | — | 1,600 | — | — | — | — | 12,800 | — | — |

* <50
‡ Not tested.

Nature of the F1 Agglutinating Principle. The finding that F1 sera would agglutinate only Coombs-positive MRBC suggested that an anti-antibody might be involved. To test this hypothesis, one positive F1 serum (serum 57) was absorbed in several ways and tested against RBC from two Coombs-positive NZB mice (E1 and C2). As indicated in Table I, this particular combination of sera and cells gave high titers of agglutination. Absorption of the F1 serum with a column of anti-mouse IgG completely removed the agglutinating activity, whereas control columns had no significant effect. This suggested that an antibody was responsible for the activity of the F1 serum. One possibility was rheumatoid factor; however, serum 57 failed to agglutinate SRBC coated either with mouse anti-SRBC antibody or coated nonspecifically with normal mouse IgG (Table I). Mouse IgG bound to Sepharose also failed to remove the agglutinating activity from another F1 serum (not shown). Similar absorption experiments with four other F1 sera gave comparable results.

Specificity of Agglutinating Principle. One of the most striking features of the agglutination phenomenon was the specificity that individual F1 sera showed for individual NZB RBC. A summary of titrations of 10 reacting F1 sera with RBC from each of 12 Coombs-positive NZB mice is shown in Table II. Although certain sera and certain
cells were more reactive than others, the overall pattern suggested individual specificities of a qualitative nature. For example, $F_1$ serum 42 agglutinated RBC C1 with a titer of 51,200, but failed to agglutinate erythrocytes J1. Conversely, serum 143 failed to agglutinate NZB RBC C1 but agglutinated RBC J1 with a titer of 12,800. It is unlikely that this individual specificity represented the reactivity of $F_1$ rheumatoid factors with selected subclasses of mouse IgG, as we found no obvious correlation of the pattern of reactivity of NZB erythrocytes with heterologous anti-subclass antisera compared with the pattern of reactivity with the (CBA × NZB)$F_1$ sera (Table III).

The individual specificity of the reactivities was confirmed by blocking experiments, as shown in Table IV. The reactivity of $F_1$ serum 96 vs. NZB E1 RBC (see Table II) could be inhibited by serum from mouse E1 but not by serum from six other Coombs-positive NZB mice, the RBC of which did not react with $F_1$ serum 96. Similarly,
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TABLE V
Agglutination of Autoantibody-coated RBC by Whole or Pepsin-digested (NZB × CBA)F1 Sera

| F1, Reagent | Uncoated MRBC | Normal MRBC coated with NZB whole sera | Normal MRBC coated with NZB Fab'2 |
|-------------|---------------|--------------------------------------|----------------------------------|
|             |               | NZB-5 | NZB-12 | NZB-22 | NZB-25 | NZB-5 | NZB-12 | NZB-22 | NZB-25 |
| Mouse 2-5 whole serum* | - | - | + | - | + | - | + | - | + |
| Mouse 2-5 F(ab')2 | - | - | + | - | + | - | + | - | + |
| Mouse 3-1 whole serum | - | - | - | - | - | + | - | - | - |
| Mouse 3-1 F(ab')2 | - | - | - | - | - | + | - | - | - |
| Mouse 4-3 whole serum | - | + | - | + | - | + | - | + | - |
| Mouse 4-3 F(ab')2 | - | + | + | + | + | + | - | + | - |
| Mouse 4-4 whole serum | - | + | - | + | - | + | - | + | - |
| Mouse 4-4 F(ab')2 | - | + | - | + | - | + | - | + | - |
| Anti α + anti λ§ | - | + | + | + | + | + | - | + | - |
| Buffer | - | - | - | - | - | - | - | - | - |

* 1:20 dilution.
§ Mixture of 1:50 dilution of monospecific goat anti-mouse α and goat anti-mouse λ sera.

TABLE VI
Comparison of F(ab')2 and Fab' Agglutination of F(ab')2 and Fab'-coated Mouse RBC

| Agglutinating fragment | F1 mouse | F(ab')2 coating reagent | Fab' coating reagent |
|------------------------|----------|------------------------|----------------------|
|                        | Saline   | NZB-5 | NZB-12 | NZB-22 | NZB-25 | NZB-5 | NZB-12 | NZB-22 | NZB-25 |
| F(ab')2*               | 2-5      | - | - | + | - | + | - | + | - |
|                        | 3-1      | - | - | + | - | + | - | + | - |
|                        | 4-3      | - | - | + | - | + | - | + | - |
|                        | 4-4      | - | + | + | + | + | + | + | + |
| Fab'†                  | 2-5      | - | - | - | - | - | - | - | - |
|                        | 3-1      | - | - | - | - | - | - | - | - |
|                        | 4-3      | - | - | - | - | - | - | - | - |
|                        | 4-4      | - | +/- | - | - | - | - | - | - |
| Anti α + anti λ§       | - | + | + | + | + | + | + | + | + |

* 1:20 dilution.
† 1:10 dilution.
§ 1:50 dilution.

serum 143 could be inhibited in its reactions with NZB J1 RBC by J1 serum, and also by serum from another NZB with which 143 reacted, D2. Thus, the individual NZB specificities seen by the F1 sera were represented consistently both on the NZB RBC and in their serum.

**Agglutination Required Fab'-Anti-Fab' Binding.** If the agglutinating phenomenon we observed was due to idiotype-anti-idiotype interactions, the binding function of both the F1 serum and the NZB Coombs antibody should be contained in the Fab portion of IgG. To demonstrate this, 45% ammonium sulphate fractions of four individual F1 sera and four NZB sera containing antibodies that reacted with at least some of the chosen F1 sera were all digested with pepsin and carefully purified free of intact IgG (see Methods). Due to the small volume involved, the resultant reagents could only be tested at a single dilution. The patterns of reactivity of the F1 F(ab')2 fragments were in general no different than those of the initial F1 sera (Table V). In addition, the F1 serum [or F(ab')2] equally agglutinated normal MRBC coated with either NZB sera or with the F(ab')2 fragments purified from these sera (Table V). Furthermore, as shown in Table VI, reduction and alkylation of the F1 F(ab')2 completely removed their agglutinating activities. Finally, normal MRBC coated with reduced and alkylated NZB F(ab')2 (i.e., Fab') were not agglutinated with F1 F(ab')2 fragments,
but still could be agglutinated with heterologous antibodies to mouse light chains (Table VI).

Discussion

The present data indicate that certain NZB F1 hybrids spontaneously produce antibodies directed against idiotypes of NZB Coombs anti-RBC antibodies. Evidence for the idiotype specificity of these antibodies derives from: (a) the selective reactivity of certain F1 sera with only certain NZB RBC (Table II); (b) the inhibition of F1 sera-NZB RBC agglutination by only those NZB sera from mice whose RBC were recognized by the F1 sera (Table IV); (c) the fact that F1 (F(ab')2 fragments agglutinated RBC coated with F(ab')2 fragments of the Coombs antibody, which indicated that Fab-Fab interactions were sufficient (Table V); and (d) the failure of F1 sera to agglutinate MRBC coated with NZB Fab', which indicated that the F1 sera recognized determinants near the antigen-binding site of the Coombs antibody. That the observed agglutination might be due to rheumatoid factor was ruled out by the patterns of serum reactivity, the failure to absorb out activity with mouse IgG in a variety of forms, and the specificity for the F(ab')2 portion of the Coombs antibody molecule. In sum, then, all our data are consistent with the interpretation that the F1-agglutinating activity is an anti-(anti-RBC)-idiotypic antibody. An unlikely alternative explanation would be that the F1 sera recognize V-region subgroups (12). This possibility cannot be formally excluded, because it has been shown that such markers on anti-Rh antibodies can be blocked by antigen binding (13, 14). However, we believe it improbable that such anti-V-region subgroup antibodies would spontaneously arise in the F1, and that they would show the high degree of specificity for individual NZB Coombs antibodies (Tables II and IV).

The observation of anti-idiotype antibody to an autoantibody is in keeping with several experimental reports (15-17) of the formation of spontaneous anti-idiotype antibodies directed against exogenous antigens. (NZB X CBA)F1 mice and other NZB hybrids develop hemolytic disease less severe than that of their NZB parent (10). Furthermore, it has been reported that the Coombs antibody found in certain NZB hybrid mice is allotype restricted and therefore probably oligoclonal (18). The small amount and clonal restriction of NZB hybrid anti-RBC autoantibody may be the result of anti-idiotype regulation (albeit imperfect) of these self-reactive specificities. It is of interest that we have not detected anti-idiotype antibody against Coombs antibody either in NZB mice or in normal strains.

That a hemagglutination assay is capable of detecting anti-(anti-Coombs) idiotypic antibody may reveal something of the nature of the Coombs antibody. It is likely that one of the two Fab regions of the Coombs antibody must be relatively free for the anti-idiotypic antibody to recognize it. The fact that (anti-RBC)Fab'-coated MRBC can be agglutinated by anti-light chain antisera, but not by F1 anti-idiotype sera is in accord with this interpretation. Such a mechanism argues for substantial univalent attachment of the Coombs antibody to the RBC surface, a hypothesis consistent with low avidity and the presence of free antibody in the serum. It also implies that the epitopes recognized by the Coombs antibody are widely spaced on RBC. Similar conclusions were reached in studies of the reactivity of idiotypic, V-region framework,
and constant-region antisera with RBC coated with F(ab')2 or Fab' fragments of anti-Rh antibodies (13, 14).

The data thus far collected allow only a rough estimation of the size of the repertoire of the NZB Coombs antibody. Cross-reactivity using anti-idiotype containing F1 sera suggests several shared idiotypes; however, the complex pattern of reactivity (Table II) implies more than one anti-idiotypic specificity in some F1 sera and more than one Coombs idiotype on some NZB RBC. It will be important to quantitate anti-idiotypic antibody and Coombs antibody levels over time in individual mice to define the relationship between RBC autoantibody levels and the appearance of anti-idiotype. With the availability of anti-idiotypic reagents to the NZB Coombs antibody, direct functional studies will also be possible to elucidate the role of such antibodies in regulation of autoantibody levels and autoimmune disease.

Summary

The F1 hybrids of NZB and several normal mouse strains are known to produce less anti-erythrocyte (Coombs) autoantibody and develop a milder hemolytic anemia than their NZB parents. We have found that serum from some (NZB × CBA)F1 mice agglutinated erythrocytes from certain Coombs-positive NZB mice, often in extremely high titer, whereas other (CBA × NZB)F1 sera agglutinated erythrocytes from different individual NZB mice. The agglutination was due to antibody, but was not due to rheumatoid factor activity. Because F(ab')2 fragments of the F1 sera agglutinated erythrocytes coated with F(ab')2 fragments of the appropriate NZB sera, the observed reactivity was probably caused by idiotype-anti-idiotype interactions. In addition, because F1 sera could not agglutinate mouse erythrocytes coated with monovalent NZB Fab' fragments, the recognized idiotype probably involved the antigen-binding site. Anti-idiotypic antibodies against anti-erythrocyte autoantibodies may play an important role in the regulation of autoantibody formation.

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