Crossreactivity and Inheritance of Idiotypes Restricted to Human Anti-Tetanus Toxoid Antibodies

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Abstract

The presence of cross-reactive idiotypes on human IgG antibodies of tetanus toxoid (TT) antigen was assessed by examining the capacity of two anti-idiotypic (ID) antisera raised against IgG (Fab')2 anti-TT (idiotype) from two subjects to bind radiolabeled “idiotype” and to inhibit the binding of radiolabeled TT to IgG from unrelated subjects and from family members of the idotype donors. Idiotypic crossreactivity with unrelated individuals was infrequent and weak but was frequent and stronger among siblings. The strongest idiotypic crossreactivity was seen between identical twins in studies using four anti-ID raised against the anti-TT idiotypes of two sets of twins.

The results of the present study suggest that idiotypic determinants restricted to human anti-TT antibodies are, at least in part, encoded by inherited genes, which are infrequently shared among unrelated individuals.

Introduction

Human and mouse antibodies to self antigens have been reported to display a high incidence of cross-reactive idiotypes (CRI)1 (1–9). This could reflect the presence of conserved germ line V regions that encode variable regions that bind to self antigens. CRI have often been detected on antibodies to foreign antigens made in inbred mouse strains. In contrast, antibodies to most foreign protein antigens made by outbred animals express individual idiotypes, and rarely express CRI unless specifically induced with anti-idiotypic (anti-ID) antibodies (9–14). The data on the prevalence of CRI on antibodies made by unrelated human subjects in response to foreign antigens is conflicting (15–22).

In the present study we have examined the presence of CRI on human antibodies to tetanus toxoid (TT) using rabbit anti-ID antisera. These antisera were raised against idiotypes restricted to the anti-TT antibodies of single individuals. Two different assays were used to detect CRI: direct binding of radiolabeled IgG anti-TT antibodies by anti-ID and inhibition of TT binding to IgG anti-TT by anti-ID. Crossreactivity of idiotypes restricted to anti-TT antibodies was found to be infrequent and weak with unrelated subjects, significantly stronger and more frequent among siblings, and strongest among identical twins.

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1. Abbreviations used in this paper: anti-ID, anti-idiotypic; CRI, cross-reactive idiotyp; DT, diphtheria toxoid; TT, tetanus toxoid.

Methods

Antigens. TT and diphtheria toxoid (DT) antigens were obtained from Massachusetts Biological Laboratories (Jamaica Plain, MA) and fractionated over Sephadex G-200 columns. The second peak eluted from the column was used.

Antisera. Rabbit anti-human IgG serum was obtained from a rabbit immunized with human immune serum globulin. Fab-specific and Fc-specific rabbit antisera to human IgG were obtained from Calbiochem-Behring Corp. (La Jolla, CA). Goat anti-rabbit IgG was produced by immunization with normal rabbit IgG and was extensively absorbed with normal pooled human IgG crosslinked to Sepharose 4-B before use.

Immunosorbents. Proteins were dialyzed against 0.1 M NaHCO3, pH 8.3, and crosslinked to cyanogen bromide–activated Sepharose 4-B beads (Pharmacia Fine Chemicals, Piscataway, NJ) at a ratio of 15–25 mg per 1 ml packed beads, by following the instructions of the manufacturers.

Radioiodinations. Proteins were radiolabeled with 125I by the chloramine T method (23) to a specific activity of 2,000–3,000 cpm/ng. TCA precipitability and the biological activity of the radiolabeled material exceeded 80% in each case.

Human sera. IgG immunosorbent purified antibodies, and pooled IgG. IgG was isolated over DEAE columns from the sera of subjects who had been recently immunized and boosted with TT (24). The IgG samples were centrifuged for 2 h at 100,000 g before use in the experiments to remove any immune complexes present. Immunosorbent purified anti-TT and anti-DT IgG were isolated by passage of serum IgG over Sepharose 4-B columns crosslinked to the appropriate antigens and by subsequent elution of the bound material with 3 M NaSCN.

Pooled human IgG was obtained from Massachusetts Biological Laboratories (Boston, MA) in the form of Cohn Fraction II prepared for intramuscular gammaglobulin therapy. For its use as an adsorbent, pooled IgG was thoroughly depleted of TT reactivity by exhaustive absorption by passage over TT-Sepharose. The reactivity-depleted IgG failed to agglutinate TT-coated red cells (25) and contained no detectable (< 2 ng/ml) IgG anti-TT as determined by radioimmunoassay (23).

Production of anti-TT “idiotypes.” All anti-ID sera used in the present study were newly prepared and have not been reported previously. Two adult subjects aged 22 and 28 yr with previous histories of immunization with TT and DT were boosted by intramuscular injection of 5 floculation units of TT antigen (Massachusetts Biological Laboratories) and plasmapheresed at days 7 and 10 after immunization. These subjects are designated as subjects 1 and 2 in Tables I and II and in Figs. 1–4. IgG F(ab')2 anti-TT was prepared from their plasma as described previously (22–24). Briefly, the plasma was clotted, precipitated with 50% (NH4)2SO4, and the precipitate was dissolved in normal saline, dialyzed against 0.01 M phosphate buffer, pH 6.8, fractionated over a DEAE column, digested with pepsin with acetate buffer at pH 4.5 (protein to enzyme ratio 50:1, 37°C, 48 h), and passed over a G-150 column to obtain IgG F(ab')2 fragments. This preparation was adsorbed against packed Staphylococcus aureus (IgG sorb; The Enzyme Center, Inc., Boston, MA) and did not show any reactivity by Ouchterlony analysis with rabbit antiserum specific for Fe of human IgG. The F(ab')2 IgG was passed over a Sepharose 4B-TT column and the material that bound to the column was eluted with 3 M NaSCN and designated “idiotype.” The material that did not bind to the column was further passed over Sepharose-TT until it was completely devoid of anti-TT activity as assessed by hemagglutination out of radioimmunoassay. This material was designated TT-nonreactive IgG F(ab')2, or “nonidiotype.”
A similar protocol was used to immunosorbent purify IgG (Fab')₂ anti-DT over a Sepharose 4B-DT column.

Production of rabbit anti-ID IgG. Albino rabbits (Pine Acre Rabbiterary, East Douglas, MA) were immunized with 0.5 mg of F(ab')₂ anti-TT (idiotype) in 0.5 ml of phosphate-buffered saline (PBS) emulsified in an equal volume of complete Freund's adjuvant (Difco Laboratories, Detroit, MI). Three booster immunizations with the same amount of the material in incomplete Freund's adjuvant were given at weekly intervals and the rabbits were bled 2 wk after the last booster dose. The serum was separated from clotted blood and the IgG fraction was obtained by 50% (NH₄)₂SO₄ precipitation. Preimmune and postimmune rabbit IgG (50--100 mg) was adsorbed on 1,000 mg TT-nonreactive IgG F(ab')₂ (coupled to Sepharose 4-B) from the idiotype donor and on 1,000 mg of pooled human immune serum globulin depleted of anti-TT activity. The adsorptions were repeated (usually two times) until the immune rabbit IgG showed no more crossreactivity with TT-nonreactive IgG F(ab')₂ derived from the idiotype donor as determined by radioimmunoassays. Furthermore, the rabbit IgG at 10 mg/ml contained no detectable anti-DT reactivity as determined by hemagglutination and by radioimmunoassays (< 2 ng/ml).

Binding of radiolabeled IgG F(ab')₂ anti-TT and TT-nonreactive IgG (Fab')₂ to rabbit anti-ID IgG. 100 μl of 125I-labeled IgG F(ab')₂ anti-TT idiotype, or TT-nonreactive IgG F(ab')₂ nonidiotypic (= 100,000--200,000 cpm and 50 ng), were mixed in Beckman microfuge tubes with different amounts of rabbit anti-ID IgG, rabbit preimmune IgG, or rabbit antihuman IgG (Fab specific) serum. After incubation at 37°C for 1 h, normal rabbit IgG and goat anti-rabbit IgG were added in quantities to achieve a uniform content of total rabbit IgG in each tube and to achieve equivalence. The tubes were incubated at 37°C for an additional hour at 4°C overnight, then were centrifuged in a Beckman microfuge (13,000 rpm), the supernatants were discarded, and the formed precipitates were washed three times in washing buffer (PBS) containing 0.5% bovine serum albumin and 0.5% Tween-20. The radioactivity of the precipitates was counted in a gamma spectrometer (Model 1185; Tracer Analytic, Inc.; Elk Grove Village, IL). All determinations were made in triplicates and all experiments were done at least twice. In all calculations the background, i.e., number of counts obtained with the appropriate amount of rabbit preimmune IgG, was subtracted from the obtained values. The amount of 125I-idiotype precipitated by anti-ID IgG was expressed as a percentage of the amount of 125I-idiotype precipitated by rabbit antihuman Fab.

Inhibition of binding of radiolabeled IgG F(ab')₂ anti-TT to rabbit anti-ID IgG. Half a milligram of rabbit anti-ID IgG was incubated for 1 h at 37°C and 1 h at 4°C with different preparations of human IgG before adding 125I-labeled IgG F(ab')₂ anti-TT. The rest of the experiment was performed as described above.

The inhibitory effect of TT and DT on 125I idiotype binding to anti-ID IgG was assessed by preincubating 125I-idiotype with TT or DT antigen for 1 h at 37°C and 1 h at 4°C. The rest of the experiment was performed as just described. The percentage of inhibition of 125I-idiotype binding to IgG was calculated as:

\[
1 - \left( \frac{cpm \text{ bound by anti-ID IgG in the presence of inhibitor}}{cpm \text{ bound by anti-ID IgG without inhibitor}} \right) \times 100.
\]

Inhibition of 125I-TT and 125I-DT binding to IgG by rabbit anti-ID sera. The dilution of human IgG necessary to precipitate 70% of the total amount of precipitable radiolabeled antigen was determined for each source of IgG from the following experiment. A set of Beckman tubes received 100 μl of serial dilutions of human IgG. Human myeloma IgG devoid of anti-TT or anti-DT activity was added to achieve a total amount of 30 μg of IgG in each tube. Radiolabeled antigen (100,000--200,000 cpm and 50 ng) was added in 100-μl vol for 1 h at 37°C, rabbit anti-human IgG was added at equivalence, and the incubation was continued at 37°C for 1 h, then at 4°C overnight. The tubes were then centrifuged in a Beckman microfuge and the formed precipitate were washed three times in washing buffer and counted in a gamma spectrometer as described above. The amount of IgG that bound 70% of the total amount of precipitable radiolabeled antigen was used in all further inhibition experiments.

The amount of IgG determined above was incubated without or with rabbit anti-ID IgG or with preimmune IgG for 1 h at 37°C, after which radiolabeled antigen was added and the experiment was carried as described above. The background in this experiment was cpm bound by myeloma IgG alone, and it was subtracted before any calculations. Percent inhibition of binding was calculated:

\[
1 - \left( \frac{cpm \text{ of 125I-TT/DT bound by IgG in the presence of inhibitor}}{cpm \text{ of 125I-TT/DT bound by IgG without inhibitor}} \right) \times 100.
\]

Evaluation and calculation of crossreactivity. A significant crossreactivity was considered to be present when binding of F(ab')₂ anti-TT by anti-ID exceeded by threefold or more the binding of F(ab')₂ anti-DT from the same donor, and when inhibition by anti-ID of 125I-TT binding to IgG exceeded by threefold the inhibition of 125I-DT binding to IgG from the same donor. In each of the two assays used the degree of idiotypic crossreactivity between the idiotype donor and the test subject was calculated as:

\[
100 \times \left( \frac{cpm \text{ of specific activity of anti-ID with IgG of test subject}}{cpm \text{ of specific activity of anti-ID with IgG of idiotype donor}} \right)
\]

Statistical analysis. The statistical analysis was performed by the Student's t test.

Family studies. Members from the families of subject 1 (family 1) and subject 2 (family 2) were all adults (> 18 yr) and gave history of immunizations with TT. The last of those immunizations was always > 1½ yr before the present study. IgG from those family members was obtained without prior boosting with TT.

Twins studies. Two sets of adult identical twins (age 22 and 32, respectively) were studied. They were boosted with TT and idiotype and rabbit anti-ID were prepared as described for donors 1 and 2.

Results

Characteristics of the anti-ID antisera. The capacity of the two anti-ID antisera used in the present study to bind the IgG (Fab')₂ anti-TT or idiotype used for immunization is shown in Fig. 1. 1 mg of anti-ID IgG bound, respectively, 48 and 40% of 125I-idiotype but < 1% 125I nonidiotypic (TT-nonreactive IgG F(ab')₂) derived from the donor of the anti-TT antibody used for immunization. Preimmune rabbit IgG bound < 1% of 125I the idiotype used for immunization (data not shown).

![Figure 1](http://www.jci.org)

**Figure 1.** Idiotype binding by rabbit anti-ID IgG. Approximately 50 ng of 125I-labeled IgG F(ab')₂ anti-TT idiotype and of TT-nonreactive IgG F(ab')₂ nonidiotypic were added to various amounts of rabbit anti-IgG raised against the idiotype of the individual subject. The percent bound was calculated as the ratio of the precipitated by the anti-ID to the cpm precipitated by rabbit anti-human Fab × 100.
Figure 2. Inhibition of the binding of 125I-labeled TT and DT to IgG by rabbit anti-ID IgG raised against the IgG F(ab')2 anti-TT (idiotype) of each of the two subjects. The cpm of 125I-TT precipitated in the absence of the inhibitor were 180,000 cpm for subject 1 and 210,000 cpm for subject 2. The cpm of 125I-DT precipitated in the absence of inhibitors were 30,000 cpm for subject 1 and 60,000 cpm for subject 2.

The capacity of the two anti-ID antisera used in the study to inhibit the binding of 125I-TT to IgG obtained from the idiotype donor is depicted in Fig. 2. When 1 mg of anti-ID IgG was used this inhibition was 42 and 38%, respectively, for the two donors studied. This inhibition was antigen specific because the anti-ID antisera did not inhibit binding of IgG from the two subjects studied to 125I-labeled DT (< 5% inhibition). The inhibition of TT binding to IgG by anti-ID was not due to the presence of anti-TT antibodies in the rabbit antisera because these antisera contained no detectable anti-TT IgG (< 2 ng/ml) and absorption of the anti-ID over TT-Sepharose did not alter its capacity to inhibit TT binding to human IgG.

The majority of the idiotypic determinants recognized by the two anti-ID antisera appeared to be closely related to the TT binding site because in each case TT antigen, but not DT antigen, inhibited the binding of anti-ID to 125I-labeled idiotype to an extent close to that obtained with cold idiotype (Fig. 3).

Cross-reactive idiotypes assessed by direct binding of idiotype to anti-ID. We directly examined the reactivity of each of the two anti-ID antisera with 125I-labeled IgG (Fab')2 anti-TT and IgG (Fab')2 anti-DT obtained from 10 adult (18-42 yr) subjects (subjects 3–12) unrelated to the two idiotype donors and from family members of the idiotype donors. None of the subjects was booster immunized with TT for at least 1½ yr before the study. Table I shows that in only four of 22 instances tested was there evidence of crossreactivity with unrelated subjects (11, 17, 20, and 25%). There was no evidence of crossreactivity in this assay with two pools of human IgG tested (data not shown).

Table I. Anti-ID Binding to Radiolabeled Idiotypes from Different Subjects

<table>
<thead>
<tr>
<th>Source of idiotype</th>
<th>Reactivity of idiotype</th>
<th>Binding of idiotype by:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Anti-ID No. 1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>%</td>
</tr>
<tr>
<td>Subject 1 TT</td>
<td>40.2</td>
<td>36.1</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>4.1</td>
</tr>
<tr>
<td>Subject 2 TT</td>
<td>3.2</td>
<td>-0.8</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>4.0</td>
</tr>
<tr>
<td>Subject 3 TT</td>
<td>6.1</td>
<td>4.1 (11)</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>0.6</td>
</tr>
<tr>
<td>Subject 4 TT</td>
<td>6.1</td>
<td>2.2</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>3.6</td>
</tr>
<tr>
<td>Subject 5 TT</td>
<td>8.2</td>
<td>3.7</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>4.5</td>
</tr>
<tr>
<td>Subject 6 TT</td>
<td>6.5</td>
<td>2.8</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>3.7</td>
</tr>
<tr>
<td>Subject 7 TT</td>
<td>4.8</td>
<td>2.3</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>2.5</td>
</tr>
<tr>
<td>Subject 8 TT</td>
<td>7.0</td>
<td>2.4</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>4.6</td>
</tr>
<tr>
<td>Subject 9 TT</td>
<td>13.4</td>
<td>9.0 (25)</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>4.4</td>
</tr>
<tr>
<td>Subject 10 TT</td>
<td>6.4</td>
<td>1.2</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>5.2</td>
</tr>
<tr>
<td>Subject 11 TT</td>
<td>7.9</td>
<td>4.6</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>3.3</td>
</tr>
<tr>
<td>Subject 12 TT</td>
<td>9.5</td>
<td>3.7</td>
</tr>
<tr>
<td></td>
<td>DT</td>
<td>5.8</td>
</tr>
</tbody>
</table>

Equivalent amounts (50 ng) of F(ab')2 anti-TT and F(ab')2 anti-DT 125I-labeled idiotypes were added to 1 mg of rabbit anti-ID. The percentage bound idiotype was calculated as described in the legend of Fig. 1. All experiments were done in triplicate and the mean value is shown. Standard deviations were always < 16% of the mean values. Specific binding of anti-ID to IgG F(ab')2 anti-TT was obtained by subtraction of the binding of 125I-DT idiotype from that of 125I-TT idiotype. Values denoting significant crossreactivity, i.e., TT idiotype binding more than threefold DT idiotype binding, are underlined. Percent crossreactivity was calculated as described in the text, and is shown in between parentheses for individuals exhibiting significant crossreactivity.
The capacity of each of the two anti-ID antisera to bind $^{125}$I-F(ab')$_2$ anti-TT derived from the immediate members of the family of the idiotype donors (donors 1 and 2) is shown in Table II. In the first family, the one parent available for study and three of four siblings showed significant idiotypic crossreactivity with the idiotype donor. In the second family, both parents and two of four siblings showed significant idiotypic crossreactivity with the idiotype donor.

Cross-reactive idiotypes assessed by the capacity of anti-ID to inhibit TT binding to IgG. We next examined idiotypic crossreactivity by studying the capacity of each of the anti-ID to inhibit the binding of $^{125}$I-radiolabeled TT antigen to IgG from 10 subjects unrelated to the two idiotype donors (subjects 3–12) and from the family members of the idiotype donors. The inhibition of DT binding to IgG was used as a control, was always <5%, and was subtracted from the value of inhibition of TT binding to IgG. The results of these experiments are shown in Fig. 4. The same four subjects who showed crossreactivity in the direct idiotype binding assay also showed crossreactivity in the inhibition of TT antigen binding assay, i.e., 4 of 22 instances. The pattern of crossreactivity was similar to that seen with the direct binding assay shown in Table IV, i.e., anti-ID No. 1 crossreacted with anti-TT antibodies from subjects 3 and 9 and anti-ID No. 2 crossreacted with anti-TT antibodies from subject No. 6 and No. 12. This idiotypic crossreactivity was weak (13, 15, 20, and 21%, respectively).

Among family members the same pattern of idiotypic crossreactivity seen with the idiotype binding assay was observed in the present assay, i.e., all parents and the same five (out of eight) siblings showed idiotypic crossreactivity with the two idiotype donors.

Table III summarizes the incidence of cross-reactive anti-TT idiotypes based on our results with both the idiotype binding assay and the inhibition of antigen binding assay. In both assays crossreactivity was more prevalent and significantly ($P < 0.05$) more substantial with siblings than with unrelated subjects.

There was no correlation between idiotypic crossreactivity and sharing of HLA antigens within family members (data not shown). In both families the study of Inv and Gm allotypic markers was not informative.

CRI among identical twins. Two sets of genetically identical twins were studied for the presence of CRI on anti-TT antibodies. Individual rabbit anti-ID antisera were prepared against the anti-TT idiotype from each member of the two sets of twins, i.e., four antisera were prepared. The results of tests for CRI are shown in Table IV. Mean idiotypic crossreactivity was 59±6% in the idiotype binding assay and 57±4% in the inhibition of TT binding assay. Idiotypic crossreactivity among identical twins significantly ($P < 0.05$) exceeded that seen among siblings of the two families we studied here. There was no crossreactivity between anti-ID No. 1 or anti-ID No. 2 with IgG anti-TT from twins A and B (data not shown).

**Discussion**

The data presented in this study indicate that IgG anti-TT antibodies from unrelated subjects exhibit infrequent and limited cross-idiotypic reactivity. In contrast, cross-idiotypic reactivity was more frequent and stronger among siblings and strongest among identical twins.

### Table II. Anti-ID Binding to Radiolabeled Idiotypes from Family Members

<table>
<thead>
<tr>
<th>Source of anti-ID</th>
<th>Source of idiotypes</th>
<th>% Binding by anti-ID of IgG (Fab')$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Anti-TT</td>
</tr>
<tr>
<td></td>
<td></td>
<td>%</td>
</tr>
<tr>
<td>Anti-ID 1</td>
<td>Family 1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Subject 1</td>
<td>40.2</td>
</tr>
<tr>
<td></td>
<td>Mother</td>
<td>20.2</td>
</tr>
<tr>
<td></td>
<td>Sibling 1</td>
<td>20.4</td>
</tr>
<tr>
<td></td>
<td>Sibling 2</td>
<td>11.3</td>
</tr>
<tr>
<td></td>
<td>Sibling 3</td>
<td>10.2</td>
</tr>
<tr>
<td></td>
<td>Sibling 4</td>
<td>14.5</td>
</tr>
<tr>
<td></td>
<td>Family 2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Subject 2</td>
<td>38.7</td>
</tr>
<tr>
<td></td>
<td>Mother</td>
<td>18.5</td>
</tr>
<tr>
<td></td>
<td>Father</td>
<td>12.0</td>
</tr>
<tr>
<td></td>
<td>Sibling 1</td>
<td>8.1</td>
</tr>
<tr>
<td></td>
<td>Sibling 2</td>
<td>13.8</td>
</tr>
<tr>
<td></td>
<td>Sibling 3</td>
<td>7.2</td>
</tr>
<tr>
<td></td>
<td>Sibling 4</td>
<td>19.5</td>
</tr>
</tbody>
</table>

See legend to Table I.
The anti-ID antisera we have used were exhaustively absorbed against TT-nonreactive IgG obtained from the idiotype donor as well as from a pool of unrelated subjects. Thus, they were presumably directed against idioptic determinants restricted to anti-TT antibodies. Thus, these anti-ID would lack reactivity against “framework” idioptic determinants that could be shared by antibodies to TT and antibodies to other antigens. Our anti-ID antisera effectively inhibited the binding of TT antigen to IgG from the idiotype donor (Fig. 2). Also, TT antigen inhibited ID-anti-ID binding (Fig. 3), suggesting that the majority of the sites recognized by the anti-ID antisera were related to the antigen binding site. There was no evidence, however, that the anti-ID antisera contained antibodies bearing the “internal image” of TT antigen because, in the vast majority of the instances, the anti-ID antisera failed to compete with 125I-TT in binding to IgG anti-TT derived from unrelated subjects (Fig. 4) as well as from rabbits (data not shown). The lack of anti-ID antibodies bearing the internal image of TT did not simply result from denaturation of the TT binding site during the course of preparing the immunogen IgG (Fab)2 anti-TT, because there was virtually complete recovery of TT binding activity in the immunizing idiotype fraction eluted from Sepharose TT column, compared with the starting IgG (data not shown).

It is important to note that in the present study the anti-ID antisera were absorbed with TT-nonreactive IgG obtained from the donor of the immunizing idiotype and from pooled IgG. Such absorptions would have removed potential CRI shared between anti-TT antibodies and antibodies of other specificity. The recognition by our anti-ID antisera only of idioptic determinants restricted to TT and not of framework idiotypes, together with the absence from these antisera of anti-ID antibodies bearing the internal image of antigen, may have contributed to the relative infrequency in which CRI were detected in unrelated subjects. It is likely that the lack of CRI was related to the high degree of outbreeding among humans. In a highly outbred species like man it is not unexpected to find substantial differences in the TT binding polypeptide genes from unrelated individuals, which result from variable degrees of mutations and/or recombination events affecting the encoding V region genes. In this respect it is known that a substitution of one amino acid can result in major changes in the profile of idiotypic reactivity (26).

It is interesting to note that in the two assays for CRI used, the same pattern of crossreactivity was seen with the two anti-ID sera, which suggests that each anti-ID was recognizing separate CRI, each of which was present in 2 out of 11 unrelated subjects tested. The relative lack of CRI on human antibodies to the foreign antigen TT contrasts with the generally agreed upon observation of high incidence of CRI of human antibodies to self antigens (1–8). Based on the very strong homologies in the amino acid sequence analysis of the V region polypeptides of many of these autoantibodies (27–29), it appears that autoantibodies are encoded for by germ line genes that remain highly conserved in the outbred individuals of the species.

Lack of CRI on antibodies to foreign antigens made by individuals of outbred species was evident in the early observations of Oudin and Michel on the idiotypes of antibodies made by outbred rabbits immunized with the bacterial antigens (10). More recently the idiotypes of murine antibodies to alprenolol have been found to generally lack CRI (14). The data on the frequency and extent of CRI antibodies to foreign antigens in humans is conflicting. Atevogt and Wigzell (15) found that a rabbit anti-ID antiseraum raised against IgG (Fab)2 anti-TT reacted with IgG from only one of four unrelated subjects. Natvig et al. (17) detected CRI on anti-Rh antibodies in only 2 of 22 subjects. More recently, Emmrich et al. (16) found CRI in antibodies to Streptococcus polysaccharide antigens from the majority of unrelated subjects, whereas Cheung et al. (18) demonstrated CRI on anti-casein antibodies of 12 out of 16 IgA-deficient patients. Bose et al. (20) detected CRI on antibodies to rye antigen in the majority of patients receiving rye antigen immunotherapy, and Kennedy et al. (19) detected CRI on antibodies to hepatitis B surface antigen in the majority of subjects. In the anti-TT system Saxon et al. (22) presented evidence for CRI at the T cell level in subjects repeatedly and vigorously immunized with TT antigen, whereas Hoffman et al. (21) demonstrated a high frequency of CRI present on a variable percentage (1–50%) of anti-TT antibodies of unrelated subjects. The reasons for the conflicting results on the incidence of CRI to foreign antigens in human species are many. First, the nature of the antigen studied may be important. Caseins represent a family of small molecular weight (12,000–24,000) proteins also present in man and potentially presenting a limited number of immunogenic determinants. Second, the immunization schedule used may predispose to the predominance of CRI as occurs in repeatedly immunized mice (30). In this respect, studies showing CRI in man have involved antibodies made in response to repeated immunizations, e.g., repeated injections of TT (22), or rye antigen (20); infection with a recurrent bacterium such as streptococcus (16) or with a replicating virus such as hepatitis B (19); and repeated injection of the antigen casein (20). Third, the population of subjects studied may be important. The highest cross-idioptic reactivity of anti-casein antibodies occurred among IgA-deficient patients of Finnish descent, a group known to be genetically less heterogeneous than North Americans. Most importantly, two critical technical considerations previously alluded to may underlie the conflicting results obtained in the different studies on CRI in man. First, unless the anti-ID antisera are exhaustively absorbed they would contain antibodies to framework determinants, which may be represented on TT as well as non-TT binding antibodies. These determinants tend to be conserved among unrelated subjects. Second, the presence in the anti-ID antisera of anti-ID antibodies bearing the internal image
of antigen will result in the detection of spurious CRI. In this respect, none of the studies demonstrating a high frequency of CRI in humans have examined the anti-ID reagents used for the presence of antibodies bearing the internal image of antigen (18–22).

Perhaps the most important observation of the present study is the high frequency and the strength of idiotype reactivity among family members (Tables II and III, Fig. 4). Crossreactivity among family members is the more significant in view of two observations. First, it is known that the idiotype profile of anti-TT antibodies fluctuates during the course of the antibody response to TT (31). The effect of such fluctuations was however minimized in our study because none of the family members had been recently immunized with TT and because the idiotypes used to raise the anti-ID were taken 7–10 d after immunization and would have not been affected by the auto anti-ID-associated fluctuations in idiotype profile (31). Second, the study of CRI among identical twins showed less than total cross-idiotype reactivity (Table IV), which suggests that uninherited factors such as somatic mutations, differential use of various genes, and ID–anti-ID interactions may influence idiotype expression.

Inheritance of strain-specific idiotypes in mice is well known (9, 32, 33). The inheritance of idiotypes in humans has been described for an individual rheumatoid factor idiotype (34). Our study suggests that this also occurs for idiotypes expressed on antibodies to a foreign antigen, such as TT. The exact interpretation of inherited CRI among family members is somewhat difficult in the absence of primary sequence data. Although in most cases serologically detected idiotype crossreactivity has been found to be due to similar amino acid v region sequences, there have been exceptions in which it was found associated with homology of only a very small region, with homology of carbohydrate groups or with no sequence homology at all (35, 36). It seems unlikely, however, that similar variable regions as detected by our anti-ID antiserum could be generated in family members exclusively by somatic diversification. The results, rather, suggest the inheritance of antibody genes related to the idiotypic determinants in humans. These genes appear not to be linked to those encoding HLA antigens. The lack, in the present study, of informative allotypic markers did not allow us to establish or to rule out linkage between genes encoding CRI and constant immunoglobulin chain genes.

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