Evidence Supporting the Identity in Graves' Disease of Thyroid-stimulating Antibody and Thyroid Growth-promoting Immunoglobulin G as Assayed in FRTL5 Cells

Margita Zakarija, Shixin Jin, and J. Maxwell McKenzie
Department of Medicine, University of Miami School of Medicine, Miami, Florida 33101

Abstract

This paper addresses the question: in Graves' disease is there a thyroid-growth stimulating IgG (TGI) separate from thyroid-stimulating antibody (TSAb)? Using the functioning rat thyroid line (FRTL5) cells for TGI (incorporation of [3H]thymidine into DNA) and TSAb (increase in cAMP concentration) assays, we tested IgG from 30 Graves' patients. Positive TGI assay occurred only if cAMP increased in the cells and responses correlated, i.e., $r = 0.95$, $P < 0.001$. With one very potent TSAb-IgG we showed that Fab was active as TGI and TSAb, IgG with pl of 8.5–9.0 was the most potent fraction in both systems and an inhibitory IgG prevented the action of both TSAb-IgG and TSH in both the TSAb and TGI assays. In the last example, the action was on the cell membrane and not on the TSH or IgG. These data are entirely compatible with the view that in Graves' disease, at least as tested in FRTL5 cells, the same IgG is active in stimulating both growth and adenylyl cyclase.

Introduction

In recent years the possibility has been raised that there may be a specific thyroid growth-stimulating IgG (TGI), distinct from the thyroid-stimulating antibody (TSAb) of Graves' disease, circulating in patients with simple goiter (1–4) or Graves' disease (3, 5). In addition, it was suggested that stimulation of thyroid growth may not be mediated by cAMP (1, 3, 6) unlike much of the rest of thyroid function (7), and despite the fact that such mediation was well documented for growth effects of TSH in canine thyroid cells (8–11). Recently, we addressed the question of whether TSH-stimulated growth (of the nontransformed functioning rat thyroid line, FRTL5 cells) was mediated by cAMP, and obtained data (12), as have others (13, 14), in support of that concept. We now describe experience with another aspect of the topic, viz., is it necessary to hypoth-

1. Abbreviations used in this paper: TGI, thyroid growth-promoting immunoglobulin G; TSAb, thyroid-stimulating antibody.

2. This IgG was prepared from the serum of a patient who, after giving birth to a hyperthyroid neonate, was treated with $^{131}$I and subsequently maintained on $T_4$. As this is the most potent TSAb in our experience and without a biphasic effect in cAMP assays, we have been using it as our laboratory standard (reference 12, TSAb; reference 15, DeL IgG). There has been no change in potency in this patient's IgG over 4 yr. IgG from 9/82 and 8/86 were tested simultaneously at $10 \mu g/ml$ (human thyroid cells); results were 570 and 571% increase in cAMP.

Address reprint requests to Dr. Zakarija, Department of Medicine (R-93), University of Miami School of Medicine, PO Box 016760, Miami, FL 33101. Dr. Jin's present address is the First Teaching Hospital, Xian Medical College, Xian, People's Republic of China.

Presented in part at the 61st Annual Meeting of the American Thyroid Association, Phoenix, AZ, September 1986.

Received for publication 7 July 1987 and in revised form 14 September 1987.

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0021-9738/88/03/0879/06 $2.00
Volume 81, March 1988, 879–884

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Table I. Thyroid Gland Size, Therapy, and Assay Data for 12 Patients with Graves’ Disease

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Thyroid Therapy*</th>
<th>IgG mg/ml</th>
<th>cAMP %</th>
<th>[3H]Tdr pmol/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Nil</td>
<td>0.2-4</td>
<td>105-75</td>
<td>103-70</td>
</tr>
<tr>
<td>2</td>
<td>PTU</td>
<td>0.2-4</td>
<td>98-85</td>
<td>66-88</td>
</tr>
<tr>
<td>3</td>
<td>PTU</td>
<td>0.2-4</td>
<td>96-96</td>
<td>100-72</td>
</tr>
<tr>
<td>4</td>
<td>PTU</td>
<td>0.2-4</td>
<td>96-95</td>
<td>95-100</td>
</tr>
<tr>
<td>5</td>
<td>PTU</td>
<td>0.2-4</td>
<td>106-125</td>
<td>105-92</td>
</tr>
<tr>
<td>6</td>
<td>PTU</td>
<td>0.2-4</td>
<td>846-1203</td>
<td>870-820</td>
</tr>
<tr>
<td>7</td>
<td>PTU</td>
<td>0.2-4</td>
<td>863-1208</td>
<td>872-870</td>
</tr>
<tr>
<td>8</td>
<td>PTU</td>
<td>1.0-4</td>
<td>282-1402</td>
<td>510-969</td>
</tr>
<tr>
<td>9</td>
<td>PTU</td>
<td>1.0-4</td>
<td>930-3788</td>
<td>985-936</td>
</tr>
<tr>
<td>10</td>
<td>PTU</td>
<td>1.0-4</td>
<td>451-1732</td>
<td>704-960</td>
</tr>
<tr>
<td>11</td>
<td>PTU</td>
<td>0.2-1</td>
<td>697-2850</td>
<td>963-962</td>
</tr>
<tr>
<td>12</td>
<td>PTU</td>
<td>0.2-4</td>
<td>1100-3462</td>
<td>970-951</td>
</tr>
</tbody>
</table>

*At time blood was obtained; PTU, propylthiouracil; T4, thyroxine, prescribed for hypothyroidism that developed 5 wk after subtotal thyroidectomy (patient 6) or 3 doses of 131I (patient 12). Blood for TGI and TSAb assays was obtained shortly before (patients 1-3) or after (patient 6) subtotal thyroidectomy. Where it was from patients under therapy with PTU all had suppressed TSH so that the thyroid weight was presumably not influenced by therapy in all except patient 12. Thyroid weight: *confirmed at operation; *clinical estimate. Assay data are the means of triplicate observations obtained with the two concentrations of IgG listed; in most instances intermediate concentration of IgG was also tested. cAMP-% and [3H]Tdr refer to values expressed as percentage of control (12); in both systems IgG 1-5 were not stimulatory. In the TSB assay the maximum response to TSH (5 mU/ml) was 6596% and in TGI assay (200 μU/ml) 973%. As discussed in the text, the TGI assay response is maximal at lower concentrations of TSH and IgG than occurs in the TSAb system.

Both systems. To exclude the possibility that the inhibitor interacted directly with TSH and TSAb-IgG, some assays were carried out by preincubating cells with the inhibiting IgG, then washing them with buffer before incubation with test materials. In this report the laboratory standard TSB-β-IgG is referred to as TSB or TSAb-IgG and the inhibiting IgG as IgG-β. TSH used was NIH-TSH-B-10, a gift from National Institute of Diabetes, Digestive and Kidney Diseases, NIH.

TSAb assay with human thyroid and FRTL5 cells in culture. FRTL5 cells were generously provided by Dr. L. D. Kohn and Dr. W. A. Valente of NIH and were used as described (12). The procedure for human thyroid cells, in hypotonic medium, was also reported previously (15). In both instances cAMP was measured by radioimmunoassay kit (Immuno Nuclear Corporation, Stillwater, MN); with the FRTL5 cells an ethanol extract of cells was processed (12) and with human cells aliquots of medium were used directly after dilution and acetylation (15). The incubation time with IgG or TSH was 2 h with FRTL5 cells and 4 h with human cells.

TGI assay. FRTL5 cells were used as described earlier (12) with increase in DNA and in the incorporation of [3H]Tdr into DNA as indices of growth. The incubation time with IgG or TSH was 72 h.

Statistical methods. Analysis of variance was used to establish the statistical significance of assay data. Correlation coefficient was calculated by linear regression analysis.

Results

Assays in human thyroid cells. These were carried out at two or three concentrations of IgG, typically 0.05, 0.5, and 1.5 mg/ml hypotonic medium, as previously described (15, 20). All 30 samples were positive at one or more concentrations. Illustrative data with IgG from 12 patients are given in Fig. 1. As we have emphasized in an earlier publication (20), the response to the three doses of IgG tested may vary from straight-line positive slope (e.g., IgG 8), through biphasic (e.g., IgG 3), to a straight-line negative slope (e.g., patient 6). Also shown in the figure is the fact that the least potent IgG (1-5) were negative in the FRTL5-cAMP assay and all others were positive. This aspect of the study is enlarged below.

Assays with FRTL5 cells. Of the 30 IgG, 17 were positive in terms of both cAMP production and [3H]Tdr incorporation; 4 were low positive (<180%) as TSAb but negative in the TGI assay; 8 were negative in both procedures; data for the 12 most recently studied IgG are in Table I. We had the opportunity to obtain repeatedly IgG from 1 patient and her data are detailed below.

There was no correlation of the data from any of the three assay systems, specifically the TGI procedure, with estimated goiter size (Table I). On the other hand, as indicated in Table I, TGI negative IgG were also negative in the FRTL5-cAMP assay and were the least potent in the human thyroid cell assay. A correlation coefficient of 0.95, P < 0.001, resulted when the values obtained with the lowest IgG concentrations in the 12 duplicate sets of data were used to compare results of TSAb versus TGI assays. The values with the highest IgG concentrations were not as satisfactory for analysis since some represented data after the maximum responsiveness of the TGI assay was reached.

Studies with individual IgG. Fig. 2 provides data with IgG collected over time from a patient initially treated with propylthiouracil (from 3/84) and then subjected twice to subtotal thyroidectomy (in June 1985 and June 1986 after relapse of hypothyroidism with goiter regrowth), before requiring thyroxine therapy for hypothyroidism. The patterns of assay response (human thyroid cells) illustrate a change from negative to positive slopes over the concentrations tested. The figure also depicts assay results using FRTL5 cells. No IgG was available from the blood of 3/84 but the four other IgG were assayed for effects on both cAMP concentration and [3H]Tdr incorporation. Only samples from 1/85 and 7/85 enhanced
cAMP concentration, as shown; IgG from 8/86 and 12/86 were negative as tested at a maximum of 4 mg/ml. In the TGI assay (data not shown) all IgG were negative except that of 1/85, which at 4 mg/ml gave a response of 463%.

Assay of fractions obtained by isoelectric focusing. Fig. 3 offers a comparison of increasing concentrations of the IgG fractions, obtained by isoelectric focusing, that were tested in the TSAb and TGI assays with FRTL5 cells. The following features merit emphasis. In the TSAb assay there is a progressive increment of response to 10–200 µU TSH/ml; none of the IgG fractions stimulated the system to the degree obtained with 200 µU TSH/ml. For each concentration of IgG the greatest effect was obtained with fraction 5, i.e., with pI of 8.5–9.0. With the TGI assay, as judged by the response to TSH, a maximal response of ~ 650–700% was achieved with fractions 4–6 at 50 µg/ml and fractions 3–7 at 200 µg/ml. However, with 10 µg/ml the greatest response was clearly obtained with fraction 5 (pI 8.5–9.0). As reported previously (12), it is characteristic of these assays that the maximum effect is readily reached with lower quantities of either IgG or TSH in the TGI system than in the TSAb assay. A similar pattern of response relative to the pI of the fractions was obtained in the TSAb assay with human thyroid cells except that there was greater sensitivity in that system; when IgG was tested at 1, 5, and 50 µg/ml, a maximum effect was observed with 5 µg of fraction 5 (data not shown).

Table II provides data from the same experiment but with the growth effect shown as micrograms of DNA/well; there was confirmation of a true growth effect with 50 and 200 µg IgG/ml. Using the lower concentration, the greatest increase in DNA was observed with fraction 5 but with 200 µg/ml all responses, except for that of fraction 2, were to the apparent maximum of the system, as was the case with the [3H]Tdr incorporation.

TSAb and TGI assays of Fab fragment of TSAb-IgG. As shown in Fig. 4 in the TSAb assay the Fab fragment of the standard TSAb-IgG was, by weight, more potent than the parent IgG; we have previously documented that on a molar basis the Fab component is approximately equipotent to the parent IgG (15). We have no explanation why, in this instance, the Fab fragment was apparently of significantly greater molar potency than the parent IgG. In the TGI assay the maximal effect (as judged by the response to 200 µU TSH/ml, not shown) was achieved by 10–50 µg Fab/ml but only by 50–200 µg IgG. A similar pattern (not shown) was obtained when an increase in DNA/well was used as the index of response. Thus in stimulation both of cAMP accumulation and growth the Fab fragment of the parent IgG had full potency.

Inhibition of TSAb, TGI, and TSH. IgG-i was previously shown to be a potent inhibitor, in the TSAb assay, of both TSH and TSAb-IgG (15). It was therefore of interest to test it in the TGI assay and the data are given in Fig. 5. IgG-i had a slight negative effect by itself, and TSAb and TSH were stimulatory at 20 µg and 20 µU per ml; these effects of TSAb and TSH were completely inhibited by addition of IgG-i to the system.

The mechanism of inhibition of TSAb-IgG and TSH by IgG-i in the TSAb assay was explored as shown in Fig. 6. In this experiment cells were pre-incubated with 1 mg IgG-i or normal IgG per ml buffer for 1 h before they were washed and used for the TSAb assay. The effect of only preincubation with IgG-i was to inhibit both stimulators by >90%; there was complete inhibition when TSH and TSAb were assayed with IgG-i included in the second incubation. Similar results were obtained when another inhibitory IgG was tested in like fashion (data not shown).

Discussion

These experiments were aimed at testing the hypothesis that there may be different IgG molecules in Graves' disease that are active in TSAb and TGI assays. Our findings are compatible with the view that the same IgG is active in both systems.

As emphasized previously we consider it important, when using human thyroid cells, to assay samples for TSAb at sev-
The fraction (Fr.) number refers to those identified by pl in Fig. 3. The DNA values in this table are those used to calculate the response ([3H]Tdr/μg DNA) in Fig. 3.

* Mean±SD, n = 3.

† P vs. control: < 0.001.

The table shows the mean and standard deviation (Mean±SD) of three triplicates for each group. The groups include Control, TSAb-IgG, Fr. 2, 3, 4, 5, 6, and 7. The DNA concentration is given in μg DNA/well at various IgG concentrations: 10 μg/ml, 50 μg/ml, and 200 μg/ml.

Comparison of responses to IgG in the three assay systems is fraught with difficulty. In general it is clear that, with FRTL5 cells, only if there is a positive TSAb effect of a certain magnitude, i.e., in our hands, an increase in cAMP accumulation of at least 180%, is there stimulation of [3H]Tdr incorporation. Secondly, overall in the two TSAb assay systems, the response of human thyroid cells is greater than that of FRTL5 cells, although direct comparison is complicated by the biphasic responses (Fig. 1) that occur more frequently with the former. In our experience, a response in the human system of at least 400% increase in cAMP with one concentration of IgG was obtained if that IgG was positive in the TGI system. Furthermore, as illustrated by the data in Fig. 2 and Table I, there was no correlation between the clinical assessment of thyroid growth (Fig. 2) or goiter size (Table I) and any of the assay data. The degree to which an inhibitor of TSAb, referred to above and characterized previously (15), enters into divergence of TSAb assay data (human cells) and thyroid size is unclear. On the other hand, overall our data indicate that an IgG that is strongly positive in the human thyroid cell system will be positive in both FRTL5 cell assays.

Turning to more detailed studies with a single IgG, we aimed to compare several qualitative aspects of the IgG as tested in the TSAb and TGI assays. Evidence was obtained that the greatest stimulatory fraction of IgG, separated by isoelectric focusing, was the same, at least within the analytical limitations of the technique used, for both assays. As already emphasized in the results section, the lower maximal responsiveness of the TGI procedure makes comparisons of this nature less than straightforward; recognizing this difference in the assays, and the inevitable cross-contamination of fractions by large-scale preparative isoelectric focusing, the pl of the most active IgG for stimulation of growth was 8.5 to 9.0, similar to that with TSAb (also reported with other TSAb assay techniques (19)). This implies, at a minimum, that the membrane component to which IgG binds for the TGI effect has a pl similar to that for the TSAb effect, viz., acidic and complementary to the alkaline pl of the IgG.

The second point is that growth-stimulation, like the TSAb effect, does not require a divalent IgG molecule. Antibodies to the insulin and acetylcholine receptors have been shown to act only as whole IgG or Fab2, but not as Fab (21, 22). This is unlike TSAb that is active as Fab (15). The implications of both growth and cAMP increase being positively affected by a monovalent fragment suggest that cross-linking of the TSH receptor is not necessary for either effect. However, for the present purpose one may merely accept that the findings make more likely the identity of TSAb and TGI in the patient's IgG.

The final point is that TSAb and TGI activities of both
TSH and the patient’s IgG were inhibited by the action of a single IgG. Again, one interpretation of the data is that this inhibition reflects an action on a common pathway, i.e., involving adenylate cyclase. The mode of such inhibition is not established but from the study depicted in Fig. 6 it seems clear that the action is on the membrane, presumably on the TSH receptor itself, and not on the stimulating molecules, i.e., TSH or TSAb. An antibody to TSAb, i.e., an antidiotypic, was suggested by others (23) who studied the same patient’s IgG; our data clearly refute this possibility. Since the TGI assay requires at least 48 h (12), it is not possible similarly to test the mode of inhibition in that procedure, but it may be reasonable to assume that the same mechanism is involved. These inhibition data do not, of themselves, speak for a single IgG acting as both TSAb and TGI but are contrary to the view that not only are these separable IgG, but that they act through distinct pathways (3).

To summarize, we have reported data from both a survey of 30 patients’ IgG tested in three assay systems, and detailed studies of the characteristics of a potent TSAb-IgG that is active in these TSAb and TGI assay procedures. Our interpretation is that in Graves’ disease the IgG that stimulates adenylate cyclase is the same as that which stimulates growth. These conclusions are contrary to those reached by others who, using FRTL5 cells, studied IgG from a series of patients and found divergence in assay results, TSAb versus TGI (5). The latter findings were in keeping with the results of other experiments apparently showing that stimulation of growth and adenylate cyclase in FRTL5 cells was due to separate bioeffects of TSH (6). However, we and others (8–14) have reported that CAMP is indeed a mediator of the growth-promoting action of TSH and therefore probably also of TSAb.

Clearly our findings need have little bearing on claims that there may be a specific “TGI” in other goitrous states (1–4) or that there are growth pathways mediated by other than cAMP, indeed data have accumulated (10, 11, 24–30) in accord with the existence of a non-cAMP route for thyroid growth stimulation. Consequently, we restrict interpretation of our current observations to their being in keeping with the view that in Graves’ disease, TSAb action, through adenylate cyclase, is sufficient to explain thyroid growth. A role in vivo for other IgG or growth factors is possible. Regarding correlates of goiter size with assay data, it is probable that more than a single bioactive substance will influence thyroid growth in an individual patient. Apart from the coexistence of TSAb and other antibodies that affect the TSH receptor (reviewed in 31), it is now recognized that γ-interferon may reduce TSH-stimulated thyroid cell metabolism (32) and growth (33) in vitro. It is thus feasible that in vivo, particularly in thyroids with marked lymphocytic infiltration, such action of γ-interferon might modulate the size of the goiter.

An important caveat is that in these studies we, and others, speculate on the basis of data obtained with model systems, e.g., FRTL5 cells; a significant achievement will be development of a human thyroid cell line retaining all differentiated functions, including responsiveness to growth stimuli.

Acknowledgments

Appreciation is expressed to Mrs. Jean Zegadlo for her technical assistance. J. M. McKenzie is the Kathleen and Stanley Glaser Professor of Medicine.

Supported by U. S. Public Health Service grant AM-31391.

References

18. Zakariaj, M. 1980. Thyroid-stimulating antibody (TSAb) of...


