Detection of Thyrotropin Binding Inhibitory Activity in Neonatal Blood Spots*

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ABSTRACT

Recent studies have suggested that maternal TSH receptor-blocking antibodies might be of primary etiological importance in some cases of transient congenital hypothyroidism (CH). Because these antibodies are extremely potent, we evaluated the feasibility of identifying babies at risk by using readily available newborn blood spots. Blood spots obtained from 84 normal babies (group 1) and from 354 infants whose initial T₄ was less than the tenth percentile for the assay and whose TSH was 40 mU/L or more (group 2) were studied without knowledge of the diagnosis. Blood was eluted from spots overnight and evaluated for [¹²⁵I]TSH binding inhibition (TBI) to solubilized porcine thyroid membranes. Four spots obtained from 3 group 2 babies, but none of those from the group 1 infants, exhibited TBI activity greater than 3 SD above the normal mean (33.9%). Four additional hypothyroxinemic infants whose mothers had Graves’ disease were also negative. Subsequent follow-up revealed that all 3 positive babies had transient CH, and all 3 mothers had primary myxedema. Potent TBI activity was confirmed in the serum of all 3 mothers and in the 2 babies in whom it was evaluated at birth.

We conclude that newborn blood spots can be used to detect potent maternal TBI activity, and that this identifies a baby likely to have transient, rather than permanent, CH. Because of their stability and ease of collection and handling, newborn blood spots should offer a convenient tool for future studies aimed at defining in more detail the incidence and clinical characteristics of this unique syndrome. (J Clin Endocrinol Metab 77: 1005-1008, 1993)

A SYNDROME of transient congenital hypothyroidism (CH) has been described in babies born to mothers with autoimmune thyroid disease and potent TSH receptor-blocking antibodies in serum (1-3). Infants with this disorder are indistinguishable at birth from those with thyroid dysgenesis, because they frequently have severe hypothyroidism, no goiter is palpable either in the baby or mother, and thyroid uptake is significantly reduced or absent (4, 5). Unlike sporadic thyroid dysgenesis, however, babies with blocking antibody-induced CH tend to have transient disease, and the disorder has a high recurrence rate in subsequent offspring due to the tendency of these antibodies to remain elevated for a long time in maternal serum (1-5).

The prevalence of TSH receptor-blocking antibodies as a cause of CH is unknown. Because only those babies born to mothers with the most potent blocking antibodies develop hypothyroidism (3, 6-8) and the binding inhibitory effect correlates well with inhibition of TSH-induced stimulation of thyroid growth and function (6, 9, 10), we hypothesized that it might be possible to screen for TSH receptor-blocking antibodies by evaluating TSH binding inhibition (TBI) using readily available newborn blood spots. The present studies were performed to evaluate the feasibility of this method.

Materials and Methods

Study population

Serum was initially obtained from two patients with known primary myxedema (1'M) and two normal individuals for use in the pilot studies to be described. Patient 1 has had two babies with transient CH, and her immunoglobulin G (IgG) results have been previously reported (10). Patient 2 was a 10-yr-old girl who presented with severe 1'M and was known to have TSH binding inhibitory IgG.

Blood spots were obtained from the New York State Hypothyroid Screening Program between 1987 and 1988 and included 84 blood spots from normal babies (group 1) as well as 465 samples from 354 infants whose initial T₄ was less than the tenth percentile for the assay and whose TSH was greater than or equal to 40 mU/L (group 2). Of the group 2 babies, 150 were considered to have confirmed CH because their TSH levels had remained elevated on retesting. In the remaining 204 unconfirmed babies, thyroid function tests were normal on repeat exam. This miscellaneous patient group included babies with borderline values and those who had been screened before 4 days of age because of early discharge. As an additional control, 4 spots obtained from hypothyroxinemic infants whose mothers had Graves’ disease were studied as well. All spots had been stored desiccated at -4°C and were studied without knowledge of the diagnosis.

The study protocol was approved by the Committee on the Protection of Human Subjects in Research at the University of Massachusetts Medical Center.

Methods

In the initial studies, dried blood spots were prepared by mixing washed cells from normal heparinized blood with serum containing the desired test hormone or IgG in a ratio of 1:1, spotting the mixture onto
filter paper, and allowing the blood to dry for at least 18 h. Disks were punched using a standard 1/8th-in. paper punch and incubated in 200 μL 50% Coon's Modified Ham's F-12 culture medium overnight at 4°C. Fifty microliters of eluted blood were then assayed for [125I]bovine TSH (bTSH) binding inhibitory (TBI) activity according to the method of Southgate et al. (11). Each sample was studied in duplicate. The assay used Triton X-100-solubilized porcine thyroid membranes and receptor-purified bTSH, kindly supplied by Dr. John Pierce, University of California-Los Angeles, as tracer. In a few of the preliminary studies, a commercial kit (Kalibre TSH Receptor Antibody, Kronus, Dana Point, CA) was used. Both methods gave equivalent results. Human pituitary TSH (NIDDK hTSH-I-6) was provided by the National Hormone and Pituitary Program; recombinant human TSH (Genzyme) was the gift from Dr. Lewis Braverman.

Results were corrected for nonsaturable binding, and the percent inhibition of [125I]bTSH binding was expressed as a TBI index, computed as follows: 1 - (specific binding of [125I]bTSH in the presence of test sample/specific binding of [125I]bTSH in presence of control samples) X 100; unknowns were compared with the mean value of 6-10 normal spots evaluated in the same assay.

All spots were studied in duplicate. Any spot with a result 1.75 SD greater than the normal mean or higher was repeated.

Results

Pilot studies

Initial studies demonstrated that the TBI activity of blood spots obtained from the two 1°M patients whose IgG was known to have strong and intermediate potency, respectively, could be clearly distinguished from normal, and that the previously identified relative difference in potency could be demonstrated (Fig. 1). A dose-response curve was observed; potency was dependent on the number of filter paper disks used for elution. From these results, elution of four blood spot disks was selected for routine assay, as TBI activity was readily detectable with this quantity of blood and this amount of blood was obtainable from the majority of spots. Initial studies indicated that elution with either 50% Coon's F-12 or 10 mmol/L Tris-50 mmol/L NaCl gave equivalent results, and both were superior to 20% sucrose. Elution with 2 mol/L NaCl completely blocked [125I]bTSH binding of both normal and 1°M samples (results not shown). For the studies to be described, 50% Coon's F-12 medium was used.

Because the infants to be studied were expected to be hypothyroid, experiments were performed to evaluate the possible cross-reactivity of human TSH in this heterologous RRA. In three separate experiments, 1 U/L human pituitary TSH added to serum produced only 17.6 ± 7.7% (mean ± SEM) inhibition of [125I]bTSH binding. Results with recombinant hTSH were similar and are shown in Fig. 2. When added to blood, spotted onto filter paper, and eluted, as much as 2 U/L hTSH, the highest concentration tested, did not produce significant inhibition of tracer binding.

TBI activity in blood spots

The TBI activity of 84 blood spots obtained from 84 normal babies (group 1) and 465 blood spots obtained from 354 infants with either confirmed (n = 150) or unconfirmed (n = 204) CH (group 2) was studied (Fig. 3). The mean blood spot TBI activity in normal babies was 0.9 ± 11.0% (mean ± SD). Four spots obtained from 3 group 2 babies had TBI activity greater than 3 SD above the normal mean (61.4%, 71.9%, 41.7%, and 54%, respectively) and remained positive on retesting (60.2%, 40.9%, 48.1%, and 58.0%, respectively). All 3 babies had confirmed CH. Two other normal spots and 8 other CH spots had activity greater than 2 SD above the normal mean on initial testing, but these results were not confirmed. No borderline spots, defined as 1.75-2.0 SD greater than the mean, were positive on retesting. The interassay coefficient of variation of a positive control (patient 1) studied in 8 assays was 26.7%; the intrassay coefficient of variation of this sample was 12.0%.

As thyroid-stimulating antibodies in patients with Graves' disease could conceivably inhibit TSH binding and be de-
TABLE 1. Clinical and laboratory data of three infants with significant TBI-positive activity in blood spots and in their mothers

<table>
<thead>
<tr>
<th>Spot no.</th>
<th>Clinical course</th>
<th>% TBI (serum)</th>
<th>Diagnosis</th>
<th>% TBI/TRAb (serum)</th>
</tr>
</thead>
<tbody>
<tr>
<td>157</td>
<td>Transient CH</td>
<td>89.1</td>
<td>1°M</td>
<td>85.5/-</td>
</tr>
<tr>
<td>394</td>
<td>Transient CH</td>
<td>82.8</td>
<td>1°M</td>
<td>77.4/-</td>
</tr>
<tr>
<td>320</td>
<td>Transient CH</td>
<td>ND</td>
<td>1°M</td>
<td>—/382 U/L</td>
</tr>
</tbody>
</table>

TBI, TSH binding inhibitory Ig, determined at Nichols Institute (San Juan Capistrano, CA; normal value, <10%). TRAb, TSH receptor antibody; this assay, similar to the TBI assay, is based on the method of Southgate et al. (11) (normal value, <10 U/L). %TBI/TRAb was performed using a commercial kit (Kallire, Kronus). ND, Not done.

Our findings, thus, correlate remarkably well with those of previous studies, which demonstrated that half-maximal TBI activity in babies of 1°M mothers who became hypothyroid was obtained with a 1:3 to 1:5 dilution of serum. Four spots obtained from three babies exhibited the most potent (>3 SD) blocking activity, and all three babies had transient hypothyroidism. All were in the confirmed CH group, in contrast, no spots in the 84 normal subjects or 204 unconfirmed CH babies exhibited TBI activity of this magnitude. Preliminary studies indicated that a serum concentration of hTSH such as might occur in a baby with CH did not interfere significantly with [125I]hTSH binding, and that results using eluted blood from blood spots were dose dependent and mirrored results using serum.

The presence of potent TSH receptor-blocking antibodies was confirmed in the serum of all three mothers and in the two babies evaluated at birth. Similar to previously reported cases, all three mothers had 1°M; in one patient, hypothyroidism was preceded by radioactive iodine therapy for severe Graves’ disease. It is of interest that TPO antibodies, commonly used as a marker of autoimmune thyroid disease, were present in just two of the three mothers identified. This is consistent with previous reports suggesting that measurement of TPO antibodies is not an adequate screen for the prediction of transient CH (10).

Because bioactivity was not evaluated specifically, screening for TBI activity could conceivably have identified babies...
of mothers with Graves’ disease and high titers of thyroid-stimulating antibodies or even those with a mixture of TSH receptor antibodies of varying bioactivity (21). For this reason, spots obtained from four known hypothyroxinemic infants of Graves’ disease mothers were studied as well; none contained significant TBI activity. These results suggest that the hypothyryoxinemia observed in such babies, therefore, is more likely to be due to maternal thiouracil therapy (22). It is of interest that studies from several laboratories have demonstrated that stimulating antibodies do not inhibit as well as blocking antibodies (8, 13), probably because they interact at a different domain on the receptor (14, 15).

We conclude that newborn blood spots can be used to detect potent maternal TBI activity, and that this identifies a baby likely to have transient, rather than permanent, hypothyroidism. Because of their stability and the ease of collection and handling, newborn blood spots should offer a convenient tool for future studies aimed at defining in more detail the incidence and clinical characteristics of this unique syndrome.

Acknowledgments

Drs. Robert Klein and Marvin Mitchell, New England Congenital Hypothyroidism Collaborative, provided the blood spots from two of the hypothyroxinemic babies born to mothers with Graves’ disease.

References

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