A Case of Idiopathic Hypothalamic Hypothyroidism

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The availability of thyrotropin releasing hormone (TRH) has made it possible to determine whether tropic hormone deficiency is caused by pituitary or hypothalamic dysfunction. A case of hypothalamic hypothyroidism was described in a 17 year old woman. This patient was admitted for the evaluation of hypothyroidism and secondary amenorrhea. Her T3 and T4 were decreased, with an undetectable level of base line thyrotropin. The TRH test revealed normal but delayed response of TSH. Her base line prolactin and its response to TRH were normal. Adenocorticotropic hormone (ACTH), cortisol, growth hormone (GH), and urinary 17-hydroxysteroids were also normal. ACTH response to metyrapone was normal. Evaluation of the pituitary-gonadal axis revealed a normal increase in both lutenizing hormone (LH) and follicle stimulating hormone (FSH) following the intravenous administration of lutenizing hormone releasing hormone (LHRH). These results suggest that she had hypothalamic hypothyroidism as an isolated disturbance in the hypothalamic-pituitary axis. A deficiency of TRH is probably caused by a disorder of hypothalamic function of unknown etiology since the extensive studies did not reveal any secondary causes. It is recommended that patients with amenorrhea and hypothyroidism be evaluated for possible hypothalamic hypothyroidism.

Key Words: Thyrotropin releasing hormone, Thyroid stimulating hormone, Secondary amenorrhea, Lutenizing hormone releasing hormone, Hypothalamic-pituitary axis.

With the availability of synthetic TRH it became possible to assess the site of lesion in patients with hypothyroidism secondary to pituitary or hypothalamic disease1-2). This differentiation can be made by the response of TSH to TRH. Pittman and his associates were the first to describe a case of hypothalamic hypothyroidism due to hypothalamic origin3). The case was characterized by decreased thyroid hormones, and detectable serum thyrotropin (TSH) levels, with a normal response to TRH which occurred following a head injury. They suggested the term "hypothalamic hypothyroidism" for TRH deficiency as a cause of hypothyroidism. Since then similar cases have been reported. However, in most cases there are usually such secondary causes as trauma, tumor, irradiation, or stalk section4-7). Furthermore, TSH deficiency is usually observed as a part of multiple pituitary deficiencies8-9).

We report here a case of hypothalamic hypothyroidism of unknown etiology where TSH deficiency was an isolated phenomenon.

CASE REPORT

A 17 year-old high school girl, was referred to our medical service for the evaluation of hypothyroidism. She was born of an uncomplicated pregnancy and delivery, and her early growth and development were considered normal. She underwent menarche at the age of 12 but her menstrual cycle never became regular, and at the age of 14 she became completely amenorrheic.

In March 1979, she began to be easily fatigued and developed cold intolerance. At that time she was seen by her physician...
and the diagnosis of hypothyroidism was made. She was started on thyroid tablets with the gradual improvement of her symptoms. She continued to take the drug for several months but then there was no follow up. Three months before her admission to our hospital, she again noticed easy fatigability, cold intolerance and constipation, which became gradually worse. She gained 9 kg during the last several months. She also withdrew from many of her school activities. She was found to have decreased levels of thyroid hormones: T<sub>4</sub>, 2.1 μg/dl; TSH, less than 2 μu/ml. She was referred to our hospital. There was no history of headache, visual difficulties, skin dryness or deepening of her voice. There was no history of head trauma. Her family history was unremarkable.

Physical examination revealed that she was tall and well developed but appeared inactive in general. Her height was 170 cm and weight, 64 kg. Blood pressure was 110/50 mmHg, pulse, 44/min, regular, and respiration, 16/min. Her temperature was as low as 35.4°C. Her skin and hair were of normal consistency. The movement of eyes was normal and funduscopic examination was within normal limits. Her visual fields were intact. The thyroid gland was palpable, but not enlarged. The cardiac examination revealed no abnormalities except for sinus bradycardia. The lungs were normal. There was no hepatosplenomegaly. Pelvic examination disclosed no abnormalities. The neurological examination revealed that deep tendon reflexes were generally diminished with no lateralization. There were no pathological reflexes. Results of the routine laboratory studies were normal, including a complete blood count, urinalysis, and blood chemistry. The skull series showed a normal sella turcica. A computed tomographic scan of the brain revealed no abnormalities.

**EVALUATION OF ENDOCRINOLOGICAL FUNCTION**

**METHODS**

Jap J Med Vol 20, No 3 (July 1981)
was greater than those obtained at 15 and 30 min. However, prolactin response was normal. LH and FSH showed normal response to LHRH. ACTH reserve test was also normal (Fig. 3). These findings are consistent with diagnosis of isolated defect of hypothalamus-pituitary-thyroid axis.

Treatment was begun with desicated thyroid 20 mg daily which was gradually increased. She improved clinically. The level of T4, increased to 7.3 μg/ml two weeks after the initiation of therapy. Three months later, she returned to her usual school activities and her menses did not resume until 3 months after thyroid replacement.

Fig. 1. Prolactin and TSH response to TRH administration. Prolactin response was normal following injection of TRH (500 μg), and TSH response was normal but delayed. The shaded area indicates the normal range of response.

Fig. 2. FSH and LH response to LHRH administration. The intravenous administration of LHRH (100 μg) caused normal response of both FSH and LH.

Fig. 3. ACTH reserve test. Metyrapone (2.0 g) was given orally at 12:00 pm. ACTH response was normal.
Hypothalamic Hypothyroidism

DISCUSSION

Our patient described here had hypothyroidism and decreased serum TSH level with delayed response to TRH. The response of prolactin levels to TRH was normal. These findings are all consistent with hypothalamic hypothyroidism. Furthermore, it is interesting to note that our patient seems to have abnormality of only the hypothalamic-pituitary-thyroid axis. As shown in Fig. 2, basal FSH and LH values were normal, as was the response to LHRH. The function of pituitary-adrenal axis was also normal.

The TSH response to TRH administration has been well studied in normal as well as in various endocrinological disorders\(^{10\text{-}14}\). Hall et al have studied TSH response after intravenous injection of TRH in 77 patients with diseases of the pituitary or hypothalamus\(^{15}\). Of 15 patients with various hypothalamic lesions, 8 had an impaired or absent response to TRH; 3 patients who responded normally to TRH had other evidence of hypothyroidism. A delayed response to TRH, where the twenty-minute TSH level was less than the sixty-minute value, occurred in 13 of 15 in this group. They concluded that a normal TSH response to TRH, but a delayed response was characteristic of hypothalamic lesions. It appears that the TSH-producing cells of pituitary gland are intact, but certain time interval should elapse before the initiation of TSH synthesis.

As far as the etiology is concerned in our patient, it is probably idiopathic on the basis of normal skull X-ray and computed tomographic scan of the brain. Although, in the majority of cases, intracranial lesions could be found as the cause of hypothyroidism, idiopathic cases have rarely been reported\(^{1\text{-}5,16}\). A search of the English literature has revealed only five cases of idiopathic hypothalamic hypothyroidism which was an isolated phenomenon\(^{15,17}\). In Japan, two similar cases have been described\(^{18,19}\). Since an enzymatic mechanism of the synthesis of a hypophysiotropic hormone from aminoacid precursors was demonstrated in the hypothalamic tissue\(^{20}\), it may be possible that the enzyme “TRH synthetase” is absent or deficient in some idiopathic cases.

Hypothyroidism has been associated with menorrhagia or irregular menses. However, amenorrhea is rare\(^{21}\). Our patient exhibited secondary amenorrhea. Woolf has reported four women who had hypothyroidism due to hypothalamic insufficiency, although one patient did not show the typical pattern of TSH release to TRH\(^{10}\). These four patients were similar to our patient in which the etiology was probably idiopathic and other hormonal systems except for pituitary-thyroid axis were normal. It is interesting to note that all four patients had secondary amenorrhea. It has been suggested that an acquired abnormality within the hypothalamus was a probable cause of amenorrhea in view of their normal LH and FSH response to exogenous LHRH administration.

The cause of amenorrhea is unclear. It is at least in part causally related to thyroid deficiency on the basis of normal values of FSH, LH and prolactin. It should be noted, however, that there is considerable evidence from both animal experiments and studies in hypothyroid patients that thyroid deficiency may lead directly to impairment in gonadotropin secretion\(^{22\text{-}24}\). The menstrual response to thyroid hormone replacement is variable. In all four patients described by Woolf, the menstruation did not occur until 4 to 6 months after beginning treatment. By contrast, Kramer et al\(^{17}\) reported patients resumed menstrual cycle 2-3 months after treatment. In our patient, menses did not resume until 3 months after thyroid replacement.

Young women presenting secondary amenorrhea and hypothyroidism may have hypothalamic hypothyroidism. In such a clinical setting, TRH and LHRH tests are recommended.

ACKNOWLEDGMENTS: We are indebted to Dr Satoru Sekoguchi for referring the patient described in this report.
REFERENCES


