TSH neurosecretory dysfunction (TSH-nd) in Down syndrome (DS): low risk of progression to Hashimoto’s thyroiditis

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ABSTRACT

Introduction: Patients with Down syndrome (DS) often have elevated TSH (hypothalamic origin), which is called TSH neurosecretory dysfunction (TSH-nd). In these cases, there is slight elevation in TSH (5-15 μUI/mL), with normal free T4 and negative thyroid antibodies (AB). Objective: To recognize the risk of progression to Hashimoto’s thyroiditis (HT). Subjects and methods: We retrospectively analyzed 40 DS patients (mean age = 4.5 years), followed up for 6.8 years. Results: HT was diagnosed in 9/40 patients, three early in monitoring, and six during evolution. In 31/40 patients, TSH-nd diagnosis remained unchanged over the years, with maximum TSH values ranging from 5 to 15 μIU/mL. In this group, free T4 also remained normal and AB were negative. There was a significant TSH reduction (p = 0.017), and normal TSH concentrations (< 5.0 μUI/mL) were observed in 29/31 patients, in at least one moment. No patient had TSH > 15 μUI/mL. Conclusion: DS patients with TSH-nd present low risk of progression to HT (10% for females and 6% for males). Arq Bras Endocrinol Metab. 2011;55(8):628-31

Keywords
Down syndrome; Hashimoto’s thyroiditis; TSH neurosecretory dysfunction; isolated TSH elevation

RESUMO

Introdução: Pacientes com síndrome de Down (SD) geralmente apresentam TSH elevado (de origem hipotalâmica), uma desordem chamada de disfunção neurossecretora de TSH (TSH-nd). Nesses casos, há uma leve elevação do TSH (5-15 μUI/mL), com T4 livre normal e anticorpos antitireoide (AB) negativos. Objetivo: Reconhecer o risco de progressão para a tireoidite de Hashimoto (HT). Sujeitos e métodos: Analisamos retrospectivamente 40 pacientes com SD (idade média = 4,5 anos), acompanhados por 6,8 anos. Resultados: A HT foi diagnosticada em 9/40 pacientes, três logo no início da avaliação e seis durante a evolução. Em 31/40 dos pacientes, o diagnóstico de TSH-nd permaneceu estável durante os anos, com valores máximos de TSH variando de 5 a 15 μUI/mL. Neste grupo, T4 livre também permaneceu normal e os AB foram negativos. Houve uma redução significativa do TSH (p = 0.017), e concentrações normais de TSH (< 5,0 μUI/mL) foram observadas em 29/31 pacientes, em pelo menos um momento. Nenhum paciente apresentou TSH > 15 μUI/mL. Conclusão: Pacientes com SD e TSH-nd apresentam baixo risco de progressão para a HT (10% para o sexo feminino e 6% para o sexo masculino). Arq Bras Endocrinol Metab. 2011;55(8):628-31

Descritores
Síndrome de Down; tireoidite de Hashimoto; disfunção neurossecretora de TSH; elevação isolada de TSH
INTRODUCTION

Down syndrome (DS) is a common chromosomal abnormality and the most common genetic cause of mental retardation (1). Autoimmune diseases, especially Hashimoto’s thyroiditis (HT), are frequently observed in patients with DS. In these situations, autoantibodies (AB) directed against the thyroid are found in 13% to 34% of patients (2). Therefore, it is suggested that thyroid function, including measurement of TSH, free T4 and AB, is performed as part of routine monitoring of DS patients (3-4). Thyroid hormones have important functions in the central nervous system (CNS). They are involved in neuronal migration and differentiation, neurotransmitter synthesis and secretion, myelination, and regulation of gene expression in neuronal cells (5). Hypothyroidism is a potentially aggravating factor of neurological abnormalities of DS patients (5).

In addition to primary thyroid disease, patients with DS may present inadequate dopaminergic regulation of pituitary TSH secretion (6). These changes determine an abnormal secretion pattern, resulting in isolated TSH elevation without changes in thyroid hormones (7). In TSH neurosecretory dysfunction (TSH-nd), TSH concentrations are slightly elevated (5-10 μUI/mL), free T4 values are normal, and AB are negative. In most cases, there is no detectable anatomic abnormality, and the etiology is not identified. Frequency presentation is similar in males and females (7).

Despite the absence of alteration in thyroid hormones, it is questionable whether TSH-nd may reduce growth velocity in these children (8). In these cases, there is no consensus on levothyroxine replacement therapy (9).

Cutler and cols. suggested that isolated TSH elevation could be a sign of risk of progression to primary autoimmune hypothyroidism, even in the absence of detectable anti-thyroid AB (10).

The aim of this study was to identify a risk of progression to HT, by means of a follow up of DS patients with TSH-nd.

SUBJECTS AND METHODS

This study was based on the retrospective analysis of 40 patients with DS (mean age = 4.5 years), both females and males, followed up at the Pediatric Endocrinology Department Unit of ISCMSP, referred to us because of previous elevated TSH and/or positive anti-thyroid AB. The study was approved by the Ethics Committee in Human Research of the ISCMSP.

The studied variables were: weight (kg), height (cm), pubertal characteristics (Tanner score); T4, free T4, TSH, anti-thyropperoxidase and anti-thyroglobulin concentrations and, for some patients, thyroid ultrasound. Weight was measured on an analog scale (Filizola®), and height, on a wall-mounted stadiometer (Gomes & Tonelli®).

Height and BMI were expressed as standard deviation z scores (SDS), calculated from reference values of SD patients (11).

T4, free T4 and TSH were measured by fluorometric assays (AutoDELFIA automatic immunoassay system). Anti-thyropperoxidase and anti-thyroglobulin AB were quantified by an immunometric assay (IMMULITE 2000). Negative results were below 35 IU/mL and 40 IU/mL, respectively.

TSH-nd diagnosis was established when TSH values were slightly elevated (5-15 μUI/mL), T4 and free T4 concentrations were normal, and thyroid AB were negative.

Statistical analysis used SigmaStat for Windows, v.2.03 (SPSS, Chicago, Il, USA). Comparisons of initial and final values for height SDS, BMI SDS, T4, free T4 and TSH of the same patient were performed by Paired T test. Statistical significance was determined by p < 0.05.

RESULTS

Forty DS patients (mean age = 4.5 years) were considered for retrospective evaluation. Three patients (3/40) were initially excluded because they had symptoms compatible with primary hypothyroidism already at diagnosis.

During follow-up, 6/37 patients (4F: 2M) initially considered as having TSH-nd presented progression to HT diagnosis, without hypothyroidism. Anti-thyroid AB were positive after a follow-up period ranging from 8 to 28 months.

TSH-nd diagnosis remained unchanged in thirty one individuals (12F: 19M). Median follow-up period of this subgroup was 6.8 years (1.0 - 20.2). Mean age at first visit was 3.4 years (SDS = 3.7), ranging from 0.1 to 14.3 years. Table 1 summarizes initial and final descriptive data, including anthropometric measurements and laboratory results.

In relation to anthropometric measurements, there was no reduction in growth during the observation period. Despite a significant increase in BMI in this subgroup, final BMI SDS values were not consistent with obesity diagnosis.
When analyzed individually, TSH peak values were slightly elevated (mean = 9.7 μU/mL, ranging from 5.4 to 15.0). There was a wide variation in TSH, with 29/31 patients (93.5%) showing normal values in at least once during follow-up. Eleven patients (11/31) had at least one TSH measurement between 10 and 15 μUI/mL. Only one patient had TSH = 15 μU/mL, although this value was observed only once during the follow-up period of 5.4 years. Figure 1 represents TSH evolution in the subgroup of DS patients with TSH-nd during follow-up.

Also in the TSH-nd subgroup, no patient developed hypothyroidism. By analyzing T4 and free T4 concentrations, the lowest mean individual values were respectively: 7.6 mg/dL (5.3 to 10.9) and 1.1 ng/dL (0.7 to 1.7).

Thyroid ultrasound was performed in 16/40 patients. In 5/9 patients diagnosed with HT, ultrasound results showed only heterogeneous texture. In 11/31 patients with TSH-nd, all results were considered normal.

### DISCUSSION

The close relationship between thyroid disorders and DS is well-documented (12). Considering the total universe of patients with DS, the frequency of autoimmune thyroid disease is described to be between 20% and 66%, depending on the type of study, sample size, region, age group and the inclusion of hypo-or hyper-thyroidism cases (13-14). It is possible that previous studies also included DS patients with TSH-nd, without TH and without hypothyroidism, producing a large number of false positive results.

In this study, long-term monitoring showed positive AB in only 6/37 patients (16%) in a period ranging from 8 to 28 months of observation. From this total, about 10% were represented by females and 6% by males. These results corroborate literature data, which show a higher frequency of HT in girls than in pre-pubertal boys (2:1) (15).

For most patients (31/37) analyzed in this study, there were only intermittent and isolated elevations of TSH without thyroid disease progression. For all these patients, anti-thyroid AB were not detected, values of free T4 remained normal, and there was no reduction in longitudinal growth over a mean period of 6.8 years. These long-term observations suggest that isolated elevation of TSH in DS does not seem to be a condition that predisposes to the development of thyroid disease, since in these cases, normal TSH values were often observed at some point of monitoring.

The results of this study are consistent with those of the literature. Dias and cols. (16) analyzed retrospectively 169 children with Down syndrome, with 46 patients selected for cross-sectional analysis. The authors observed a high prevalence of isolated TSH elevation (39.6%), with slightly elevated TSH. Mean TSH values were similar to those observed in our study. These authors also observed a high frequency (67.4%) of normal TSH values in at least one time point during the clinical follow-up. Similar to our results, they also detected a low progression to thyroid disease, and hypothyroidism
was detected in only 4/169 of the DS patients. Thus, they concluded that isolated TSH elevation in DS children may be a transitional condition in these patients.

Other studies have also observed transient elevation of TSH in DS patients. Selikowitz (17) suggested a less active form of TSH. However, Konings and cols. (18) showed normal TSH bioactivity in DS children who had subclinical hypothyroidism.

Other studies have suggested that the genesis of isolated TSH elevation in DS patients may be related to increased dopaminergic tone, determining increased secretion of TSH, which is related to down-regulation of thyroid TSH receptors and may maintain normal free T4 values (6).

Dopamine is a catecholamine that acts both on the hypothalamus and on D2 receptors of pituitary trophs, inhibiting the secretion of TSH. Therefore, dysfunctions of the Hypothalamic-Pituitary-Thyroid axis in patients with DS may be associated with primary thyroid diseases or with disorders of TSH secretion dependent on insufficient dopaminergic control of pituitary secretion (19).

Previous studies conducted on patients with DS reinforce the hypothesis of reduced dopamine secretion in the CNS, and describe atrophy and reduction in the number of dopamine-producing cells in the substantia nigra at the base of the brain, and ventral tegmental area. These findings are common in individuals with DS who are over 40 years old, and occur as dementia similar to Alzheimer’s disease. Just like the substantia nigra and the ventral tegmental area (important dopamine-producing areas in the brain), the arcuate nuclei of the hypothalamus (dopamine producers that act as inhibitory hormone upon TSH secretion) could also be compromised (19-21).

We concluded that patients with DS may have elevated TSH, even when thyroid hormone values are normal. Thus, caution is recommended in relation to the indication of levothyroxine replacement therapy (22), since monitoring of these patients showed long-term, low risk of progression to HT. Repeated clinical and laboratory evaluation is necessary for correct indication of treatment in DS patients who have TSH-nd.

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