Hypothyroidism in children: diagnosis and treatment

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Abstract

Objective: To present relevant and updated information on the status of hypothyroidism in the pediatric population (newborn infants to adolescents).

Sources: Original and review articles and books containing relevant updated data.

Summary of the findings: This review addressed data on the etiopathogeny of hypothyroidism and on the importance of screening for congenital hypothyroidism to assure early diagnosis and treatment of the newborn. We point out the difficulties experienced in the handling of subclinical hypothyroidism; we also address the importance of diagnosing autoimmune Hashimoto’s thyroiditis, the high incidence of the disease among adolescents, mainly females, and the occurrence of a severe neurological condition, Hashimoto’s encephalopathy. We indicate situations in which severe hypothyroidism may lead to puberty disorders (precocious or delayed puberty) and describe the importance of transcription factors in thyroid embryogenesis. Diagnostic and therapeutic criteria are also addressed.

Conclusion: Thyroid hormones are necessary for normal growth and development since fetal life. Insufficient production or inadequate activity on the cellular or molecular level lead to hypothyroidism. These hormones are necessary for the development of the brain in the fetus and in the newborn infant. Neonatologists and pediatricians deal with child development issues in their practice, and many of these issues start during intrauterine life. Currently, with neonatal screening, neonatologists and pediatricians can prevent irreversible damage through early treatment. They should also be alert for dysfunctions such as subclinical hypothyroidism and Hashimoto’s thyroiditis, which may provoke damage not only to growth, but also to the neurological and psychological development of these children and adolescents.

Introduction

Deficiency in the production or in the activity of thyroid hormones (TH) leads to hypothyroidism, one of the most frequent hormone diseases in children. The first known description of this syndrome dates back to 1874, by Gull; the name myxedema was defined by Ord in 1878. The term myxedema was used for several years to refer to the disease, although Haliburton, in 1893, emphasized the fact that many patients did not present that sign.¹ Clinical conditions resulting from TH deficiency will depend on the degree and duration of the deficiency, and will affect basically all tissues to a lower or greater extent. However, it is during intrauterine life that the lack of adequate TH production determines more damaging consequences, since these hormones have a fundamental role in normal fetal brain development.² The advent of molecular biology brought significant advances regarding information on the disease, including elucidations regarding its etiology, which may have an origin during intrauterine life. In the past 3 decades, knowledge about the otogenesis, pathophysiology and early diagnosis of hypothyroidism have grown strongly, and early diagnosis has allowed for intervention on the first days of life of newborn infants (NB), thus preventing damage to neuropsychomotor development. For an adequate TH production, it is important that the hypothalamic-pituitary-thyroid axis be maintained whole so as to ensure the sequence of activities of the hypothalamic releasing hormone (thyrotropin-releasing hormone – TRH) over the pituitary gland, producing thyroid-stimulating hormone (TSH), which in turn acts on the thyroid, producing TH. Deficiencies in these stages lead to tertiary (hypothalamic), secondary (pituitary) or primary (thyroid) hypothyroidism.³,⁴
Physiology

Iodine is an essential element for the synthesis of TH, the only substances in our body that contain iodine in their configuration. Dietary sources of iodine include bread, iodized salt and dairy products. The recommended daily intake of iodine is of at least 75 µg/day, which corresponds to 10 g of iodized salt, according to recommendations of the World Health Organization (one part of sodium iodide in 100,000 parts of NaCl).5

Inorganic iodine present in circulation enters the thyroid follicular cells, where it is organified. This transport depends on the TSH and on a sodium-iodide symporter (NIS), which is located in the membrane of the thyroid cells. In general, an increase in the organic iodine content inside the follicular cells decreases iodide transport; in addition, transport can also be inhibited by some anions, such as perchlorate and thiocyanate. Human NIS has been already identified in breast, colon and ovary cells, and tissues such as salivary glands and gastric mucosa are also capable of concentrating iodide.3 Pendrin, a protein of the Pendred syndrome gene, was described after the performance of studies with patients carrying the syndrome, which consists of an association between hypothyroidism and hearing and speaking impairment. Pendrin also acts in the transport of iodide into follicular cells. Once inside the cell, iodide binds to tyrosine, a thyroglobulin residue. Such iodization is catalyzed by hydroxygen peroxid or peroxidase, whose source is unknown. This passage may be inhibited by thiocarbamides and cyanates.

The thyroid has a limited capacity to use iodides.3 In normal conditions, thyroid iodide clearance rates are higher than organification rates (iodide incorporation into amino acids). Progressively higher concentrations of extracellular iodide increase its transport into the cell until organification reaches its maximum rate; then a sudden decrease is observed, a short-duration phenomenon known as Wolff-Chaikoff effect.

After tyrosine organification, the formation of monoiodotyrosines (MIT) and diiodotyrosines (DIT) already incorporated into thyroglobulin will take place. These hormones will couple to form two main TH: triiodothyronine (T3) and tetraiodothyronine (T4). Thyroglobulin is a large soluble protein with a molecular weight of 660 kDa that is present in the light of the thyroid follicle (colloid). Only three to four T4 molecules are formed in each thyroglobulin molecule, and the thyroid gland usually produces a significantly greater quantity of T4 than T3. The T4 to T3 ratio is 15:1 in normal thyroglobulin. MIT and DIT formation may be inhibited by sulfamides (Figure 1).

Thyroglobulin releases TH by the action of lysosomal proteases inside the follicular cell. Colloid droplets then form on the apical surface of the cell via endocytosis, stimulated by TSH; finally, lysosomes release proteolytic enzymes which will in turn release TH. Considerable amounts of circulating thyroglobulin are only found when the thyroid cell has been damaged. Excess iodide inhibits the release of TH. The treatment of severe hyperthyroidism usually benefits from this effect.3

The TH released in the circulation will bind to carrier molecules: globulin (thyroxine-binding globulin – TBG); transthyretin (TTR), previously called prealbumin (thyroxine-binding prealbumin – TBPA); and albumin. TBG binds 70% of T4 and 80% of T3.5,7

Reverse T3 (rT3) derives from the peripheral monodeiodination of T4.

Mechanism of action of thyroid hormones

Most part of the biological effects associated with TH are determined by interactions between T3 and their specific nuclear receptors. The binding of TH with their nuclear receptors allows the transcription of specific mRNA (nuclear receptors are transcription factors). Nuclear receptors have a high affinity for T3, and their affinity for T4 is 15 times lower. In the normal animal, about 85% of the total iodothyronine that is bound to the nucleus of hepatic and renal cells is of the T3
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type, and only 15% are T4. TH stimulate Na⁺, K⁺ -ATPase in the cell membrane, increasing the consumption of oxygen.

TH may actually be considered a growth factor, and TH deficiency impairs child growth and development, even when the growth hormone (GH) is present. TH act in practically all tissues of the body and influence enzyme concentration and activity, the metabolism of substrates, vitamins and mineral salts, basal metabolism or calorigenesis; they also stimulate the consumption of oxygen and act in other endocrine systems.¹,³

TH stimulate the synthesis and degradation of proteins. The influence of TH on growth is related to its activity in protein synthesis. When TH reach significantly high levels, they accelerate protein catabolism and increase nitrogen excretion.

TH alter the metabolism of carbon hydrates. By increasing the action of epinephrine, they stimulate glycogenolysis and neoglucogenesis and also improve insulin action in glycogen synthesis and glucose use. Low levels of TH increase glycogen synthesis in the presence of insulin, whereas high levels stimulate glycogenolysis. TH also increase the rate of intestinal glucose absorption and its uptake in the adipose and muscular tissues.

TH act on lipid metabolism. In cases of TH insufficiency, a decrease in cholesterol synthesis and its metabolic conversion is observed; however, since the degradation is more affected than the synthesis, blood cholesterol levels become high. The opposite is observed in cases of excess TH, when cholesterol, phospholipid and triglyceride levels are low. One mechanism that may contribute to an increase in cholesterol metabolism in response to TH is the ability of TH to increase the number of low-density lipoprotein receptors on the cell surface. By increasing lipolysis in the adipose tissue, TH affect the metabolism of fatty acids.¹,³

TH are essential for the development of the central nervous system, and deficiency of these hormones during fetal and newborn life extends tissue immaturity, leads to hypoplasia of cortical neurons, delayed myelination and reduced vascularization. If hormone replacement therapy is not carried out soon after birth, lesions will become irreversible, and the child's neuropsychomotor development will be damaged.

**Effects of TH deficiency: clinical features of hypothyroidism**

The early recognition of clinical features in a case of TH deficiency is of fundamental importance and is considered a pediatric emergency in newborn care. Early signs include: prolonged or recurrent jaundice, delay in umbilical cord separation and umbilical hernia. Crying is hoarse, and sounds emitted are low. In the first months of life, other signs become present: feeding difficulty, insufficient weight gain, noisy breathing, nasal congestion, respiratory disorders, obstipation, lethargy, dry, cold and pale skin, with *livedo reticularis.* However, these signs and symptoms are not always evident, and a precious time can sometimes be wasted before treatment is started. This is why the performance of laboratory tests in the nursery room is so important.

Delayed neuropsychomotor development and growth are observed, body proportions are abnormal, and the lower limbs are short if compared to the trunk.

When hypothyroidism is acquired at a later stage, mental retardation may be less evident, but growth will be affected, and these children will present a delay in bone maturation or bone age. In adolescents, hypothyroidism clinical features may show a slower evolution, with tiredness, difficulties at school, intestinal obstipation, dry skin and hair, hair loss, brittle nails, intolerance to cold weather and decreased appetite (it is important to emphasize that obesity is not a characteristic of hypothyroidism). Girls may present menstrual irregularities, and an increase in menstrual cycle periods are more common than amenorrhea.¹

When hypothyroidism remains untreated, more significant physical alterations may be observed in the long term. The skin becomes cereous, pale or yellowish due to carotene impregnation. Myxedema may occur due to the high concentration of mucopolysaccharides in the subcutaneous cell tissue and in other tissues. Movements and bone-tendon reflexes are slow. Some children with severe muscle myxedema show muscular pseudo-hypertrophy and slow muscle action. Myxedema may affect the cardiac musculature, possibly increasing its volume and finally causing stroke.¹

Other endocrine alterations may also be observed in hypothyroidism. Some adolescents may present sexual infantilism, and paradoxically, some others may present precocious puberty. In the long term, thyrotroph hypertrophy may be observed, with an increase in the pituitary gland and in the sella turcica.⁸

**Differential diagnosis**

Differential diagnosis should include Down syndrome, Beckwith syndrome, mucopolysaccharidoses, chondrodystrophies, hypopituitarism, and obesity. It is always important to take into consideration that hypothyroidism is very rarely associated with obesity.

**Congenital hypothyroidism: classification**

Table 1 presents the classification and prevalence of congenital hypothyroidism according to Fisher.⁹

Hypothyroidism manifestations in practically all tissues do not depend on its etiology, but rather on the degree of hormone deficiency.

The causes of thyroid agenesis remain unknown, but there is evidence suggesting an association with mutations in some transcription factors, such as TTF1, TTF2 and PAX8, which are important in thyroid gland embryogenesis.¹⁰-¹⁴
Neonatal screening

Neonatal screening (heel prick test) should be performed in the nursery room, ideally between 3 and 5 days after birth. Many mothers are discharged from hospital before the third day after delivery; dosages performed before the ideal time increase the prevalence of NB with high levels of TSH, due to the physiological increase of this hormone, and may lead to false-positive results. One drop of blood is collected on a filter paper card. Currently, T4 and TSH dosages may also be carried out. TSH values are considered significant when around 20 to 25 µU/mL. Taking into consideration that primary hypothyroidism is the most frequent manifestation of the disease, elevated TSH values allow for early diagnosis and treatment. Normal male NB may present low total T4 levels and normal TSH levels. In these cases, free T4 and TBG carrier values should also be assessed. If free T4 values are normal in the presence of TBG deficiency, this means that the boy is normal, or that the diagnosis of congenital hypothyroidism is discarded. The prevalence of TBG deficiency is 1:5,000 to 1:12,000, and its genetic transmission is related to the X gene.15

Dysgenesis

Agenesis
Hypogenesis
Ectopia

Dishormonogenesis
Absence of TSH response
Defect in iodide transport or uptake
Defect in organification
Defect in thyroglobulin synthesis
Defect in iodotyrosinase deiodinase

Hypothalamic-pituitary
Hypothalamic-pituitary anomaly
Pan-hypopituitarism
TSH deficiency alone
Resistance to TH

Transitory hypothyroidism
Drug-induced
Induced by maternal antibodies
Pregnant women treated with antithyroid or irradiation drugs
Idiopathic

Table 1 - Classification and prevalence of congenital hypothyroidism9

<table>
<thead>
<tr>
<th>Type</th>
<th>Prevalence</th>
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<tbody>
<tr>
<td>Dysgenesis</td>
<td>1:4,000</td>
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<tr>
<td>Agenesis</td>
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<tr>
<td>Hypogenesis</td>
<td></td>
</tr>
<tr>
<td>Ectopia</td>
<td></td>
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<tr>
<td>Dishormonogenesis</td>
<td>1:30,000</td>
</tr>
<tr>
<td>Absence of TSH response</td>
<td></td>
</tr>
<tr>
<td>Defect in iodide transport or uptake</td>
<td></td>
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<tr>
<td>Defect in organification</td>
<td></td>
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<tr>
<td>Defect in thyroglobulin synthesis</td>
<td></td>
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<tr>
<td>Defect in iodotyrosinase deiodinase</td>
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<tr>
<td>Hypothalamic-pituitary</td>
<td>1:100,000</td>
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<tr>
<td>Hypothalamic-pituitary anomaly</td>
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<tr>
<td>Pan-hypopituitarism</td>
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<tr>
<td>TSH deficiency alone</td>
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<tr>
<td>Resistance to TH</td>
<td></td>
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<tr>
<td>Transitory hypothyroidism</td>
<td>1:40,000</td>
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<tr>
<td>Drug-induced</td>
<td></td>
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<tr>
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<td></td>
</tr>
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<td></td>
</tr>
<tr>
<td>Idiopathic</td>
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</table>

TH = thyroid hormone; TSH = thyroid-stimulating hormone.

Neonatal screening

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Dishormonogenesis

Inborn errors of metabolism correspond to about 15% of the causes of congenital hypothyroidism and are associated with enzyme defects of autosomal recessive genetic transmission. When the cascade reactions to TH synthesis are analyzed, it is possible to observe that each inefficient enzyme action alters the cascade, causing deficient hormone production and hypothyroidism. Except for the absence of TSH response, all other forms progress with goiter, which may or may not be present from birth. Several types of hypothyroidism have similar clinical features, and their distinction is only possible based on laboratory tests. The only exception is Pendred’s syndrome, an organization defect associated with both hypothyroidism and hearing and speaking impairment.16,17

Hypothalamic-pituitary hypothyroidism

Central hypothyroidism is relatively rare among NB. Its prevalence is between 1:50,000 and 1:150,000. Up to the 1990s, the dysfunction was considered to be a consequence of trauma associated with delivery. The finding of TH deficiency and image exams revealing posterior pituitary ectopia started to suggest that central hypothyroidism might be part of a broader set of pituitary hormone deficiencies, linked to gene mutations of transcription factors involved in hypothalamic-pituitary embriogenesis. POU1F1 (previously Pit1) gene mutations are associated with a subtype of pan-hypopituitarism that evolves with GH, prolactin and TSH deficiency. In spite of the severity resulting from the presence of these multiple hormone deficiencies, they are rarely diagnosed in the neonatal period. Diagnostic suspicion may occur based on neonatal screening tests showing low levels of T4 and TSH.14,18

Resistance to thyroid hormones

Resistance to TH (RTH) may reveal two different conditions: hypothyroidism, in which all tissues are affected,
also known as generalized RTH syndrome; and hyperthyroidism, which affects the pituitary more severely, also known as pituitary RTH syndrome. There is consensus in that the phenotype of these two defects does not correspond to two different syndromes, but rather reflects a continuum spectrum of a similar molecular defect with variable tissue resistance. TH receptor proteins are coded by two genes: α gene, located in chromosome 17, and β gene, located in chromosome 3. The molecular defect of the cases studied so far involves the β1 receptor of chromosome 3. Inheritance is considered to be autosomal dominant, with 15 to 20% of sporadic cases. RTH patients usually present increased serum levels of T3 and T4 and normal or increased TSH results. Newborn screening programs that primarily assess TSH may detect the condition, since TSH can be slightly or moderately high, and increased T3 remains high).19

Subclinical hypothyroidism

This denomination applies to asymptomatic patients presenting normal T3 and T4 levels and slightly high TSH levels. This type of hypothyroidism is considered to be mild and to represent a risk factor for evolution to overt hypothyroidism and other dysfunctions. Diagnostic implications start with the definition of normal TH levels, more specifically TSH levels. The accepted cut-off point for normal TSH levels is 4 to 5 mU/L, which has been conventionally used to diagnose high concentrations of TSH.20 Some studies have considered lower cut-off points, of 2 to 2.5 mU/L; however, justifications for adopting such numbers were considered insufficient, and as a result it has been recommended that normal TSH levels be maintained at 0.4-4 mU/L. The classification of results between 2 and 4 mU/L as abnormal and the consequent introduction of medication in these cases would most likely have more disadvantages than advantages.

Subclinical hypothyroidism is considered a risk factor for some cardiovascular diseases, hypothyroidism, alterations in lipid and carbohydrate metabolism, neuromuscular symptoms, and decreased energy metabolism. When limits considered to be normal are exceeded, patients should be assessed for signs that justify treatment with levothyroxine: goiter, presence of antiperoxidase and thyroglobulin antibodies, manic-depressive disorders, fertility problems, pregnancy or anticipation of delivery, autoimmune thyroiditis patients (risk for progression of thyroid dysfunction) and children and adolescents with or without goiter (to avoid possible side effects on growth and development).

TSH may return to normal levels spontaneously, without medication, in about 40% of the cases, which explains the origin of controversies about the treatment of subclinical hypothyroidism: cardiovascular risk factors have not been totally proved; there is not a defined standard for TSH normalization; treatment cost and noncompliance are relevant issues; and T4 overdoses may worsen osteoporosis. TSH levels should be monitored carefully to prevent them from going below normal, since T3 and T4 stimulate bone resorption and increase the number of osteoblasts.

If parameters contraindicating treatment are found, it is recommended that clinical and laboratory assessments be carried out every 6 months.

Transient hypothyroidism

In this situation, hormone levels behave similarly as in primary hypothyroidism, that is, low T4 levels and high TSH levels will be observed. The prevalence of transient hypothyroidism varies according to different geographical regions, is related to the intake of iodine and is higher at lower gestational ages. Premature newborns require higher levels of iodine than term newborns in order to maintain a positive balance of iodine and an adequate production of T4 in extrauterine life; therefore, in areas that are geographically poor in iodine, newborns may develop neonatal iodine deficiency. Transient hypothyroidism manifests itself in the first or second week of life, usually associated with transient hypothyroxinemia of prematurity. Treatment is recommended, as this form of hypothyroidism may persist for several months.

Transient hypothyroxinemia

These patients are usually premature newborns with clinical features similar to those of tertiary or hypothalamic hypothyroidism. The condition is transient and resolves spontaneously by the 10th week of life. Treatment is not necessary, except if TSH levels are high.15

Other causes

These can be iatrogenic situations, such as the ones resulting from surgical interventions, antithyroid drug therapy, or radioactive iodine. Hypothyroidism provoked by excessive intake of drugs containing iodine is rare, but should also be considered.

Chronic lymphocytic thyroiditis or autoimmune/Hashimoto’s thyroiditis

In 1912, Hashimoto first described chronic lymphocytic thyroiditis in women with asymptomatic goiter. After surgical
removal of the gland, the author classified them as \textit{struma lymphomatosa}. Later on, in 1938, diagnosis was made in children presenting goiter with lymphocytic infiltrate. Up to 1956, when antibodies were detected, it was considered a rare disease in pediatrics, but since then incidence numbers have been increasing. Currently, Hashimoto’s thyroiditis is considered to be the most frequent thyroid disease in pediatric patients when compared to other autoimmune thyroid diseases.

\textbf{Etiopathogeny}

Chronic lymphocytic thyroiditis (CLT) is basically determined by immunological mechanisms and can be detected in the blood by the presence of antithyroglobulin and peroxidase antibodies. CLT and Graves’s disease are controlled by altered autoimmune processes, and sometimes it is difficult for pathologists to differentiate between both conditions. Cases of patients with classic histological manifestations of CLT and classic clinical manifestations of Graves’s disease have been described. Both processes may appear in the same family and share HLA haplotypes. Major histocompatibility complex (MHC) genes are responsible for different immunological responses, including thyroid auto-antigens. The high incidence of CLT in the female sex at any age group suggests the participation of mutant dominant X chromosome genes, or even an influence by the absence of chromosome Y, with changes in genetic susceptibility potentially associated with chromosomes X and 21. This could explain the high incidence of CLT in Turner and Down syndromes (trisomy 21). There have been reports of families with homozygote twins where one child has CLT and the other Graves’s disease. Such families very frequently are found to carry autoimmune diseases, and sometimes cases of diabetes mellitus, pernicious anemia, myasthenia gravis, rheumatoid arthritis and Addison’s disease are also detected. Although genetic predisposition plays its part in CLT etiopathogeny, few patients evolve to a clinically evident manifestation, and the great majority of patients will most likely remain in a subclinical status, which the authors denominate immunological surveillance status.

\textbf{Incidence}

CLT is considered to be the most common thyropathy among children and adolescents, and it is recognized as the main cause of nontoxic goiter. In an American population with age between 11 and 18 years, five new cases were detected out of 1,000 adolescents screened every year. The incidence is higher among girls, varying from 4:1 to 8:1 depending on the geographical region covered. The disease is rare before 4 years of age and is frequent between 10 and 11 years.

\textbf{Clinical features}

The presence of goiter is one of the main complaints. The gland presents a diffuse increase in volume (two to five times its normal size) and is generally not nodular. The natural history of the disease is as follows: 1) toxic, transient, self-limited thyroiditis; 2) euthyroid goiter; 3) hypothyroidism with/without goiter. However, children may be in any of these phases on the first medical consultation, since there is not a fixed duration for each stage. The clinical course of toxic thyroiditis may vary from weeks to months. In this phase, laboratory data (TH and antibodies) may be confounded with hypothyroidism data. Therefore, it is often difficult to establish a clear clinical profile. Many children may remain euthyroid for some years and then present the clinical features of hypothyroidism.\textsuperscript{21} Children and adolescents with low stature or a progressively lower growth rate, delayed bone age, dry skin and other hypothyroidism-related aspects, even in the absence of goiter, may present a more severe form of hypothyroidism, in which the gland has become fibrotic. Therefore, patients with CLT should be reassessed periodically, with special attention to the finding of nodules on ultrasound, which may require a biopsy puncture to prevent the development of tumor (10 to 25% of these nodules may be carcinomas).\textsuperscript{22,23}

Hashimoto’s encephalopathy, which consists of the involvement of the central nervous system in an encephalopathy status, should be considered in cases of unknown etiology. Adolescents with a positive history for the presence of antibodies, even in euthyroid situations (normal T4 and TSH), and who present a progressive cognitive decline should be assessed. Although the etiology is unknown, a good response to steroid medications suggests an inflammatory or autoimmune dysfunction.\textsuperscript{22,23} Antibodies are considered important markers for the identification of patients who will benefit from an efficient treatment with glucocorticoids.

\textbf{Laboratory features of hypothyroidism}

The diagnosis of congenital hypothyroidism can be confirmed based on T4 and TSH dosages. In the neonatal period, i.e., between 1 and 4 weeks of life, T4 levels < 6.5 µg/dL and TSH levels > 10 mU/L are suggestive of congenital hypothyroidism.

Hypothalamic-pituitary hypothyroidism is characterized by low levels of T4 and normal or even low levels of TSH. Decreased TSH response during the TRH test suggests a diagnosis of central hypothyroidism.

Male children with low total T4 and normal TSH levels should undergo assessment of free T4 and TSH values. This situation may be related with TBG deficiency in normal children, who thus should not be treated for hypothyroidism.

Especially after 4 years of age, in addition to T4 and TSH values, antithyroglobulin and antiperoxidase antibodies should also be assessed to diagnose Hashimoto’s thyroiditis.

The presence of thyroglobulin in serum indicates parenchymal lesion and may be a tumor marker.\textsuperscript{22,23}
Thyroid ultrasound will always be an important laboratory test for the purposes of diagnosis and follow-up. Images showing an irregular texture in the parenchyma are suggestive of thyroiditis. The presence of nodules or cysts deserves special attention in order to discard the possibility of carcinomas.24,25

Two and 24-hour thyroidal radioactive isotope uptake tests with 99mTc or 123I are carried out to diagnose ectopic glands, agenesis or thyroid dysgenesis.26

**Treatment**

In the nursery room, screening can ensure early diagnosis and treatment (in the first 3 to 4 weeks of life), thus guaranteeing an adequate neuropsychomotor development for the NB.

TH replacement is the simplest among all hormone replacement therapies. The drug of choice is levothyroxine (L-T4 sodium salt), which allows measuring serum T4 levels to assess the efficacy of treatment and adjusting doses. Levothyroxine has a mean life of 7 days, and the maximum response is reached in the second week of treatment, when great part of T3 will have been converted. It is administered once a day in the morning. Table 2 shows the recommended doses for different age groups.

These doses may change according to laboratory variations. They should be adjusted whenever signs of overdose are observed: irritability, inability to sleep, red areas on the skin, diarrhea, tachycardia, and sweating. Breastfed infants submitted to high doses of levothyroxine may develop cranioesthesia.

Since these children may present some degree of psychomotor disorder, they should be followed by professionals from the areas of speech therapy, physical therapy and psychopedagogy.27

**References**


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**Table 2 - Recommended levothyroxine doses for children and adolescents**

<table>
<thead>
<tr>
<th>Age</th>
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<td>10 to 15</td>
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<tr>
<td>3 to 12 months</td>
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</tr>
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<td>10 to 16 years</td>
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