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Status for Congenital Hypothyroidism at Advanced Ages

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1. Introduction
Congenital hypothyroidism is one of the most frequent causes of growth and developmental delay and preventable mental retardation. It’s incidence of approximately 1 in 4,000 births (Fisher, 2008). Recently in some countries higher incidences were 1/1800, 1/2759 reported (Skordis et al., 2005; Henry et al., 2002).

2. Classification and etiology
Congenital hypothyroidism is usually classified as primary, secondary and tertiary or as permanently and transient hypothyroidism. Primary congenital hypothyroidism accounts for 90% of all cases. Approximately 85% of cases are sporadic, while 15% are hereditary. Besides dietary iodine deficiency, the causing reasons of permanently congenital hypothyroidism are embryologic and anatomical defect of the thyroid gland, biosynthetic defect in the production of thyroid hormones in sporadic cases. The most common causes are thyroid gland dysgenesis associated with ectopic, hypoplastic and absent gland (athyreosis). The next most common cause of permanent congenital hypothyroidism is dyshormonogenesis (MacGillivray, 2004). Affected patients have normally located and normally shaped thyroid gland but are enlarged due to thyroid-stimulating hormone chronic and hyperstimulation.

The pathogenesis of dysgenesis is largely unknown, some cases are now discovered to be the result of mutations in the transcription factors PAX-8 and TTF-2. Loss of function mutations in the thyrotropin (TSH) receptor have been demonstrated to cause some familial forms of athyreosis. The most common hereditary etiology is the inborn errors of thyroxine (T4) synthesis. Recent mutations have been described in the genes coding for the sodium/iodide symporter, thyroid peroxidase (TPO), and thyroglobulin. Transplacental passage of a maternal thyrotropin receptor blocking antibody (TRB-Ab) causes a transient form of familial congenital hypothyroidism (Brown, 2009).

3. Actions of thyroid hormone and clinical findings in congenital hypothyroid
Thyroid hormone has multiple effects in cells, including stimulation of thermogenesis, water and ion transport and acceleration of substrate turnover and amino acid and lipid metabolism. Thyroid hormone also potentiates the action of catecholamine’s (Brown, 2009).
Whereas thyroid hormone-mediated effects in the pituitary, brain and bone can be detected prenatally, thyroid hormone-dependent action in brown adipose tissue, liver, heart, skin and carcass are apparent only postnatally.

The most prominent finding in view of the rough face of delayed diagnosis. In some cases, depending on the TSH stimulation can be found enlarged thyroid gland (fig 1a-1b). During the perinatal period, brown adipose tissue is essential for non-shivering thermogenesis. Thyroid hormone stimulates transcription of thermogenin that protein that uncouples nucleotide phosphorylation and the storage of energy as ATP. As the child matures, shivering thermogenesis assumes greater importance and brown adipose tissue disappears (Brown, 2009). In children with thyroid hormone deficiency is impaired temperature regulation. Thus the skin is rough and cold skin, and is evidence "cutis marmorata" signs. Hypertrichosis in children with delayed diagnosis and treatment of hypothyroidism can be seen in the back (fig 2).

Fig. 1a, 1b. A 19-year-old girl with congenital primary hypothyroidism. Enlarged thyroid gland, coarse facial, umbilical hernia, thick hair, outpouring of the medial parts of the eyebrows, abdominal distansion and delayed pubertal findings are seen (Nobel Med, 2010; 6(1):74-77).
Other important thyroid hormone target in the perinatal period is bone, as evidenced by the striking growth retardation, decreased growth velocity and delayed ossification of the epiphyseal growth plate characteristic of long-standing untreated hypothyroidism in infancy and childhood. Thyroid hormone-mediated bone maturation involves both direct and indirect actions. The indirect action mediated by regulation of growth hormone gene expression and the IGF system (Robson et al., 2000, 2002). T3 regulates endochondral ossification and controls chondrocyte differentiation in the growth plate both in vitro and in vivo as a direct (Robson et al., 2000; Ball et al., 1997). Osteoblasts and growth plate chondrocytes both express TRs and several T3-specific target genes have been identified in bone (Stevens et al., 2003). T3 also stimulates closure of the skull sutures in vivo, the basis for the enlarged anterior and posterior fontanelle characteristic of infants with congenital hypothyroidism (Akita et al., 1994). Due to delayed bone maturation, there were skeletal deformed, kyphoscoliosis, on thoraco lumbar vertebrae in a 21 year old female with delayed diagnosis of hypothyroidism (fig 3). Bone age is retarded in hypothyroidism almost always exceeds the retardation in linear growth. Tooth eruption may be delayed, and in rare cases stippled epiphyses are evident radiographically.

In the brain, thyroid hormone provides the induction signal for the differentiation and maturation of neural system, and a critical window of brain development. These processes include neurogenesis and neural cell migration (occurring predominantly between 5 and 24 weeks), neuronal differentiation, dendritic and axonal growth, synaptogenesis, gliogenesis (late fetal to 6 months postpartum), myelination (second trimester to 24 months postpartum) and neurotransmitter enzyme synthesis. The thyroid hormone- deficient patient usually exhibits slowing of the deep tendon reflexes, with a delayed relaxation phase. (Brown, 2009). The sella turcica may be enlarged (Oatridge, 2002).
Fig. 3. A 21-year-old female with congenital primary hypothyroidism. Short stature, coarse facial, kyphoscoliosis on her back and absent of pubertal findings are seen (Nobel Med, 2010; 6(1):74-77).

The absence of thyroid hormone appears to delay rather than eliminate the timing of critical morphological events or gene products, resulting in a disorganization of intercellular communication. TRs are found in highest concentration in developing neurons and in multiple areas of the fetal brain, including the cerebrum, cerebellum, auditory and visual cortex. Consistent with a nuclear receptor-mediated mode of action, thyroid hormone stimulates numerous developmentally regulated genes, including genes for myelin, neurotropins and their receptors, cytoskeletal components, transcription factors, extracellular matrix proteins and adhesion molecules, intracellular signaling molecules, as well as mitochondrial and cerebellar genes. In addition, thyroid hormones regulate some genes at the level of mRNA stability or mRNA splicing (Brown, 2009).

Sexual development of most hypothyroid children is delayed in approximate proportion to the retardation of skeletal maturation (fig 1a-3). However, rare children with severe hypothyroidism present with signs of precocious puberty, the Van Wyk-Grumbach syndrome, (Van Wyk & Grumbach, 1960; Hemady et al., 1978; Chattopadhyay et al., 2003). Girls manifest precocious menstruation, breast development, and galactorrhea. In boys, this syndrome is associated with excessive enlargement of the penis and testes. Most of these patients lack pubic hair, and bone age is retarded in keeping with the duration of the hypothyroid state. Serum prolactin and TSH levels are elevated in some children, but the molecular mechanism of precocious puberty is not clear. The increased serum prolactin levels are probably explained by the fact that TRH stimulates TSH and prolactin release from the pituitary. A paracrine action of the hyperstimulated thyrotropic cells on
gonadotrope cells may explain the increased gonadotropin secretion. It is also possible that these patients have genetic variants of the gonadotropin receptors that can be stimulated by the increased TSH levels (Anasti et al., 1995). Similar findings have been reported for TSH and FSH receptor variants stimulated by HCG (Rodien et al., 1998; Montanelli et al, 2004). When the hypothyroid state is alleviated, the manifestations of sexual precocity regress—and normal puberty ensues when the general level of maturity has progressed appropriately.

4. Newborn screening for congenital hypothyroidism

The morbidity of congenital hypothyroidism can be reduced to a minimum by early diagnosis and therapy. Thus mental retardation in affected infants is eliminated completely with treatment. Unfortunately, usually the disease may become evident after many symptoms of the condition leads to an irreversible brain damage. It was reported that, during the first month of birth, only 10% of the congenital hypothyroidism cases were diagnosed by clinical findings while 35% were diagnosed within 3 months after labor and 70% within a year and 100% only within 3-4 years of age, before screening for hypothyroid (Klein, 1972). Diagnosis of hypothyroidism has been delayed in the countries not applied national newborn screening yet (Malik & But, 2008; Tahirović & Toromanović, 2005).

The process involves measurement of T4 and/or TSH on dried blood spots obtained from skin puncture done in first days after birth. In most center, only TSH is used to screen newborn infant, because of primary hypothyroidism is most common causes of congenital hypothyroidism. The cutoff for reporting an elevated TSH is a level above 20-25 U/L in most screening programs.

4.1 Diagnostic criteria

Diagnosis of congenital hypothyroidism has essentially based levels of serum TSH and frees T4. In affected infants presenting with very low serum free T4 and very high TSH levels. Rarely, some infants have only a moderate elevation of serum TSH and normal T4 levels.

Congenital hypothyroidism is usually diagnosed during the neonatal period or early infancy. Sometimes, the diagnosis may be delayed in families with low level of socio-economic, and if the birth at home is frequent in population (Yuca et al. 2010). The newborn screening programs for early diagnosis and treatment is vital in congenital hypothyroidism.

5. Treatment

The primary aim of treatment for congenital hypothyroidism is begin adequate thyroid hormone replacement as early as possible to optimize the prognosis for intellectual development. L-thyroxin is preferred to triiodothyronine, because T4 to T3 convert locally in brain and peripheral tissues.

The starting oral dose of L-thyroxin is 10 to 12 μg/kg/day. The target range for serum is T4 to 10-16 μg/dl. The clinical responses vary among infants even on the standardized dose regimen. Adjustment dose is based on the serum T4 levels and the clinical examination. The patient should be follow at regular intervals.
6. Prognosis in delayed diagnosis and treatment

At 2-year-old and over children may refer to hospital due to uncertain growth and developmental retardation. Untreated congenital hypothyroidism cases may display different levels of mental retardation and delayed linear growth and bone maturation. Infants with delayed treatment may demonstrate neurological disorders such as spasticity and corrupted walking patterns, dysarthria or mutism and autistic behavior (Delong, 1996). Patients receiving treatment with delayed diagnosis is under an obvious target height, but can show some physical growth. These are may gain the skill and awareness of their daily functions, and if they does not speak, will be start talking or improve of talking. If they not walking, are start walk, and have more active movements. It returns to the findings of thermogenesis and skin disorders. The findings of the skeleton will have been better by support therapies such as vitamin D₃ and calcium. In the patients had goiter with pressure symptoms, in fact the thyroid gland is nonfunctional, must be thyroidectomy. Eventually, puberty is developed in the patients with congenital hypothyroid, and they may be fertile. But they cannot reach the mental development accordance with their own age, which is easy for patient with early diagnose and treatment (Oerbeck et al., 2003; Kempers et al., 2006; Josef et al., 2008).

7. Conclusion

Today, there is sensitive radioimmunoassay to measure serum T₄ and TSH using a blood spot made it possible to initiate newborn thyroid screening programs. Affected patients have get out of permanent mental retardation by early diagnosis and treatment with adequate dose of L-thyroxine. Hypothyroidism not only the brain but also the other tissues affect and lead to functional and developmental abnormalities there. Some of these functions are recovered a small amount with long-term treatment. Unfortunately, in the patients diagnosed after the completion of the development of the brain, mental retardation is severe and irreversible despite appropriate therapy.

8. References


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This book aims to provide readers with a general as well as an advanced overview of the key trends in endocrine disorders. While covering a variety of topics ranging from thyroid carcinogenesis and pituitary adenomas to adrenal tumors and metabolic bone disease, this book also focuses on more specific issues not yet fully elucidated (e.g. the molecular pathways involved in thyrotropin beta gene regulation or monogenic phosphate balance disorders). Readers of different fields and background will have the opportunity to update their knowledge and more importantly to clarify areas of uncertainty and controversies in several topics of endocrine disorders.

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