

EDITORAL

Ain Shams University

The Egyptian Journal of Medical Human Genetics

www.ejmhg.eg.net



Subclinical hypothyroidism in children with Down syndrome: To treat or not to treat???



In general, hyperthyrotropinemia and whether to treat or not, is still an unresolved issue. There are no clear guidelines.

Elevated TSH at screening, even when of very short duration, may be a clinically relevant marker of thyroid abnormalities [1]. Data from the Italian National Registry of Infants with Congenital Hypothyroidism reported an association between defects of thyroid development and mild increase of b-TSH at screening. Specifically, 19.6% of babies with a mild increase of b-TSH at screening had defects of thyroid development (thyroid hypoplasia, hemiagenesis, and ectopy) [2].

However, the European Society for Paediatric Endocrinology Consensus Guidelines on Screening, Diagnosis, and Management of Congenital Hypothyroidism, did not reach a solid and clear way of management. They stated that "If venous TSH concentration is ≥ 6 to 20 mU/l beyond 21 days in a well baby with a FT4 concentration within the limits for age, we suggest (a) investigation, which should include diagnostic imaging, to try to obtain a definitive diagnosis; (b) consideration, in discussion with the family, of either initiating thyroxine supplementation immediately and retesting, off treatment, at a later stage; or withholding treatment but retesting two weeks later" [3].

In children with subclinical hypothyroidism, The American Thyroid Association in 2014 declared that treatment is generally not recommended when the TSH is 5-10 mIU/L. For patients with a TSH > 10 mIU/L with signs and symptoms consistent with primary thyroid disease and/or risk factors associated with progression, L-T4 replacement may be reasonable. [Weak recommendation Low quality evidence] [4].

As regards Down syndrome (DS), the therapeutic management of subclinical hypothyroidism remains also object of debate. Very few randomized control trials are presented in this field.

The most important question is whether the TSH elevation reflects mild hypothyroidism that could harm brain growth and development in the youngest children with DS and contribute to the ever present mental retardation. An often-used argument against this hypothesis is the finding that accompanying plasma thyroid hormone concentrations generally are within the age-specific normal range. In addition, it is unknown whether thyroid hormone concentrations within the low-normal range always guarantee optimal thyroid hormone provision of the developing brain [5].

A wait and see policy with frequent thyroid function screening could be considered adequate and may be the preferred strategy, thus avoiding chronic hormonal therapy at least in Down syndrome patients in whom TSH levels tend to spontaneously normalize. A positive anti-TPO antibody test is a key factor in the follow-up of these patients because of its potential risk of progression to manifest thyroid disease [6].

Iughetti et al. (2014), suggest that antithyroid antibodies might represent a marker of deteriorating thyroid function unlike 1st year TSH levels that do not predict future thyroid dysfunction requiring therapy, in subjects with DS [7].

Refetoff (2014) would treat hyperthyrotropinemia in DS only if there is evidence for superimposed autoimmunity [8].

In the last paragraph of the editorial, Elsayed mentioned that "L-thyroxine administration will improve growth, hypotonia and psychomotor functions." I would rather use the word "may" instead of "will" improve.

Recently, Bongers-Schokking et al. (2013) reported that congenital hypothyroidism overtreatment during the first 2 years leads to lowered cognitive outcomes at 11 years, whereas undertreatment, if not complicated by overtreatment, results in a normal cognitive development [9].

In conclusion, more evidence is required regarding the optimal course of treatment for subclinical hypothyroidism in DS. Such evidence may be best obtained by conducting a prospective randomized control trial.

References

- Calaciura Francesca, Motta Rosa Maria, Miscio Giuseppe, Fichera Graziella, Leonardi Daniela, Carta Anna, et al.. Subclinical hypothyroidism in early childhood: a frequent outcome of transient neonatal hyperthyrotropinemia. J Clin Endocrinol Metab 2002;87:3209–14.
- [2] Olivieri Antonella, Corbetta Carlo, Weber Giovanna, Vigone Maria Cristina, Fazzini Cristina, Medda Emanuela. The Italian study group for congenital hypothyroidism congenital hypothyroidism due to defects of thyroid development and mild increase of TSH at screening: data from the Italian national registry of infants

http://dx.doi.org/10.1016/j.ejmhg.2014.10.005

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Peer review under responsibility of Ain Shams University.

with congenital hypothyroidism. J Clin Endocrinol Metab 2013;8:1403-8.

- [3] Léger Juliane, Olivieri Antonella, Donaldson Malcolm, Torresani Toni, Krude Heiko, van Vliet Guy. The congenital hypothyroidism consensus conference group. European society for paediatric endocrinology consensus guidelines on screening, diagnosis, and management of congenital hypothyroidism. J Clin Endocrinol Metab 2014;99:363–84.
- [4] Jonklaas J, Bianco AC, Bauer AJ, Burman KD, Cappola AR, Celi FS, et al. Guidelines for the treatment of hypothyroidism prepared by the American thyroid association task force on thyroid hormone replacement. Thyroid [doi: 10.1089/thy.2014.0028 ahead of print].
- [5] Van Trotsenburg ASP, Vulsma T, Van Santen HM, Cheung W, De Vijlder JJM. Lower neonatal screening thyroxine concentrations in Down syndrome newborns. J Clin Endocrinol Metab 2003;88:1512–5.
- [6] Rubello D, Pozzan GB, Casara D, Girelli ME, Boccato S, Rigon F, et al.. Natural course of subclinical hypothyroidism in Down's

syndrome: prospective study results and therapeutic considerations. J Endocrinol Invest 1995;18(1):35–40.

- [7] Iughetti L, Predieri B, Bruzzi P, Predieri F, Vellani G, Madeo SF, et al.. Ten-year longitudinal study of thyroid function in children with Down's syndrome. Horm Res Paediatr 2014;82(2):113–21.
- [8] Samuel Refetoff: personal communication, 2014.
- [9] Bongers-Schokking Jacoba J, Resing Wilma CM, de Rijke Yolanda B, de Ridder Maria AJ, de Muinck Keizer-Schrama Sabine MPF. Cognitive development in congenital hypothyroidism: is overtreatment a greater threat than undertreatment? J Clin Endocrinol Metab 2013;98:4499–506.

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Available online 1 December 2014