Hypothyroidism

Series N. 15b

Patient’s Guide

Average Readability Leaflet
Hypothyroidism - Series 15b (Revised August 2006)

This leaflet was produced by Fernando Vera MSc and Prof Gary Butler at the Institute of Health Sciences, University of Reading, Reading, UK (August, 2006). Some portions of the text were extracted or modified from the Growth and Growth Disorders Booklet Series (Third edition, 2000)* and may be used in conjunction with these as they provide a choice of leaflets providing the same information, but for people of different ages and reading abilities. The numbering sequence in each series is the same for easy cross-reference. The original leaflet series can be also obtained from the links given at the end.

All illustrations were created and produced by Fernando Vera MSc.

This leaflet is part of the Hormone Disorders Leaflet Series. The following are also available:

Series N 4. Precocious Puberty
Series N 5. Emergency Information for Children with Cortisol and GH Deficiencies and those Experiencing Recurrent Hypoglycaemia.
Series N 6. Congenital Adrenal Hyperplasia
Series N 7. Growth Hormone Deficiency in Young Adults.
Series N 10. Constitutional delay of growth and puberty
Series N 11. Multiple Pituitary Hormone Deficiency
Series N 12. Diabetes Insipidus
Series N 13. Craniopharyngioma
Series N 14. Intrauterine Growth Retardation or Small Gestational Age
Series N 15.a. Hyperthyroidism
Series N 15.b. Hypothyroidism
Series N. 16. Type 2 Diabetes and Obesity

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Introduction

The aim of this leaflet to provide information about hypothyroidism in babies, children and adults. We hope this information will give you a better understanding of this condition and form the basis for discussions with your GP, Paediatrician and Nurse.

What is the thyroid gland?

The thyroid gland is a small gland found in the neck below the larynx (Adam's apple). It makes and releases thyroid hormones to help regulate body growth and metabolism.

What do the thyroid hormones do?

Hormones are messengers used around the body to produce an effect. The main hormone produced by the thyroid gland is thyroxine. This hormone controls the amount of energy used by the body to maintain vital processes such as breathing, circulation and digestion.
Too much thyroxine makes the body work too fast, whereas too little allows the body to slow down. The thyroid hormones also affect brain growth and metabolism in babies in the womb and up to the age of about two years.

What is Hypothyroidism?

Normally, the thyroid gland should be fully developed in all babies by the 22nd week of pregnancy. However, this development can go wrong and lead to underdevelopment or absence of the thyroid gland.

When the thyroid gland is poorly developed or absent, it will not produce thyroxine. This condition is called congenital hypothyroidism. In children, hypothyroidism usually appears as either congenital (from birth) or acquired. There are important differences between these two forms of hypothyroidism, which will be described.

What is Congenital Hypothyroidism?

Congenital hypothyroidism is caused when the thyroid gland does not reach its proper position during development in the womb. In other occasions, the thyroid gland grows normally but fails to produce thyroxine. This condition is present from birth and it’s likely to be passed in families (inherited).

What are the symptoms of congenital hypothyroidism?

Common symptoms in babies include the following:

- Slow feeding
- Sleepiness
- Constipation
- Prolonged jaundice after birth (yellowish skin coloration)

Unfortunately, these are very common symptoms in babies, even when they do not have hypothyroidism. For this reason, all babies are now screened at birth for congenital hypothyroidism with the heel prick test. This test is done in the first week of life of all newborns. If this suggests the existence of this condition, additional testing will be needed to confirm diagnosis.

How is diagnosis confirmed?

A blood sample from the vein will be taken to confirm diagnosis. This test will examine if the levels of thyroxine and TSH are within normal range. Other tests that may be performed include an X-ray of the knee and a thyroid scan.
How is congenital hypothyroidism treated?

Once hypothyroidism has been diagnosed, treatment will start and it will often remain for life. Thyroxine will be given in tablet once a day by mouth. These tablets are available in 25, 50 and 100 micrograms (mcg).

Most doctors will regularly check your baby’s/child’s levels of thyroxine in the blood. This is to make sure that they are on the correct dose. If the dose is much too high, symptoms of hyperthyroidism may develop, with restlessness, weight loss and mild diarrhoea.

Most doctors believe that children with congenital hypothyroidism should remain under the care of a paediatrician with experience in treating the condition.

What is Acquired Hypothyroidism?

Acquired hypothyroidism occurs when the thyroid gland becomes under-active in later childhood. Often this results from a condition in which the body attacks its own tissues. It also may occur because the gland gradually stops working over time.

What are the symptoms of acquired hypothyroidism?

The most common symptoms include:

**Growth problems**: Children may have slow growth and a tendency to put on weight. Puberty may be late, very slow or absent

**Physical changes**: Children may develop a gradual change in facial appearance. The face may develop a rather pale, puffy appearance. Other changes may include constipation, a slow heart rate, some hair loss and slow reflexes.
How is acquired hypothyroidism treated?

Treatment of acquired hypothyroidism is with thyroxine. This is given as a once daily tablet and the dose will vary according to an individual needs.

Generally, a child will be given 50 to 100 mcg a day. Your doctor will review the dose from time to time as it may take sometime to build up to the full replacement dose. The outcome for your child should be excellent. Usually, the symptoms disappear within the first 3 months of treatment.

It’s very important that treatment is taken every day. This can become difficult with older children as they become responsible for taking their own treatment. So, as children become young adults, the importance of taking their treatment should be explained to them.

Who is at risk of developing hypothyroidism?

As already mentioned, thyroid problems may appear in families. Around 40% of children with acquired hypothyroidism have relatives who also have some type of thyroid gland problem. Girls and women are much more commonly affected than boys or men. In addition, the following conditions are considered a risk factor for hypothyroidism:

- Down’s Syndrome
- Turner Syndrome
- Metabolic and Blood Conditions
- Pituitary Gland Problems

What are other sources of useful information?

The goal of this leaflet was to provide a basic overview of hypothyroidism. Further information can be found in the following sources:

- **European Society for Paediatric Endocrinology**  
  ESPE Secretariat, BioScientifica  
  Euro House 22 Apex Court Woodlands, Bristol BS32 4JT - UK  
  Telephone No: + 44 (0) 01454 642208  
  Internet: [http://www.eurospe.org/](http://www.eurospe.org/)

- **British Society for Paediatric Endocrinology and Diabetes**  
  BSPED Secretariat, BioScientifica  
  Euro House 22 Apex Court Woodlands, Bristol BS32 4JT - UK  
  Telephone No: + 44 (0) 01454 642208  
  Internet: [http://www.bsped.org.uk/](http://www.bsped.org.uk/)

- **Child Growth Foundation**  
  2 Mayfield Avenue, Chiswick London W4 1PW UK.
Telephone +44 (0) 20 8995 0257
Internet: http://www.childgrowthfoundation.org/

You can also consult your doctor or nurse for additional information in your local area.