A new TRβ mutation in resistance to thyroid hormone syndrome

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ABSTRACT

Thyroid hormones (TH) exert their actions by binding nuclear receptors alpha (TRα) and beta (TRβ1 and TRβ2). Resistance to thyroid hormone (RTH) is a clinical syndrome with various clinical manifestations, its hallmark being decreased tissue sensitivity to the action of thyroid hormones. We report the case of a family harbouring a novel TRβ mutation. Sequencing of the TRβ gene revealed a single nucleotide substitution-C to G in codon 340: glutamine was replaced by glutamic acid. The clinical picture and biochemical and hormonal panel showed significant differences within the family, despite their sharing the same mutation. We also present the result of low-dose antithyroid treatment in one member of the family diagnosed with this rare condition.

Key words: Mutation, Resistance to thyroid hormone, Single nucleotide substitution, Thyroid hormone receptors

INTRODUCTION

Resistance to thyroid hormone (RTH) is a rare condition with subtle clinical manifestations, characterized by decreased sensitivity of the target tissues to thyroid hormone action.1 This leads to elevated levels of thyroid hormones (TH), accompanied by normal or high thyroid-stimulating hormone (TSH) values, in the absence of intercurrent illness or drug use. At the same time, TH feedback on TSH expression is preserved, but higher doses are required to produce the expected response at both a pituitary and periphery level. The resistance is always partial and the underlying genetic defect is usually, but not always, a mutation in the TH receptor (TR) β gene.2

Given the rarity of the syndrome, diagnosis may be overlooked and subtle abnormalities in lab tests may be considered technical mistakes. Patients are often clinically euthyroid, the most frequent clinical finding being goiter: when this is present, such signs and symptoms as tachycardia, hyperkinesia and emotional disturbances are quite common complaints.3,4

CASE REPORT

A 37-year old Caucasian woman, presenting the characteristics described above, sought medical at-
attention at our Department for recurrent episodes of supraventricular tachycardia. In the absence of a specific cardiological condition, she was treated empirically with beta-blockers and propaphenone, but with limited efficacy. She was referred to our clinic for abnormal thyroid function tests: elevated free thyroxine (FT4) and normal TSH [TSH=1.1 μUI/ml (N=0.5-4.5), FT4=31.8 pmol/l, (N=12-22)].

The initial evaluation revealed normal intelligence, normal body mass index, a grade 2 nodular goiter but no signs and symptoms of hyperthyroidism except mild extrasystolic activity. The hormone panel was repeated using a different assay at another laboratory which confirmed the initial results. Thyroid ultrasound revealed a multinodular goiter (Figure 1), while scintigraphy showed diffuse increased uptake (Figure 2).

The differential diagnosis included two main entities, thyrotopinoma and RTH, for which dynamic testing was required. A thyrotropin-releasing hormone (TRH) test was performed by administrating a 200 μg TRH iv bolus which led to an increase of the TSH level to 12.95 μUI/ml at 30’. Further, the patient was given 87.5 μg/day triiodothyronine (T3) orally for 10 days, which suppressed the TSH level to 0.05 μUI/ml.

This lower dose of T3 was chosen to be given as the outcome of discussion of the patient’s higher cardiac risk at a multidisciplinary team meeting. However, the suppression of TSH at a lower than the standard T3 total dose, together with the positive response of TRH, supported the suspicion of RTH.

The patient also underwent pituitary magnetic resonance imaging which showed a microadenoma of 4.3/4.1/3 mm. The rest of her biochemical pituitary panel was within normal limits. Considering all of the results it was labeled as an incidentaloma.

Further tests were performed to establish the degree of peripheric resistance, and they showed a normal bone turnover (normal osteocalcin and alkaline phosphatase despite elevated T3 and T4) and also hepatic resistance demonstrated by the normal sex hormone-binding globulin (SHBG) level.

The immediate family was investigated and it was discovered that the patient’s sister and niece both shared the hormonal pattern of elevated TH and normal TSH. The patient’s only son had normal thyroid function tests.

We contacted Prof. S. Refetoff (University of Chicago) whose lab performed the genetic tests on this family.

**Thyroid function tests**

Total T4 (TT4), total T3 (TT3) and TSH were measured by chemiluminescence immunometric assays using the Elecsys Automated System (Roche Diagnostics, Indianapolis, IN, USA). Total rT3 (re-
verse T3 or 3,3',5'-triiodothyronine) was measured by a commercial radioimmunoassay (RIA) (Adaltis Italia, Bologna, Italy) and thyroglobulin (TG) by an in-house RIA. The free T4 index (FT4I) and free T3 index (FT3I) were calculated as the product of the total serum concentrations of each iodothyronine and the normalized resin T4 uptake ratio. Antibodies (Ab) against TG and thyroperoxidase (TPO) were measured by passive hemaglutination (Fujirebio, Inc., Tokyo, Japan).

**Molecular analysis**

Genomic DNA was extracted from peripheral blood leukocytes using the QIAamp DNA Blood Mini Kit (QIAGEN, Valencia, CA, USA) in accordance with the manufacturer’s instructions. For sequencing, coding regions (exons 7, 8, 9 and 10) and flanking introns of the TRβ gene were amplified using primers and a thermocycler setting similar to those previously described.7 The resulting polymerase chain reaction products were visualized on 1.8% agarose gel before direct automated sequencing.

**RESULTS**

Sequencing of the TRβ gene revealed a single nucleotide substitution-C to G in codon 340, with glutamine replaced by glutamic acid. This precise mutation has not been reported to date, although a mutation (Q340H) in the same codon has been documented.8

The genetic tests confirmed the diagnosis and highlighted the variability of clinical findings and laboratory tests of this condition; our patient’s sister and her 4-year old daughter were both asymptomatic, but the patient’s niece showed a greater increase in T3 levels, which may be accounted for by decreasing thyroid hormone levels with age. Moreover, the patient’s sister had a thyroid autoimmune disease positive for TG Ab.

Results are illustrated in Figure 3.

We performed an *in silico* analysis using PolyPhen 2 software (Polymorphism Phenotyping version 2 - http://genetics.bwh.harvard.edu/pph2/) to predict the

![Figure 3. Family Mcon (January 2011).](image-url)
pathogenicity of the mutation. The result showed that it is probably damaging with a high score of 0.998 (sensitivity: 0.27, specificity: 0.99).

DISCUSSION

The family we studied fell into the broad category of patients with RTH harbouring a mutation in the TRβ gene.9

RTH represents the main component of a larger group of genetic disorders which includes defects of the transmembrane transporter and of thyroid hormone metabolism.3 Recently, mutations of the SLC16A2 gene have been described, which encodes MCT8, a protein responsible for the transmembrane transport of TH and which is associated with the clinical picture of severe psychomotor disturbance10 and mutations of the SECISBP-2 gene implicated in the synthesis of deiodinases, altering the intracellular metabolism of TH.11

Abnormalities of thyroid hormone-binding proteins (thyroxine-binding globulin and albumin) may resemble the laboratory pattern of RTH, but using equilibrium dialysis to determine T3 and T4 levels12 establishes the diagnosis.

The following refers to the most frequent form of RTH, caused by TRβ gene mutations.

The general features of RTH are elevated serum levels of free T4 and T3, normal or slightly increased TSH level that responds to RTH and increased doses of thyroid hormone and absence of the usual clinical consequences of TH excess and goiter.3,13 Depending on the symptoms, these patients have initially been classified as having selective pituitary resistance to TH if they appeared to be hypermetabolic, or as having generalized resistance to TH if the defect seemed to be compensated by the high levels of TH, with no signs of hyperthyroidism. However, since individuals sharing the same mutation, even within the same family, seem to tolerate differently the elevation in thyroid hormone, it was concluded that the two forms are the result of subjectivity rather than being two separate entities.14

The defect in the RTH syndrome lies in the thyroid hormone receptor, which is a nuclear receptor encoded by two genes, designated alpha and beta, on chromosome 17 and 3. They each generate TRα and TRβ molecules that have substantial structural and sequence similarities. The primary transcript for each gene can be alternatively spliced, generating different alpha and beta receptor isoforms. Of the four different thyroid hormone receptors (alpha-1, alpha-2, beta-1 and beta-2) only 3 are able to bind thyroid hormones. To a certain degree, TRβ and TRα are interchangeable, but some thyroid hormone effects are TR isoform specific. The receptors have specific organ distribution: TRα1 is widely expressed, especially in cardiac and skeletal muscles, while TRα2 is also widely expressed, but unable to bind hormone; TRβ1 is predominately expressed in brain, liver and kidney, while TRβ2 expression is limited to the hypothalamus and pituitary.15

There are three functional domains of the thyroid hormone receptor. A ligand-binding and dimerization domain at the carboxy-terminus, a DNA-binding domain that binds to the sequences of the DNA promoter known as the hormone response elements and a transactivation domain at the amino terminus that interacts with other transcription factors to form complexes that repress or activate transcription. Linking the DNA and ligand-binding domains is the hinge region that contains the signal for nuclear localization.

Mutations have been described almost exclusively in the beta isoform and the vast majority are point mutations. Only one family was reported to have a TRβ gene deletion.16 A few years ago, a mutation was described in the TRα gene in association with a hypothyroid phenotype.17

All TRβ gene mutations are localized in the carboxyl terminus of the TRβ, mostly contained within three CpG-rich “hot spots”, in the ligand-binding domain and adjacent hinge domain. Three mutational clusters have been identified with intervening cold regions. With the exception of the family with the TRβ gene deletion, in all others, to the best of our knowledge, inheritance is autosomal dominant. Mutant TRβ molecules are characterized by reduced affinity for T3 abnormal interaction with cofactors involved in thyroid hormone action and interference with the function of the wild type (WT) receptor.18
In a growing number of individuals, RTH occurs in the absence of mutations in the TRα or TRβ genes (non-TR-RTH).

The precise molecular etiology of these cases is not yet known, one of the hypotheses being a defect in one of the cofactors involved in the mediation of TH action.\(^{14}\)

The more common form of RTH is characterized by minor defects in one allele of the TRβ gene, in contrast to individuals who lack one allele of the TRβ that do not exhibit the RTH phenotype. These findings indicated that RTH is not simply the consequence of a reduced amount of a functional TR (haploinsufficiency) but is caused by the interference of the mutant TR with the function of the WT-TR (dominant negative effect).\(^{19}\)

The variability of the RTH phenotype is explained both by individual tolerance to supraphysiological levels of thyroid hormone and by the distribution of receptor isoforms. The majority of untreated subjects maintain a normal metabolic state at the expense of high levels of TH, but the degree of this compensation of tissue hyposensitivity to the hormone is also variable amongst individuals as well as in different tissues. As a consequence, clinical and laboratory evidence of TH deficiency and excess often coexist.\(^{2}\)

Aside from diagnosis, another issue with regard to RTH is establishing a course of treatment or deciding if one is necessary at all. Considering the disturbing symptomatology in our patient despite the antiarrhythmic treatment, we decided in favour of a low-dose (10 mg/d) thiamazole therapy. Five months later her condition improved, allowing the cardiologist to stop propafenone and to reduce by half the beta-blocker dose, decreasing FT4 at the cost of a small increase in TSH that remained within normal limits.

Since the patient’s sister and niece shared no abnormalities except for an abnormal antibody level, they will remain under our observation, especially for growth monitoring of the 4-year old girl. The absence of any clinical signs or symptoms in their case underlines the differences that may occur even within a family that shares the same mutation in the ligand-binding domain, allowing for variability of signal transduction due to both coexistence of the wild type receptor and intricate corepressor/coactivator mechanisms.\(^{20}\)

Overall, RTH is a rare condition with only slightly more than 3,000 cases described, which may raise difficulties both in determining the diagnosis and in establishing an individualized management: studies show that up to one third of affected patients receive inappropriate treatments (thyroidectomy, radioiodine) before diagnosis is made,\(^{21}\) since genetic testing is relatively expensive and not easily accessible.

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CONFLICT OF INTEREST STATEMENT

No competing financial interests exist.

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A new TRβ mutation in resistance to thyroid hormone syndrome

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