Thyroxin-binding globulin deficiency in a boy with fragile X syndrome: a case report

Abstract:

Fragile X syndrome (FXS) is the most common known genetic cause of male intellectual disability. A wide variety of medical problems has been reported in FXS syndrome including seizures, facial abnormalities, macroorchidism, and autistic disorders. Here we reported a 9-year-old boy with fragile X syndrome who confirmed through karyotyping and mental retardation. Initially he was diagnosed as hypothyroidism when he was 15 months old. However due to unusual clinical presentation, we re-evaluated the patient according to his history and clinical findings. Subsequently, targeted laboratory tests were done and the results were indicative for thyroxin-binding globulin (TBG) deficiency in our patient. Therefore, levothyroxine was discontinued and one month later, laboratory tests were repeated and his diagnosis confirmed. As inherited TBG deficiency might be also X linked, FXS and TBG deficiency may be associated or coincidental findings in the patient.

Key words: Fragile X Syndromes, Thyroid Function Test, Hypothyroidisms, TBG Deficiency

Introduction:

Fragile X syndrome (FXS) is a common genetic disorder which is considered as the most common cause of male mental retardation. It might be presented in various clinical manifestations including mental retardation, cognitive and behavioral disorders, macroorchidism, joint hyperlaxity, and some facial abnormalities such as large forehead, narrow face, prominent chin, large and anteverted ears [1-3]. Approximately, the FXS incidence is about one out of 2000-5000 individuals [4] and it is more prevalent in males than females [1].

FXS is a chromosome X-linked disorder caused by a dynamic mutation in exon 1 of the FMR-1 (Fragile X-linked Mental Retardation type 1) gene. This gene is located on the band q27.3 of the chromosome X. This dynamic mutation occurs as a result of an expanded CGG (cytosine-guanine-guanine) trinucleotide repeats [5]. However 25-30% of these patients do not present the typical characteristics of FXS [1]. Several studies have investigated the role of thyroid dysfunction in FXS [6-8]. It has been hypothesized that thyroid dysfunction may contribute to macroorchidism in males with FXS [9, 10].

Thyroxin-binding globulin (TBG), a 54 kilodalton glycoprotein, is the major transporting protein of thyroid hormone in human serum [11]. TBG consists of 395 amino acids and it is located on the long arm of the X chromosome between bands q21 and q22 (q22.2) [12]. Therefore most of the TBG abnormalities are inherited in x-linked manner [13]. Whereas estrogen excess is
among the causes of increased serum TBG concentration, corticosteroids administration decrease the serum TBG concentration independently [14,15].

TBG deficiency is either acquired or inherited through X-linked inheritance. The cause of both TBG deficiency forms stems in TBG gene defect(s). Mutations in this gene may result in partial or complete TBG deficiency, which is sometimes accompanied by other genetic defects such as carbohydrate deficient glycoprotein syndrome type 1 (CDG1), an autosomal recessive disease. Some of the TBG deficiency acquired causes include hyperthyroidism, chronic renal or liver failure, malnutrition, HIV/AIDS, and severe systemic illness [16].

Case presentation:

A 9-year-old boy was referred to our pediatric endocrinology clinic, Imam Reza Hospital, Mashhad, Iran with an initial diagnosis of hypothyroidism. According to his past medical history, he was born via normal vaginal delivery with a normal Apgar score at birth. His birth weight was 3450 grams. His mother stated that he had weakness, delayed crying, and poor feeding in infancy. His growth pattern seemed to be normal, whereas his development was retarded as he was unable to sit before 15 months. In addition, his speech was retarded, he was easily irritated, and frequently cried and knocked his head. He was not toilet-trained (voiding control) before 4 years old.

In his familial history, he was the second child in a family of four. The parents were not related. His mother and elder sister were receiving levothyroxine for hypothyroidism.

At 15 months a total T4 of 3.1 µIU/ml (normal range: 5.1-12.5 µIU/ml for 7-16 y) and a TSH of 6.8 µIU/ml (normal range: 0.6-5.2 µIU/ml for 7-12 y) were noted and accordingly, the patient was treated with levothyroxine. Periodic laboratory results are listed in Table 1. In addition, it should be noted that the patient previously received clonidine, risperidone, biperiden, and carbamazepine due to his behavioral disorders. No abnormalities were reported in his brain CT scan.

At the time of admission to pediatric endocrinology, the patient’s height was 150 cm (Height Z-score: 2.54), his weight was 42 kg (weight Z-score: 1.6), and his body mass index was 18.6. Physical examination revealed macroorchidism, as the testes size with orchidometer was more than 3cc (normal size in prepubertal boys is 1-3 cc), and puberty was in Tanner stage one. Based on his initial examination and past medical history, TBG deficiency was suspected. Therefore, levothyroxine was discontinued and laboratory tests were repeated after one month. The results were the following: TSH: 3.2 µIU/ml, Total T4: 4.8 µIU/ml, T3RU: 42%, Free T4: 24 pmol/L.

Thyroid laboratory profiles in the patient’s mother and sister also indicated TBG deficiency. Following confirmation, their levothyroxine was discontinued as well. One year after cessation of therapy, the mother’s laboratory results were as the following: TSH: 2 µIU/ml, Total T4: 2.9 µIU/ml, T3RU: 29%, Free T4: 9.1 pmol/L. Considering the probable diagnosis of FXS based on
mental retardation and macroorchidism, genetic evaluation requested for the patient and FXS 46, XY, Fra(x) (q27.3) was reported.

Table 1: Thyroid functional tests in the patient

<table>
<thead>
<tr>
<th>Date</th>
<th>TSH (µIU/ml)</th>
<th>T4 (µg/dl)</th>
<th>T3 (µg/dl)</th>
<th>T3RU (%)</th>
<th>Free T4 Index (ratio)</th>
<th>Free T4 (pmol/l)</th>
</tr>
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<tr>
<td>2002 Aug</td>
<td>6.8</td>
<td>3.1</td>
<td>67</td>
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<tr>
<td>2002 Sep</td>
<td>1.2</td>
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<td>110</td>
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<tr>
<td>2002 Oct</td>
<td>1.5</td>
<td>5.2</td>
<td>170</td>
<td></td>
<td></td>
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<tr>
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<td>4.2</td>
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<tr>
<td>2003 Nov</td>
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<td>2</td>
<td>24(H)</td>
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<td>&lt;2</td>
<td>44(H)</td>
<td>15.4</td>
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</tr>
</tbody>
</table>

In the latest follow-up, when he was 10 years old, the patient’s height was 152 cm and his weight was 60 kg. The thyroid profile was TSH: 3.7 µIU/ml, Total T4: <2 µIU/ml, T3RU: 44%, FT4: 15.4 pmol/l. He was under observation by pediatric psychologists for his developmental retardation.
Peripheral blood samples were collected from the patient in order to karyotyping evaluation. With GBG banding (G-bands after trypsin and Giemsa) in 100 metaphases in a folate deficient medium, fragile site was observed (see figure 1).

Figure 1. The karyotype of the patient, 46, Y, fraX(q27.3) with X chromosome showing fragile site at Xq27.3 (arrow shows this site on terminal end of X chromosome long arm).

Discussion:

FXS is a common chromosomal disorder and understanding new aspects of this syndrome is likely to be useful for diagnoses and treatment. It is responsible for about 30% of all mental retardations and is associated with diverse clinical manifestations [1, 2]. However 25-30% of patients with FXS do not present typical clinical signs [1]. The presented patient was a 9-year-old boy with macroorchidism and TBG deficiency and a karyotyping result compatible with FXS.

Several studies have investigated the role of thyroid dysfunction in FXS [6-8], but up to our best knowledge, this is the first report of TBG deficiency accompanied by FXS syndrome.

Coincidence or association of fragile X syndrome with TBG deficiency is quite rare. It seemed that as the inherited TBG deficiency may be X linked, FXS and TBG deficiency might be associated. However, it remained controversial whether they are coincidental or associated and requires further investigation.

Some investigators have evaluated thyroid function in FXS [9, 10]. Bregman and colleagues [9] evaluated thyroid profiles in twelve 3-28 years old males with FXS. Their study demonstrated that T4, TBG, and EFT (estimated free thyroxin) values were within normal ranges. No correlations between age with serum T4 and EFT levels were observed. TSH response to thyrotropin releasing hormone (TRH) was blunted and baseline prolactin was elevated in these subjects. These results supported hypothalamic-pituitary abnormalities in FXS [9]. In our case, EFT, TSH, T3RU were normal, but due to TBG deficiency total T4 was low.

FMR1 and TBG genes are located on Xq27.3 and Xq22.3, respectively. Togetherness of these conditions would be possible if they were part of contiguous gene syndrome due to a large deletion on this site. But, the gene sites are not contiguous. Moreover, FXS is a triplet repeat (allelic expansion) disorder, not a deletion syndrome. There is no reported similar case so far. Therefore, coincidence is the most probable explanation for this togetherness.

Conclusion:
In summary, we presented a 9-year-old boy with FXS and TBG deficiency in himself and his mother and sister. Because TBG deficiency may be inherited X linked, FXS and TBG deficiency might be associated. However, it is controversial whether they are coincidental or associated and requires further investigations. In many syndromes with mental retardation in boys such as Down syndrome, fragile X syndrome and Klinefelter syndrome, thyroid functional tests are abnormal. We should interpret thyroid functional tests carefully because misinterpretation of tests in mental retarded boys may lead to delayed or overlooked diagnosis of hypothyroidism.

Consent

Written informed consent was obtained from the patient’s father for the publication of this report and any accompanied images.

Conflict of Interest: None declared.

Acknowledgment: None.

References:


