Novel splice-affecting variants in CYP27A1 gene in two Chilean patients with Cerebrotendinous Xanthomatosis

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Abstract

Cerebrotendinous Xanthomatosis (CTX), a rare lipid storage disorder, is caused by recessive loss-of-function mutations of the 27-sterol hydroxylase (CYP27A1), producing an alteration of the synthesis of bile acids, with an accumulation of cholestanol. Clinical characteristics include juvenile cataracts, diarrhea, tendon xanthomas, cognitive impairment and other neurological manifestations. Early diagnosis is critical, because treatment with chenodeoxycholic acid may prevent neurological damage. We studied the CYP27A1 gene in two Chilean CTX patients by sequencing its nine exons, exon-intron boundaries, and cDNA from peripheral blood mononuclear cells. Patient 1 is a compound heterozygote for the novel substitution c.256-1G > T that causes exon 2 skipping, leading to a premature stop codon in exon 3, and for the previously-known pathogenic mutation c.1183C > T (p.Arg395Cys). Patient 2 is homozygous for the novel mutation c.1185-1G > A that causes exon 7 skipping and the generation of a premature stop codon in exon 8, leading to the loss of the crucial adrenoxin binding domain of CYP27A1.

Keywords: Cerebrotendinous Xanthomatosis, splicing, mutation, exon skipping.

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Introduction

Cerebrotendinous xanthomatosis (CTX) [MIM #213700] is an autosomal recessive disease caused by mutations in the CYP27A1 gene, which encodes the mitochondrial enzyme 27-sterol hydroxylase (EC 1.14.13.15) involved in bile acid synthesis. The primary bile acids cholic acid (CA) and chenodeoxycholic acid (CDCA) are synthesized by two complementary chemical pathways: the classic “neutral pathway”, considered as the main producer of bile acids (approximately 50% CA and CDCA) and an alternative “acidic pathway” leading to CDCA synthesis. The 27-sterol hydroxylase encoded by CYP27A1 catalyzes the initial step in the acidic pathway, with the C27-hydroxylation of cholesterol oxidation of the side chain sterol intermediates in bile acid formation (Sundaram et al., 2008). As a consequence of inherited genomic defects in CYP27A1, there is an alteration of the production of bile acids, mainly decreasing CDCA production. The immediate consequence of this genetic defect is the compensatory increase in the activity of the rate-limiting enzyme in the neutral bile acid synthesis pathway, cholesterol 7α-hydroxylase, leading to the accumulation of 7α-hydroxylated bile acid precursors, in particular 7α-hydroxy-4-cholestene-3-one. This molecule is a precursor of cholestanol, which is elevated in the plasma of CTX patients, as well as an increase in
plasma cholestanol/cholesterol ratio (Björkhem and Hanson, 2010). Consequently, these molecules are abnormally deposited in different tissues such as central nervous system (CNS) and crystalline lens (Khan et al., 2013).

CTX diagnosis is based on clinical grounds. Recently, an Italian group of experts in CTX has proposed a suspicion index for early diagnosis (Mignarditi et al., 2014). Elevated levels of cholestanol levels in serum, high bile alcohol levels in urine and genetic testing, confirm the diagnosis. According to the literature, there is no established genotype-phenotype correlation pattern in CTX. (Lee et al., 2001).

At early stages of CTX, symptoms are childhood diarrhea and bilateral cataracts, which are usually diagnosed during the first or second decade of life (Cenedella, 1996; Moghadasi et al., 2002; Pilo-de-la-Fuente et al., 2011). In more advanced stages, the disease is characterized by tendon xanthomas as well as the presence of progressive neurologic dysfunction signs that usually occur during the second and the third decade of life. Less frequently, the presence of seizures, coronary heart disease and early atherosclerosis has been also described (Dotti et al., 2000).

Although the prevalence of the disease has been roughly estimated in 1/50,000 (Salen et al., 1983), this figure varies by ethnicity and geographic location. In populations of Sephardic Jews of Moroccan origin, a prevalence of 1/108 has been described (Berginer and Abeliovich, 1981), while in Caucasian populations of the United States of America, the estimated prevalence is approximately 3-5/100,000 (Lorincz et al., 2005). There are, so far, more than 50 different CYP27A1 mutations published (http://www.hgmd.org). In Chile, two CTX patients have been previously reported (Filippi et al., 2009; Huigen et al., 2012). The aim of the present report is to describe the genetic, molecular and clinical characteristics of two Chilean CTX patients.

Subjects and Methods

Clinical description of patients

This study was approved by the Ethical Review Board of the School of Medicine of the Pontificia Universidad Católica de Chile. Written informed consent was signed by patients and relatives.

Patient 1: female, 42 years old. No history of consanguinity. History of unexplained chronic diarrhea and bilateral cataracts diagnosed at age of 5 and operated in childhood. Since age of 28 she developed paraparesis and progressive difficulty to walk. Neurological examination in that moment showed bilateral pyramidal syndrome, cerebellar syndrome and deep sensory deficit in lower extremities. Brain magnetic resonance imaging (MRI) showed involutive signs incerebellar and bilateral frontal regions, with white matter involvement bulb and cerebellum. An electrophysiological study of the lower extremities showed signs of chronic sensorymotorpolyneuropathy. Laboratory tests showed negative results for HIV, HTLV-1, and VDRL. Plasma B12 and folic acid levels were normal. Thyroid function was normal. The patient showed a normal lipid profile except for slightly elevated high density lipoprotein cholesterol (HDL-C) (74.6 mg/dL). At age of 31, CTX was suspected, although without presenting typical tendinous xanthomas. This patient showed a concentration of 64 μmol/L of plasma cholestanol reported from an external laboratory (normal values: 2-12.6 μmol/L). At the moment of recruitment for this study, the patient was prostrated, with a severe spastic tetraparesis, flexor pattern of the four limbs, severe dysphagia that required installation of a percutaneous endoscopic gastrostomy. There is no clinical data of her bone condition at the moment of diagnosis. Due to her clinical condition, it was not possible to perform a bone densitometry at the moment of the recruitment.

Patient 2: male, 17 years old. Patient was adopted at age of 5 months and no data of his biological parents are available. He was hypotonic at infancy, without feeding or respiratory impairments. He had motor and developmental delay, with independent walking after 2.5 years old and language (phrases) after 2 years old. Mother refers frequent stools (twice daily with low consistency by periods) since toddler. Difficulties in visual acuity were noticed at the age of 10 years. Miopic astigmatism and bilateral cataracts were diagnosed at 13 years old. On neurologic examination, the following signs were evident: macrocephaly, dysmorphic signs with mild facial hypomimia, distal hypotrophy in lower limbs; xanthomas on Achilles tendons (both) and one on triceps tendon. He had speaking, reading, writing and drawing difficulties and distal paresis with hyperreflexia with no spasticity in lower limbs. Fine hands tremor; difficulty in walking on heels and subtle instability in turns were evident. Brain MRI revealed T2 signal abnormalities on cerebellar hemispheric white matter. Thyroid function was normal. Bone densitometry showed low bone mineral mass in both hips and normal spine values (Left femoral neck Z-score -3.3; right femoral neck Z-score -3.2). Levels of 25(OH) vitamin D were 14.2 ng/mL (normal values: 20-60 ng/mL). Plasma cholestanol levels were analyzed in an external laboratory yielding a result of 2.6 mg/dL (normal values 0.17 ± 0.12 mg/dL, for children 3-16 years), thus confirming the diagnosis.

Molecular and bioinformatic methods

Total DNA and RNA were extracted from peripheral blood in EDTA tubes using Mini blood kits (Qiagen) and Trizol (Ambion) respectively. Genomic DNA was amplified and sequenced (Macrogen, Korea) using primers designed to amplify the 9 intron/exon boundaries, as well as the 3' UTR of the CYP271A1 gene (NG_007959.1). Primer sequences and thermal profiles are available under request from authors. RNA (NM_000784.3) was reverse transcribed with random primers (Applied Biosystem).
Genomic sequence variation was described according to the guidelines of the Human Genome Variation Society (Den Dunnen and Antonarakis, 2000).

To assess the possible impact of the amino acid substitution p.Arg395Cys on protein structure and function, we used the Polyphen-2 and PROVEAN (v1.1) programs (Adzhubei et al., 2010; Choi et al., 2012). The two novel intronic mutations found in our patients (Table 1) disrupt the canonical splicing 3' acceptor site. Subsequently, we used the Human Splicing finder (HSF) software (Desmet et al., 2009), which predicts the effect of the splice affecting variants with high accuracy based on position weight matrices. Natural splice sites (5' and 3'), branch points and exonic splice enhancers among others are assigned with consensus values. A delta Consensus Value (ACV) of the splice sites strength > 10% is likely to have an impact on splicing (Desmet et al., 2009). When mutations affecting intron positions -1,-2, +1 or +2 occur, the probability of exon skipping, the use of pseudo 3' or 5' splice sites, retention of the mutated intron or cryptic splice sites, among others, is higher (Faustino and Cooper, 2003).

Results

We found three different mutations in the CYP27A1 gene, listed in Table 1. Patient 1 is a compound heterozygote harboring the novel mutation c.256-1G > T and the previously described missense mutation c.1183 C > T (p.Arg395Cys), shown in Figure 1A. Patient 2 is homozygous for the novel mutation c.1185-1G > A, shown in Figure 1B. We have registered the new CYP27A1 mutations in the Leiden Open Variation Database with accession numbers as following: c.256-1G > T: # DB_ID:CYP27A1_00069, and c-1185-1G > A: # DB_ID:CYP27A1_00068. The third mutation c.1183C > T or p.Arg395Cys (patient 1) has accession number: # DB_ID:CYP27A1_00004 (http://www.lovd.nl/3.0/home), is shown in Figure 1A.

The analysis of the novel mutation c.256-1G > T on the splicing process, using the HSF program, predicted that the nucleotide T lowers the strength (ACV -32.9%) of the natural splicing acceptor 3' site (AG > AT) and also seems to swap the 3' site to a 5' donor site with a ΔCV -14.0. This would predict a highly plausible skipping of exon 2 (191 nucleotides long). This imprecise splicing may shift the reading frame creating a new amino acid sequence of 30 residues containing a premature stop codon (GRTPLPAAAPGSEPVAEAASGSALYGCFQ*).

In order to demonstrate the effect of the mutations found in patient 1, we reverse transcribed total RNA extracted from leukocytes of peripheral blood. The cDNA was amplified and sequenced in two segments: exon 1 to 4, and exon 5 to 9. The sequence was concordant with the compound heterozygote genotype found in the genomic DNA, as both mutations were present. The cDNA sequence
of exon 1 to 4 of showed the extended haplotype sequence containing the intronic mutation c.256-1G > T and the normal haplotype. The main peaks correspond to the nucleotide sequence of exon 2, while the weak background peaks of the sequence correspond to the nucleotide sequence of exon 3 (Figure 1C). This result indicates that exon 2 is incorrectly spliced and exon 1 is joined to exon 3 in the extended haplotype carrying the c.256-1G > T mutation. Thus, we have demonstrated the existence of exon skipping, which was previously predicted by the HSF program. The mutation c.256-1G > T has been named as p.Val86Glyfs30Ter (Table 1).

Patient 1 cDNA also carry the c.1183C > T (p.Arg395Cys) variant, which is located in the cDNA sequence corresponding to a TGT codon, codifying for cysteine, as shown in Figure 1D. The first two nucleotides of this codon come from exon 6 and the third nucleotide comes from exon 7, with the c.1183C > T mutation thus being located in the penultimate site of exon 6. The cDNA sequence analysis showed that there is no splice defect derived from this mutation, given that exon 6 is correctly joined with exon 7. On the other hand, the c.1183C > T mutation generates a change in the amino acid sequence (p.Arg395Cys) previously known to be pathogenic. According to bioinformatics predictions, Polyphen-2 showed that p.Arg395Cys was “probably damaging” with the highest score possible of 1.0 (range 0.0 to 1.0). The tridimensional model of CYP27A1 (±A) shows that Arg395 interacts with Phe438 by pi-cationic interaction and is also involved in h-bond interactions with the carboxyl group of Glu392 and the main-chain nitrogen from Pro441 (Figure 2B upper panel).

We found the novel mutation c.1185-1G>A in the “-1” site of intron 6 in homozygous state in patient 2 (Figure 1B). The analysis of this mutation using HSF software showed a decrease in the strength of the 3’ splice site (ΔCV -30.7%) and MaxEnt revealed a ΔCV of -119.7% in strength variation. Both values can be considered strong evidence to predict a splice defect. Following the same procedure used in Patient 1, we reverse transcribed total RNA and amplified and sequenced the cDNA from patient 2. The cDNA sequence of exon 5 to exon 9 of CYP27A1 of patient

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Figure 1 - Genomic and cDNA sequences showing case 1 and case 2 mutations. A) Case 1 genomic sequence showing heterozygote mutations. Black arrows point to mutated nucleotides. B) Case 2 genomic sequence. Homozygous mutation shown by arrow. C) cDNA sequence of case 1 (Exons 1-3). One haplotype has the normal sequence of exon 1 joined to exon 2, while the second haplotype shows exon 1 incorrectly joined to exon 3 (small peaks) (RefSeq NG_007959). D) cDNA sequence of case 1 showing exon 6 and 7 with the mutation at codon 395. E) cDNA sequence of case 2 showing the join of exon 6 to exon 8, skipping exon 7.
2 shows that exon 6 is joined to exon 8 (Figure 1E), demonstrating that the homozygous mutation did affect normal splicing of exon 7. We expect that the mutated protein loses 136 amino acids of the original sequence and only preserves sequence identity up to Arg395, after this position there is a new sequence of 28 amino acids (PSLCSATMWCPTPLPSLSLKASSPTAG*), which finalizes with a premature stop codon, generating a predicted truncated protein of 423 amino acids. The nomenclature for this mutation is p.Leu396ProfsTer28 (Table 1).

Discussion

As a consequence of the metabolic impairments observed in Cerebrotendinous xanthomatosis, it has been proposed that the primary treatment of this disease would be oral CDCA administration, which is reported to decrease plasma cholestanol levels (Sundaram et al., 2008). Statins may also produce a synergistic effect by lowering the cholesterol pool and improving the clinical response (Watts et al., 1996; Verrips et al., 1999; Federico and Dotti, 2003).

Patient 1 is a compound heterozygote for two CYP27A1 variants: c.256-1G > T and the missense mutation c.1183C > T (p.Arg395Cys). The c.256-1G > T mutation has not been previously reported in other case series. We show here molecular evidence based on cDNA analysis that such a mutation leads to a splice defect, also supported by bioinformatics analysis. In this context, the c.256-1G > T mutation causes exon 2 skipping and a premature stop codon in exon 3 (p.Val86Glyfs30Ter). The second mutation found in patient 1 (c.1183C > T), alters the protein sequence (p.Arg395Cys). This mutation (also termed p.Arg362Cys) is the most common mutation found in CTX patients of Spanish origin (Pilo de la Fuente et al., 2011). It is very likely that its presence in Chile is derived from migrations from Spain, one of the main population contributors to the current Chilean population. It was shown by Cali et al. (1991) that the p.Arg395Cys mutation dramatically reduces enzyme activity. In our study, we provide support to this observation using bioinformatics tools. According to our tridimensional protein model, position 395 is located within the ERR triad, which may act as a folding motif, stabilizing heme binding and redox partner binding (Prosser et al., 2006). It is predicted that mutation in Cys395 may increase distances and the loosened interactions may favor misfolding, thus affecting adrenoxin (ADX) binding (Figure 2B lower panel).

It is worthy of note that the mutation c.Arg395Ser is located in the same genomic position (penultimate nucleotide of exon 6) as the c.Arg395Cys (codon CGT to TGT; c.1183C > T) (Chen et al. (1998)). Given the special location of this mutation in the exon-intron boundary, Chen et al. (1998) proposed that the c.Arg395Ser mutation may affect normal splicing and gene expression efficiency, implying that this specific genomic DNA position may also affect gene expression in other mutations located in the same genomic site. As far as we know, there are no reports assessing the possible impact of the mutation c.1183C > T on gene expression or the splicing process. Although we have not determined whether c.1183C > T affects gene expression efficiency, we show here that the sequence containing the c.1183C > T mutation is fully transcribed (Figure 1D).
Therefore, it is very likely that the sole effect of the amino acid change generated by the amino acid change c.Arg395Cys is sufficient to impair enzyme activity and cause CTX (Cali et al., 1991).

Regarding the c.1185-1G > A mutation found in homozygous state in patient 2, we also show bioinformatic and molecular evidence supporting a role in the CYP27A1 splicing process. We show that this mutation causes exon 7 skipping and the generation of a premature stop codon in exon 8. As a consequence, the protein lacks the two binding domains for its cofactors adrenodoxin (ADX) and ADX reductase, as well as relevant residues for HEM coordination such as Cys476. It is also possible that this patient lacks the CYP27A1 protein, as the stop codon generated by exon skipping is located > 55 nucleotides upstream of the most 3’ exon-exon junction, possibly leading the activation of the Nonsense-Mediated mRNA Decay (NMD) surveillance pathway (Karam et al., 2013).

It is necessary to emphasize the importance of clinical and biochemical CTX screening in young patients with early disease signs, such as neonatal cholestatic jaundice, childhood diarrhea, and congenital or juvenile bilateral cataracts, given that a treatment with CDCA is available. In the absence of national screening programs, ophthalmologic findings in patients with CTX are interesting since they are one of the first signs of the disease, often before diagnosis of CTX is made. It is important to emphasize that early treatment during childhood and before CNS deposits appear, may avoid progression of the disease.

In conclusion, we have reported herein the genetic, molecular and clinical characteristics of two Chilean CTX cases, showing two novel splice-affecting variants in the CYP27A1 gene. Since CTX is a treatable disease, we propose to screen new CTX cases among children with early disease signs, such as bilateral cataracts, followed by testing their plasma cholestanol levels and searching mutations in the CYP27A1 gene in the patients who are positive for biochemical screening.

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