Introduction

Cerebrotendinous Xanthomatosis (CTX) is a rare autosomal recessive disorder, caused by a deficiency of the mitochondrial enzyme 27α-sterol-hydroxylase, which is encoded by the CYP27A1 gene (2q33-qter) [1] is characterized by an abnormal accumulation of cholesterol and cholestanol affecting many tissues including the brain [2]. The most common clinical manifestations include xanthomas, cataracts, and progressive neurological dysfunction [1]. The first case of CTX was described in 1937 by van Bogart [3], since then, more than 300 cases have been reported around the world [4]. CTX is rarely reported in Latin American countries, where biochemical and genetics testing have limited availability [5]. We present a Peruvian CTX that reveal the clinical and biochemical features revealing the diagnosis.

Case presentation

A 49-year-old male (III-1 individual, Figure 1), experienced 10-year history of gait disturbances and progressive postural instability with frequent falls. He also developed slurred speech over the last 3 years. According to his parents, he had learning difficulties at school and quit during elementary school. He had several sporadic and temporary jobs until mid-thirties. He stopped working collaborating in household chores.
The neurological examination revealed moderate dysarthria, spasticity in all extremities with increased tendon reflexes, cerebellar ataxia, dysmetria and bilateral dysdiadochokinesia, as well as multidirectional nystagmus in primary gaze. Physical examination revealed xanthomas in both Achilles’s tendons (Figure 2). The ophthalmological evaluation revealed bilateral sub capsular cataracts. Cognitive assessment demonstrated significant cognitive decline scoring 9/30 on MoCA test. Neuropsychological battery found IQ of 43 points affecting verbal fluency, attention, concentration, abstraction, calculation and working and visuospatial memory.

Nerve conduction study revealed mixed mononeuropathy in the right median nerve. MRI neuroimaging showed hypointensities surrounded by external hyper intensities in both cerebellar dentate nucleus and global brain atrophy (Figure 3). Laboratory tests found both serum cholesterol and triglycerides in normal range. Cholestanol levels were measured in external laboratory and found it markedly elevated 951 ng/ml (NV: <25 ng/ml). Genetic testing was not performed. Patient evolution was stationary for the next 6 months. Unfortunately, chenodeoxycholic acid was not available locally, and then he was put on atorvastatin 20 mg/day. Patient get lost on follow-ups.

Discussion

We presented a case of a normolipemic adult patient with Achilles tendinous xanthoma and progressive neurological symptoms. Head MRI studies and plasma sterol analysis support the diagnosis of Cerebrotendinous xanthomatosis. CTX is a rare and underreported lipid storage disorder. In Peru, a case with a late clinical diagnosis has been published [5]. The accumulation of cholestanol in various tissues (crystalline, tendons, brains, blood vessels, etc.) determines its clinical manifestations, our patient presented sub capsular cataracts of early onset, it often appear during the first decade of life [6], also, we observe in the Achilles tendon, which appear in the second or third decade. Neurological symptoms of our patient was slow progression, such as signs of the pyramidal tract, cerebellar ataxia and peripheral neuropathy. The learning difficulty of our patient is described since childhood, however, the cognitive deterioration was evidently progressive after the age of 30, typically cognitive decline has been described around the age of 20 [7]. The diagnosis is often overlooked at initial presentation due to the lack of a classic clinical picture, additionally routine biochemical tests of blood, urine, and cerebrospinal fluid are often normal, except plasma sterol analysis. In our case, the cholestanol dosage could be performed, obtaining 951 ng/ml (NV: 25 ng/ml), this biochemical result being compatible with the disease. The brain MRI findings support the diagnosis in our patient; the most characteristic neuroimaging lesions have seen in the basal ganglia, cerebral peduncles and dentate nucleus [7]. In case of our patient, we can observe bilateral low intensity of the dentate nuclei surrounded by hiperintensity lesions, as well as global atrophy of the cerebellum (Figure 3). This disorder can be treat with Chenodeoxycholic Acid (CDCA), which provides negative feedback for the bile acid biosynthesis pathway therefore the treatment consists in suppressing the production of cholestanol and bile alcohols [8]. Starting treatment at an early age can improve or previse neurological injurie and cognitive decline. Unfortunately, the drug is not available in Peru, so in this case we choose atorvastatin 20 mg/day.

Conclusion

We present a case with clinical and neuroimaging criteria characteristic of CTX, which was confirmed with a biochemical study of cholestanol dosage. The diagnosis of this case was very late, with a high risk of permanent sequelae despite being a disease that, treated early, can avoid many complications. The possibility of CTX should be considered in all patients with xanthomas, juvenile cataracts, and neurological manifestations. Biochemical and genetic diagnostic studies should be implemented that allow a timely diagnosis of these cases.
Declarations

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Conflicts of interest: The authors report no disclosures relevant to the manuscript.

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