Bilateral Xanthomas of Tendoachilles in A Patient of Cerebrotendinous Xanthomatosis - A Rare Case Report and Review of Literature

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Abstract

**Introduction:** Cerebrotendinous xanthomatosis is a very rare disorder of cholesterol metabolism with autosomal recessive inheritance. It can present to the out-patient clinics of orthopaedic department with painful and swollen tendoachilles on one or both sides. It is commonly mistaken for a local disorder and excision may be recommended without realizing its systemic nature.

**Case Report:** We find such a rare and perplexing case in a 19 year boy who presented with painful swellings of both achilles tendons causing signification limitation of walking distance. This was initially interpreted as a localized benign disorder and was offered surgical treatment. Excision of the swollen achilles tendon followed by reconstruction using peroneus brevis tendon was done, first on the more symptomatic right side. The diagnosis of cerebrotendinous xanthomatosis was made retrospectively after histopathological as well as biochemical analyses and appropriate medical therapy was initiated.

**Conclusion:** Cerebrotendinous xanthomatosis commonly manifests with bilateral swollen tendoachilles, which is actually a part of systemic pathology with poor prognosis. High suspicion is warranted and surgical excision should not be the primary option without the confirmation of the cause. A detailed history with clinical assessment followed by the laboratory investigations certainly leads to the correct diagnosis. The treatment is more of medical means and appropriate medication can even retard the disease progression, if initiated early in life.

**Keywords:** Cerebrotendinous xanthomatosis; bilateral xanthomas; tendoachilles

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Website: www.jocr.co.in

DOI: 10.13107/jocr.2250-0685.223

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Introduction

Cerebro-tendinous xanthomatosis (CTX) is a rare disorder of cholesterol metabolism with an autosomal recessive inheritance. It is characterized by the lack of production of primary bile acids from cholesterol and accumulation of a byproduct called cholestanol in various tissues like brain, spinal cord, peripheral nerves, eyes, lungs, liver, kidneys, tendon xanthomas and bile [1]. The disease starts with chronic diarrhea and cataract in the early decades of life. The painful swellings of tendoachilles usually appear in the second decade. Full blown disorder soon follows resulting in low intelligence, convulsions and abnormal movements. Out of all, the central nervous symptoms are more disabling and the situation continues to worsen with the advancing age making them completely dependent. Here we describe a classical presentation of this rare disorder that has to be considered in the differential diagnosis of commonly occurring tendon xanthomas secondary to conditions [2,16] like familial hypercholesterolaemia and other hyperlipidaemias in whom surgical excision is the correct option. Whereas the present condition is a different one with systemic nature and has poor prognosis if the diagnosis is missed. Postoperative recurrence is common if surgical excision is carried out for tendon xanthomas as happened in this case.

Case Report

A 19 year old boy presented with bilateral slow growing and painful swellings of his achilles tendons of 4 year duration. On clinical examination each swelling was of size 5x2 cm, firm, nodular, tender and localized to the distal portion of the tendoachilles just above its insertion point to the calcaneal tuberosity (Fig. 2). He was more symptomatic on the right side. He was initially advised rest, analgesics and foot wear modification. However, the symptoms did not subside and the situation worsened resulting in significant disability with limitation of his walking distance.

Initial radiographs of both ankles showed homogenous soft tissue shadow in the lower halves of tendoachilles (Fig. 3). Magnetic resonance imaging showed localized homogenous hyper intense signals with fusiform swellings of tendoachilles (Fig. 4). Surgical excision was offered and the right sided one was selected first as it was more symptomatic. The swollen tendinous portion measuring 6x3 cm was excised (Fig. 5) followed by reconstruction using the ipsilateral peroneus brevis tendon. Immediate and early post-operative period was uneventful. Biopsy of the excised specimen revealed it to be a xanthoma characterized by the accumulation of mononuclear cells with foamy cytoplasm and multinucleated giant cells with high concentration of cholestanol. Thus a diagnosis of cerebro-tendinous xanthomatosis was made.

On detailed retrospective inquiry, he had surgery for bilateral juvenile cataracts at the age of 8, along with history of chronic intractable diarrhea. He was coherent and co-operative but situation worsened resulting in significant disability with limitation of his walking distance. Out of all, the central nervous symptoms are more disabling and the situation continues to worsen with the advancing age making them completely dependent.

Here we describe a classical presentation of this rare disorder that has to be considered in the differential diagnosis of commonly occurring tendon xanthomas secondary to conditions [2,16] like familial hypercholesterolaemia and other hyperlipidaemias in whom surgical excision is the correct option. Whereas the present condition is a different one with systemic nature and has poor prognosis if the diagnosis is missed. Postoperative recurrence is common if surgical excision is carried out for tendon xanthomas as happened in this case.

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Discussion

Cerebrotendinous xanthomatosis (CTX) is a rare genetic metabolic disorder of cholesterol and bile acid metabolism that results in systemic and neurologic abnormalities [1,4] The primary enzymatic defect is in mitochondrial sterol 27-hydroxylase, a key enzyme in the complicated process of bile acid synthesis from cholesterol [2,7]. Defects in this enzyme result in decreased synthesis of primary bile acids, this, in turn disrupts feedback regulation on cholesterol 7-alpha-hydroxylase, which is the rate limiting step in bile acid synthesis [1,6]. Therefore, bile acid precursor – cholestanol accumulates in various tissues. Deposition of cholestanol in the central nervous system (the brain and spinal cord), muscles (including the heart and eye-lens), tendons, blood vessels and other tissues results in a degenerative process that worsens over time unless treated [3,4,5].

The disease was first described in 1937 by Van Bogaert and his colleagues [3]. In 1968, Menkes et al described the accumulation of cholestanol, the primary metabolite found in elevated concentrations in cerebrotendinous xanthomatosis, in tissues of the CNS [4]. In 1971, Salen found that chenodeoxycholic acid (CDCA), an important bile acid, was virtually absent in patients with clinical symptoms of the disease [5]. This led to successful trials of therapy with CDCA replacement therapy by Salen and colleagues in 1975 [6]. In 1980, defects in mitochondrial 27-hydroxylase were implicated in the biochemical pathophysiology of the disease by Oftebro et al [7]. In 1991, mutations in the gene CYP27A1 were discovered as causative [8,9,10]. Since then, more than 50 mutations have been implicated.

The clinical manifestations are due to lack of bile acid synthesis and accumulation of bile acid precursors. Morbidity begins with intractable diarrhea, with onset typically in infancy due to defect in the synthesis of bile acids. The diarrhea continues through adulthood if left untreated [11]. Neonatal or infantile hepatitis and prolonged jaundice have been described [12]. Juvenile cataracts seen as early as 5-6 years of age, may be the presenting sign [13], resulting in vision abnormalities due to accumulation of cholestanol in lens tissues. Xanthomas are rarely seen before the second decade. They are usually found on the achilles tendons but may also be found on the patella, elbow, hand, and neck tendons. They have also been reported on the parenchyma of the lungs and brain, as well as in the bones.

Musculoskeletal xanthomas can cause motor restriction and joint deformities, resulting in ambulation disabilities. Vascular abnormalities such as premature atherosclerosis (especially in the carotid and coronary vessels) due to excess cholestanol deposition can lead to stroke and myocardial infarction. Significant neurologic impairment with low intelligence, seizures, dementia and extra pyramidal dysfunction typically begins in the third decade of life and progress until death, if left untreated. This is because of the accumulation of high serum cholestanol levels for a prolonged period which have a toxic effect on the neurons and myelinated axons leading to atrophy and secondary demyelination [14]. If CTX is not diagnosed, the life expectancy is into the fifth and sixth decades; however, deaths have been reported even in the second and third decades. This is a progressive and terminal disease if left untreated. Treated patients may have a normal life span.

The differential diagnosis of tendon xanthomatosis includes familial hypercholesterolemia, leukodystrophy and sitosterolemia. These patients have accelerated atherosclerosis with tendon xanthomata, but absence of neurological symptoms and diarrhea differentiates them from CTX [15]. Our patient showed all the typical features of this rare disorder like infantile diarrhea, juvenile cataracts, tendoachilles xanthomata, low intelligence and short stature except seizures and symptoms of extra pyramidal dysfunction. The serum cholestanol level was typically elevated to 4.37 mg/dL (the normal levels range from 0.02-0.12 mg/dL).

The treatment of choice in cerebrotendinous xanthomatosis is chenodeoxycholic acid (CDCA) replacement therapy. The recommended dose is 15 mg/kg/day orally in 3 divided doses would normalize the cholestanol levels after as little as 4 months of treatment. Neurological, gastrointestinal and musculoskeletal symptoms improve gradually along with the feeling of general well-being.

The common order of presentation of this rare disease, as explained, is infantile intractable diarrhea, juvenile cataracts, adolescent tendon xanthomata followed by adulthood neurological manifestations. Appearance of tendon xanthomas is an important clinical hallmark of the disease. If tendon xanthomas are not present, a diagnosis of CTX often will not be

Figure 4: MRI scan T1 weighted image showing homogenous hyper intense signals with fusiform swelling of both the tendoachilles.

Figure 5: Excised piece of the affected tendon during surgery on right side, measuring 6x3 cm.

Figure 6: Post-operative follow up after 2 years revealing painless and minimal enlargement of the operated tendon on the right side.
made, unless biochemical tests are performed. So all the patients with achilles xanthomas need to be evaluated with a detailed history, clinical examination, followed by serum cholestanol estimation. Surgical temptation in the form of excision of the swollen achilles tendons may miss a dreadful, yet treatable condition.

**Conclusion**

There are different causes of tendoachilles swellings, of which one of the rare but a medically treatable condition is cerebrotendinous xanthomatosis. Early recognition of this entity is crucial as it has got an effective medical therapy that can prevent further progression and even reverse the disease manifestations. Surgery should not be the option for the tendon xanthomata without the confirmation of the cause as it could be a part of a systemic pathology.

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