Case Report

Ptosis as the Main Presenting Sign in a Patient with Leigh Syndrome

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INTRODUCTION

This case report highlights a rare but lethal disease which can have ophthalmic signs as the presenting signs. Leigh Syndrome (LS, subacute necrotizing encephalopathy) is a progressive neurodegenerative disorder with onset usually in infancy or early childhood [1-2]. It is characterized by basal ganglia, brainstem and thalamus changes (symmetrical hypo densities in the basal ganglia on CT or hyperintense lesions on T2-weighted MRI) but with considerable clinical and genetic heterogeneity [3-4]. The estimated incidence of LS is 1:40,000 live births. LS is associated with defects in mitochondrial energy production [5]. Defects of the mitochondrial respiratory chain are associated with a diverse spectrum of clinical phenotypes, and may be caused by mutations in either the nuclear or the mitochondrial genome (Mt-DNA)[5]. Frequently encountered clinical signs include motor and/or intellectual delay and signs of brainstem dysfunction like respiratory abnormalities [6]. Other neurological manifestations include ataxia and dystonia. Ophthalmologic manifestations are nystagmus, ophthalmoparesis, strabismus, pigmentary retinopathy, optic atrophy and ptosis [7]. Typically, symptoms begin in the first months and progress to death within 2 years, but later onset and/or slower progression can occur [6].

REPORT OF A CASE

A 16-month-old girl presented with a history of progressive drooping of the left upper eyelid over a 5-month period which worsened after a head concussion 3 months prior to presentation. Her primary care physician diagnosed developmental delay. There was no discernable eyelid ptosis on photographic review of the patient’s early infancy however there was an esotropia.

On ophthalmologic evaluation, visual acuity was decreased with fixation but only brief following behavior in both eyes. Her pupils were reactive to light with no relative afferent pupillary defect. On external exam, bilateral ptosis (left upper eyelid more than the right upper eyelid), asymmetric Bruckner’s reflexes and a 15 Prismatic Diopters (PD) exotropia were noted (Figure 1).

Bilateral optic nerve pallor was noted on fundus examination. Cycloplegic retinoscopy revealed mild ‘against the rule’ hyperopic astigmatism. Due to findings on our examination including variable strabismus during infancy, bilateral optic nerve pallor, developmental delay and acquired bilateral ptosis, neurologic and metabolic evaluation were recommended.

Neurologic exam revealed: upper extremity hypotonia with occasional head titubation and normal symmetrical reflexes. Brain MRI revealed “Symmetric reduced diffusion [and T2/FLAIR hyperintense lesions] were noted in bilateral medial thalami, cerebral peduncles, periaqueductal grey, ventral pons and medulla, substantia nigra and inferior cerebellar peduncles” (Figure 2). This pattern of involvement was concerning for an underlying metabolic etiology, in particular Leigh syndrome. Genetic testing was performed. Mitochondrial-DNA (Mt-DNA) sequencing revealed ND5 gene 13094T > C Val253Ala mutation.

A few days after the MRI scan, the patient developed a rapid and progressive deterioration systemically due to her mitochondrial disease. She quickly developed abnormal breathing, associated with congenital stenosis of her pulmonary valve and brainstem dysfunction caused by the LS, as well as paralysis of the eye muscles due to a progressive ophthalmoplegia. Dyspnea led to hospitalization where the patient unfortunately passed after a few days secondary to cardiopulmonary arrest, as was consistent with the natural history of her newly diagnosed disease. Exome sequencing was not able to be completed due to her sudden worsening status.

DISCUSSION

LS is a little known but fatal disease that can present with ophthalmic signs. Eyelid ptosis can be the main presenting sign in patients with LS as in other mitochondrial disorders. These patients can initially be misdiagnosed as having juvenile myasthenia gravis because the presentation of ptosis in LS is variable. Han et al. [7]
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reported ophthalmologic manifestations in 44 patients with childhood onset LS. Among them, only seven patients (15.9%) had a ptosis phenotype while ptosis was the initial sign in four patients. Sudo et al. [5] reported on 6 patients with LS caused by ND5 gene mutation in the Mt-DNA. Among these 5/6 (83.3%) patients had eyelid ptosis which was the initial sign in most (4/5) of these patients. Rahman et al. [8] found the ptosis phenotype in 6/35 patients (17.1%), almost the same percentage of Han et al. [7].

Therefore, LS should be included in the differential diagnosis of patients who develop eyelid ptosis or gaze-palsy, and these patients should be followed to check for developmental delay, gait disturbance, or other neurologic symptoms [7]. In conclusion, acquired eyelid ptosis is a clinical manifestation that an ophthalmologist must examine in depth and carefully. Multidisciplinary management with a pediatric ophthalmologist, pediatric neurologist, geneticist, neuroradiologist and possibly a metabolic specialist is fundamental in patient assessment due to the lethality of the disease.

REFERENCES