Leber’s (Plus?) Hereditary Optic Neuropathy: A Case Report

Arife CA1*, Cansu S2* and Ufuk E1

1Department of Neurology, Istanbul Education and Research Hospital, Istanbul, Turkey
2Istanbul Education and Research Hospital, Istanbul, Turkey

Corresponding author: Arife CA, Department of Neurology, Istanbul Education and Research Hospital, Istanbul, Turkey, Tel: 905338141817; E-mail: cimenatar@yahoo.com.tr

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Abstract

Leber’s hereditary optic neuropathy (LHON) is a maternally inherited genetic disorder of young males and characterized by severe progressive vision loss with other neurological and systemic symptoms. Here we present a young male with subacute progressive vision loss and Parkinsonism symptoms like right arm rigidity and endocrine abnormalities like hypoparathyroidism as a probable LHON-plus case.

Introduction

Leber’s hereditary optic neuropathy (LHON) is an important cause of progressive painless visual loss among young male patients [1]. It is characterized by bilateral subacute loss of central vision owing to focal degeneration of the retinal ganglion cell layer and optic nerve [2]. Recently new case reports of subacute visual failure and additional prominent neurological features like dystonia, myoclonic jerks, ataxia, multiple sclerosis like symptoms are described and defined as LHON-PLUS cases [3,4]. Here we present a case with bilateral subacute visual loss accompanied by different neurological examination and cranial magnetic resonance imagining (MRI) findings as a probable LHON-plus case.

Case Report

17 years old male patient applied to an outpatient ophthalmology clinic with bilateral progressive painless blurred vision starting one month ago. There was bilateral optic disc pallor in fundoscopy and his visual accurate was diminished to counting fingers from one meter. The patient was diagnosed as optic neuropathy and referred to our clinic by ophthalmologist for neurological assessment. His personal background was unremarkable. His uncle had a history of bilateral optic atrophy, the pupils were reactive to light and there was an afferent pupillary defect. The orbital MRI results were normal. In the cranial MRI there were bilateral symmetrical hypertensins lesions in T2 and FLAIR and hypointens lesions without contrast enchancement in T1 sequences at mesencephalon (Figure 1). In visual evoked potentials (VEP) there were bilateral loss of P100 responses. The organic aminoacid levels for detecting mitochondrial cytopathies in urine were in normal ranges. Laboratory results of parathormone was 3.2 pikogram (pg)/millileter/ml (normal ranges are 10-65 pg/ml) and 25–hydroxyl vitamin D: 7.8 ng/ml (normal ranges: >30 ng/ml). In cerebrospinal fluid (CSF) examination the protein levels were slightly elevated, the IgG index, lactic acid and piruvic acid levels were significantly high. The CSF lactat/piruvate ratio was normal. Serum copper level (70 microgram/desyleter), ceruloplasmin level (20.8 mg/dl) and 24 hour-urine copper levels (3.0 micrograms per day) were performed for differential diagnosis of Wilson disease and all results were normal. The electroencephalopy (EEG) examination showed generalized bioelectric disorganisation. Maculray optic coherans tomography and fundus angiography results were normal. We planned genetic testing for mtDNA analysis for confirmation but the patient declined.

As a result the presence of subacute bilateral optic neuropathy with left arm rigidity, apathy, positive family history, the MRI findings, and exclusion of other reasons of optic neuropathy lead us to LHON-plus diagnosis.

Discussion

LHON is a maternally inherited disorder characterized by acute or subacute visual loss leading to a central scotoma [5]. It is the most common cause of blindness in young males [6]. Over 95% of LHON cases are primarily the result of one of three mitochondrial DNA point mutations [7]. First in 1988 at the 11787/ND4, then in the following years the 14484/ND6 and 3460/ND1 mitochondrial DNA mutations were defined [2,3]. These three mutations involve genes encoding complex I subunits of the respiratory chain [2]. In 5-10% of the cases different types of mutations were defined [4]. Central visual loss is the only symptom of the disease in majority of the LHON cases, but recently new cases are defined in which optic neuropathy is with different neurological symptoms like dystonia, parkinsonism, cerebellar ataxia and myoclonus [2-4]. These cases are called LHON-plus. Among these abnormalities multiple sclerosis (MS) like symptoms, spinal cord disease, brainstem and basal ganglia involvement, Charcot-Marie-Tooth and skeleton abnormalities, progressive hearing neuropathy can be mentioned [8]. In Niskoskelainen et al. study 46 LHON patients were included and 59% had different types of additional neurologic symptoms like tremor, parkinsonism, dystonia, epilepsy, migraine, dementia, polyneuropathy, MS like disease, brainstem involvement, toracal kifosis and pes cavus [8]. Funalot, et al. reported a three patient series of LHON symptoms accompanied by a Leigh-like encephalopathy [9]. All these patients had symmetrical bilateral brainstem lesions like LHON in their cranial MRI and these lesions significantly regressed at
the follow-up [10]. Also different cases with bilateral vision loss and MS like lesions in the white matter (LHON-MS) and dystonic syndrome with pediatric onset ‘with striatal necrosis sites in cranial MRI where 14459 mtDNA mutation is positive, are described. None of these cases had brainstem lesions [8,11]. Other neurological symptoms accompany to LHON are summarized at Table 1.

![Image](image.jpg)

**Figure 1:** Hyperintens symmetrical pathologic signal changes in bilateral mesencephalon at FLAIR and T2 sequences in cranial MRI.

<table>
<thead>
<tr>
<th>Age/Gender</th>
<th>Neurologic Symptoms</th>
<th>MRI findings</th>
<th>Mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Funakawa I,1995</td>
<td>Cerebellar ataxia, dysarthria</td>
<td>Cerebellar atrophy</td>
<td>Mitochondrial DNA 11778 mutation</td>
</tr>
<tr>
<td>Murakami T,1996</td>
<td>Cerebellar ataxia</td>
<td>Cerebellar atrophy</td>
<td>Mitochondrial DNA 11778 mutation</td>
</tr>
<tr>
<td>Shoffner JM</td>
<td>Dystonia</td>
<td>Bilateral basal ganglia lesions</td>
<td>ND6 subunit complex mitochondrial DNA LDYT14445A point mutation</td>
</tr>
<tr>
<td>Parry-Jones 2008</td>
<td>32/female</td>
<td>Periventricular white matter spinal cord</td>
<td>Mitochondrial DNA,T14484C mutation</td>
</tr>
<tr>
<td>B ceranic, 2004</td>
<td>45/female, 59/male</td>
<td>Hearing loss</td>
<td>Mitochondrial DNA 11778 mutation</td>
</tr>
<tr>
<td>W kucker, 2007</td>
<td>36/female</td>
<td>Spastic paraparesis</td>
<td>Bilateral periventricular white matter involvement</td>
</tr>
<tr>
<td>W leuzzi, 1992</td>
<td>20/female 11/male</td>
<td>Dystonia hypokinesia, bradykinesia myoclonic jerks hypokinesia bradykinesia bilateral putaminal right n.caudatus head involvement bilateral putaminal involvement</td>
<td>mitochondrial DNA 11778 mutation</td>
</tr>
</tbody>
</table>

**Table 1:** Neurological symptoms accompany to LHON.

We finally considered the patient as LHON-plus with progressive bilateral vision loss, right arm rigidity and apathy in neurological examination and bilateral, symmetrical pathological signal changes at mesencephalon in cranial MRI. We searched the literature and could not find any other case report with similar MRI findings. The patient had an uncle with the same vision loss history but we could not
perform genetic study to other family members. Our patient also had primary hypo-parathyroidism which supports the mitochondrial disease diagnosis. There are some case reports of mitochondrial disease in literature where hypo-parathyroidism is presented as the initial symptom [12,13].

As a result in a male patient with subacute progressive optic neuropathy the presence of other neurological symptoms can suggest a LHON-plus phenotype. Clinicians must be aware of these ‘plus’ phenotypes to ensure early diagnosis, and patients in this high-risk group are best served by a multidisciplinary team minimize the functional consequences of these systemic complications.

References