Pelizaeus–Merzbacher Disease: the First Genetically Approved Case Report from Iran

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Abstract

Background: Classic Pelizaeus-Merzbacher disease is a rare x-linked disorder of proteolipid protein expression first described clinically in 1885. This disease is characterized by abnormal eye movements, very slow motor development and involuntary movements. The causative gene is PLP1.

Case Presentation: A 1-year-old boy was referred to our clinic due to abnormal eye movements. He had horizontal and flickering eye oscillation, psychomotor retardation, hypotonia and head nodding. We found hypomyelination in brain MRI.

Conclusion: The possibility of Pelizaeus-Merzbacher disease should be considered in boys with abnormal eye movements, psychomotor retardation and hypotonia.

Key Words: Pelizaeus-Merzbacher Disease; Eye Movements; Hypotonia; Head Nodding

Introduction

Classic Pelizaeus-Merzbacher disease is a rare x-linked disorder of proteolipid protein expression that was described clinically by Pelizaeus in 1885 and later in 1910 the clinical and pathologic features of the same family were documented by Merzbacher [1].

The clinical features of this disease are slowly progressive and consist of peculiar pendular eye movements, head nodding, hypotonia, mental retardation, abnormal movements (dystonia, chorea) and pyramidal signs. Also, optic atrophy and seizures may occur in this disease.

Typically, the progression of the disease slows towards the middle or end of the first decade, with death occurring in the early adult ages [2].

This disease results from point mutations that result in amino acid substitutions or complete deletion/duplication of the PLP1 gene. These gene
products play a major role in the development of oligodendrocytes for myelination [3]. The PLP gene is linked to the Xq21-22 region [4].

Brainstem auditory-evoked responses and somatosensory-evoked potentials are consistently abnormal [5]. In some cases, visual-evoked potentials are abnormal [6]. Brain MRI is helpful in demonstrating the hypomyelination that is characteristic of this disorder [6].

**Case Presentation**

This 1-year-old boy is the first child of non-consanguineous parents. He was born at term following an unremarkable pregnancy and delivered by cesarean section. Apgar score at 5 minute was 9. Birth weight 2900 gram, head circumference 34 centimeter. At birth, he had axial hypotonia and limbs hypertonicity. At 2 months, binocular nystagmus was noted, which stabilized by 6 months of age. At 8 months of age, he presented with head nodding and at times, he was noted to have markedly increased tone of both lower extremities. At present time, he has global developmental delay and progressive limbs spasticity with abnormal eye movements and axial hypotonia.

Visual and brainstem evoked potentials are abnormal.

MR study reveals poor myelination and dysmyelination of periventricular and sub cortical white matter in T1W and T2W images comparing normal myelination process for the patient’s age. Similar finding in infra-tentorial white matter is also seen (Fig. 1 and 2).

DNA investigated for mutations and deletion/duplication by sequencing revealed duplication in the PLP1. This confirmed the diagnosis Pelizaeus-Merzbacher disease.

**Discussion**

Pelizaeus-Merzbacher Disease (PMD) is a rare x-linked inherited disorder characterized by hypomyelination. In fact this disease is a hypomyelinating leukoencephalopathy. In PMD, normal myelination either never occurs or is incomplete.

In 1885, Pelizaeus and later in 1910, Merzbacher described this disorder for the first time[7].
PMD has traditionally been divided into four categories: classic, connatal, transitional and adult form (spastic paraplegia)\[7\].

The most common form is classic PMD (our case). Symptoms in classic PMD are usually manifest during infancy. These patients present with nystagmus, head nodding, hypotonia, dystonia, ataxia and cognitive delay. Hypotonia typically evolves into spastic para paresis over time. Prior to the progressive phase, the disease is often misdiagnosed as cerebral palsy\[7\].

Proteolipid protein 1 gene (PLP1) mutations are responsible for PMD. This gene encodes the major myelin components in the CNS, the two proteolipid proteins, PLP and its spliced isoform, DM20. PLP is produced by mature oligodendrocytes while DM20 is produced earlier in myelin development \[8\].

The most common mutation of PLP1 gene is duplication (70%) and the most common clinical phenotype of PMD caused by PLP1 duplication corresponds to the classic form of PMD (our case)\[9\]. More than 60 point mutations in the PLP1 gene coding region have been identified \[9\].

Brain MRI images in PMD typically demonstrate a pattern of hypomyelination with a reversal or discordance of white matter signal intensity on T2 weighted images. T1 weighted images demonstrate normal or isointense white matter\[10\].

In patients with PLP1 duplication, the brain MRI had some degree of myelination, while no myelination was noted in most patients with PLP1 point mutations\[11\].

In this study, our case suffered from manifestations of classic PMD and had no new or unusual signs or symptoms. He had global developmental delay and progressive spasticity with abnormal eye movements and axial hypotonia with increased peripheral hypertonicity.

**Conclusion**

The possibility of PMD should be considered in boys with abnormal eye movements and hypotonia in early infancy. Also with early diagnosis, we can determine prognosis based on genetic study.
References


