Conflict of Interest

All authors have no financial disclosures to report.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author on reasonable request.

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Supporting Data

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.

Both Heterozygous and Homozygous Loss-of-Function JPH3 Variants Are Associated with a Paroxysmal Movement Disorder

Recently, a single patient with a homozygous truncating variant in JPH3 and a neurodevelopmental disorder involving paroxysmal dystonia was described.1 We report both a second individual with recessive disease and a family with a milder phenotype and autosomal dominant inheritance (Fig. 1A).

Patient one, now 15 years old, was born to first-cousin parents with no relevant family or perinatal history. Her mother was asymptomatic, but the medical history of her father was unavailable. Early infant milestones were normal, but she did not walk until 3 years old. At school, she was identified as having moderate intellectual disability and behavioral difficulties. From 6 months old, her mother noticed episodes of abnormal body stiffening. Over time, these evolved into two clear episode types. “Minor” episodes consisted of brief right or occasionally left hemidystonia, which occurred several times each week. “Major” episodes happened two or three times per month and lasted up to 30 minutes. After a similar onset, they progressed to involuntary flailing movements of all limbs, impaired responsiveness, dysarthria, aphasia, and drooling, sometimes associated with upward eye deviation. Although “major” episodes tended to be triggered by fatigue, there was no obvious trigger for “minor” episodes. A clinical diagnosis of alternating hemiplegia of childhood (AHC) was made. Neurological examination between episodes was unremarkable. There was no disease progression over time. There was no clinical response to levodopa, oxcarbazepine, flunarizine, trihexyphenidyl, topiramate, or gabapentin, although prolonged episodes could be terminated with buccal midazolam. Extensive investigation including brain magnetic resonance imaging, electroencephalogram (EEG) with capture of a typical episode, and cerebrospinal fluid neurotransmitters was normal. Whole genome sequencing (WGS) identified a novel homozygous frameshift in JPH3 (NM_020655.3:c.1310delG; p.Arg437Leufs*34). Parental DNA was not available for testing. The only other variant of interest was a homozygous NDUF5 variant (NM_024120.5:c.524A>G; p.His175Arg), which was considered unlikely to be relevant because the phenotypic fit for mitochondrial complex I deficiency was poor and in silico tools predicted that it was probably benign.

Patient two, now 10 years old, experienced episodes of limb, facial, and orolingual dyskinesia at least weekly from the age of 6 months (Video S1). These usually started in the right upper limb before becoming generalized and lasted for several hours. No abnormal eye movements were evident during the episodes, and there was no obvious trigger. Between

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episodes, very mild resting dyskinesia was evident in the upper limbs. She had moderate learning difficulties, but no behavioral issues. Again, extensive investigation was unremarkable, including EEG with capture of episodes. Several different medications—carbamazepine, valproate, topiramate, acetazolamide, and trihexyphenidyl—proved ineffective. Her maternal grandmother had experienced lifelong episodes of orofacial dyskinesia and limb posturing and had never learned to read. Her mother reported episodes of orofacial dystonia triggered by alcohol and had a slight resting tremor. She had completed compulsory education. The only relevant variant identified on WGS was a novel heterozygous nonsense variant in \textit{JPH3} (NM020655.3:c.1014C>G; p.Tyr338*) shared by mother and daughter: grandmaternal DNA was unavailable (Fig. 1B).

\textit{JPH3} encodes junctophilin-3, a component of the junctional complex linking the plasma membrane with the endoplasmic reticulum in excitable cells. Expression is highest in the brain. Pathological heterozygous triplet-repeat expansion variants (believed to cause toxic intracellular accumulation of abnormal protein)\cite{3,4} occur in Huntington-like disease type 2, characterized by adult-onset chorea, dementia and atrophy of the cortex, and basal ganglia.\cite{5} However, we believe that the disorder from either biallelic or heterozygous truncating variants is a distinct condition, mediated by loss of protein function rather than toxic accumulation, and so far, appears non-progressive in reported cases (Fig. 1C). The gene is predicted to be highly intolerant of loss-of-function,\cite{6} and in mice, both haploinsufficiency and knockout of \textit{JPH3} result in an abnormal motor phenotype.\cite{7} Although the association of the variably penetrant, heterozygous phenotype will require further confirmation in additional patients, \textit{JPH3} variants should be considered in patients with undiagnosed complex paroxysmal movement disorders.

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Data Availability Statement

We are unable to make patients’ full genomic data available under the terms of our ethical approval but are happy to share details of methods and laboratory or clinical results on request.

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Supporting Data

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