Does SCA45 Cause Very Late-Onset Pure Cerebellar Ataxia?

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Spinocerebellar ataxias (SCAs) are a large group of genetically and phenotypically heterogeneous autosomal dominant, neurodegenerative disorders manifesting with progressive cerebellar ataxia usually with adult-onset. Currently, 48 subtypes of SCAs are described; of which, for 40 SCAs, the genes have been identified. The most frequent types are related to coding repeat expansions including SCA1, SCA2, SCA3, SCA6, and SCA7. However, for unusual SCA types, whole exome sequencing (WES) is necessary to identify the genetic cause. Late-onset cerebellar ataxia can be caused by several genetic mutations, but a large percentage of patients remain undiagnosed after WES.

In this article, we describe 2 siblings who presented with very late-onset cerebellar ataxia, mild cerebellar atrophy, and whose genetic investigation disclosed a novel heterozygous c.10906T>G, p.Tyr3636Asp FAT2 (SCA45) missense variant. In addition, given that the variant is novel, affects a conserved amino acid, and segregates with disease in combination with the previous report on missense variants in FAT2, we may classify this variant of unknown significance as likely pathogenic.

Case Reports

Patient 1
An 86-year-old man presented with a progressive ataxia that started at 70 years of age. At 81 years, he developed dysarthria and dysphagia and started using a wheelchair at the age of 84 years. Family history was remarkable for cerebellar ataxia (figure A). Neurologic examination showed normal ocular movements and deep tendon reflexes, slurred speech, head titubation, and global cerebellar ataxia (video 1, links.lww.com/NXG/A403). Cognition was normal. Brain MRI (MRI) showed mild cerebellar atrophy (figure, B and C). Given the dominance inheritance pattern of the disease within the family, the SCA1, 2, 3, 6, 7, and 10 genes were tested for repeat expansions but was negative. WES was performed using a routine procedure including exome capture with Agilent Clinical Research Exome v1, followed by sequencing using the Illumina NextSeq platform. WES identified a novel heterozygous missense variant c.10906T>G, p.Tyr3636Asp in the SCA45 gene, FAT2 (ENST00000261800.5).

Patient 2
The sister of patient 1, 79 years old, presented with a 6-year history of progressive cerebellar ataxia. She also suffered of mild dysarthria, slurred speech, and dysphagia. Neurologic examination showed global ataxia (video 1). Cognition and ocular movements were normal, as well as deep tendon reflexes. Brain MRI showed mild cerebellar atrophy (figure, D and E). WES identified the same c.10906T>G, p.Tyr3636Asp variant in FAT2.
Discussion

SCA45 (MIM #617769) related to the FAT2 gene variants is an unusual conventional SCA type. SCA45 was reported in 2017 in a WES-based study that reported 2 different missense variants c.10758G>C, p.Lys3586Asn and c.10946G>A, p.Arg3649Glu affecting highly conserved amino acids in the last cadherin repeat and linker region between the cadherin repeat and the laminin A-G of FAT2. The first variant was identified in a patient who presented with pure cerebellar ataxia, downbeat nystagmus, dysarthria, a positive family history, and disease onset after the age of 40 years. The second case also had slowly progressive ataxia and a disease onset after the age of 50 years, but no family history. Moreover, a recent study including patients with CANVAS phenotype and negative for the RFC1 repeat-expansion disclosed one subject with the heterozygous c.4370T>C, p.Val1457Ala variant affecting the 13th cadherin domain of FAT2.

The p.Tyr3636Asp variant described in our report is novel and absent in control databases (gnomAD, ABraOM) and literature. Although this may seem a variant of unknown significance, there are some relevant points to be considered: (1) the variant is located close to the previously reported p.Lys3586Asn variant in a well-established functional domain, (2) cosegregates with disease, and (3) the phenotype is highly similar to cases described in previous studies of SCA45—we considered this variant likely pathogenic and the cause of the ataxia.

FAT2, previously named MEGF1, belongs to the family of human FAT genes. AT2 is a large transmembrane adhesion molecule specifically expressed in the cerebellar granule cells, playing an important role in postnatal cerebellum development. FAT genes are the human homologous of Drosophila fat. Recently, the depletion of fat in Drosophila was reported to cause neurodegeneration of the fly eye, very likely to deficits in autophagy. The question remains whether FAT2 in humans is also implicated in autophagy and thereby mediates neuronal homeostasis in the cerebellum needs to be established.

So far, only one family with a few affected members and one sporadic case carrying likely pathogenic FAT2 variants have been described. All cases presented with a slowly progressive late-onset ataxia. Therefore, additional cases with likely pathogenic variants presenting with late-onset ataxia and future functional studies on how variants in FAT2 cause late-onset ataxia will further add to link between FAT2 and SCA45. In conclusion, SCA45 seems characterized by a very late-onset cerebellar ataxia and cases presenting with such features without repeat expansions in the most common SCAs should be screened for variants in FAT2 using WES.

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**References**

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