NOVEL AMPD2 MUTATION IN PONTOCEREellar HYPOPLASIA, DYSMORPHISMS, AND TEETH ABNORMALITIES

Pontocerebellar hypoplasias (PCHs) are a genetically and clinically heterogeneous group of autosomal recessive inherited neurodevelopmental disorders. In the past decade, whole-exome sequencing (WES) has led to the identification of new genes, allowing the recognition of at least 10 different PCH types with broad and overlapping phenotypes.1

PCH type 9 (PCH9) (MIM 615809) is caused by homozygous mutations in the adenosine monophosphate deaminase 2 (AMPD2) gene.

AMPD2 plays an evolutionary conserved role in purine and nucleotide metabolism by regulating the guanine nucleotide biosynthesis and protein translation.2 Patients typically present with severe developmental delay, microcephaly, axonal neuropathy, and epilepsy. Neuroimaging in patients with PCH9 shows a typical midbrain “figure of 8” appearance, callosal hypoplasia, and periventricular white matter involvement.3

Previous reports have demonstrated the pivotal role of AMPD2 during neurogenesis and showed a potentially rescue therapy in vitro by administration of purine precursors.2 Since the original report of 5 PCH9 families,2 only 6 additional patients from 2 families have been described.3,4 Furthermore, a homozygous frameshift mutation in the AMPD2 gene has been identified in 2 members of a consanguineous family affected by spastic paraplegia without PCH.5

We report a novel homozygous AMPD2 mutation in 3 siblings with severe PCH9 phenotype.

Clinical description. The probands are 2 girls (IV:1, aged 9 years and IV:3 aged 8 years) and 1 boy (IV:2, aged 7 years) born to first-cousin parents from the Middle East (figure e-1 at Neurology.org/ng). Family history was unremarkable. All individuals were born at term by normal delivery, following an uneventful pregnancy. Since the first months of life, developmental delay without any motor skills acquisition, drug-resistant focal and tonic-clonic seizures, and progressive spasticity were noticed. All patients shared microcephaly and common dysmorphic features, including sloping forehead, large and posterior rotated ears, upper lateral incisor agenesis, mandible hypoplasia, and clinodactyly of the V fingers (figure 1, A–C). In all individuals, brain MRI showed PCH, severe callosal hypoplasia, leukoencephalopathy, and basal ganglia involvement (figure 1, D–F). Diffusion tensor imaging revealed marked thinning of the corticospinal tracts (figure e-2), and MR spectroscopy showed low N-acetyl aspartate in the thalamic region (data not shown) in patient II-3. Electrophysiologic studies performed in individual IV-2 did not reveal signs of axonal neuropathy. At the final follow-up, all patients showed cortical blindness and were using wheelchair or were bedridden. Their clinical and imaging details are summarized in table e-1.

Genetic study. We first excluded copy number and structural DNA variations by karyotyping and array-comparative genomic hybridization. We then performed WES on DNA samples of all family members, using a standard Illumina pipeline (e-Methods, tables e-2 and e-3) and identified a novel homozygous frameshift mutation c.495delG (p.R165fs*21) in the gene encoding for the adenosine monophosphate deaminase 2 enzyme (AMPD2, NM_001257360.1). Sanger sequencing showed that the mutation segregates according to a recessive model of inheritance (figure e-2). The institutional review board approved the use of human samples for this study.

Discussion. AMPD2 encodes 1 of 3 adenosine monophosphate (AMP) deaminase enzyme homologs, which convert AMP to IMP. AMPD2 deficiency results in accumulation of adenosine nucleotides and depletion of guanine nucleotide, impairing the guanosine-5′-triphosphate–dependent initiation of protein translation, similarly to what was observed in transfer RNA splicing endonuclease complex genes, linked to other PCH types.2,6 Recently, AMPD2 mutant mice and Ampd2−/− mice have been associated with nephrotic syndrome and proteinuria in the absence of any brain abnormality.7 A neurodegenerative phenotype has been observed when both Ampd2 and Ampd3 are knocked out, suggesting a functional redundancy among AMP
deaminase homologs. However, it remains to be elucidated whether humans carrying AMPD2 mutations may have renal involvement.

To date, 14 patients with PCH9 have been reported. All but 1 AMPD2 mutation reside within the conserved catalytic AMP deaminase domain\(^2\)-\(^4\) (table e-1). The novel identified homozygous AMPD2 mutation (c.495delG p.R165RfsX21) lies outside the catalytic domain and is associated with a very severe phenotype as described for c.751C\(^T\) (p.R251W).\(^4\) Neuroimaging revealed the involvement of the basal ganglia and thalami and marked hypoplasia/atrophy of the corticospinal tracts, unraveling the anatomical basis of the characteristic midbrain “figure of 8” appearance and pontine flattening observed in PCH9 individuals. However, our patients showed peculiar facial dysmorphisms and teeth abnormalities that might be an additional feature associated with PCH9. Genetic studies did not show any additional homozygous variant that could readily explain these findings (table e-3). Further studies will clarify the phenotypic spectrum associated with AMPD2 mutations.

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