Recurrent Stroke Due to a Novel Voltage Sensor Mutation in Ca$_2$.1 Responds to Verapamil

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Background and Purpose—Familial hemiplegic migraine is characterized by recurrent migraine, hemiparesis, and ataxia. Causes may be mutations in calcium and sodium channels or in a subunit of the Na/K-ATPse. Migraine treatment with calcium channel blockers was only successful in some patients.

Summary of Case—We describe a 6-year-old girl with recurrent ischemic strokes after minor head trauma associated with seizures, hemiparesis, fever, and altered consciousness. Genetic analysis revealed a spontaneous, novel dominant CACNA1A mutation (c.4046G→A, p.R1349Q) that removed a highly conserved arginine of the voltage sensing region of the P/Q-type Ca$_2$.1 channel. Because a homologous mutation in the tottering-5J mouse increased open probability of the channel as well as calcium influx, we treated the patient with the calcium channel blocker verapamil during characteristic prodromi after head trauma. Treatment was instantly effective and prevented a new stroke.

Conclusion—CACNA1A mutations should be considered in the diagnostic workup of childhood stroke, especially if associated with ataxia and migraine. (Stroke. 2011;42:00-00.)

Key Words: calcium channels ■ genetics ■ pediatric neurology ■ pediatric stroke ■ stroke in children

Familial hemiplegic migraine (FHM) is a rare, childhood-onset, autosomal-dominant subtype of migraine with aura. It is characterized by hemiparesis during the aura phase and may also be associated with cerebellar ataxia, delay of motor development, and other neurological symptoms such as cerebral seizures, mental retardation, brain edema, fever, and coma.1,2 Attacks may be provoked by typical migraine triggers such as stress, exertion, and minor head trauma. Some small open pharmacological studies tested the efficiency of acetazolamide and verapamil to suppress aura symptoms, but not all patients improved and the relation between therapeutic effect and molecular defect had remained unexplored.3 Gene mutations leading to FHM comprise defects in CACNA1A (FHM1, MIM #141500), ATP1A2 (FHM2, MIM #602481), and SCN1A (FHM3, MIM #609634). On the reverse, mutations in CACNA1A may cause various clinical phenotypes, which include FMH1, episodic ataxia Type 2 (MIM #108500), and spinocerebellar ataxia Type 6 (MIM #183086). There exists a certain, albeit restricted, genotype–phenotype relation: gain-of-function mutations cause FHM1, truncating mutations or mutations affecting folding of the channel cause episodic ataxia Type 2, and triplet extension mutations around Q84 lead to spinocerebellar ataxia Type 6.4 We report the case of a young girl in whom FHM Type 1 manifested itself with repeated cerebral infarctions due to a novel human gain-of-function mutation in the voltage-gated calcium channel Ca$_2$.1. The child responded promptly to therapy with the calcium channel blocker verapamil, thus linking a defined molecular defect in FHM1 with a potential pathogenesis-based therapy for the first time.

Case History
The 6-year-old girl is a single child of healthy nonconsanguineous parents. At 3 months, she presented with a severe convergent squint. Motor development was retarded (head control 3 months, turns from supine to prone 12 months, sitting freely 14 months, pulls to standing from supine 18 months). At the age of 2 years, she had 2 episodes of cerebral seizures and coma for 2 weeks after a fall from a height of 1 m from a bed. An MRI excluded intracranial hemorrhage and detected mild cerebellar atrophy and delay in myelination. From this time, her motor development was stagnant and until now she is unable to walk freely. At 3.5 years of age, she was admitted again with encephalopathy and left-sided weakness, this time a few hours after a backward fall from a hobby-car. MRI revealed a massive ischemic lesion with prominent perifocal edema in the right parietal and occipital lobe (Figure 1A–B). Despite an extensive search for emboli, including transesophageal cardiac ultrasound and coagulation...
studies, no obvious cause was found. Sequence analysis of all mitochondrial tRNAs ruled out mitochondrial encephalomyopathy, lactic acidosis, and stroke-like episodes (MELAS) syndrome and related disorders. The girl recovered slowly, needed long-term rehabilitation, and a left-sided hemiplegia remained. Absence-like seizures were treated with levetiracetam. At 5 years of age, after falling from the lap of a grandfather, she was again admitted to the hospital with loss of consciousness, now with fever, right-sided focal seizures, and weakness. Serial MRI scans revealed increased signal intensity on T2 and fluid-attenuated inversion recovery images and hypointensity in apparent diffusion coefficient maps of the left hemisphere that were not restricted to vascular territories and showed variance in the temporal progression of the lesions (Figure 1C–H). Digital subtraction angiography excluded an arterial or venous occlusion. Over the next 3 months, vigilance and motor capabilities of the child improved slowly.

Genetics

The clinical symptoms and the MRI diagnostic findings led us to suspect a severe form of FHM and prompted sequence analysis of \textit{CACNA1A}. Genomic DNA was isolated from peripheral blood cells by salt extraction and all 47 exons of \textit{CACNA1A} (GenBank NM_001127222), including flanking intronic regions, were polymerase chain reaction-amplified and analyzed by automatic sequencing. We identified a heterozygous c.4046G→A transition in \textit{CACNA1A} (Figure 2). The heterozygous c.4046G→A transition leads to the exchange of arginine for glutamine and was absent in both parents. Segregation analysis of the mutation. The c.4046G→A mutation generates a novel BstN I restriction site, thus cleaving the 241-bp wild-type fragment into 171/70 bp (mut).

Figure 1. A, Cerebral MRI at the age of 3.5 years. Axial fluid-attenuated inversion recovery (FLAIR) image demonstrates edema in the right parietal and occipital lobe, which is not restricted to vascular territories. Corresponding hypointense signal changes on the apparent diffusion coefficient (ADC) map confirm an ischemic lesion. B–D, Serial cerebral MRI at the age of 5.5 years. B, At the day of symptom onset, the axial FLAIR image shows mild cortical edema in the left frontal and parietal lobe, whereas the ADC map is still unremarkable at this time, suggesting vasogenic rather than cytotoxic edema. At the right parietal and occipital lobe, the cortex is atrophied in the area of the old infarction. C, At the second day, cytotoxic edema with increasing hyperintense signal changes on FLAIR images and signal reduction on the ADC map corresponding to an ischemic lesion can be seen in the left frontal lobe. A vascular occlusion of the left posterior cerebral artery was ruled out by conventional angiography (data not shown). D, Ten days after symptom onset, cerebral MRI depicts progressive subcortical edema in the left parietal lobe on FLAIR images without signs of new ischemic tissue damage on the ADC map.

Figure 2. Molecular genetic findings in the patient’s family. A, The heterozygous c.4046G→A transition leads to the exchange of arginine for glutamine and was absent in both parents. B, Segregation analysis of the mutation. The c.4046G→A mutation generates a novel BstN I restriction site, thus cleaving the 241-bp wild-type fragment into 171/70 bp (mut).
To confirm the c.4046G→A mutation and to screen normal controls, we polymerase chain reaction-amplified the DNA sequence of exon 25 using the oligonucleotides (forward) 5‘-GGACCACTCTTACCCAGGAA-3‘ and (reverse) 5‘-CCACCCTCCTCCATCTCAC-3‘. The resulting 241-bp band was cut into 171±70 bp by BstNI only in the presence of the mutation (Figure 2B). On the protein level, the mutation causes the substitution of arginine to glutamine at codon 1349. The de novo mutation was absent in her parents (Figure 2) and in 200 normal alleles excluded a common polymorphism. The alleles of normal control subjects (data not shown). To exclude uniparental disomy or a complex rearrangement and to confirm paternity, we investigated 20 microsatellite markers flanking the CACNA1A locus on chromosome 19 (D19S424, D19S1034, D19S586, D19S906, D19S221, D19S558, D19S226, D19S923, D19S885, D19S429, D19S925, D19S931, D19S255, D19S875, D19S433, D19S882, D19S414, D19S871, D19S223, D19S178) by polymerase chain reaction amplification with fluorescent dyes (D19S433, D19S882, D19S414, D19S871, D19S223, D19S178) by polymerase chain reaction and subsequently with the GeneMapper Version 3.7 software (data not shown).

Clinical Course and Verapamil Treatment

Shortly after we had found the CACNA1A mutation, at the age of 5.5 years, the girl was again admitted to the emergency department after another backward fall from a chair and hitting her head on the floor. She showed the characteristic prodromi that had anticipated her last ischemic strokes (all secondary to a minor head trauma), which comprised severe headaches, yawning, truncal unsteadiness, and progressive loss of consciousness. Based on the molecular genetic results, we decided to initiate immediately an intravenous therapy with the calcium channel blocker verapamil at a single dose of 0.1 mg/kg body weight over 20 minutes. Already during infusion the girl regained consciousness and started to talk to her mother. The dramatic clinical improvement was sustained even after the infusion had finished. As a long-term prophylactic treatment, we prescribed oral verapamil at a daily dosage of 3 mg/kg body weight under which the child did not develop any signs of stroke for 1 year despite 2 episodes of head trauma of the same order of magnitude as the ones causing stroke before.

Discussion

We report on a young girl who had several ischemic strokes due to a spontaneous dominant mutation in her CACNA1A gene (c.4046G→A, p.R1349Q). Ischemic stroke has not been reported in FHMI before. These findings expand the clinical spectrum of CACNA1A mutations and place our patient at one extreme of a clinical continuum between simple migraine aura and the most severe vascular complications with manifest stroke.

The CACNA1A gene encodes the α1-subunit of the highly conserved P/Q-type voltage-gated calcium channel Ca2.1. Analysis of the c.4046G→A sequence variant with Mutation Taster, an evaluation tool of the pathogenic potential of sequence variants,5 predicted pathogenicity with a high probability (R=0.988). Moreover, the absence of this variant in 200 normal alleles excluded a common polymorphism. The mutation replaces an evolutionary highly conserved alkaline arginine by a neutral glutamine. R1349 contributes to a motif of 4 arginines that form the voltage sensor of various calcium as well as of sodium and potassium channels (Figure 3).6 Missense mutations removing such arginine residues have been shown to alter Ca2.1 channel function in a gain-of-function manner. This was confirmed by in vitro and ex vivo studies on human CACNA1A FHMI1 mutations (p.R192Q and p.R583Q) and for the p.R1252Q mutation of the tottering-5j (Tg-5j) mouse (this mutation corresponds to p.R1300Q on NP_031604.3; Figure 3) that led to a shift to more negative potentials for both Cav2.1 channel activation and inactivation.7–9 Interestingly, the p.R1300Q mutation of the Tg-5j mouse is exactly homologous to the human p.R1349Q mutation (Figure 3). The functional consequences of this mutation were grave, because homozygous mice died soon after birth.9 We thus have reason to assume that in humans, the p.R1349Q mutation may also lead to a gain-of-function with enhanced Ca2.1 channel activity and subsequent increased intracellular calcium concentrations leading to cortical spreading depression.
Spreading depression of cortical electric activity is thought to be the pathophysiological correlate of migraine aura and the result of near-complete and sustained depolarization of neurons above the inactivation threshold for their action-potential generating ion channels. Spreading depolarization can be observed as a large change of the slow potential, whereas spreading depression is characterized by electric silence in the higher frequency band of the electrocorticogram in humans and animals. Resistance vessels change their vascular tone in response to spreading depolarization, causing either (1) transient hyperperfusion (physiological hemodynamic response) in healthy tissue; or (2) severe hypoperfusion (inverse hemodynamic response) under pathological conditions leading to spreading ischemia, which in itself may cause widespread cerebral infarctions in animals and which contributes to lesion progression in humans. Mild head trauma is a well-known trigger for such spreading depolarizations, and missense mutations in CACNA1A may decrease their threshold, an effect presumably due to increased glutamatergic neurotransmission. The findings by Cav2.1 channels. Because Cav2.1 channels also seem to be present on the smooth muscle layer of resistance vessels, they may mediate vasoconstriction secondary to calcium influx. Interestingly, another L-type calcium channel blocker, nimodipine, was able to almost completely reverse spreading ischemia in a rat model.

We therefore hypothesize that the mild head trauma in our patient was sufficient to induce spreading depolarization, which, on top of an inverse hemodynamic response, led to widespread brain infarctions in a superadditive manner. This hypothesis deserves further study in animals, for example, in the tottering-5j (Tg-5J) mouse with the homologous mutation.

We are aware that analysis of a single case cannot prove the therapeutic efficacy of verapamil in this condition. Further controlled clinical studies in combination with molecular genetic analyses are thus necessary to substantiate such an effect. Beyond that, our findings suggest that sequence analysis of CACNA1A should be considered during diagnostic workup of childhood stroke associated with migraine and ataxia; all the more, it seems to be a treatable condition.

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Disclosures
None.

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