Identification of a novel CACNA1A mutation in a Chinese family with autosomal recessive progressive myoclonic epilepsy

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Background: Progressive myoclonic epilepsy (PME) is a heterogeneous neurodegenerative disorder, which is commonly manifested with refractory seizures and neurologic deterioration. The prognosis of PME is poor, so early diagnosis of PME is critical. The aim of our study is to identify the novel pathogenic gene in a Chinese family with PME, which may be helpful in future.

Subjects and methods: A three-generation consanguineous Chinese Han family with PME was recruited. A novel homozygous variant was identified by the genetic technique of exome sequencing and certificated by Sanger sequencing and functional prediction.

Results: A novel homozygous variant, c.6975_6976insCAG, in the CACNA1A was identified in the PME family. The novel variant encoding the alpha-1A subunit of the calcium channel Cav2.1 was found in two siblings in the Chinese family and was absent in 50 normal controls. Our research indicates that the homozygous c.6975_6976insCAG might be the pathogenic mutation for PME.

Conclusion: As a molecular diagnostic strategy, our research explores the mutation gene spectrum of PME and has resulted in significant predictions for genetic counseling.

Keywords: CACNA1A, PME, exome sequencing, myoclonus, muscular hypotonia

Introduction

The CACNA1A gene has been reported to encode the pore-forming alpha-1A subunit of the calcium channel Cav2.1, which acts as a voltage sensor and an ion pore.1,2 Several mutations within this gene lead to variable clinical phenotypes such as familial hemiplegic migraine type 1, spinocerebellar ataxia type 6 and episodic ataxia type 2 (EA2), which have been recognized widely. There are only two reports on a CACNA1A-related generalized epileptic seizure3 and a photoparoxysmal response (PPR).4 Muscular hypotonia or muscular atrophy has also been reported to be associated with CACNA1A gene.5

Progressive myoclonic epilepsy (PME) is generally manifested with refractory myoclonus seizures, cognitive decline and neurologic deterioration, caused by metabolic, genetic and neurodegenerative diseases, which has an unfavorable course and poor prognosis. A wide range of specific etiology contributes to the PME, such as neuronal ceroid lipofuscinoses (NCLs), Tay–Sachs, myoclonic epilepsy with ragged red fibers (MERRF), POLG1 Mutation, Lafora Disease and Unverricht–Lundborg disease (ULD).

Herein, we describe two PME patients. Those gene mutations recognized widely in the PME were screened with negative results. On the other hand, a causative novel
mutation in the CACNA1A gene was found in the PME by exome sequencing (ES), and then confirmed by Sanger sequencing subsequently.

**Subjects and methods**

**Subjects**

A three-generation consanguineous Chinese Han family with PME was studied, and four members of the family agreed to participate in our study. PME has been found in the two siblings (II:1 and II:3; Figure 1) with the same phenotype of refractory myoclonic seizures, neurologic deterioration and cognitive decline, who have the same problems of refractory myoclonic seizures, neurologic deterioration and cognitive decline. However, their parents (I:1 and I:2; Figure 1) and their children (III:1–3; Figure 1) were normal persons. Fifty healthy Chinese Han subjects (age 32.5±4.5 years) were enrolled as controls with neurologic physical and clinical assessments. After obtaining written informed consent from the participants, we collected the peripheral blood samples from April 19 to October 30, 2015.

**Clinical evaluations**

Clinical and electrophysiologic assessments were performed on the PME family in the First Hospital of Jilin University. Both patients were afraid of light. Electroencephalogram (EEG) was conducted in both patients with PPR during the process of light stimulation frequently. Magnetic resonance imaging was also carried out.

**Whole ES and relevant variant analysis**

Genomic DNA was extracted from peripheral blood samples by using standard phenol–chloroform extraction method.6 ES was performed according to relevant guidelines and regulations. At least 1.5–2 μg of genomic DNA was obtained from the proband (II:1 and II:3; Figure 1), and then following the manufacturers’ procedures, the samples was sheared, enriched, hybridized and captured on the Agilent SureSelect. The captured library was enriched with Illumina HiSeq 2000 sequencing instruments and provided enough depth to identify variants at 97.4% of targeted exome exactly.7

All variants were screened with the single nucleotide polymorphism (SNP) database version 142 (dbSNP142), 1000 Genomes Project (version 2014 October) and NHLBI ES Project (ESP) 6500. The clean reads were mapped to the human reference genome (UCSC hg19, [http://genome.ucsc.edu](http://genome.ucsc.edu)) using Burrows–Wheeler Alignment tool.8,9 SNPs and insertions/deletions were identified by the Sequence Alignment/Map tools (SAMtools),10 and duplicate reads were marked by Picard tools. Functional prediction was carried out by Sorting Intolerant from Tolerant and Polymorphism Phenotyping version 2 (PolyPhen-2). Candidate variants were analyzed by the ANNOVAR (Annotate Variation) software.11

**Sanger sequencing and functional prediction**

Potential causative variants were confirmed by direct Sanger sequencing using the ABI3500 sequencer system (Thermo Fisher Scientific, Waltham, MA, USA).12 Primers for the mutation of RS753460234 in the CACNA1A gene were designed as follows: RS753460234-F: TCCCGTGTCTCC TTGATTT; R: GGTAAGCGCGTGGAGT. MutationTaster was applied to predict the functional changes in the CACNA1A variant.13

**Ethics approval and consent to participate**

This study has been approved by the Research Ethics Board of Jilin University First Hospital. Written informed consents were obtained from the patients and patients’ family members for publication of this study and any accompanying images. A copy of the written consent is available for review by the editor of this journal.

**Results**

**Clinical findings**

Both patients (II:1 and II:3; Figure 1) presented with refractory myoclonus at around 50 years of age. Their previous history was noncontributory. The seizures could be described as symmetrical and rhythmic jerk. The frequency
increased to dozens of times a day. They had much more myoclonus in day than at night. Besides this, they were afraid of light and sound before seizure onset. And 2 years later, they gradually presented with muscle weakness, muscular atrophy and cognitive decline. Neurologic examination showed both proximal and distal muscle atrophy of lower limbs, muscle weakness and progressive ataxia. Mini-Mental State Examination scores were 20/30 and 22/30, respectively, which revealed that they had trouble in memory and locating. EEG on admission showed generalized polyspike waves or generalized spike-slow complex wave without previous EEG available (Figure 2). Also, the electromyogram results were normal. Light stimulation caused photosensitivity frequently. Magnetic resonance imaging was normal. According to the clinical history, presentation and EEG features, we made a diagnosis of PME. After the administration of antiepileptic drug valproic acid, no improvement was seen. The clinical picture of PME family is summarized in Table 1.

Common pathogenic gene of PME screening
A wide range of specific gene mutation contributes to the PME, such as NCLs, Tay–Sachs, MERRF, POLG1 mutation, Lafora Disease and ULD. However, we obtained negative results with the common pathogenic gene screening.

Exome sequencing
A total of 162.565 million reads were generated from ES with an average read length of 150 bp, and 162.308 million reads (99.84%) were mapped to the human genome. Also, 16,041.78 Mb of effective sequences was mapped to the target region. Moreover, 180.973 million bases covered the target region and occupied about 99.7% of the target region. The sequence at 10× or greater covered 98.7% of the target region. Also, 64,952 SNPs in the exon regions and 7,001 SNPs in the splicing sites were identified in total. Five hundred and sixty indels in the exon regions and 402 in the splicing sites were identified, respectively.7 We used a prioritization scheme to eliminate common known variants reported in public databases, such as dbSNP137 (Minor Allele Frequency >1%), 1,000 genomes project with a frequency >0.5% and NHLBI ESP6500, in order to detect the possible pathogenic variant. Sorting Intolerant from Tolerant and PolyPhen-2 analyses were utilized to predict the functional changes of the mutation. As a result, a homozygous CACNA1A c.6975_6976insCAG variant was observed in the proband (II:1 and II:3; Figure 1) as a possible pathogenic mutation.

![Figure 2 EEG showed generalized polyspike-slow complex wave mixed with some muscle artifacts, which indicated a myoclonic seizure.](https://www.dovepress.com/)

**Notes:** Red mark: photic stimulation onset; blue mark: muscle artifacts; yellow mark: polyspike-slow complex wave.

**Abbreviation:** EEG, electroencephalogram.
Confirmation of pathogenic mutation

A homozygous CACNA1A c.6975_6976insCAG in the exon 47 identified in the proband was further confirmed by direct Sanger sequencing. The same homozygous variant was found in the affected sibling (II:1 and II:3; Figure 1), but not in both unaffected offsprings (III:1–3; Figure 1). The identical variant was absolutely absent in 50 ethnically matched normal controls (Figure 3). The homozygous CACNA1A c.6975_6976insCAG variant relevant with the PME phenotype, leading to an insertion at amino acid position 2326, was predicted to change alanine to another quisqualic acid, which may trigger neurotoxicity. CADD (http://cadd.gs.washington.edu/score) predicts the variant to be harmful with a functional CADD_Phred of 14.13 (≤10); the mutation may be the disease-causing mutation in the PME family.

Discussion

PME is an epilepsy syndrome characterized by progressive myoclonus, cognitive impairment, ataxia as well as other neurologic deficits. It includes several specific etiologies. Cystatin B gene mutations cause ULD, a rare form of PME. Mutation of the nuclear lamin gene LMNB2 causes PME with early ataxia. CLN6 mutation is reported to cause NCLs, another form of PME. Also, PME encompasses a wide spectrum of mitochondrial and metabolic diseases such as MERRF and Lafora disease. MERRF is the common form with the underlying mutation in the tRNALys gene, and Lafora was reported to be associated with mutation in EPM2A gene. However, in our cases, none of the screened mutation genes are positive. By whole ES, we identified CACNA1A mutation in our subjects. This mutation cannot be seen in normal controls, but has been found in our PME family. Only homozygous mutation can cause PME, although the offsprings may show heterozygous mutation, while they are normal people without PME presentation. Therefore, one good allele is enough in our family.

The CACNA1A gene encodes the alpha-1A subunit of the Cav2.1 P/Q-type voltage-gated calcium channel, which is in connection with the transmembrane performing.

Table 1  Phenotypes and genotypes of the PME family

<table>
<thead>
<tr>
<th>Subjects</th>
<th>Age, years</th>
<th>PME</th>
<th>MRI</th>
<th>CACNA1A c.6975_6976insCAG</th>
</tr>
</thead>
<tbody>
<tr>
<td>II:1</td>
<td>56</td>
<td>+</td>
<td>Normal</td>
<td>Homozygous</td>
</tr>
<tr>
<td>II:3</td>
<td>54</td>
<td>+</td>
<td>Normal</td>
<td>Homozygous</td>
</tr>
<tr>
<td>III:1</td>
<td>30</td>
<td>Normal</td>
<td>Normal</td>
<td>Heterozygous</td>
</tr>
<tr>
<td>III:2</td>
<td>31</td>
<td>Normal</td>
<td>Normal</td>
<td>Heterozygous</td>
</tr>
<tr>
<td>III:3</td>
<td>34</td>
<td>Normal</td>
<td>Normal</td>
<td>Heterozygous</td>
</tr>
</tbody>
</table>

Abbreviations: MRI, magnetic resonance imaging; PME, progressive myoclonic epilepsy.

Figure 3  Sequence analysis of a PME family.

Notes: CACNA1A c.6975_6976insCAG was confirmed in II:1 and II:3 by Sanger sequencing. Also, the variant was not identified in both unaffected offsprings and 50 normal controls.

Abbreviation: PME, progressive myoclonic epilepsy.
Several heterozygous mutations in this gene have been reported to cause three allelic autosomal dominant diseases, such as EA2, familial hemiplegic migraine type 1 and spinocerebellar ataxia type 6. Among these mutations, nonsense, missense and cytosine–adenine–guanine repeats are commonly observed. However, an association with idiopathic epilepsy and a link with epileptic syndrome have only been recently recognized. Reinson et al describe in their report that two patients, a sister and brother, developed daily refractory seizures with extreme muscular hypotonia and global developmental delay. A missense mutation c.4315T>A in CACNA1A was identified. Similar clinical symptoms of refractory seizures, muscular hypotonia and progressive ataxia are seen in our cases. Kaido et al report two cases of EA2 with PPR in EEG. A novel single base deletion (c.3575delA) in CACNA1A was uncovered. Special and rare PPR response to light stimulation is also presented in our cases.

In consistent with our clinical findings, mice with loss of Cav2.1 channel exhibit a severe phenotype of chronic dystonia and ataxia. Inborn errors of CACNA1A, an early P/Q-type voltage-gated calcium release defect, produces a stable generalized epilepsy phenotype. The various phenotypes caused due to mutation in CACNA1A have been found in both human and mice. Thus, CACNA1A is likely a novel disease-causing gene for the PME.

The mutation of polymorphic CAG insertion in exon 47 was located in an important functional region of the human a1A subunit of the neuronal Cav2.1 (P/Q-type) Ca2+ channel, which is associated with dysfunction of the intracellular C-terminus closely (Figure 4). Furthermore, mutation p.A2326delinsQA in CACNA1A changes alanine to another quisqualic acid, which may contribute to convulsions. Chase reports that l-quisqualic acid is an excitatory amino acid agonist that may trigger neurotoxicity.

**Conclusion**

In our study, the homozygous CACNA1A c.6975_6976insCAG mutation was identified in two affected siblings of PME family, but was absent in the unaffected children and 50 normal controls. The mutation is associated with PME phenotype closely and might be disease causing. The c.6975_6976insCAG mutation in CACNA1A in the PME family is a novel finding. This mutation of the base-insertion was responsible for the function of amino acid and protein significantly.

In summary, ES reveals a novel disease-causing CACNA1A gene for PME as a powerful molecular diagnostic technique.

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**Author contributions**

All authors participated in drafting the article or revising it critically for important intellectual content. All authors participated in conception and design, acquisition of data, or analysis and interpretation of data. All authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work. Also, all authors agreed and gave final approval of the version to be published.

**Disclosure**

The authors report no conflicts of interest in this work.

**References**


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