SHORT REPORT

Rare KCNJ18 variants do not explain hypokalaemic periodic paralysis in 263 unrelated patients

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ABSTRACT

Objective To examine rare KCNJ18 variations recently reported to cause sporadic and thyrotrophic hypokalaemic periodic paralysis (TTP).

Methods We sequenced KCNJ18 in 474 controls (400 Caucasians, 74 male Asians) and 263 unrelated patients with periodic paralysis (PP), including 30 patients with TTP without mutations in established PP genes.

Results In 10 patients without TTP, we identified 9 heterozygous, novel variations (c.-3G>A, L15S, R81C, E273X, T309I, I340T, N365S, G394R, R401W) and a questionable heterozygous causative R399X stop variant. Studies on 40 relatives of these 10 patients showed that none of the variants were de novo in the patients and that R399X occurred in 3 non-affected relatives. Most affected amino acids lacked conservation and several clinically affected relatives did not carry the patient’s variant. T309I, however, could be pathogenic under the pre-requisite of strongly reduced penetrance in females. Of the controls, 12 revealed a novel rare variants including the heterozygous E273X stop variant in three individuals.

Conclusions Our study shows many different, rare KCNJ18 alterations in patients as well as controls. Only perhaps one meets the requirements of a disease-causing mutation. Therefore, KCNJ18 alterations are seldom pathogenic. Additional studies are required before patients with PP can be genetically diagnosed on the basis of a KCNJ18 variant alone.

INTRODUCTION

Hypokalaemic periodic paralyses (PP) are a group of diseases characterised by episodes of flaccid muscle weakness associated with hypokalaemia. These episodes usually begin in the first or second decade of life, occur spontaneously and can be triggered by serum potassium reduction due to insulin (following carbohydrate-rich meals), glucocorticoids (stress, infection) and muscle reuptake at rest after strenuous exercise. The genes responsible for hypokalaemic PP are CACNA1S encoding the calcium channel Cav.1.1 (HypoPP1) and SCN4A coding for the sodium channel Nav.1.4 (HypoPP2). Both are voltage-dependent channels of the skeletal muscle fibre membrane. A mutation-induced aberrant current leads to a paradoxical membrane depolarisation that renders muscle fibres unexcitable. A third gene responsible for PP with concomitant arrhythmia and dysmorphia is KCNJ2, encoding the inwardly rectifying potassium channel Kir.2.1 of skeletal and cardiac muscle. The typically observed hypokalaemia might result from a reduced outward conductance of this channel.

The most frequent form of hypokalaemic PP is thyrotrophic periodic paralysis (TTP). It resembles HypoPP1/2 with respect to provocative factors. However, the hypokalaemia is more pronounced in TTP (between 1.0 and 2.5 mM) and therefore, often alters the ECG. Patients with TTP only experience paralytic attacks in the hyperthyroid state whereby clinical signs of hyperthyroidism may not be obvious. Although hyperthyroidism is much more frequent in females, the male-to-female ratio for TTP in Asians is about 6:1 and the onset is usually after the age of 20 years. Paralytic attacks cease when the euthyroid state is restored.

Mutations in KCNJ18 gene encoding an inwardly rectifying potassium channel (Kir.2.6) cause TTP and sporadic, that is, non-familial cases of HypoPP. Mutant Kir.2.6 proteins form a heterotetrameric Kir channel complex with Kir.2.1 and thereby reduce cell surface expression of the complex.

The aim of our work was to identify potentially disease-causing KCNJ18 variants (defined as <1% in the normal population) in 263 unrelated patients with PP, in whom mutations in the established PP genes have been excluded. Then the identified variants were analysed by (1) conservation of KCNJ18 variants, (2) segregation studies, (3) prediction programmes and (4) comparison with non-synonymous variants of a control group of same origin without neuromuscular disease.

METHODS

Patients and volunteers

Samples of DNA were collected from a total of 263 unrelated patients with a history of at least two episodes of quadriparesis associated with hypokalaemia. The phenotype was defined as mild if the episodes were majorly paretic (60% of cases) and severe if the episodes were majorly plegic (40% of cases). Thirty of these patients (17 Caucasians, 13 Asians, all males) were diagnosed as TTP according to accepted criteria.

Forty relatives of the 10 patients with novel variants were also studied. Additionally 474 DNA samples from individuals without muscle disease were examined (400 Caucasians, 74 Asians). Genomic DNA was isolated from EDTA blood using the QIAamp DNA Blood Kit (Qiagen, http://www.qiagen.com) according to the instructions of the producer. Informed consent was obtained from patients, relatives and volunteers with no evidence of muscle disease.
Analysis of KCNJ18 and evaluation of rare variants

Amplification, nested PCR, sequencing and our reference sequence are described elsewhere (see online supplementary file S1). Sequence analyses were evaluated using software SeqPilot of JSI (http://www.jsi-medisys.de/). Predictions regarding missense changes were made with PolyPhen-2 (http://genetics.bwh.harvard.edu/pph2/) and Mutations Taster (http://www.mutationtaster.org/). Splicing behaviour was predicted with MaxEnt (http://genes.mit.edu/burgelab/maxent/Xmaxentscan-). Alignments of several sequences of these species were carried out using the program ClustalW (http://www.genome.jp/tools/clustalw/) to check conservation.

RESULTS

In our 263 unrelated patients with PP we found the known amino acid substitutions R66Q, Q39R, H40R, V100I, H192Q, V249I, F281L, and Y338F which are not considered to be pathogenic.7 We also identified eight synonymous, presumably non-pathogenic changes. Of greater interest are the following nine novel alterations (table 1): one prestart base change (c.–3G>A), seven non-synonymous amino acid substitutions (L15S, R66Q, Q39R, H40R, V100I, H192Q, V249I, F281L, and Y338F), and a novel E273X mutation and a known questionably causative R399X stop mutation.7 These 10 variants were heterozygous in 10 index patients with normal thyroid function and were studied more deeply according to conservation, concordance of predictions on disease causality and segregation:

- Base exchange c.–3G>A is considered ‘improbable’ by MaxEnt to generate a splice site and is carried by an unaffected brother of the male patient.
- L15S is not conserved and occurs in two controls and for KCNJ12 in 5%, but does not occur in an affected family member of the patient.
- R81C is conserved and is predicted as pathogenic; however, R81P is found in one control.
- T309 is conserved and T309I is pathogenic according to the prediction programmes and two non-affected female relatives are carriers (mother and daughter).
- I340 is different in four species and I340T is predicted as a benign polymorphism.
- N365 is conserved, but predictions on N365S are discordant and the index patient’s daughter is affected although she is hypothyroid.
- G394R is not conserved and is predicted as benign by both programmes and two non-affected carriers are carriers.
- R401 is not conserved and the substitution R401W is excluded in three affected family members (see online supplementary figure 1S).
- E273X is found in three unrelated controls.
- R399X is identified in a 10-year-old boy whose mother and paternal grandfather are R399X carriers without PP history. As previously described, R399X also occurred in 1 of 100 controls.7

| Table 1 Novel variants and known questionably causative mutations of 263 patients with PP and 474 controls |
|---|---|---|---|---|---|---|---|
| Exchange nucleotide amino acid | Index, n | Localisation | PolyPhen2 | Mutation taster | Conservation | dbSNP (KCNJ12) | Severity/ segregation |
| Patients with PP | | | | | | | |
| –3G>A | – | 1 | Intron 2–3, no splice site predicted | | | | |
| 44T>C | L15S | 1 | N | – | + | – | No entry | –/– |
| 241C>T | R81C | 1 | N | + | + | + | 50 (1089) | –/– |
| 759insT | E273X | 1 | C | – | – | – | No entry | –/– |
| 926C>T | T309I | 1 | C | + | + | + | No entry | –/– |
| 1019T>C | I340T | 1 | C | + | + | + | No info | –/– |
| 1094A>G | N365S | 1 | C | – | + | + | No entry | –/– |
| 1180G>A | G394R | 1 | C | – | – | – | No info | +/+ |
| 1195C>T | R399X* | 1 | C | + | + | + | No info | –/– |
| 1201C>T | R401W | 1 | C | (+) | – | – | No entry | +/+ |
| Controls | | | | | | | |
| –7C>T | L15S | 2 | N | – | + | – | 50 (1089) | NA |
| 109G>A | G34S | 1 | N | (+) | + | + | No entry | NA |
| 242G>C | R81P | 1 | N | + | + | – | No entry | NA |
| 578C>T | T193M | 1 | C | + | + | + | No info | NA |
| 754G>A | D252N | 3 | C | – | – | – | No info | NA |
| 782G>A | R261H | 1 | C | + | + | + | No info | NA |
| 759insT | E273X | 3 | C | – | – | – | No entry | NA |
| 1037A>G | H346R | 1 | C | + | + | + | No entry | NA |
| 1137C>A | N379K | 1 | C | + | + | + | No entry | NA |
| 1153A>C | S385R | 1 | C | – | + | + | No entry | NA |
| 1219C>T | Q407X* | 1 | C | + | + | + | No entry | NA |
| 1228C>T | H410Y | 1 | C | (+) | – | – | No entry | NA |

PolyPhen2: benign –, possibly damaging (+), probably damaging ++; mutation taster: disease-causing +, benign polymorphism –; conservation: 100% conserved (16/16) +, <100% conserved –; dbSNP (KCNJ12): alignment was performed with KCNJ12 because KCNJ18 data are not available and the identity of the two coding sequences is 98.7%; severity: mild –, severe ++; segregation: no segregation –, no available relatives 0.

*Published as potential causative mutations previously.7
In summary, only T309I fulfills the criteria of a disease-causing mutation—but only if the two female carriers without PP history are explained as reduced penetrance, as sometimes reported for HypoPP.1,3 This interpretation cannot be excluded since the phenotype in the index case is clinically mild.

In the 474 controls, we found the known amino acid substitutions R6Q, Q39R, H40R, V100I, H118R, L156P, H192Q, V249I, F281L, and Y348F which are not considered to be pathogenic. Additionally, we identified seven synonymous, presumably non-pathogenic changes, the earlier reported questionably causative mutation Q407X, a prestart base change (c.–7C>T), the E273X stop mutation and 10 novel non-synonymous heterozygous changes (L15S, G34S, R81P, T193M, D252N, R261H, H346R, N379K, S385R, H410Y). Again, we evaluated the novel changes according to the above criteria (table 1):

- The prestart exchange c.–7C>T is considered ‘improbable’ to generate a splice site by MaxEnt.
- L15S, D252N, S385R and H401Y are not conserved, and the predictions on disease causality are discordant.
- G34S, T193M, R261H, H346R and N379K are perfectly conserved and concordantly predicted to be damaging, but all occurred only in healthy controls.

**DISCUSSION**

**KCNJ18 mutations have been reported to cause TPP.** A requirement for 
KCNJ18 being a responsible TPP gene in the presence of hyperthyroidism would be a mutation-specific change in T3-induced expression or translocation of the *mutant* product. Previous functional studies showed that only L156P translocates to the cell surface.9 No such effect has been demonstrated for any of other *KCNJ18* variants.

*KCNJ18* mutations have also been reported to cause sporadic periodic paralysis (SPP).8 A sporadic disease on a monogenic basis like SPP is due to an autosomal-dominant gene defect, arising by a new mutation transmitted through a non-penetrant or very mildly affected parent, or by a clinically unaffected parent who carries a mosaic germ line mutation.12 To clarify whether the 10 index patients have de novo variants, we have clinically studied and genotyped their parents and siblings. For each patient, a parent (or at least a sibling) carried the variant or was clinically affected. Therefore, we conclude that *KCNJ18* variants, if disease-causing at all, are neither frequent de novo mutations nor mosaic germ line mutations. Surprisingly, family members have not been studied in the article on SPP.8 Assuming that *KCNJ18* is a PP gene, it also remains unclear why both gain-of-function and dominant-negative mutations should have the same clinical effects, that is, weakness episodes.7 8

The identified variants, here, except for T309I, do not meet the requirements of a disease-causing mutation. All heterozygous stop mutations, such as E273X, R399X and Q407X (as well as early framshift mutations) should be excluded as relevant *KCNJ18* alterations. The missing carboxyterminus prevents the assembly with wild type proteins.11 All tetrameric channel complexes are, therefore, normal—no matter whether the incomplete RNA is unstable and immediately destructed or not. Often the other allele is overexpressed so that the number of normal channels is not reduced. For example, we are aware of healthy controls carrying in their CACNA1S gene a heterozygous stop mutation (c.C3709del) without showing a defect in muscle excitation-contraction coupling. With these individuals provocative stimuli do not elicit bouts of paralysis.

A functional change brought about by an ion channel variants is usually an important criterion for disease causality; but this criterion is not sufficient to prove causality, especially when the functional defects do not explain the phenotype. In previous studies, Kir2.6 mutations have been shown to lead to loss of function defects and suppression of the main rectifier Kir2.1.7 8 However, the Kir2.6 channel itself is already shown to have a dominant negative effect on Kir2.1 function.9 Therefore, a loss of function of the dominant negative suppression should actually lead to a gain of function of Kir2.1 which does not fit with the known depolarisation-induced paralysis pathogenesis.9 Testing for functional defects in a channel with unclear significance for muscle membrane potential would not be interpretable. Also, regardless of the outcome, it would not change the fact that the genetic criteria are not met for any of the variants but perhaps T309I. Finally, any functional effect could simply be the result of the variants being polymorphisms, such as sodium channel Nav1.4 variant S906T14 as well as others reviewed previously.9

In summary, it remains unclear whether *KCNJ18* is a PP gene. Without doubt, Kir2.6 contains functional units that could make it a PP gene. However, the large homology with *KCNJ12* leaves us with some problems concerning the interpretation. *KCNJ18* might be a duplication of *KCNJ12*, because the SNPs that occur in *KCNJ18* are predominantly related to amino acid positions that discern *KCNJ18* and *KCNJ12* (eg, L15S, Q39R, H40R, V100I, H118R, L156P, H192Q, V249I). Similarly the high percentage of rare variants in patients and controls (10/263=3.8% vs 17/474=3.6%) additionally questions the pathogenicity of these variants taken all together. Up to the final clarification neu rologists should not consider *KCNJ18* as an established periodic paralysis gene and should continue the genetic analysis of the three other known causative genes. If *KCNJ18* can be confirmed as PP gene by additional studies, the number of identified mutations9 8 and our results demonstrate its rarity.

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**Contributors** MK established and performed gene analysis and assisted with manuscript revisions; KJ-R and FL-H designed the study, collected patient DNA and data, and authored the manuscript.

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**Competing interests** None.

**Patient consent** Obtained.

**Ethics approval** These studies were approved by the Institutional Review Board of Ulm University (IRB Study #30/12_Pathogenesis of hypokalaemic periodic paralysis).

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**REFERENCES**


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Supplementary File S1. Analysis of KCNJ18 and our reference sequence

*KCNJ18* has one coding exon (exon 3). This exon was amplified using polymerase chain reaction (PCR) under the following conditions: 94 °C for 5 min, followed by 35 cycles with 94°C for 30 s, 65°C for 1 min and 72°C for 1.5 min as well as a final elongation at 72°C for 10 min. For this procedure the primers of Ryan et al. (2010) were slightly modified for higher specificity (CCAGACATGCTGTCCTCTCTGTC/ GGGCCTCTCCCGGCA) and amplified by means of AmpliTaq Gold with GeneAmp (Applied Biosystems by Life Technologies, Carlsbad, CA, USA). For each preparation of 50 µl we used 50 ng DNA.

For some products, a second amplification was done using a nested PCR with the primers (ATGCTGTCCCTCTCTGTCC/ CGGCCAGGGGTGGATGCTGACATGCTGACATG) under the following conditions: 94 °C for 5 min, followed by 35 cycles at 94 °C for 30 s, 65 °C for 1 min und 72 °C for 1.5 min as well as a final elongation at 72 °C for 10 min. The resulting products were sequenced using BigDye terminator v1.1, v3.1 mix and the forward-primers (CCAGACATGCTGCTCCTCTGTCC/CTGGCGCTACATGCTGCTCATC/ CGCCGTGTTGCCCCCTGCGAGAC/GCCAATGAGATCCTGTGGGGTCAC) on the capillary sequencer ABI 3130xl (Applied Biosystems by Life Technologies, Carlsbad, CA, USA).

Our wild type sequence differed from the published *KCNJ18* sequence in 11 bases and from the known *KCNJ12* sequence in 17 bases. Dr. Ryan’s actual *KCNJ18* sequence (personal communication) differed from our most frequent *KCNJ18* sequence in one base (c.473 T>C) that did not show up in our study. Our reference sequence for *KCNJ18* is:

**Exon 3**

ATG ACC GCG GCC AGC CGG GCC AAG CCC TAC AGC ATC GTG TCA TTG  
GAG GAG GAC GGG CTG CAT CTG GTC ACC ATG TCG GCC GCC AAC GGC  
TTC GGC AAC GCC AAG GTG CAC ACG CAG CAC AGG TGC CGC AAC CGC  
TTC GTC AAG AAG AAT GGC CAG TGC AAC ATT GCG TTC GCC AAC ATG
Supplementary Figure 1S: Lack of segregation in a PP family with KCNJ18-p.R401W.

Ten affected family members (and one clinically equivocal) are carriers as well as one non-affected relative. In contrast, three affected relatives do not carry R401W.