Novel mutations in the KCNQ2 gene link epilepsy to a dysfunction of the KCNQ2-calmodulin interaction

M C Richards, S E Heron, H E Spendlove, I E Scheffer, B Grinton, S F Berkovic, J C Mulley, A Davy

Key points

- We set out to ascertain KCNQ2 mutations in 10 families with benign familial neonatal seizures (BFNS), and to determine if any mutations from the C-terminal region affected the interaction between KCNQ2 and the ubiquitous calcium sensing protein, calmodulin (CaM).
- Eight novel mutations were found: two missense start codon mutations and a frameshift mutation in the N-terminal region and a truncation, two putative splice site mutations and two missense mutations in the C-terminal region.
- Using a yeast two hybrid assay, we determined that the KCNQ2 C-terminal missense mutations R353G and L619R both affect normal CaM binding. The R353G mutation lies within one of the two putative CaM binding regions and the L619R lies within close proximity of the other.
- On the basis of our data we postulate that dysfunction of CaM binding may be one of the pathogenic mechanisms associated with benign familial neonatal seizures.

Mutations in the voltage gated potassium channels KCNQ2 (OMIM 602235) and KCNQ3 (OMIM 602232) are associated with an autosomal dominant idiopathic epilepsy syndrome of newborns, benign familial neonatal seizures (BFNS) (OMIM 121200). BFNS is characterised by unprovoked partial seizures typically beginning when the infant is around three days old. BFNS associated genes were mapped to human chromosomes 20q13.3 and 8q24, which led to the identification by positional cloning of KCNQ2 as the chromosome 20 gene. KCNQ3 was subsequently identified as the 8q24 BFNS gene, based on genomic location and homology with KCNQ2.

The potassium channels of the KCNQ gene family consist of four subunits, each with a 6 transmembrane topological organisation. KCNQ subunits, comprising KCNQ1–5, have an intracellular amino terminus, a single pore loop (P-loop) that forms the selectivity filter of the channel, a positively charged, voltage sensing fourth transmembrane domain (S4), and a large intracellular carboxy terminus (C-terminus). All five known KCNQ proteins can form homomeric channels, but the association of specific subunits to form heteromeric channels is restricted to certain combinations.

KCNQ2 and KCNQ3 are mostly expressed in the central nervous system, where they form a heteromultimeric channel that mediates the neuronal muscarinic regulated current (M-current), also known as an M-channel or M-type channel that mediates the neuronal muscarinic regulated current (M-current). The M-current is a slowly activating, non-inactivating potassium conductance known to regulate neuronal excitability by determining the firing properties of neurones and their responsiveness to synaptic input. Because it is active at voltages near the threshold for action potential initiation, the M-current has a major impact on neuronal excitability.

Since the KCNQ2/KCNQ3 ion channel plays a pivotal role in the regulation of neuronal excitability, it is not surprising that several mutations in the gene have been associated with epilepsy. The first K+ channel mutations associated with epilepsy were reported in 1998 and more have subsequently been reported in both KCNQ2 and KCNQ3.

Whilst it is generally hypothesised that mutations in either KCNQ2 or KCNQ3 may result in a mild reduction of the M-current and subsequent dysfunction of the neuronal response to synaptic input, for the most part no significant dominant negative effects or alterations in channel gating or ion selectivity have been observed.

In one BFNS family, a mutation in the voltage-sensing S4 domain of KCNQ2 was shown to have a dominant negative effect on channel activation; however this was associated with the occurrence of myokymia in patients.

KCNQ2 C-terminal mutations account for over half of all known mutations associated with BFNS and to date there have been no functional studies elucidating the pathogenic effects of such mutations. Recently, the interaction between the C-terminal region of KCNQ2 and the ubiquitous calcium binding protein calmodulin (CaM) has been reported, suggesting a possible pathogenic mechanism of action for C-terminal KCNQ2 mutations.

Here, we report the association of eight novel KCNQ2 mutations in families with benign familial neonatal seizures. Specifically, we show that two missense mutations affect the binding affinity of CaM and we discuss the involvement of an aberrant KCNQ2-CaM interaction in epileptogenesis.

MATERIALS AND METHODS

Clinical data

Detailed clinical histories and pedigrees were obtained from families where two or more individuals had neonatal seizures, without a known extraneous cause. Venous blood was sampled for DNA extraction. The study was approved by the Austin and Repatriation Medical Centre Committee on Human Ethics.

Abbreviations: BFNS, benign familial neonatal seizures; CaM, calmodulin; C-terminus, carboxy terminus; KCNQ2, Homo sapiens potassium voltage gated channel, Kv7-like subfamily, member 2; KCNQ3, Homo sapiens potassium voltage gated channel, Kv7-like subfamily, member 3; P-loop, pore loop; M-current, neuronal muscarinic regulated current; S1–6, first to sixth transmembrane domains.
Mutation screening

KCNQ2 was screened by single strand conformation polymorphism analysis or sequencing as described elsewhere.26 Primer sequences for these and all other primers from the study are available from the authors upon request.

Generation of yeast two-hybrid screen constructs

A yeast two hybrid screen was carried out using the ProQuest™ Two-Hybrid System with Gateway™ Technology (Invitrogen™, Australia) according to the manufacturer’s directions. A KCNQ2 C-terminal entry clone was generated using the pENTR Directional TOPO® Cloning Kit (Invitrogen™, Australia). Primers were designed to amplify the intracellular C-terminal region of KCNQ2 based on the sequence of human KCNQ2 (Genbank accession number NM_172107). The 1611 bp cloned fragment included exon 10a (found in all our amplified clones), corresponding to aa 373–382.1,17 The extra 30 bp (10 aa) were included in our numbering. Following sequence verification, the KCNQ2 cDNA fragment was then subcloned into pDEST™32, the DNA Binding domain (DB) Gateway™ Destination Vector (Invitrogen™, Australia).

KCNQ2 C-terminal bait mutagenesis

Primer were designed to incorporate the c.1057C>G (R353G) and c.1856T>G (L619R) changes into the pDEST™32-KCNQ2 C-terminal bait construct. Overlapping PCR products were amplified, gel extracted and purified before a second round of PCR using the initial KCNQ2 F and R primers. These products were also gel extracted before cloning into the pDEST™32 bait vector using the TOPO® system. Mutant baits were sequence verified.

Yeast two-hybrid assay

The DBLeu (empty bait vector), DB-KCNQ2 wild-type (DB-Q2 wt) and mutant (R353G, L619R) C-term baits were transformed into the yeast strain Mav203 and plated onto minimal selective media lacking leucine. A duplicate was carried out, in which the empty Activation Domain (pAD) prey vector was cotransformed with the baits and plated onto minimal selective media lacking leucine (−Leu) and tryptophan (−Trp). Yeast control strains (Invitrogen™) were included on all plates. Control 1, used as a negative control, contains empty plasmids pPC97 and pPC86. Control 2 has pPC97-RB and pPC86-E2F1, which express a relatively weak interaction. Control 3 contains plasmids encoding the Drosophila DP (pPC97) and E2F (pPC86) domains that have a moderately strong interaction, and provide a control for plasmid shuffling. Control 4 contains pPC97-Fos and pPC86-Jun which express a relatively strong interaction, and control 5 has a pCL1 plasmid encoding full-length GAL4p and empty pPC86 and is used as a positive control. The constructs were tested for self-activation of the HIS3 and β-galactosidase reporter genes according to Invitrogen™ instructions. The interaction between each DB-Q2C mutant and CaM was then tested by yeast two-hybrid assay and compared to the interaction with DB-Q2 wt. Three different PCR amplified CaM clones were introduced by gap repair20 into the prey vector (pPC86) in the yeast strain expressing either DB-Q2C wt, DB-Q2C mutants, or the empty DBLeu vector, which was used as a negative control. CaM interaction with the DB-Q2C wt and mutants was then assessed by expression of the HIS3 and LacZ reporter genes.

Yeast protein extraction

Yeast cultures were prepared for protein extraction according to the Clontech protocol. Protein extracts were prepared by the Urea and sodium dodecyl sulphate method described elsewhere.21

Western blotting

The NuPage® Novex 4–12% Bis-Tris Precast gel system (Invitrogen™, Australia) was used to analyse the yeast protein extracts. Gels were run according to the manufacturer’s instructions. A 42 kDa Gal4 (DBD-1–147) positive control (Santa Cruz Biotechnology, Inc., Santa Cruz, California) and MagicMark™ Western Protein Standard (Invitrogen™, Australia) were run on the gel for comparison. A 1:1000 dilution of the mouse monoclonal Gal4 (DBD) antibody (RK51, Santa Cruz Biotechnology, Inc., Santa Cruz, CA) was used as the primary antibody, followed by a 1:5000 dilution of the horseradish peroxidase conjugated sheep antimouse immunoglobulin secondary antibody (Silenus Labs, Pty. Ltd., Australia). The membrane was visualised with SuperSignal® West Dura Extended Duration Substrate (Pierce Biotechnology, Inc., Rockford, IL).

Assay for β-galactosidase activity in liquid culture using chlorophenol red-β-D-galactopyranoside

The chlorophenol red-β-D-galactopyranoside assay was carried out according to the ProQuest™ Two-Hybrid System with Gateway™ Technology (Invitrogen™, Australia) instructions. The units of β-galactosidase in each sample were calculated, where 1 unit of β-galactosidase is defined as the amount that hydrolyses 1 μmol of chlorophenol red-β-D-galactopyranoside to chloramphenicol red and D-galactose in one minute:

$$\text{β-galactosidase units} = \frac{1000 \times \text{OD}_{574}}{V \times \text{OD}_{600}}$$

where $V$ = volume of culture used in the assay (ml); OD$_{574}$ = optical density at 574 nm, which is equal to the absorbance by chloramphenicol red (and light scattering by cell debris); OD$_{600}$ = optical density at 600 nm, which is equal to the cell density at the start of the assay. Data were analysed with Microsoft® Excel 2000 and for statistical evaluation, a two-tailed, unpaired Student’s t test was applied.

RESULTS AND DISCUSSION

Novel KCNQ2 mutations were found in eight out of ten BFNS families in our study. All of the mutations were found on one allele only (—that is, patients were heterozygous) and were shown to segregate within the families. None of the mutations were detected in at least 50 unaffected control samples. The novel mutations that we found comprised two mutations of the start codon (M1V, M1T), one frameshift (K69fsX119), one truncation (R430X), two putative splice site mutations (E491 and R570) and two missense mutations (R353G and L619R) (table 1 and fig 1). The one base insertion at aa 69 creates a premature termination of the protein within the N-terminal domain of KCNQ2 and, along with the two mutations of the start codon, would act as an allelic knockout, likely to result in decreased KCNQ2 subunit expression and a concomitant reduction in neuronal M-channels. The subsequent reduction in M-current in the brain is predicted to result in reduced inhibition of neuronal excitability, resulting in hypereexcitability as observed in benign familial neonatal seizures. The other five mutations are all located within the KCNQ2 C-terminal domain. All known mutations are shown in table 1 and fig 1.

To date, most functional studies have been on mutations located within transmembrane domains. In particular mutations associated with BFNS and found in the voltage sensing S4 domain of KCNQ2 were shown to alter the gating properties of the M-channels, displaying slower opening and faster closing kinetics and a decreased voltage sensitivity.
with no concomitant changes in maximal current or membrane expression.

Effects of mutations in the KCNQ2 C-terminal domain are less well understood, although the cytoplasmic C-terminal domain of KCNQ2 has been shown to be crucial for normal activity of the channel.12 13 19 30 31 The C-terminus accounts for 63% of the KCNQ2 protein and, in common with other KCNQ subunits, contains a conserved ‘A domain’ thought to be involved in subunit interactions and another distal short domain, encompassing the A domain and the putative assembly domain, was recently found to determine the subunit specificity of KCNQ channel assembly.32 33 and consequently essential for expression of functional homomeric or heteromeric channels. Heteromeric KCNQ2/KCNQ3 channels are expressed pre and postsynaptically, where they are likely to be associated with the cytoskeleton by means of scaffold proteins. KCNQ2, because it is more resistant to detergent proteins have a binding site for calcium, and it has been suggested that some ion channel proteins have a binding site for calmodulin.34 When this site is not occupied, the channel is in its inactivated (closed) conformation, but in response to an influx or internal release of calcium,calcified CaM binds to the channel, resulting in an activated (openable) form. Indeed, CaM was then tested by yeast two hybrid assay and compared with the interaction with DB-Q2 wt and the control interactions. Fig 3B shows the growth of transformed yeast and controls on –Leu–Trp selection after 24 h. Yeast can grow on –Leu if they contain the DB plasmid, and –Trp if they have the AD plasmid. Fig 3C shows growth of transformed yeast and controls on –Leu–Trp–His + 40 mM 3AT selective plate after 48 h. Only yeast cells where the expression of the His3 reporter gene is activated by interaction between the bait and prey plasmids can grow on this selective plate. Figs 3D–F show a LacZ filter assay for interaction between bait and prey plasmids after 2, 7, and 24 h, respectively. A change in colour from cream to blue indicates activation of the β-galactosidase reporter gene by interaction of the bait and prey plasmids. The Q2C R353G mutant did not interact with CaM, as seen by no growth on HIS3 selective plate (fig 3C) and no blue readout in the LacZ filter assay (figs 3D–F). On the other hand, the DB-Q2C L619R mutant was shown still to interact with CaM, as seen by growth on an HIS3 selective plate (fig 3C) and the blue readout in the LacZ filter assay (figs 3D–F). Interestingly, the DB-Q2C L619R mutant showed an even greater growth level on HIS3 selective plate than the DB-Q2C wt and also appeared to stain faster and more intensely blue in the LacZ filter assay, suggesting a stronger interaction between CaM and this mutant.

To quantify β-galactosidase activity more accurately, a second assay was carried out using the high sensitivity substrate chlorophenol red-β-D-galactopyranoside in liquid culture. The affinity of the DB-Q2C/AD-CaM interaction was measured in terms of units of β-galactosidase activity, with a zero value indicating no expression of the LacZ reporter gene, and hence no interaction. In the chlorophenol red-β-D-galactopyranoside assay, a value of 0.05 units β-galactosidase activity (fig 4) was significantly different from the empty bait vector replicate (p<0.01, Student’s t test), confirming the interaction of the DB-Q2C wt with CaM. As observed in the LacZ filter assay, the chlorophenol red-β-D-galactopyranoside assay showed a significant difference in the interaction between the Q2C R353G mutant and CaM as compared to the wild-type replicate (p<0.01, Student’s t test, fig 4).

These results suggest that the R353G mutation alters the structural conformation of the KCNQ2 C-terminal domain such that it is no longer able to bind to CaM. Indeed, the R353G missense mutation is within one of the CaM binding domains previously identified in KCNQ235 36 (fig 1). By abolishing CaM binding, the R353G mutation could lead to an impairment of M-current in vivo, through decreased opening of the channel. This might be predicted to have a similar effect on neuronal excitability to truncation and frameshift BFNS mutations that disrupt the structure of the KCNQ2 protein.

This is supported by experiments where KCNQ2 mutants deficient in CaM binding co-expressed with KCNQ3 in Chinese hamster ovary cells did not give rise to detectable currents.15 The membrane targeting of the mutants was not altered, neither was the ability to co-assemble with KCNQ3, thus demonstrating that CaM binding is not necessary for membrane biogenesis of KCNQ2 or KCNQ3 channels. Furthermore, it has been suggested that some ion channel proteins have a binding site for calmodulin.37 When this site is not occupied, the channel is in its inactivated (closed) conformation, but in response to an influx or internal release of calcium, calcified CaM binds to the channel, resulting in an activated (openable) form. Indeed, CaM was shown in two studies to be tethered constitutively to the M-channel, suggesting that the C-terminal domain of KCNQ channels may fold in a manner that allows CaM to induce M-channel opening.14 15

### Table 1

<table>
<thead>
<tr>
<th>Variant</th>
<th>Amino Acid change</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>c.1A→G</td>
<td>M1V (no start)</td>
<td>N-ter</td>
</tr>
<tr>
<td>c.2T→C</td>
<td>M1T (no start)</td>
<td>N-ter</td>
</tr>
<tr>
<td>c.204insC</td>
<td>K69X119</td>
<td>N-ter</td>
</tr>
<tr>
<td>c.585insT</td>
<td>H194PfsX262</td>
<td>Loop II–III</td>
</tr>
<tr>
<td>c.587C→T</td>
<td>A166V</td>
<td>S4</td>
</tr>
<tr>
<td>c.590T→C</td>
<td>L197P</td>
<td>S4</td>
</tr>
<tr>
<td>c.619C→T</td>
<td>R207W</td>
<td>S4</td>
</tr>
<tr>
<td>c.640C→T</td>
<td>R214W</td>
<td>S4</td>
</tr>
<tr>
<td>c.740C→G</td>
<td>S247W</td>
<td>S5</td>
</tr>
<tr>
<td>c.749T→C</td>
<td>V250G</td>
<td>S5</td>
</tr>
<tr>
<td>c.847insGT</td>
<td>K283x319</td>
<td>P-loop</td>
</tr>
<tr>
<td>c.851A→G</td>
<td>Y284C</td>
<td>P-loop</td>
</tr>
<tr>
<td>c.926G→A</td>
<td>A306T</td>
<td>S6</td>
</tr>
<tr>
<td>c.1016T→G</td>
<td>L339R</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1057C→G</td>
<td>R353G</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1221T→G</td>
<td>Splice donor site (K407)</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1288C→T</td>
<td>Truncation (R430X)</td>
<td>C-ter</td>
</tr>
<tr>
<td>Unknown</td>
<td>R458X</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.14711G→A</td>
<td>Splice donor site (E491)</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1501delC</td>
<td>S504delS506</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1577G→A</td>
<td>Splice donor site (C526)</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1630insGCCCT</td>
<td>Y546insG549</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1688G→A</td>
<td>R563Q</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1711delT</td>
<td>Exonic splice site (R570)</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1856T→G</td>
<td>L619R</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.1870delTTG</td>
<td>Y626fsX910</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.2073delCTT</td>
<td>P284X</td>
<td>C-ter</td>
</tr>
<tr>
<td>c.2543delG</td>
<td>G848fsX911</td>
<td>C-ter</td>
</tr>
</tbody>
</table>

*Our data; †Numbering according to authors; ‡Phenotype not compatible with typical benign familial neonatal seizures.

...
In contrast, the chlorophenol red-β-D-galactopyranoside assay for the L619R Q2C mutant showed a significantly higher level of β-galactosidase activity (0.26 units) than the wild-type replicate (p < 0.001, Student’s t test, fig 4). This finding indicates that the L619R mutation alters the conformation of the protein in a manner that increases CaM binding to the KCNQ2 C-terminal domain approximately fivefold.

More recent work by Gamper and Shapiro 17 is consistent with a role for CaM in Ca²⁺ sensing rather than in channel assembly (as found by Wen and Levitan 15 and Yus-Nájera et al16), or at least a dual role for CaM in KCNQ channel physiology. Their work suggests that bradykinin modulation of the M-current, at least in sympathetic neurones, uses CaM in concert with rises in calcium concentration ([Ca²⁺]), over the physiological range of <10–400 nM.

KCNQ2 and KCNQ3 channels are strongly modulated by several distinct signalling pathways in neurones that are mediated by second messengers, and these include M1-type muscarinic receptors and activators of G-protein coupled receptors, such as bradykinin B₂ and purinergic P₂Y₂. The signal provoked by the latter two receptors involves the phospholipase C/inositol triphosphate (PLC/IP₃) pathway and release of Ca²⁺ from internal stores to produce rises in intracellular [Ca²⁺].

Gamper and Shapiro 17 show that bradykinin induced rises in intracellular [Ca²⁺] act on M-channels via Ca²⁺-CaM to suppress the M-current. In the case of the KCNQ2 L619R

---

**Figure 1** Model of the KCNQ2 channel protein showing the known mutations associated with BFNS, numbered according to the mutated residue. Numbering is based on the long form of KCNQ2 including exon 10a. * numbering according to the authors (see table 1). Grey regions of the C-terminal loop indicate CaM binding regions. 15 16

**Figure 2** Pedigrees of families with benign familial neonatal seizures. *, members carrying a KCNQ2 mutation that have been tested; -, members who were tested and are negative for the mutation.

---

In contrast, the chlorophenol red-β-D-galactopyranoside assay for the L619R Q2C mutant showed a significantly higher level of β-galactosidase activity (0.26 units) than the wild-type replicate (p < 0.001, Student’s t test, fig 4). This finding indicates that the L619R mutation alters the conformation of the protein in a manner that increases CaM binding to the KCNQ2 C-terminal domain approximately fivefold.

More recent work by Gamper and Shapiro 17 is consistent with a role for CaM in Ca²⁺ sensing rather than in channel assembly (as found by Wen and Levitan 15 and Yus-Nájera et al16), or at least a dual role for CaM in KCNQ channel physiology. Their work suggests that bradykinin modulation of the M-current, at least in sympathetic neurones, uses CaM in concert with rises in calcium concentration ([Ca²⁺]), over the physiological range of <10–400 nM.

KCNQ2 and KCNQ3 channels are strongly modulated by several distinct signalling pathways in neurones that are mediated by second messengers, and these include M1-type muscarinic receptors and activators of G-protein coupled receptors, such as bradykinin B₂ and purinergic P₂Y₂. The signal provoked by the latter two receptors involves the phospholipase C/inositol triphosphate (PLC/IP₃) pathway and release of Ca²⁺ from internal stores to produce rises in intracellular [Ca²⁺].

Gamper and Shapiro 17 show that bradykinin induced rises in intracellular [Ca²⁺] act on M-channels via Ca²⁺-CaM to suppress the M-current. In the case of the KCNQ2 L619R
mutation that results in increased binding of CaM to the M-channel, it can be inferred that the effect of bradykinin modulation of M-channels would be amplified because of a likely increased sensitivity to $[\text{Ca}^{2+}]$. Thus, in the case of the L619R mutant channel, in response to a rise in bradykinin-induced intracellular $[\text{Ca}^{2+}]$, the KCNQ2/KCNQ3 derived M-current would be suppressed to a greater extent than for the wild-type channel. Suppression of the M-current would lead to decreased inhibition of neuronal excitability and could potentially be associated with BFNS pathogenicity.

CONCLUSION

Our results implicate CaM in the pathogenesis of epilepsy and specifically in the BFNS syndrome. Whilst further work will be required to elucidate fully the involvement of the KCNQ2-CaM interaction in neuronal excitability and its correlation with idiopathic epilepsy, these data suggest that dysfunction of this interaction leads to aberrant neuronal excitability in some BFNS patients.

ACKNOWLEDGEMENTS

We thank the families for their participation and Allison J Hall for technical assistance. Family A was kindly referred by Dr Gregory Holmes (Children's Hospital, Boston) and family B by Dr Jeremy Freeman (Royal Children’s Hospital, Melbourne).

Authors' affiliations

M C Richards, S E Heron, H E Spendlove, J C Mulley, Department of Laboratory Genetics, Women’s and Children’s Hospital, North Adelaide, South Australia, Australia
I E Scheffer, B Grinton, S F Berkovic, Epilepsy Research Institute and Department of Medicine (Neurology), University of Melbourne, Austin and Repatriation Medical Centre, Heidelberg, Victoria, Australia
J C Mulley, Department of Molecular Biosciences, University of Adelaide, Adelaide, Australia
A Davy, Bionomics Ltd, 31 Dalgleish St, Thebarton, South Australia, Australia

The study was supported by the Australian National Health and Medical Research Council and Bionomics Ltd.

Bionomics Ltd funded part of the study.

Correspondence to: Dr A Davy, Bionomics Ltd, 31 Dalgleish St, Thebarton, SA, Australia, 5031; adavy@bionomics.com.au

Received 1 September 2003
Revised version received 21 October 2003
Accepted 28 October 2003

REFERENCES


Novel mutations in the KCNQ2 gene link epilepsy to a dysfunction of the KCNQ2-calmodulin interaction

M C Richards, S E Heron, H E Spendlove, I E Scheffer, B Grinton, S F Berkovic, J C Mulley and A Davy

*J Med Genet* 2004 41: e35
doi: 10.1136/jmg.2003.013938