The CACNA1B R1389H variant is not associated with myoclonus-dystonia in a large European multicentric cohort

Niccolo E. Mencacci1, Léa R’bibo1, Sara Bandres-Ciga1,2, Miryam Carecchio3,4, Giovanna Zorzi3, Nardo Nardocci3, Barbara Garavaglia4, Amit Batla5, Kailash P. Bhatia5, Alan M. Pittman1, John Hardy1, Anne Weissbach6, Christine Klein6, Thomas Gasser7, Ebba Lohmann7 and Nicholas W. Wood1,*

1Department of Molecular Neuroscience, Institute of Neurology, University College London, London WC1N 3BG, UK, 2Department of Physiology and Institute of Neurosciences Federico-Olóriz, Centro de Investigaciones Biomedicas (CIBM), University of Granada, Granada 18071, Spain, 3Neuropediatrics Unit, IRCCS Istituto Neurologico Carlo Besta, Milan 20133, Italy, 4Molecular Neurogenetics Unit, IRCCS Istituto Neurologico Carlo Besta, Milan 20133, Italy, 5Sobell Department of Motor Neuroscience and Movement Disorders, UCL Institute of Neurology, London WC1N 3BG, UK, 6Institute of Neurogenetics, University of Lübeck, Lübeck 23538, Germany and 7Department of Neurodegenerative Diseases, Hertie Institute for Clinical Brain Research, University of Tübingen, and German Center for Neurodegenerative Diseases (DZNE), Tübingen 72076, Germany

*To whom correspondence should be addressed at: Department of Molecular Neuroscience, UCL Institute of Neurology, Queen Square, WC1N 3BG London, UK. Tel: +44 2034484255; Fax: +44 2072785616; Email: n.wood@ucl.ac.uk

Abstract

Myoclonus-dystonia (M-D) is a very rare movement disorder, caused in ~30–50% of cases by mutations in SGCE. The CACNA1B variant c.4166G>A; (p.R1389H) was recently reported as the likely causative mutation in a single 3-generation Dutch pedigree with five subjects affected by a unique dominant M-D syndrome and cardiac arrhythmias. In an attempt to replicate this finding, we assessed by direct sequencing the frequency of CACNA1B c.4166G>A; (p.R1389H) in a cohort of 520 M-D cases, in which SGCE mutations had been previously excluded. A total of 146 cases (28%) had a positive family history of M-D. The frequency of the variant was also assessed in 489 neurologically healthy controls and in publicly available data sets of genetic variation (1000 Genomes, Exome Variant Server and Exome Aggregation Consortium). The variant was detected in a single sporadic case with M-D, but in none of the 146 probands with familial M-D. Overall, the variant was present at comparable frequencies in M-D cases (1 out of 520; 0.19%) and healthy controls (1 out of 489; 0.2%). A similar frequency of the variant was also reported in all publicly available databases. These results do not support a causal association between the CACNA1B c.4166G>A; (p.R1389H) variant and M-D.
Introduction

Myoclonus-dystonia [M-D (MIM 159900)] is a rare familial movement disorder, which classically features a variable combination of non-epileptic myoclonic jerks and dystonia (1). Heterozygous loss-of-function mutations in the maternally imprinted ε-sarcoglycan gene [SCGE, DYT11; (MIM 604149)] represent a major cause of autosomal dominant M-D (2). However, up to 50–70% of familial cases with M-D lack mutations in SCGE (3–5), suggesting that disease-causing mutations in other genes are responsible for this syndrome.

Recently, Groen et al. (6) identified the missense variant c.4166G>A; (p.R1389H) (rs184841813) in CACNA1B [MIM 601012] as the likely causative mutation in a Dutch pedigree with five subjects affected by autosomal dominant M-D lacking mutations in SCGE. Unique features in the pedigree were lower limb orthostatic high-frequency myoclonus, attacks of limb painful cramps and cardiac arrhythmias in three of the affected subjects (7). Sanger sequencing of the CACNA1B exons coding for the protein portion spanning from III-S5 to III-S6 failed to reveal other mutations in a further 47 M-D cases.

CACNA1B encodes neuronal voltage-gated calcium channels CaV2.2, which have a key role in controlling synaptic neurotransmitter release (8). Furthermore, CACNA1A [MIM 600111] mutations in the homologous region of the gene cause familial hemiplegic migraine [MIM 141500] (9) and episodic ataxia type 2 (MIM 108500) (10).

The CACNA1B p.R1389H substitution represents an excellent candidate as a disease-causing mutation for M-D. However, in the absence of identification of CACNA1B mutations in other unrelated pedigrees, the implication of mutations in this gene as a cause for M-D is not confirmed.

In this study, we assessed the frequency of the CACNA1B c.4166G>A; (p.R1389H) variant in a multicentric cohort of M-D cases without mutations in SGCE (both point mutations and copy number variants).

Results

A total of 520 M-D cases (28% were familial) were screened for the presence of the c.4166G>A; (p.R1389H) variant. Additionally, we assessed the frequency of the variant in whole-exome sequencing data from 489 white healthy controls of UK and US origin and in European cases listed in publicly available data sets of genetic variation (1000 Genomes, Exome Variant Server and Exome Aggregation Consortium).

None of the 146 probands with familial M-D carried the CACNA1B c.4166G>A; (p.R1389H) variant. The variant was detected only in a single female case of UK origin with sporadic M-D (see chromatogram of the mutation in the Supplementary Material, Fig. S1). This case presented in her mid-30s with tremulous cervical dystonia and myoclonic jerks in the upper limbs. She had no family history for M-D or any other movement disorder. No other family members were available for segregation analysis of the variant.

The total carrier frequency in our M-D cohort, including familial and sporadic cases, is 0.19% (1/520 cases). The variant is present at a similar frequency in our healthy controls (0.2%; 1 out of 489 individuals). The control carrier of the variant is a 38-year-old male without any neurological disease and with no relevant family history of movement disorders.

The CACNA1B c.4166G>A; (p.R1389H) variant is reported at comparable frequencies in the 1000 genome project (0.26%; 1/379 individuals) and Exome Variant Server (0.28%; 12/4203 individuals) databases. In the Exome Aggregation Consortium database, c.4166G>A; (p.R1389H) is present in 0.11% (38 out of 33,367) of the European subjects (difference to M-D cases not significant; Fisher’s exact test \( P = 0.4 \)).

Discussion

The advent of next generation sequencing has led to an extraordinary acceleration in the discovery rate of rare genetic variants, the majority of which are of uncertain clinical significance. Hence, a close scrutiny is necessary before causally linking a candidate variant to a disease. To avoid false assignment of pathogenicity, MacArthur et al. (11) have recently proposed guidelines for implicating causality of rare variants in human disease.

In family-based studies, assessment of co-inheritance of a candidate variant with the disease status within family members represents the first requirement to prove causality.

The c.4166G>A; (p.R1389H) variant was identified by Groen et al. through a combination of whole-exome sequencing and linkage analysis (13 chromosomal regions identified, with a maximum LOD score of 1.2) in a single dominant M-D pedigree. Notably, two other rare missense changes, c.10355A>G; (p.Q9452R) in VPS13D [MIM 6068877] and c.5308C>T; (p.R1770C) in SPTAN1 [MIM 182810], were found to perfectly co-segregate with the disease in the family. De novo mutations in SPTAN1 have been shown to cause a neurological phenotype (West syndrome with severe cerebral hypomyelination, spastic quadriplegia and developmental delay) (12) and more recently a microdeletion encompassing SPTAN1 was detected in a child with epileptic encephalopathy and severe dystonia (13).

Given the clinical presentation pointing towards a possible channelopathy, the authors assumed that the causative variant was the one in CACNA1B (6).

However, co-segregation of a variant with disease in a single pedigree does not establish with certainty its pathogenic role, especially if other co-segregating coding variants and the possibility of a separate undetected pathogenic variant in linkage disequilibrium cannot be convincingly ruled out.

In addition, a candidate variant responsible for a rare disease should be found at a low frequency in population controls, consistent with the proposed model of inheritance and disease prevalence.

M-D is a very rare disorder with a suggested prevalence of around two per million in Europe (14). We would therefore anticipate highly penetrant mutations causing dominant forms of M-D to be absent or extremely rare in the general population. Yet, this is not the case for p.R1389H, which is present at a considerable frequency in our healthy controls and all publicly available databases (~0.1–0.3%). According to the Exome Aggregation Consortium database, the carrier frequency of this variant in Europeans is ~4 times higher than the TOR1A [MIM 605204] c.904_906delGAG deletion (0.026%), which is by far the most common single mutation responsible for dystonia described to date (15). Given this frequency, if c.4166G>A; (p.R1389H) were a pathogenic variant, we would expect it to be responsible for a large proportion of familial M-D cases. However, in our cohort, not only was the variant not identified in any of the probands with familial M-D, but the overall frequency of the variant did not differ between M-D cases and healthy controls. This does not support a pathogenic effect of the variant even assuming a reduced penetrance.

In conclusion, our study suggests that the role of the CACNA1B variant c.4166G>A; (p.R1389H) as a cause for M-D is questionable.
Further genetic evidence is needed before designating CACNA1B mutations as a cause for dominant M-D.

Materials and Methods

A total of 520 M-D cases of British, German and Italian origin were recruited in four tertiary movement disorders centers (London, Lübeck, Tübingen and Milan). All selected cases fulfilled the proposed diagnostic criteria for M-D (2). A total of 146 cases (28%) had a positive family history of M-D. All participants provided written informed consent.

M-D cases were screened by direct Sanger sequencing for mutations in exon 28 of CACNA1B (RefSeq NM_000718.3), which contains the c.4166G>A; p.(R1389H) variant. Each reaction was performed in a 20 µl volume containing 10 µl of FastStart PCR master mix (Roche), 5 µl of water, 2 µl of each primer (5pmol/µl) and 30 ng of genomic DNA. After purification, PCR products were sequenced in both forward and reverse directions using BigDye Terminator v3.1 sequencing chemistry and then were loaded on the ABI3730xl genetic analyser (Applied Biosystems, Foster City, CA, USA). The sequences were analysed with Sequencher software (version 4.9; Gene Codes).

Whole-exome sequencing data from 488 white healthy controls of UK and US origin were provided by the International Parkinson’s Disease Genomic Consortium (IPDGC). In short, prior to sequencing, DNA templates were bridge amplified to form clonal clusters inside a flowcell via the cBot cluster generation process. The flowcells were then loaded into the next-generation sequencer Illumina HiSeq 2000. Paired end sequence reads were aligned with Burrows-Wheeler Aligner (BWA) against the reference human genome (UCSC hg19). Duplicate read removal, format conversion and indexing were performed with Picard (http://picard.sourceforge.net/). The Genome Analysis Toolkit (GATK) was used to recalibrate base quality scores, perform local realignments around possible indels and to call and filter the variants.

Web Resources

1000 Genomes project (URL: http://www.1000genomes.org/) [last accessed: April 2015].
Exome Variant Server, NHLBI GO Exome Sequencing Project (ESP), Seattle, WA, USA (URL: http://evs.gs.washington.edu/EVS/) [last accessed: April 2015].
Exome Aggregation Consortium (ExAC), Cambridge, MA, USA (URL: http://exac.broadinstitute.org) [last accessed: April 2015].

Supplementary material

Supplementary Material is available at HMG online.

Acknowledgements

We are grateful to all participants in this study. We thank the International Parkinson Disease Genomic Consortium (IPDGC) for providing the exome sequencing data of healthy population controls. We would also like to thank the Exome Aggregation Consortium and the groups that provided exome variant data for comparison. A full list of contributing groups can be found at http://exac.broadinstitute.org/about. The authors would also like to thank the NHLBI GO Exome Sequencing Project and its ongoing studies which produced and provided exome variant calls for comparison: the Lung GO Sequencing Project (HL-102923), the WHI Sequencing Project (HL-102924), the Broad GO Sequencing Project (HL-102925), the Seattle GO Sequencing Project (HL-102926) and the Heart GO Sequencing Project (HL-103010).

Conflict of Interest statement. None declared.

Funding

The work was undertaken at University College London Hospitals (UCLH) and University College London (UCL), who receive support from the Department of Health’s NIHR Biomedical Research Centers funding streams. N.E.M. is funded by an MRC-Wellcome Trust grant. S.B. holds an FPU fellowship from the Spanish Ministry of Education and Science jointly with a short-term stay grant by Cei-BioTic and University of Granada. A.M.P. is funded by the Reta Lila Weston Trust. C.K. is the recipient of a career development award from the Herman and Lilly Schilling Foundation. E.L. and T.G. are supported by a grant from the Dystonia Medical Research Foundation (DMRF). The funders had no role in study design, data collection and analysis, decision to publish or preparation of the manuscript. This work was supported by a Medical Research Council/Wellcome Trust Strategic Award (grant number WT089698/Z/09/Z) and a grant from the Bachman-Strauss Dystonia Parkinsonism Foundation. Funding to pay the Open Access publication charges for this article was provided by Medical Research Council/Wellcome Trust.

References

1. Nardocci, N. (2011) Myoclonus-dystonia syndrome. Hand. Clin. Neurol. 100, 563–575.
2. Kinugawa, K., Vidalinhet, M., Clot, F., Aparis, E., Grabl, D. and Roze, E. (2009) Myoclonus-dystonia: an update. Mov. Disord., 24, 479–489.
3. Ritz, K., Gerrits, M.C., Foncke, E.M., van Ruijssen, F., van der Linden, C., Vergouwen, M.D., Bloem, B.R., Vandenbergh, W., Crols, R., Speelman, J.D. et al. (2009) Myoclonus-dystonia: clinical and genetic evaluation of a large cohort. J. Neurol. Neurosurg. Psychiatry, 80, 653–658.
4. Carecchio, M., Magliozzi, M., Copetti, M., Ferraris, A., Bernar-
dini, L., Bonetti, M., Defazio, G., Edwards, M.J., Torrente, I., Pel-
legrini, F. et al. (2013) Defining the epsilon-sarcoglycan (SGCE) gene phenotypic signature in myoclonus-dystonia: a re-appraisal of genetic testing criteria. Mov. Disord., 28, 787–794.
5. Grunewald, A., Djarmati, A., Lohmann-Hedrich, K., Farrell, K., Zeller, J.A., Allert, N., Papengut, F., Petersen, B., Fung, V., Sue, C.M. et al. (2008) Myoclonus-dystonia: significance of large SGCE deletions. Hum. Mutat., 29, 331–332.
6. Groen, J.L., Andrade, A., Ritz, K., Jalalzadeh, H., Haagmans, M., Bradley, T.E., Jongejan, A., Verbeek, D.S., Nurnberg, P., Denome, S. et al. (2015) CACNA1B mutation is linked to unique myoclonus-dystonia syndrome. Hum. Mol. Genet., 24, 987–993.
7. Groen, J., van Rootselaar, A.F., van der Salm, S.M., Bloem, B.R. and Tijsen, M. (2011) A new familial syndrome with dystonia and lower limb action myoclonus. Mov. Disord., 26, 896–900.
8. Beuckmann, C.T., Sinton, C.M., Miyamoto, N., Ino, M. and Ya-
 nagisawa, M. (2003) N-type calcium channel alpha1B subunit (Cav2.2) knock-out mice display hyperactivity and vigilance state differences. J. Neurosci., 23, 6793–6797.
9. Carrera, P., Piatti, M., Stenirri, S., Grimaldi, L.M., Marchioni, E., Curcio, M., Righetti, P.G., Ferrari, M. and Gelfi, C. (1999) Genetic heterogeneity in Italian families with familial hemiplegic mi-
graine. Neurology, 53, 26–33.
10. Jen, J., Wan, J., Graves, M., Yu, H., Mock, A.F., Coulin, C.J., Kim, G., Yue, Q., Papazian, D.M. and Baloh, R.W. (2001) Loss-of-function EA2 mutations are associated with impaired neuromuscular transmission. *Neurology*, 57, 1843–1848.

11. MacArthur, D.G., Manolio, T.A., Dimmock, D.P., Rehm, H.L., Shendure, J., Abecasis, G.R., Adams, D.R., Altman, R.B., Antonarakis, S.E., Ashley, E.A. et al. (2014) Guidelines for investigating causality of sequence variants in human disease. *Nature*, 508, 469–476.

12. Saitsu, H., Tohyama, J., Kumada, T., Egawa, K., Hamada, K., Okada, I., Mizuguchi, T., Osaka, H., Miyata, R., Furukawa, T. et al. (2010) Dominant-negative mutations in alpha-II spectrin cause West syndrome with severe cerebral hypomyelination, spastic quadriplegia, and developmental delay. *Am. J. Hum. Genet.*, 86, 881–891.

13. Matsumoto, H., Zaha, K., Nakamura, Y., Hayashi, S., Inazawa, J. and Nonoyama, S. (2014) Chromosome 9q33q34 microdeletion with early infantile epileptic encephalopathy, severe dystonia, abnormal eye movements, and nephroureteral malformations. *Pediatr. Neurol.*, 51, 170–175.

14. Asmus, F. and Gasser, T. (2010) Dystonia-plus syndromes. *Eur. J. Neurol.*, 17(Suppl. 1), 37–45.

15. Valente, E.M., Warner, T.T., Jarman, P.R., Mathen, D., Fletcher, N.A., Marsden, C.D., Bhatia, K.P. and Wood, N.W. (1998) The role of DYT1 in primary torsion dystonia in Europe. *Brain*, 121, 2335–2339.