Screening of CACNA1A and ATP1A2 genes in hemiplegic migraine: clinical, genetic, and functional studies

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Abstract
Hemiplegic migraine (HM) is a rare and severe subtype of autosomal dominant migraine, characterized by a complex aura including some degree of motor weakness. Mutations in four genes (CACNA1A, ATP1A2, SCN1A and PRRT2) have been detected in familial and in sporadic cases. This genetically and clinically heterogeneous disorder is often accompanied by permanent ataxia, epileptic seizures, mental retardation, and chronic progressive cerebellar atrophy. Here we report a mutation screening in the CACNA1A and ATP1A2 genes in 18 patients with HM. Furthermore, intragenic copy number variant (CNV) analysis was performed in CACNA1A using quantitative approaches. We identified four previously described missense CACNA1A mutations (p.Ser218Leu, p.Thr501Met, p.Arg583Gln, and p.Thr666Met) and two missense changes in the ATP1A2 gene, the previously described p.Ala606Thr and the novel variant p.Glu825Lys. No structural variants were found. This genetic screening allowed the identification of more than 30% of the disease alleles, all present in a heterozygous state. Functional consequences of the CACNA1A-p.Thr501Met and ATP1A2-p.Glu825Lys, were investigated by means of electrophysiological studies, cell viability assays or Western blot analysis. Our data suggest that both these variants are disease-causing.
Introduction

Familial or sporadic hemiplegic migraine (FHM, MIM #141599, or SHM) are rare subtypes of migraine with aura (MA) characterized by paroxysmal episodes of hemiparesis generally preceding or accompanying a headache attack (International Headache Society [IHS] 2004)). FHM is considered a monogenic disorder and follows an autosomal dominant inheritance pattern (Pietrobon 2007). Both FHM and SHM are genetically heterogeneous disorders. Up to now, four causative genes have been described in hemiplegic migraine (HM): CACNA1A on chromosome 19p13 (FHM1, MIM #301011) (Ophoff et al. 1996), ATP1A2 at 1q23 (FHM2, MIM #182340) (De Fusco et al. 2003), SCN1A at 2q24 (FHM3, MIM #182389) (Dichgans et al. 2005) and, recently, PRRT2 at 16p11.2 (MIM #614386) (Riant et al. 2012). Additionally, two other loci have been reported in FHM families at 1q31 (Gardner et al. 1997) and 1q32 (Cuenca-Leon et al. 2009), although the specific genetic defects have not yet been uncovered.

Mutational screenings of HM patients have reported more than 30 mutations in the CACNA1A gene, over 60 mutations in the ATP1A2 gene, only five in SCN1A (de Vries et al. 2009; Riant et al. 2010a; Freilinger et al. 2011) and eight in PRRT2 (Cloarec et al. 2012; Dale et al. 2012; Gardiner et al. 2012; Marini et al. 2012; Riant et al. 2012). Additionally, a quantitative study that used multiple ligation-dependent probe amplification (MLPA) identified a deletion of exons 39–47 of CACNA1A in a SHM patient (Labrum et al. 2009). CACNA1A encodes the pore-forming z1 subunit of the voltage-gated neuronal Ca2.1 (P/Q-type) channel. Ca2.1 channels are located in cortical glutamatergic presynaptic terminals and play an important role in controlling neurotransmitter release. ATP1A2 encodes the z2 subunit of the Na+/K+ ATPase, is expressed in astrocytes and is involved in the clearance of extracellular K+ and production of a Na+ gradient used in the reuptake of glutamate. SCN1A encodes the z1 subunit of the neuronal voltage-gated sodium channel NaV1.1. This channel is critical in the generation and propagation of action potentials (Wessman et al. 2007). Finally, PRRT2 codes for a transmembrane protein of unknown function that is capable to bind to synosomal-associated protein 25 (SNAP25), which suggests a role in synaptic exocytosis (Lee et al. 2012).

The allelic heterogeneity displayed by the CACNA1A gene also correlates with substantial clinical variation, as mutations in this gene are also responsible for two other autosomal dominant diseases: episodic ataxia type 2 (EA2, MIM #108500) and spinocerebellar ataxia type 6 (SCA6, MIM #183086). The range of CACNA1A-linked phenotypes has even been broadened by the recent descriptions of patients presenting with alternating hemiplegia of childhood (de Vries et al. 2008), acute striatal necrosis (Carreño et al. 2011), hemiplegia–hemiconvulsion–epilepsy syndrome (Yamazaki et al. 2011), and recurrent ischemic stroke (Knierim et al. 2011). Clinical variation is also seen within the HM phenotype, a condition in which CACNA1A may sometimes be implicated as a modifier gene rather than a disease-causing gene (Serra et al. 2010). Typical attacks in HM are often associated with other aura symptoms: the clinical spectrum includes permanent cerebellar signs and, less frequently, various types of epileptic seizures, mental retardation, and coma. Furthermore, in approximately 50% of FHM1/CACNA1A families, chronic progressive ataxia occurs independently of the migraine attacks (IHS 2004). ATP1A2 has also been associated with alternating hemiplegia of childhood (Bassi et al. 2004). Also, the SCN1A gene has been associated with phenotypes other than HM, as it has been identified as a cause of generalized epilepsy with febrile seizures plus type 2 (GEFS+2, MIM #604403) (Escayg et al. 2000), severe myoclonic epilepsy in infancy (SMEI, MIM #607208), also called Dravet syndrome (Dravet 2011), childhood epilepsy with generalized tonic-clonic seizures (ICEGTC, MIM #607208), familial febrile convulsions type 3A (FEB3A, MIM #604403) (Mantegazza et al. 2005), and elicited repetitive daily blindness (ERDB) with HM (Vahedi et al. 2009). Finally, mutations in PRRT2 have been found in a number of paroxysmal disorders, including paroxysmal kinesigenic dyskinesia (PKD, MIM #128200), infantile convulsions with PKD (PKD/IC, MIM #602066), benign familial infantile epilepsy (BFIE, MIM #605751), and episodic ataxia or febrile seizures, apart from HM (Wood 2012).

At the functional level, HM and EA2 mutations typically have opposite effects on the CaV2.1 channels leading to increased or decreased Ca2+ influx, respectively (Pietrobon 2013). HM-related mutations in the ATP1A2 gene typically produce a loss of function of the pump (de Vries et al. 2009).

In a previous study, we analyzed 21 Spanish patients with HM episodes and identified three mutations in the CACNA1A gene, but no disease-causing changes in ATP1A2 (Cuenca-Leon et al. 2008). In this study we analyzed 18 additional patients with HM of Spanish and Greek origin and identified four mutations in the CACNA1A gene and two mutations in ATP1A2. The two changes that had not previously been studied at the functional level were subjected to functional analyses to establish their relevance to the disease phenotype.

Materials and Methods

Patients

All 18 patients, examined and diagnosed by specialized neurologists, fulfilled the International Criteria for...
Headache Disorders 2nd edition (IHS 2004) for FHM or SHM diagnoses except patient #157, with probable HM but only one severe HM episode at the time of diagnosis. Clinical characteristics of the sample are presented in Table 1 and the pedigrees in which mutations were identified are shown in Figure 1. One hundred Caucasian Spanish or Greek unrelated adult control individuals with no personal or family history of recurrent or disabling headache were screened for the presence of the changes identified in the CACNA1A and ATP1A2 genes. The Spanish controls were blood donors or individuals that underwent surgery unrelated to migraine at Hospital Vall d’Hebron (Barcelona), whereas the Greek ones were healthy individuals collected as controls for a mutation screening of cystic fibrosis. This study was approved by the local Ethics Committee and informed consent was obtained from all adult subjects, children, and their parents according to the Helsinki declaration.

Sampling and mutation screening

Peripheral blood samples were collected from all probands. Genomic DNA was isolated using a standard salting-out method (Miller et al. 1988). The CACNA1A and ATP1A2 genes were sequenced as previously described, including all exons, splice sites, and branch points (Cuencía-Leon et al. 2008).

The promoter and 3’UTR regions of CACNA1A were also included in the mutational screening. We covered an 894-bp region upstream from the initiation codon and a 553-bp segment including exon 48, as previously described (Veneziano et al. 2009). The extension of the mutational screening was also applied to 18 HM patients from our previous study (Cuencía-Leon et al. 2008). All mutations were assessed by bidirectional sequencing. In addition, all mutations were confirmed by restriction fragment length polymorphism (RFLP) analysis of polymerase chain reaction (PCR) products (Table 2). Control individuals were screened by Sanger sequencing, single-strand conformation polymorphism or RFLP analysis to test the possible presence of the identified changes. Mutation nomenclature follows HGVS guidelines (http://www.hgvs.org/mutnomen/recs-DNA.html) and refers to the CACNA1A cDNA sequence NM_023035.1 (protein sequence NP_000702.3 (protein sequence NP_000693.1) with nucleotide 1A (ATG) corresponding to +1 and the ATP1A2 cDNA sequence NM_000702.3 (protein sequence NP_000693.1) with nucleotide 1A (ATG) corresponding to the initiation codon.

Copy number variant (CNV) analysis of CACNA1A was performed using two complementary approaches in order to cover most exons (Fig. S1): MLPA and quantitative multiplex PCR of short fluorescent fragments (QMPSF). For the MLPA assay, we used the SALSA MLPA kit P279-A2 for CACNA1A (MRC-Holland, Amsterdam, the Netherlands), and for QMPSF we used four sets of probes that covered 16 additional exons not included in the MLPA kit design. Further information about the methods and analysis is provided in Figure S1 and PCR conditions and sample analysis are available upon request.

Paternity was assessed in the de novo mutation identified as previously described (Carreño et al. 2011).

DNA constructs and site-directed mutagenesis

Human z1A (CaV2.1) was originally cloned into a pCMV vector and mutation p.Thr501Met was introduced by site-directed mutagenesis (GenScript Corporation, Piscataway, NJ). Rabbit z2A and rat CaV1β3, CaV2.1 channel regulatory subunits were subcloned into a pcDNA3 expression vector.

A full-length human ATP1A2 ouabain-resistant cDNA clone (z2_oua_wt, with p.Gln116Arg and p.Asn127Asp changes conferring ouabain resistance), housed in pcDNA3.1 with the myc tag, was used. Mutation c.2473G>A (z2_oua_825Lys) was introduced in a z2_oua_wt clone with the QuickChange II XI Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA).

All cDNA clones used in this study were sequenced in full to confirm their integrity.

Functional analysis of CACNA1A

Heterologous expression and electrophysiology

HEK 293 cells were transfected using a linear polyethyleneimine (PEI) derivative, the polycation ExGen500 (Fermentas Inc., Hanover, MD) (8 equivalents PEI/3.3 μg DNA/dish) as previously reported (Fernandez-Fernandez et al. 2004). Transfection was performed using the ratio for z1A (wild type or p.Thr501Met), CaV1β3, z2δ, and EGFP (transfection marker) of 1:1:1:0.3. Recordings were done 24–72 h after transfection.

Currents through wild-type (WT) or p.Thr501Met CaV2.1 (P/Q) channels were measured using the whole-cell configuration of the patch-clamp technique (Hamill et al. 1981). Pipettes had a resistance of 2–3 MΩ when filled with a solution containing (in mmol/L): 140 CsCl, 1 EGTA, 4 Na2ATP, 0.1 Na3GTP, and 10 Hepes (pH 7.2–7.3 and 295–300 mosmoles/L). The external solution contained (in mmol/L): 140 tetraethylammonium-Cl, 3 CsCl, 2.5 CaCl2, 1.2 MgCl2, 10 Hepes, and 10 glucose (pH 7.3–7.4 and 300–305 mosmoles/L). Recordings were obtained with a D-6100 Darmstadt amplifier (List Medical, Germany), filtered at 1 kHz and corrected for leak and capacitive currents using the leak subtraction procedure (P/8). Currents were acquired at 33 kHz. The
pClamp8 software (Axon Instruments, Foster City, CA) was used for pulse generation, data acquisition, and subsequent analysis.

Peak inward Ca$^{2+}$ currents were measured from cells clamped at $-80$ mV and pulsed for 20 msec from $-60$ mV to $+70$ mV in 5 mV steps. A modified Boltzmann equation (eq. 1) was fitted to normalized current voltage (I–V) to obtain the voltage dependence of activation. The voltage dependence of steady-state inactivation was estimated by measuring peak $I_{Ca}$ currents at $+20$ mV following 30-sec steps to various holding potentials (conditioning pulses) between $-80$ and $+5$ mV. During the time interval between

| Table 1. Clinical features of 18 patients with HM and other accompanying symptoms. |
|------------------|---------|---------|-------------|-------------|--------------------|---------|-------------|---------|
| Patient | Gender | Age (years) | Age at onset (years) | Triggering factors | Diagnosis | Other ictal features | Other | Family history | Origin |
|---------|---------|-------------|---------------------|-------------------|----------|-------------------|-------|----------------|--------|
| 5C | F | 50 | 11–12 | – | FHM | – | – | MA, HM | Spain |
| A03_44 | F | 66 | – | – | FHM | – | Nystagmus, progressive ataxia | – | Spain |
| A00_100 | F | 31 | Adolescent | – | FHM | Episodic ataxia | Progressive ataxia, cerebellar atrophy on MRI | – | Spain |
| 99 | F | 40 | 8 | Fasting | SHM | MA | – | MO | Spain |
| 112 | M | 19 | 11–12 | – | SHM | MA | – | Headache | Spain |
| 157 | M | 15 | 8 | Stress | HM (1 episode) | MA | – | Headache, vestibular migraine | Spain |
| 161 | M | 20 | 11 | – | SHM | MA | – | – | Spain |
| 3228 | F | 47 | <10 | – | FHM | MA | Interictal cerebellar signs | – | Spain |
| 388A | F | 15 | 11 | – | FHM | Generalized seizure, transient cerebral edema on MRI | – | – | Spain |
| 391A | F | 11 | 5 | Stress | SHM | – | – | MO | Spain |
| 431 | F | 42 | 38 | Stress | SHM | – | – | MO | Spain |
| 475 | F | 50 | 14 | – | FHM | – | – | – | Spain |
| 1713 | M | 37 | 12 | – | FHM | MA | – | – | Spain |
| G248 | F | 43 | 17 | Stress, menses | SHM | Aphasia | – | – | Spain |
| 8873 | F | 37 | 9 | – | FHM | Aphasia, transient episodes of bilateral visual loss | – | – | Spain |
| 1310 | F | 47 | 15 | Stress, strong odors | FHM | Partial epileptic seizures | – | HM | Spain |
| 387A | M | 10 | 2 | Head injury | FHM | Febrile seizures | Tension-type headache | HM | Greece |
| 489A | F | 8 | 5 | – | FHM | Prolonged dysphasia/confusion | Nystagmus, cerebellar atrophy on MRI | MA, panic attacks | Spain |

HM, hemiplegic migraine; FHM, familiar hemiplegic migraine; SHM, sporadic hemiplegic migraine; MA, migraine with aura; MO, migraine without aura; EA2, episodic ataxia type 2; MRI, magnetic resonance imaging.
test pulses (20 msec) cells were held at −80 mV. \( I_{Ca} \) currents obtained following the conditioning pulses were normalized to maximal \( I_{Ca} \) to determine the persistent current. Half-maximal voltage (\( V_{1/2, \text{inact}} \)) was obtained by fitting the data to the following Boltzmann equation (eq. 2). All experiments were carried out at room temperature (22–24°C).

\[
I = G_{\text{max}}(V - V_{\text{rev}})/(1 + \exp(-[V - V_{1/2,\text{act}}]/k_{\text{act}})) \tag{1}
\]

\[
I/I_{\text{max}} = 1/(1 + \exp([V - V_{1/2,\text{inact}}]/k_{\text{inact}})) \tag{2}
\]
Statistics

Data are presented as the means ± SEM. Student’s t test or Mann–Whitney test was used for statistical analysis, as appropriate. The t test assumes that data are sampled from population that follow a normal distribution and with equal standard deviations. Therefore, the use of the Mann–Whitney test is justified either if at least one of the two population data that are compared fails the normality test (using the Kolmogorov–Smirnov method) or if there are significant differences among the standard deviations of the two populations. Differences were considered significant if \( P < 0.05 \).

Functional analysis of ATP1A2

Cell viability assay

HeLa cells were transfected with 2 \( \mu \)g of constructs using Lipofectamine 2000 (Invitrogen, Carlsbad, CA). Forty-eight hours after transfection, 2/3 of the cells were harvested for Western blot analysis and 1/3 of cells were seeded in petri dishes. After 24 h, the \( \text{Na}^+\text{K}^+\)-ATPase inhibitor ouabain (1 \( \mu \)mol/L) was added to the medium (Dulbecco’s modified eagle medium containing 10% fetal calf serum and 1% Penicillin-Streptomycin). After 48 h of ouabain exposition, cell viability was quantified with the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) reduction assay (De Fusco et al. 2003; Vanmolkot et al. 2003).

Western blot analysis

A volume of 25 \( \mu \)g of total protein extract (per lane) was subjected to 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis. Gels were transblotted to a nitrocellulose membrane and were incubated overnight with monoclonal antibodies against c-Myc (4 \( \mu \)g/mL). Horseradish peroxidase-conjugated anti-mouse immunoglobulins were used as a secondary antibody. Proteins were visualized with the enhanced chemiluminescence kit (GE Healthcare, Little Chalfont, Buckinghamshire, U.K.). Blots were probed with antitubulin antibodies as a loading control.

Statistics

We performed four independent experiments, each with triplicate measurements, and carried out statistical analyses by the Student’s t test, considering one-tail distribution and two-sample equal variance (homoscedastic). Statistical significance was set at \( P < 0.05 \).

Results

Clinical data

Eighteen patients with HM and other accompanying clinical features were identified and screened for mutations in the CACNA1A and ATP1A2 genes. Eleven were classified...
as FHM and seven as SHM. The clinical characteristics of all patients are summarized in Table 1.

As previously described, we observed a preponderance of female patients in our HM population (13 female patients/five male patients). Age of onset was known for 17 cases. Seven patients presented onset within the first decade; the youngest age of presentation of hemiplegia was at age 2 in one FHM case (#387A). Onset in postpuberty or adolescence was documented in nine patients while in one case (#431) an unusual late onset in the fourth decade was reported. The episodes triggering factors did not differ from those commonly associated with common migraine, except for mild head trauma-induced attacks in case #387A, bearing an ATP1A2 mutation.

Other ictal manifestations were common in this cohort of HM patients: four cases developed migraine with typical visual or sensory aura and one migraine without aura (MO) episodes; three had episodes of aphasia or dysphasia, presumably the expression of migrainous aura, in the absence of concurrent hemiplegia. Two patients developed epileptic seizures, one partial and one generalized, and one patient had a history of febrile seizures; transient visual loss and episodic ataxia were described in one case each.

Four patients had interictal, permanent neurological abnormalities, mainly a cerebellar syndrome featuring nystagmus and different degrees of chronic ataxia. Three or these harbored mutations in one of the two analyzed genes. All patients were subjected to MRI studies. The only remarkable neuroradiologic findings were chronic cerebellar atrophy in three cases and cerebral edema at the time of the episodes in one case. The remaining patients did not have any chronic disability and had normal interictal exams (Table 1).

A more detailed clinical description of patient #387A and his father, bearing a previously undescribed mutation in the ATP1A2 gene, is provided as supplementary material (see Data S1).

Genetic analysis

The extensive sequencing of the CACNA1A gene in 18 subjects with HM allowed the identification of four previously reported changes in four unrelated families. In the remaining 14 patients, a novel and a previously described mutation was detected in the ATP1A2 gene (Table 2 and Fig. 2). The presence of all these variants was confirmed by restriction analysis of the corresponding PCR products (Table 2). The identified changes were not present in 100 Spanish nonmigraineurs. The novel p.Glu825Lys mutation in the ATP1A2 gene, identified in a Greek patient, was also absent from a set of 100 healthy Greek individuals. One of the changes, p.Ser218-Leu, was present in the affected sib, but not in her parents, indicating a de novo origin (Fig. 1A). False paternity was excluded by genotyping 16 polymorphic microsatellite markers in all the family members (data not shown).

All the identified mutations led to amino acid substitutions (p.Ser218Leu, p.Thr501Met, p.Arg583Gln, and p.Thr666Met in CACNA1A; p.Glu825Lys and p.Ala606Thr in ATP1A2). One of them, p.Glu825Lys in ATP1A2, is described here for the first time. The rest of the changes had previously been reported in other patients from several countries, four of them in HM and only one, p.Thr501Met in CACNA1A, in association with another phenotype, EA2.

The CACNA1A-p.Thr501Met mutation, described only once prior to this study, and the novel missense mutation p.Glu825Lys in ATP1A2, both of them subjected to functional studies in this study (see below), alter amino acid residues that are highly conserved in evolution as shown by a comparison of their paralogous and orthologous counterparts (Fig. 3). The p.Glu825Lys variant segregates with the HM phenotype, being transmitted from the affected father to the affected sib (Fig. 1B; for a detailed clinical description see Data S1). DNA was not available from the p.Thr501Met family, where a son and a daughter of the studied proband (#A03_44) have also been diagnosed with HM.

The 12 patients with no mutations identified after extensive CACNA1A and ATP1A2 sequencing, as well as 18 HM patients negative for a mutational screening performed in our previous study (Cuenca-Leon et al. 2008), were subjected to MLPA/QMPSF quantitative analysis to seek duplications/deletions in the CACNA1A gene. However, no structural variations were found in these probands.

Functional studies

We subjected two of the identified mutations, p.Thr501Met (CACNA1A) and p.Glu825Lys (ATP1A2), to functional analyses. These changes are the only ones that had not previously been studied by other authors at the functional level.

CACNA1A (p.Thr501Met): current density and activation/inactivation voltage dependence of heterologously expressed Ca\textsubscript{v}2.1 (P/Q) channels

Mutation p.Thr501Met changes a hydrophilic amino acid to a hydrophobic one and is located in a functionally important region of the human \(\alpha\textsubscript{1A}\) subunit of the neuronal \(\text{Ca}_{\text{v}}\text{2.1}\) (P/Q-type) \(\text{Ca}^{2+}\) channel (Fig. 2A). It lies in
Figure 2. Gene structure of CACNA1A and ATP1A2 and detection of mutations. (A) 1: CACNA1A gene structure, with black boxes indicating exons. The identified mutations causing HM are indicated by colored dots: p.Ser218Leu (light green), p.Thr501Met (red), p.Arg583Gln (blue), and p.Thr666Met (purple). 2: Protein structure and location of the identified mutations. (B) 1: ATP1A2 gene structure, with black boxes indicating exons. The newly identified mutation p.Glu825Lys causing HM is indicated by a yellow dot and p.Ala606Thr with a green dot. 2: Protein structure and location of the identified mutations. Detection of the mutations by direct sequencing of PCR products: electropherograms. Cyt, cytoplasm; M, cytoplasmic membrane; ES, extracellular space.
the S1 segment of domain II (II-S1) (Fig. 2A), which makes up part of the voltage sensor (Tombola et al. 2006). Because in a previous study of a patient with HM we found that another S1 mutation, p.Tyr1245Cys in domain III of the protein mainly affected voltage dependence of both activation and inactivation of the channel (Serra et al. 2009), we focused our functional analysis of the p.Thr501Met mutation on those same parameters.

Maximum current densities resulting from expression of mutant p.Thr501Met were significantly higher than current densities of wild-type (WT) channels \((P < 0.05)\) in a physiological range of depolarized voltages (from \(-15\) to +5 mV) (Fig. 4A and B, left panel). The potential for half-maximal activation \((V_{1/2, \text{act}})\) was also significantly \((P < 0.0001)\) shifted to hyperpolarized potentials for p.Thr501Met channels (by \(-7 \text{ mV}\)) (Fig. 4B, right panel). Consistently, the maximum current amplitude was elicited by depolarizing pulses to \(+15\) mV or \(+5\) to \(+10\) mV for WT or p.Thr501Met channels, respectively (Fig. 4A and B). The half-maximal voltage for steady-state inactivation \((V_{1/2, \text{inact}})\) induced by 30-sec-conditioning prepulses between \(-80\) and +5 mV was left-shifted (\(-12 \text{ mV}\)) in p.Thr501Met channels \((P < 0.0001)\). (Fig. 4C and D).

### ATP1A2 (p.Glu825Lys): Ouabain resistance survival assays

p.Glu825Lys exchanges a negatively charged residue for a positive one and is located in the intracellular L6/7 loop of the ATP1A2 protein, between the transmembrane segments M6 and M7 (Fig. 2B). Potential pathogenicity of the mutation p.Glu825Lys was indirectly tested through ouabain resistance survival assays.

HeLa cells transfected with the \(a2_{\text{oua}}_{825}\text{Lys}\) construction showed a 10% survival rate, compared with cells transfected with \(a2_{\text{oua}}_{\text{WT}}\) \((P < 0.0003)\) (Fig. 5A).

Western blot analysis of HeLa extracts obtained after transfection with the mutant and WT constructions showed a diminished amount of 825Lys protein compared with the WT, indicating that the altered protein may be unstable (Fig. 5B).

### Discussion

We have identified six potentially disease-causing missense mutations in the \(CACNA1A\) and \(ATP1A2\) genes in a cohort of 18 unrelated probands with HM from Spain and Greece (33% of patients covered).

There is evidence supporting the finding that the identified amino acid substitutions are indeed disease-causing mutations: (1) When family material was available, the mutations cosegregated with the disease phenotype within the families, (2) the changes were not present in at least 200 chromosomes from unaffected individuals nor in the main public single-nucleotide polymorphism databases (Hapmap, dbSNP), (3) no other molecular alterations were identified within the studied genes after the analysis of the whole coding region and the exon–intron boundaries, including splice sites, branch points, the promoter, and the 3′UTR region in \(CACNA1A\), (4) for the two mutations subjected to functional studies the involved amino acid residues are strongly conserved in evolution, both at the intraspecific (\(CACNA1A\), B, C, D, E, F, S or \(ATP1A1\), A2, A3, A4) and interspecific levels (\(CACNA1A\) and \(ATP1A2\) Genes in Hemiplegic Migraine O. Carreño et al.).
or ATP1A2 subunits from human, cattle, rat, mouse and zebrafish), indicating functional/structural relevance, (5) all these mutations but one (p.Glu825Lys) have been reported previously in other HM or EA2 cohorts, and (6) functional analyses reported here (p.Thr501Met in CACNA1A and p.Glu825Lys in ATP1A2) and by other authors (p.Ser218Leu [Tottene et al. 2005; Weiss et al. 2008; Debiais et al. 2009; Adams et al. 2010], p.Arg583Gln [Kraus et al. 2000] and p.Thr666Met [Kraus et al. 1998; Tottene et al. 2002] in CACNA1A; p.Ala606Thr [Jen et al. 2007; Tavraz et al. 2008] in ATP1A2) demonstrate functional consequences of all six changes suggesting pathogenicity.

Interestingly, all but one of the patients that have HM plus additional ictal/interictal neurological features or atypical paroxysmal signs (e.g., epilepsy or cerebral edema on MRI) bore mutations in one of the two genes studied. In general, this was not the case for patients without such “extended” HM phenotype, with no mutations identified in those genes. The molecular defects associated with this extended phenotype are unlikely to mimic those present in common migraine. It is, however, possible that HM cases without known mutation and with less severe clinical presentations, which often overlap with common migraine (in our series most had MO and MA attacks in addition to HM) may represent a better model for primary headaches. In this regard, deciphering the molecular basis of this group of patients may shed new light on the global landscape of migraine genetics.

**CACNA1A screening**

Four patients presented missense mutations in the CACNA1A gene, all of them described in previous screenings by other authors: p.Ser218Leu, p.Arg583Gln, p.Thr666Met, and p.Thr501Met (Fig. 2A).

**p.Ser218Leu**

This change was identified as a de novo mutation in a SHM patient with aphasia, transient cerebral edema and generalized seizures. This transition in a CpG dinucleotide had been described in eight HM patients (Kors et al. 2001; Curtian et al. 2006; Chan et al. 2008; Debiais et al. 2009; Stam et al. 2009; Zangaladze et al. 2010) having appeared de novo in two of them (Stam et al. 2009). Previous electrophysiological studies for the p.Ser218Leu mutation revealed a lower activation threshold, an increase of channel recovery after inactivation (Tottene et al. 2005) and the reduction of the inhibitory pathway carried by G-protein-coupled-receptor activation (Weiss et al. 2008). This dramatic gain of function, compared to
other missense changes, is thought to explain the severe clinical outcome observed in the patients carrying this mutation. Studies with a mouse model provide evidence that p.Ser218Leu mutations directly affect Ca²⁺-dependent facilitation and synaptic plasticity (Adams et al. 2010).

**p.Arg583Gln**

This prevalent mutation has been previously identified in FHM families and in sporadic patients (Battistini et al. 1999; Ducros et al. 2001; Terwindt et al. 2002; Wada et al. 2002; Kors et al. 2003; Thomsen et al. 2007; Freilinger et al. 2008; Yabe et al. 2008; Riant et al. 2010a). In family #A00_100, only the proband presents HM (Fig. 1A) and the mutation cosegregates with EA2, cerebellar atrophy and/or HM. Previous functional studies for the p.Arg583Gln mutation revealed a change in the voltage dependence of activation and inactivation toward more negative potentials due to the neutralization of residue 583 that is positively charged in the WT channel, and a decrease in the recovery of inactivation (Kraus et al. 2000).

**p.Thr666Met**

This is also a prevalent mutation that has previously been found in 24 FHM or SHM probands (Ophoff et al. 1996; Ducros et al. 1999; Friend et al. 1999; Terwindt et al. 2002; Wada et al. 2002; Kors et al. 2003; Thomsen et al. 2007; Freilinger et al. 2008; Yabe et al. 2008; Riant et al. 2010a) Functional studies in heterologous systems showed a gain of function through the following mechanisms: a decrease in the recovery after inactivation (Kraus et al. 1998) and increased Ca²⁺ influx through the channel in a broad voltage range around the threshold of activation, which was also reduced (Tottene et al. 2002).

**p.Thr501Met**

This change, found in patient #A03_044 with FHM, nystagmus, and episodic and progressive ataxia, has been described in another family with EA2 (Mantuano et al. 2010) and in a case with bouts of episodic ataxia and confusion but no hemiplegia, in association with vermian cerebellar atrophy (Cleves et al. 2009). This is the second reported mutation in the S1 segment in any domain of the α1A subunit in patients with HM. As the previously described p.Tyr1245Cys (Serra et al. 2009), p.Thr501Met alters channel activation and inactivation. It promotes channel activity by shifting the current activation curve to lower voltages (~9 mV) and increasing Ca²⁺ current density to a range of voltages that neurons can encounter during action potential firing. p.Thr501Met also shifts voltage-dependent steady-state inactivation to less depolarized voltages (~15 mV). These results further support an important role of the S1 segments in the function of voltage sensors leading to channel gating (Campos et al. 2007).

The functional consequences of mutation p.Thr501Met are consistent with a causative role in the disease. In this respect, a reduction in the voltage threshold of channel activation by ~10 mV is a trait shared by all FHM-causing mutations in CACNA1A (van den Maagdenberg et al. 2004; Tottene et al. 2005). Such gain of channel function specifically promotes cortical excitatory neurotransmission and favors cortical spreading depression (CSD) initiation and propagation in FHM knock-in (KI) mouse models (Tottene et al. 2009), which has been pointed out as the cause of the aura and migraine itself (Bolay et al. 2002; Pietrobon 2007).

Mutations in CACNA1A are also associated with other autosomal dominant neurological disorders characterized

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**Figure 5.** (A) Viability of HeLa cells transfected with the mutant ATP1A2 cDNA (E825K) normalized to the viability of cells transfected with the ouabain-resistant wild-type ATP1A2 cDNA (WT). NT, untransfected cells; pcDNA3.1, cells transfected with the empty vector. Four independent experiments were performed, each with triplicate measurements. The * symbol indicates the existence of significant differences between the p.Glu825Lys and the WT ATP1A2 constructs (P < 0.0003). (B) Western blot assay of the different protein extracts using anti-myc and anti-tubulin antibodies. The molecular weight of the Na⁺/K⁺-ATPase α2 subunit and tubulin are indicated. The constructs with the ATP1A2 ouabain-resistant cDNA carry the myc tag. The clone with the mutation displayed diminished band intensity.
by cerebellar dysfunction, such as EA2 (Ophoff et al. 1996). However, contrary to FHM mutations, most EA2 mutations produce loss-of-channel function (Strupp et al. 2007). Our patient carrying the p.Thr501Met CACNA1A mutation also developed cerebellar symptoms. This also occurred with other FHM-causing CACNA1A mutations (Ducros et al. 2001). As yet, it is not clear why some CACNA1A mutations cause pure FHM and other FHM with cerebellar signs, as functional studies in vitro do not reveal any notable difference among these two groups of FHM mutations (Pietrobon 2007). Nevertheless, the study of p.Arg192Gln and p.Ser218Leu KI animals may help us to unveil the role of FHM CACNA1A mutations in EA2 (van den Maagdenberg et al. 2004, 2010). While homozygous p.Arg192Gln (RQ/RQ) and heterozygous p.Ser218Leu (SL/WT) mice did not exhibit an overt phenotype, homozygous p.Ser218Leu (SL/SL) KI animals may help us to unveil the role of FHM CACNA1A mutations in EA2 (van den Maagdenberg et al. 2004, 2010). While homozygous p.Arg192Gln (RQ/RQ) and heterozygous p.Ser218-Leu (SL/WT) mice did not exhibit an overt phenotype, homozygous p.Ser218Leu (SL/SL) KI model exhibited the main features of the severe p.Ser218Leu (SL/WT) mice did not exhibit an overt phenotype, homozygous p.Ser218Leu (SL/SL) KI model exhibited the main features of the severe p.Ser218Leu clinical syndrome, including mild permanent cerebellar ataxia (van den Maagdenberg et al. 2004, 2010).

**ATP1A2 screening**

**p.Ala606Thr**

This mutation was found in a patient with FHM and focal epileptic seizures. It has previously been reported in three FHM families (Jen et al. 2007; Riant et al. 2005). Functional assays on HeLa cells transfected with WT and mutant ATP1A2 constructs suggested a loss of function of the sodium–potassium pump (Jen et al. 2007). Same results were found for mutations p.N717K and p.P786L, also located in the L4/5 cytoplasmatic loop of the protein (Jen et al. 2007; Tavraz et al. 2008). A more detailed electrophysiological functional study showed that the Na⁺/K⁺-ATPase activity was decreased due to a lower affinity for potassium (Tavraz et al. 2008).

**p.Glu825Lys**

This novel change, located in the L6/7 loop (Fig. 2B), was identified in an individual with HM and seizures. Ouabain-resistant survival assays showed a decrease in cell viability in HeLa cells transfected with the mutant ATP1A2 construct (Fig. 5A), again supporting the idea that the mutation results in loss of function of the sodium–potassium pump. Also, Western blot analysis suggests that the mutant protein may be unstable. Other mutations in this loop that have been identified by others and were functionally tested are as follows: p.Met829Arg, p.Arg384*, and p.Arg834Gln (de Vries et al. 2007; Tavraz et al. 2008). p.Arg834* showed no cell survival in ouabain challenge assays, and electrophysiological analyses of p.Arg834Gln displayed altered affinities for extracellular cations or reduced enzyme turnover. It was suggested that negatively charged residues in loop L6/7 contribute to the initial recognition of Na⁺ or K⁺ ions and constitute the cytoplasmic cation entry port (Shainskaya et al. 2000; Jorgensen et al. 2003). Other experiments found that loop L6/7 is important in the transmission of the activation signal initiated by cation binding to the phosphorylation domain of the protein (Xu et al. 2003). These experiments included a Glu to Ala mutation in a Ca²⁺-ATPase that is equivalent to the p.Glu825Lys identified in our patient.

**Unidentified mutations/genes**

Overall, 12 of 18 HM patients included in this study do not bear mutations in either CACNA1A or ATP1A2. Considering also the HM patients screened by us in a previous study (Cuenca-Leon et al. 2008), the level of molecular identification is 9/39 (23%), suggesting the involvement of other genes that still need to be uncovered. In sum, from these 39 patients, 22 were FHM (eight bearing mutations, 36%) and 17 were SHM (only one identified mutation, 6%). These data are in line with previous studies where the coverage was around 40% in FHM patients (Riant et al. 2005) and between 7% and 16% in SHM (Terwindt et al. 2002; de Vries et al. 2007), indicating that CACNA1A and ATP1A2 are major genes in the familial forms of HM but not in the sporadic ones, where there might be a larger genetic heterogeneity and/or other contributing factors. Only in one study with SHM patients presenting an early-onset of the disease the coverage was higher when analyzing the CACNA1A, ATP1A2, and SCN1A genes (Riant et al. 2010a). It is also possible that a few pathogenic mutations have remained unidentified in the two genes studied, including changes in introns or in distant regulatory regions, or CNVs in ATP1A2 that may be undetectable by PCR. Previously, a large rearrangement in the CACNA1A gene was described in a SHM patient (Labrum et al. 2009), and so it is conceivable that deletions or duplications in ATP1A2 could also be responsible for the disorder. However, CNV analysis has only reported this single deletion in CACNA1A so far in HM, and this type of alteration is more frequently found in episodic ataxia, where the typical pathogenic mechanism is a loss of function of the channel (Riant et al. 2008, 2010b; Labrum et al. 2009; Wan et al. 2011).

Mutations in SLC4A4 have been reported in pedigrees with proximal renal tubular acidosis (pRTA) with HM and migraine (Demirci et al. 2006; Suzuki et al. 2010), but this gene was not considered in this study as none of

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the patients in our series showed renal abnormalities. Mutations in the \textit{SCN1A} gene, previously linked to FHM and involved in several forms of epilepsy, would appear to be a rather unusual cause of HM. Indeed, more than 700 mutations have been identified in \textit{SCN1A} with SMEI compared to just five in FHM (http://www.molgen.vib-ua.be/SCN1AMutations) (Claes et al. 2009). Furthermore, very recently the genetic heterogeneity of HM phenotype has been broadened with the \textit{PRRT2} gene, previously related to paroxysmal kinesigenic dyskinesia and other episodic disorders. Eight mutations have been reported in 249 screened HM cases (Cloarec et al. 2012; Dale et al. 2012; Gardiner et al. 2012; Marini et al. 2012; Riant et al. 2012). Both \textit{SCN1A} and \textit{PRRT2} may be targeted by sequencing in the future, although they are not expected to explain a substantial proportion of our unresolved HM cases.

Finally, at least one other FHM locus has been mapped out of the three known loci, at 14q32, but the underlying gene still awaits identification (Cuenca-Leon et al. 2009). It is very likely that most of the unresolved 12 HM patients from the present cohort and 18 HM patients from a previous study by our group (Cuenca-Leon et al. 2008) bear mutations in other yet unknown HM genes that need to be uncovered. Thus, future studies will use whole-exome sequencing to find new genes responsible for the disease.

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\section*{Conflict of Interest}

None declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Figure S1. CNV study design using multiplex ligation-dependent probe amplification (MLPA) and quantitative multiplex PCR of short fluorescent fragments (QMPSF) approaches. Symbols (▲) indicate the exons covered by each assay. The MLPA kit allowed inspection of 24 exons. The analysis software Coffalyser v8 was used to evaluate the possible presence of CNVs, considering a deletion when the ratio was under 0.7 and a duplication when it was over 1.3. For QMPSF, we used four sets of primer pairs covering 16 exons. A reference fragment from the RNF20 gene was coamplified in each multiplex. Furthermore, two individuals with deletions corresponding to exons 20–47 and 32–47 of CACNA1A were also included in every assay as positive controls. Data were analyzed using the PeakScannerTM v1.0 software (Applied Biosystems), and the final ratios for each exon were expressed using the following formula: (height of the peak corresponding to the tested fragment for the patient/
height of the peak corresponding to RNF20 for the patient)/(height of the peak corresponding to the tested fragment for the average of controls/height of the peak corresponding to RNF20 for the average of controls). We considered a deletion when the ratio was under 0.6 and a duplication when it was over 1.4.

Data S1. Clinical features of patient #387A and his father.