**ABSTRACT**

**Objective:** To ascertain the genetic and functional basis of complex autosomal recessive cerebellar ataxia (ARCA) presented by 2 siblings of a consanguineous family characterized by motor neuropathy, cerebellar atrophy, spastic paraparesis, intellectual disability, and slow ocular saccades.

**Methods:** Combined whole-genome linkage analysis, whole-exome sequencing, and focused screening for identification of potential causative genes were performed. Assessment of the functional consequences of the mutation on protein function via subcellular fractionation, size-exclusion chromatography, and fluorescence microscopy were done. A zebrafish model, using Morpholinos, was generated to study the pathogenic effect of the mutation in vivo.

**Results:** We identified a biallelic 3-bp deletion [p.K19del] in CHP1 that cosegregates with the disease. Neither focused screening for CHP1 variants in 2 cohorts (ARCA: N = 319 and NeuroMics: N = 657) nor interrogating GeneMatcher yielded additional variants, thus revealing the scarcity of CHP1 mutations. We show that mutant CHP1 fails to integrate into functional protein complexes and is prone to aggregation, thereby leading to diminished levels of soluble CHP1 and reduced membrane targeting of NHE1, a major Na+/?H+ exchanger implicated in syndromic ataxia-deafness. Chp1 deficiency in zebrafish, resembling the affected individuals, led to movement defects, cerebellar hypoplasia, and motor axon abnormalities, which were ameliorated by coinjection with wild-type, but not mutant, human CHP1 messenger RNA.

**Conclusions:** Collectively, our results identified CHP1 as a novel ataxia-cause gene in humans, further expanding the spectrum of ARCA-associated loci, and corroborated the crucial role of NHE1 within the pathogenesis of these disorders. *Neurol Genet 2018;4:e209; doi: 10.1212/NXG.000000000000209*

**GLOSSARY**

**ARCA** – autosomal recessive cerebellar ataxia; **CaP-MN** – caudal primary motor neuron; **cDNA** – complementary DNA; **EGFR** – epidermal growth factor receptor; **GFP** – green fluorescent protein; **HMW** – high molecular weight; **HPRT** – hypoxanthine-guanine phosphoribosyltransferase; **HSP90** – heat shock protein 90; **KO** – knockout; **LKS** – Lichtenstein-Korsakoff syndrome; **LOD** – Logarithm of the odds; **mRNA** – messenger RNA; **MO** – Morpholino; **OE** – overexpression; **RT-PCR** – reverse transcription PCR; **SEC** – size-exclusion chromatography; **WB** – Western blot; **WES** – whole-exome sequencing; **WT** – wild type.

Autosomal recessive cerebellar ataxias (ARCAs) comprise a heterogeneous group of neurodegenerative disorders associating cerebellar degeneration to a variable combination of central or peripheral neurologic or nonneurologic signs. Collectively, ARCs have an estimated frequency of 1:20,000 and often show overlapping features with spastic paraplegias and peripheral neuropathies. Approximately 40% of individuals with suspected ARCA remain genetically charge funded by the authors. This is an open access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivatives License 4.0 (CC BY-NC-ND), which permits downloading and sharing the work provided it is properly cited. The work cannot be changed in any way or used commercially without permission from the journal.
undiagnosed, implying that a large number of causative genes—usually the most rare and challenging to identify—still need to be uncovered.

Here, we report the identification and characterization of a novel ARCA causative mutation in CHPI (calcineurin homologous protein-1, NM_007236.4:c.52_54del:p.Lys19del). CHPI serves as an essential cofactor of NHE1 (sodium/hydrogen exchanger 1), a major regulator of intracellular ions and pH homeostasis. Biallelic splice mutation of mouse Chp1 leads to reduced NHE1 membrane targeting and degeneration of Purkinje cells, ultimately causing ataxia. Similarly, mouse Nhe1 (encoded by Slec9a1) knockout (KO) causes Purkinje cell loss, severe locomotor ataxia, and epilepsy, while homozygous loss-of-function SLC9A1 mutations in humans cause cerebellar ataxia and sensorineural hearing loss. Therefore, we hypothesized that a mutation in CHPI, as a crucial regulator of NHE1, could impair expression and targeting of the exchanger, resembling the pathogenesis in mice and humans. Taking advantage of the zebrafish model system for modeling ARCA,– we generated a chp1-deficient zebrafish, which not only resembled the clinical features of the affected siblings but also corroborates the pathogenicity of the CHPI hypomorphic Lys19del mutation in vitro and in vivo.

METHODS Patient consents and standard protocol approvals. Informed consent was obtained from participants in accordance with French ethics regulations (Paris Necker ethics committee approval [RBM 01-29 and RBM 03-48] to A.B. and A.D.). Zebrafish procedures were approved by the local animal protection committee LANUV NRW; reference 84-02.04.2012.A251.

Whole-exome sequencing and whole-genome linkage analysis. Whole-exome sequencing (WES) of individuals 1, 2, 6, and 8 of family AAR-087 was performed on the Illumina HiSeq 2000 as paired-end 100bp reads after library preparation with the SureSelect Human All Exon 50 Mb V4 kit (Agilent Technologies, Santa Clara, California). ARCA and NeurOmics cohort WES and bioinformatic processing details are given in supplemental data. Whole-genome linkage analysis was performed using Illumina LINKAGE_12 microarrays (6090 single nucleotide polymorphism markers). Genotypes were determined by BeadStudio (Illumina, San Diego, CA) and analyzed with MERLIN 1.0.5

Expression vector generation and site-directed mutagenesis. Full-length human CHPI complementary DNA (cDNA) (NCBI: 11261) was cloned into 2 expression vectors: pcDNA3.1/CT-V5His-TOPO and pcDNA3.1/CT-green fluorescent protein (GFP)-TOPO following the manufacturer’s instructions (Life Technologies, Darmstadt, Germany). K19del mutation was introduced by site-directed mutagenesis (see primers, table e-1 at links.lww.com/NXG/A29), following QuickChange kit instructions (Agilent Technologies, Santa Clara, CA). Clones were verified by Sanger sequencing.

Cell culture and transfection. HEK293T (human embryonic kidney), HeLa (human cervical carcinoma), PC12 (murine pheochromocytoma), and N2A (murine neuroblastoma) cells were grown, maintained, and differentiated following standard protocols. Cells were transfected with plasmid DNAs, including control empty vectors, using Lipofectamine2000 (Life Technologies) and harvested after 24, 48, or 72 hours for RNA and protein isolation.

Semiquantitative reverse transcription PCR. RNA was extracted using the RNasy kit (Qiagen, Hilden, Germany), and concentrations were determined using the Ribogreen method (Life Technologies). Three hundred nanograms of RNA were reverse transcribed to cDNA with the QuantiTect Reverse Transcription Kit (Qiagen). PCR was performed with low cycle number (N = 24) and as multiplex with the housekeeping gene hypoxanthine-guanine phosphoribosyltransferase (HPRT) (see primers, table e-1, links.lww.com/NXG/A29). Densitometric analysis was performed with ImageLab 5.2.1 (Bio-RAD).

Cellular fractionation, size-exclusion chromatography, and Western blot. After 48 hours of transfection, HEK293T cells were processed with the Subcellular Protein Fractionation Kit for cultured cells (Thermo Scientific, Schwerte, Germany) following the manufacturer’s instructions. Size-exclusion chromatography (SEC) was performed with a Superose, 10/300-GL column connected to the ÄKTA-pure FPLC system (GE Healthcare, Freiburg, Germany). Sample preparation and column calibration details are given in supplemental data. Protein lysate preparation for Western blot (WB), sodium dodecyl sulfate-polyacrylamide gel electrophoresis, and transfer were performed according to the standard protocols. Antibodies used for immunologic protein detection are listed in table e-2, links.lww.com/NXG/A29. Densitometric analysis was performed with ImageLab 5.2.1 (Bio-RAD).

Fluorescence immunostaining. N2A, PC12, and HeLa cells transiently expressing pcDNA3.1-CHPI-GFP vectors were seeded in laminin-coated coverslips (Thermo Scientific, Schwerte, Germany) to preserve neurite-like structures, if necessary. After 24 hours of transfection and 4–5 days of differentiation, cells were immunostained as described in reference 16. Antibodies used are listed in table e-2, links.lww.com/NXG/A29.

Zebrafish experiments. Experiments for caudal primary motor neuron (CaP-MN) analysis were performed with the transgenic line tgf(ngn1-GFP)ml2TG (synonym TL/EK-h69-GFP). Cerebellar analyses were performed in the wild-type (WT) TL/EK line. Control and chp1 Morpholinos (MO) (against translational start codon, table e-1, links.lww.com/NXG/A29) were designed and purchased from GeneTools, LLC (Philomath, OR). WT and mutant human CHPI cDNAs were introduced into a pcDNA3.1 plasmid vector for in vitro transcription of capped messenger RNAs (mRNAs), performed as previously described. Zebrafish protein lysates for WB were prepared following the protocol described in reference 18 with modifications, explained in e-Methods. Embryo injections and CaP-MN immunostaining were performed as previously described. For cerebellar immunostainings, ~72-hpf zebrafish larvae were fixed overnight in Dent solution and processed as described in reference 12. Antibodies
Microscopy. Images were acquired using a Zeiss AxioImager M2 microscope equipped with ApoTome2 system to mimic confocality (Zeiss, Jena, Germany). Zebrarhachis CaP-MN and cerebellum images were acquired as Z-stacks and processed using ZEN software. Colocalization analyses were performed with ZEN. Representative images were processed using ImageJ. All experiments were blinded.

Protein modeling. Protein structures of free CHP1 (2CT9) and NHE1-bound CHP1 (2E30) were acquired from RCSB Protein Data Bank (http://www.rcsb.org/pdb/home/home.do). Cartoon representations were generated using PyMOL 1.5.0.4.

Statistics. Statistical analyses were performed using GraphPad Prism 6. Two-tailed Student t tests, χ², and Fisher exact statistics tests were performed. Levels of statistical significance are given in GraphPad format (*p ≤ 0.05, **p ≤ 0.01, ***p ≤ 0.001, and ****p ≤ 0.0001). Specific tests, sample size, data representation, and p values are indicated in figure legends.

RESULTS Identification of a 3-bp deletion in CHP1 causes autosomal recessive spastic ataxia. Two of 6 siblings of a consanguineous Moroccan family (figure 1A) developed autosomal recessive spastic ataxia with onset during the first decade of life. They show gait instability with moderate cerebellar atrophy (figure 1B) and upper and lower motor neuron involvement, intellectual disability, growth retardation, slow ocular saccades, and ovarian failure in the female proband. Positive pyramidal and cerebellar signs along with additional clinical features are summarized in table e-3, links.lww.com/NXG/A29 and extendedly described in supplemental data.

WES of individuals 1, 2, 6, and 8 yielded an average of 84,556,652 sequence reads per sample, of which >98.5% could be aligned to the reference sequence. Mean coverage was 77-fold, with 78.9% of the targeted sequence covering more than 20×. Twelve variants, all located within the linked regions (multipoint logarithm of the odds [LOD] score > -2), segregated recessively.
in the family (table e-4, links.lww.com/NXG/A29, and figure e-1, A and B, links.lww.com/NXG/A30). Further filtering based on alternate allele homozygosity and weak mutation effect prediction excluded 5 variants (table e-4, links.lww.com/NXG/A29). Of these, a biallelic 3-bp deletion in CHP1 (NM_007236.4:c.52-54del:p.Lys19del), hereafter defined as K19del, was validated by Sanger sequencing (figure 1C) and further selected as a top disease candidate since (1) it is located in the largest nonexcluded region on chromosome 15, where the maximum LOD score is reached, (2) it is absent from all public databases, (3) the mutation affects an amino acid highly conserved across species (figure 1D), (4) a point mutation reported in mouse Chp1, causing aberrant splicing and reduced full-length transcripts, leads to degeneration of Purkinje cells and ataxia,6 (5) CHP1 assists posttranscriptional glycosylation and membrane localization of NHE1; a major Na+/H+ exchanger listed in the ataxia-ome,19 (6) KO of mouse Nhe1 also cause ataxia,19 and (7) mutations in NHE1 cause ataxia-deafness Lichtenstein-Knorr syndrome (LKS).11 Neither focused screening for CHP1 variants in 2 large cohorts, namely ARCA (N = 319) and NeurOms (individuals with various neuromuscular diseases N = 657) nor GeneMatcher12 interrogation revealed any additional variant in CHP1, fulfilling our pathogenic selection criteria (table e-5, links.lww.com/NXG/A29).

Endocrine dysfunction is CHP1 independent. The syndromic presentation of ataxia with hypergonadotropic hypogonadism was first described in 190821,22; nonetheless, the genetic basis of this combination, except for few successful studies,19,23 remains elusive. The female proband of this study presented with ovarian failure; however, neither Chp1 vacillator nor Nhe1-depleted female mice were reported as infertile. Our analysis of uteri from vacillator mice showed neither uterus nor ovarian impairments (data not shown). We speculated that a CHP1-independent mutation might account for this defect, further clarifying the CHP1 family pathogenic landscape. WES revealed a homozygous variant (c.859C>T, p.G258E) in Basonuclin-1 (BNC1), which encodes a protein expressed only in reproductive germ cells with an essential function in oogenesis and spermatogenesis24,25 (table e-4, links.lww.com/NXG/A29). Bnc-1 null mice are sex-differentially affected, with females presenting complete infertility and subfertile males exhibiting smaller testes and reduced sperm motility.24,25 Furthermore, a pathogenic copy number variant including the BNC1 gene was recently detected in a retrospective study of spontaneous premature ovarian failure susceptibility.26

CHP1-K19del mutation alters protein solubility. To investigate the effect of CHP1-K19del, we examined the expression of CHP1-WT and CHP1-K19del-V5-tagged proteins in transiently transfected HEK293T cells by WB. Time-course analysis revealed reduced CHP1-K19del-V5 expression, whereas CHP1-WT-V5 exhibited higher levels over 72 hours of overexpression (OE) (figure 2A and figure e-2A). Semiquantitative reverse transcription PCR (RT-PCR) showed no differences between WT and mutant CHP1 transcripts (figure 2B and figure e-2B, links.lww.com/NXG/A30), thus excluding a transcriptional impairment.

Next, we assessed the distribution of CHP1-WT and mutant in soluble and insoluble fractions. Compared with CHP1-WT-V5 levels, CHP1-K19del-V5 was ~62% reduced in the soluble fraction and increased by ~68% in the insoluble fraction, respectively (figure 2C). Similar results were seen in N2A cells (figure e-2C, links.lww.com/NXG/A30). Together, these observations hint toward defects in protein localization, solubility, or stability of the CHP1 mutant form which, alone or in combination, might explain the reduced levels of soluble protein.

Mutant CHP1 is prone to aggregation. Given the clear effect of the K19del mutation on CHP1 expression, we investigated the cellular localization of WT and mutant CHP1-GFP–tagged proteins in N2A (figure 3A), PC12, and HeLa cells (figure e-3, A and B, links.lww.com/NXG/A30) via fluorescence microscopy.

CHP1-WT-GFP staining displayed a uniform distribution throughout the cells, including neurite-like structures in N2A and PC12 cells (figure 3A and figure e-3A, links.lww.com/NXG/A30), whereas CHP1-K19del-GFP expression led to massive protein aggregates, which on detailed observation revealed 2 distinct patterns: small puncta-like structures and large amorphous accumulations with a very strong fluorescent signal. These aggregates were observed in cell body and neurite-like projections (figure 3A). Quantification of total accumulation/aggregation events in N2A cells showed that ~50% of CHP1-K19del-GFP cells presented aggregates compared with ~20% in CHP1-WT-GFP cells. Large aggregations were detected almost exclusively in CHP1-K19del-GFP cells (~22% vs ~4% in the WT; p < 0.001) (figure 3A). Similar accumulation patterns were observed in PC12 and HeLa cells (figure e-3, A and B, links.lww.com/NXG/A30).

To further characterize these abnormal protein aggregates, CHP1-WT– and K19del-GFP–expressing cells were stained for Protein Homeostasis markers such as ubiquitin and P6227–29 (figure 3, B and C). Larger aggregates displayed positive colocalization and extensive accumulation of both markers, indicating that these structures are recognized by neuronal protein quality control systems, thus explaining CHP1-K19del-reduced expression. Altogether, these
results strongly indicate that K19del causes an increased proneness of CHP1 to aggregation, potentially attributable to misfolding and/or structural protein defects.

**The K19del mutation alters CHP1 association into functional complexes and impairs NHE1 membrane targeting.** To conclusively confirm that the K19del mutation leads to formation of higher molecular size CHP1 species, we performed SEC. Elution profiles of CHP1-WT-V5 and K19del-V5 proteins were determined by V5-immunoblotting. CHP1-WT-V5 eluted in 2 regions, representing the monomeric CHP1 form (~25 KDa)\(^{30,31}\) and the high-molecular-weight (HMW) complexes (~1 MDa). By contrast, the CHP1-K19del-V5 elution profile shifted not only away from the monomeric range toward higher fractions but also appeared in HMW fractions in which CHP1-WT is absent (figure 4A and e-4A).

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in CHP1-WT HMW fractions, while 3 bands were observed for CHP1-K19del. Immunoblotting of glycerinaldehyde 3-phosphate dehydrogenase, an interacting partner of CHP1,32 did not show differences between CHP1-WT and mutant elution profiles (figure e-4A, links.lww.com/NXG/A30). These results demonstrate that additional higher molecular complexes of CHP1-K19del-V5 are formed, which result from the unspecific aggregation proneness of mutant CHP1.

Characterization studies of the CHP1-NHE1 complex demonstrate that this calcium-dependent interaction exclusively involves 8 specific residues of the aJ helix of the C-terminal region of CHP1.7,30 These findings excluded the possibility of a direct implication of K19 in the interaction between both proteins. Examination of the crystal structure of CHP130 revealed that K19 is localized on the CHP1 first N-terminal aα helix (figure 4B). Of interest, comparison of this structure with an NMR-generated CHP1-NHE1 segment complex structure33 indicated that CHP1-aα undergoes large conformational changes on NHE1 binding, forming a clamp fixating NHE1-C-terminal segment (figure 4B). Although the structure-function relationship of this feature is poorly understood, this structural analysis suggests that K19del could interfere with a fully functional CHP1-NHE1 complex formation impairing NHE1 targeting to the membrane or compromising the function of this complex.

Since membrane localization of CHP1 is dependent of NHE1,6 but also membrane expression and maturation of NHE1 is reliant on CHP1,34 we investigated the effect of CHP1-K19del on NHE1 membrane targeting. Thereby, membrane fractions from HEK293T cells

**Figure 3** Protein aggregation in N2A cells expressing mutant CHP1

(A) Representative images of differentiated N2A cells transiently expressing CHP1-WT-GFP or CHP1-K19del-GFP proteins. CHP1-WT-GFP appears uniformly expressed both in the cell body and neurite-like structures of N2A cells. Small puncta-like aggregates are visible. CHP1-K19del-GFP expression leads to formation of large protein aggregations, both in the cell body and throughout axon-like projections. Puncta-like and large aggregations are indicated with white arrowheads. GFP exposure time in milliseconds is indicated inside the panels. Scale bar: 20 μm. The graph represents aggregation quantification in GFP-positive cells. Bars show the mean ± SEM from 6 independent experiments (n ≥ 250). *** denotes statistical significance (p < 0.001, χ² test). (B and C) CHP1-K19del aggregates colocalize with protein homeostasis markers. N2A cells expressing CHP1-WT-GFP or CHP1-K19del-GFP proteins were immunostained with ubiquitin and P62 antibodies. Representative images of puncta-like and large aggregations are depicted. Costes-adjusted Pearson colocalization coefficients were calculated for large aggregates, areas depicted in open circles: ubiquitin p = 0.9 and P62 p = 0.85. Positive coefficients indicate that aggregates colocalize with ubiquitin and P62. Scale bar: 20 μm. GFP = green fluorescent protein; WT = wild type.
expressing CHP1-WT– and K19del-V5–tagged proteins were analyzed by WB. In line with reduced cytoplasmic levels, CHP1-K19del membrane expression was also markedly diminished (figure 4C). It is important that marked NHE1 reduction in membrane fractions was observed on OE of mutant CHP1 (figure 4C), implying that CHP1-K19del interferes with proper NHE1 membrane targeting. Similar results were seen in N2A cells (figure e-4B, links.lww.com/NXG/A30).

**chp1** downregulation in zebrafish causes motor axon, cerebellar, and movement defects, which are rescued by wt, but not mutant CHP1 mRNA. To analyze the pathogenic consequences of the hypomorphic CHP1-K19del
Chp1 downregulation in zebrafish leads to CaP-MN, cerebellar, and movement defects rescued with WT, but not mutant CHP1 mRNA

(A) Lateral views of whole-mount embryos stained with synaptotagmin antibody (znp1). Truncated and absent motor axons are indicated with solid arrowheads and terminally branched axons with open arrowheads. Scale bar: 25 μm. (B and C) Based on overall appearance, CaP-MNs were classified as follows: normal, short (truncated axonal projection), or absent (total axonal atrophy). Based on terminal branching, axons were classified as normal, mild (branching ventral from midline), medium (2-3 or more branches at ventral or midline), or severe (>3 branches ventral or dorsal from midline). chp1-depleted zebrafish larvae display CaP-MN truncations and increased terminal branching in comparison with control (non-targeting) Morpholino (MO) and uninjected fish. Quantitative analysis of CaP-MN demonstrates that coinjection of chp1 MO and CHP1-WT mRNAs significantly improves axonal defects. Coinjection of chp1 MO and CHP1-K19del mRNAs fails to ameliorate CaP-MN defects in comparison with CHP1-WT mRNA. Results are presented in percentages from n = 150 analyzed axons representing 3 independent experiments. * denotes statistical significance in comparison with the chp1 MO group (p < 0.05 Fisher exact test).

(D) chp1-depleted; 34 hpf morphants show an increased frequency of spontaneous contractions. For graphic representation, spontaneous contractions were categorized as follows: 0 (no contraction), 1 (1 contraction in 30 seconds), and 2 or more (2 or more contractions in 30 seconds). Results are presented as the percentage of total from n = 50 observed zebrafish larvae representing 3 independent experiments. Coinjection of chp1 MO and CHP1-WT mRNAs, but not CHP1-K19-del mRNA, significantly reduced spontaneous contractions in 30 seconds **** denotes statistical significance (p < 0.0001 Fisher exact test).

(E) Dorsal views of whole-mount embryos labeled with acetylated tubulin at 72 hpf. The structure depicted between white horizontal lines in the uninjected morphant indicates the cerebellar area of observation. Optic tectum (OT) is outlined. Severe cerebellar hypoplasia is observed in chp1-depleted zebrafish in comparison with control MO and uninjected fish. * in chp1 MO image indicates trochlear decussation. Scale bar: 25 μm. (F) Quantitative analysis of cerebellar defects demonstrates that coinjection of chp1 MO and CHP1-WT mRNAs significantly ameliorates this phenotype. Coinjection of chp1 MO and CHP1-K19del mRNAs fails to improve cerebellar defects to WT levels. Results are presented in percentages from n = 40 observed zebrafish morphants from 3 independent experiments. * denotes statistical significance (p < 0.05 Fisher exact test). CaP-MN = caudal primary motor neuron; mRNA = messenger RNA; NS = not significant; WT = wild type.
mutation in vivo, we used MO to inhibit chp1 translation in zebrafish. Different MO concentrations were injected to select a dosage with little effect on overall morphology and development, but quantifiable neurologic readout (figure e-5, A–D, links.lww.com/NXG/A30). MO efficacy was confirmed by WB (figure e-5E, links.lww.com/NXG/A30).

Chp1 deficiency resulted in CaP-MN defects, namely axonal truncations and increased terminal branching. In detail, ~23% of the analyzed CaP-MNs exhibited defects in axonal projection and ~35% showed increased terminal branching (figure 5, A–C). Furthermore, Chp1 reduction led to severe cerebellar hypoplasia in ~70% of the morphants. Trochlear decussation became visible on severe cerebellar reduction, and reduced size of the optic tectum was also observed but not included for quantification analyses (figure 5, E and F). Negligible Ca-P-MN impairments and no cerebellar defects were observed in control groups (figure e-5, B–D, links.lww.com/NXG/A30). Since individuals carrying CHPI mutation exhibit spastic paraparesis, gait instability, and ataxia (table e-3, links.lww.com/NXG/A29), we additionally examined chp1 morphants movement in detail. Chp1 deficiency led to increased spontaneous contractions (figure 5D and video e-1, links.lww.com/NXG/A31) and to abnormal spastic-like trunk movement unaccompanied of locomotion (video e-2, links.lww.com/NXG/A32).

To corroborate chp1-MO phenotype specificity and confirm CHPI-K19del pathogenicity, rescue experiments were performed using WT and mutant CHPI-mRNAs. Protein OE was confirmed by WB (figure e-5F, links.lww.com/NXG/A30), and neither Ca-P-MN nor cerebellar defects were observed on CHPI-mRNAs OE (figure e-5, B–D, links.lww.com/NXG/A30). Coinjection of chp1 MO and CHPI-WT mRNA, but not CHPI-K19del mRNA, ameliorated all neurologic and movement defects associated with Chp1 deficiency (p ≤ 0.05). Thus, CHPI-WT mRNA improved axonal truncations, terminal branching, and cerebellar hypoplasia by ~13%–20% and 25%, respectively, compared with chp1 MO alone (figure 5, A–C, E and F). Spontaneous contractions were partially restored to uninjected larvae levels (figure 5D). Rescue experiments did not fully reach control levels probably because of the limited expression of synthetic mRNAs. Together, the neurologic and movement defects observed in chp1 morphants validate CHPI reduction as the underlying cause of MN defects and ratify cerebellar hypoplasia as the central pathogenic feature underpinning CHPI pathogenesis.

**DISCUSSION** Following a combination of exome and linkage analysis in a consanguineous family, we identified a 3-bp deletion in a highly conserved domain of CHPI as a novel cause of ARCA. Neither focused CHPI screening in 2 large cohorts with ARCA and NMD nor interrogating GeneMatcher unraveled a second mutation, emphasizing the scarcity of CHPI variants. This is further underpinned by the reduced tolerability for missense mutation found in control population (Exome Aggregation Consortium z-score: 2.64) and the high degree of conservation among CHPI orthologs (99% amino acid identity in mouse and 92% in zebrafish).

Here, we demonstrated that CHPI-K19del severely affects protein expression and localization, thereby impairing NHE1 membrane expression, and consequently its function. Therefore, CHPI mutation causes ataxia, most likely, in an NHE1-dependent manner (figure e-6, A and B, links.lww.com/NXG/A30), resembling the mechanism of cerebellar degeneration described for Chp1 vacillator and Nhe1 KO mice. Albeit aggregation is inherent to misfolded proteins in neurodegenerative disease, the aggregation proneness described for CHPI-K19del should be considered as a readout of abnormal protein folding rather than a disease mechanism itself.

The affected siblings of this study present with cerebellar atrophy and motor neuropathy, which were recapitulated in zebrafish. chp1 morphants exhibited severe cerebellar hypoplasia and CaP-MN alterations, thus demonstrating that MNs are also compromised on CHPI downregulation. Moreover, Chp1 reduction in zebrafish resulted in increased spontaneous contractions and spastic-like trunk movements, supporting that CHPI deficiency underlies the spastic phenotype in humans. Although the common denominator between Chp1 and Nhe1 mouse models is Purkinje cell degeneration, other central nervous areas such as the hippocampus and hypothalamus are also sensitive to NHE1 depletion. We speculate that abundant NHE1 expression in MN and MN precursors might regulate the proclivity of this population to CHPI imbalance. In contrast to the CHPI-deficient associated phenotype, NHE1 deficiency was presumptively associated with hearing loss, despite that neither Nhe1 nor Chp1 models presented deafness. More strikingly, depletion of mouse Nhe1 results in ataxia and epilepsy, while patients with LKS do not present with episodic seizures. This phenotypic variability strongly suggests that differential NHE1 dosages finely tune different pathogenic outcomes. Notably, CHPI-K19del reduces but not abolishes CHPI expression, thereby NHE1 membrane targeting is reduced, but not fully compromised. We hypothesize, in conformity with others, that reduced NHE1 certainly leads to ataxia, but remaining levels are sufficient to protect patients with CHPI mutations from epilepsy or deafness as in patients...
with LKS. Moreover, tissue-specific expression of other CHP or NHE proteins could modulate phenotypic compensations, further explaining the incomplete phenotypic translation from mouse to humans and the variability among human phenotypes.

AUTHOR CONTRIBUTIONS
Designing the study, drafting the manuscript, data interpretation, and study supervision: N.M.-F. and B.W. Functional experiments, data acquisition, processing, and analysis: N.M.-F. with help from E.J., S. H., H.L., S.S., J.M., M.R., C.P., L.T.-B., and M.H. Clinical evaluation and follow-up of patients: A.D. and A.B. Genetic and linkage analysis and data interpretation: M.C., M.K., G.S., A.S., and S.Z. Revising and approving the manuscript: all authors.

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DISCLOSURE
N. Mendoza-Ferreira co-holds patent 17172826.4-1401 (European patent office) for Calcineurin B homologous protein 1 inhibitors and therapeutic and non-therapeutic uses thereof. M. Coutelier has received research support from Fonds National de la Recherche Scientifique (F.R.S.-FNRS). E. Janzen and S. Hosseinibarkooie co-hold patent 17172826.4-1401 (European patent office) for Calcineurin B homologous protein 1 inhibitors and therapeutic and non-therapeutic uses thereof. H. Lühr, S. Schneider, J. Milbradt, and M. Kazakaya report no disclosures. M. Rieselband patented the use of specific antisense oligonucleotides to block NCALD expression to treat SMA. C. Pichilo and L. Torres-Benito report no disclosures. A. Singleton has received travel funding from 23andMe; serves on the editorial boards of Annals of Neurology, Lancet Neurology, Neurogenetics, Neurdegenerative Diseases, Brain, and the Journal of Parkinson’s Disease (no financial compensation); has a patent pending for panel of markers to diagnose stroke; and has received research support from NIH Intramural funding and Department of Defense. S. Zuchner reports no disclosures. A. Brice has served on the scientific advisory boards of FWO (Research Foundation—Flanders), ERC (European Research Council), and BMBF (Bundesministerium für Bildung und Forschung—Berlin, Germany); has served on the editorial boards of Neurology and Clinical Neuroscience, Parkinsonism & Related Disorders, Brain, Neurdegenerative Diseases, The Cerebellum, and Neurogenetics; and has received research support from French Research Agency (EU), the French program, FP (France Parkinson Association), RDS (Roger de Spoelberch Foundation), FDF (Fondation de France), and FRM (Fondation pour la Recherche Médicale). A. Durr has served on the scientific advisory boards of INSERM (National Institute for Health and Research), the Helmholz Initiative on Personalized Medicine—Germany, the Grenoble Neuroscience Institute of Grenoble, and the France Board of Directory of the ENP; has served on the editorial boards of Neurology® Genetics and Journal of Huntington’s Disease; has served on the International Advisory Committee of Archives of Neurology; co-holds a patent and receives royalty payments for Anaplerotic therapy of Huntington disease and other polyglutamine diseases (B1006353 Mochel/Durr [EP 0629187A5, April 12, 2006]); has been a consultant for Roche and Brainvictis; and has received research support from CoEn 2015: WIN–HD—Encoding Presymptomatic white matter changes in Huntington Disease 2015–2018. M. Hammerschmidt has served on the editorial boards of Development, Genes and Evolution and Molecular Genetics and Genomics; and has received research support from the National Institute of General Medicine Sciences and Deutsche Forschungsgemeinschaft. G. Stevanin has received travel funding from the Movement Disorder Society; holds patents for Diagnosis of hereditary spastic paraplegias (HSPI) by identification of a mutation in the KIAA1840 gene or protein; and has received research support from the European Community’s Seventh Framework Program, French Agency for Research, Erare Programme 2013–2018, Erare Programme 2016–2019, the program “Investissements d’Avenir,” Association Strümpell-Lorrain France, and Association Connaitre les Syndromes Cérébelleux. B. Wirth has served on the scientific advisory board of SMA Europe; has received travel funding and speaker honoraria from Biogen; co-holds patent 17172826.4-1401 (European patent office) for Calcineurin B homologous protein 1 inhibitors and therapeutic and non-therapeutic uses thereof; and has received research support from the European Community’s Seventh Framework Program. Go to Neurology.org/ng for full disclosure forms.

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