New spinocerebellar ataxia subtype caused by \textit{SAMD9L} mutation triggering mitochondrial dysregulation (SCA49)

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Spinocerebellar ataxias consist of a highly heterogeneous group of inherited movement disorders clinically characterized by progressive cerebellar ataxia variably associated with additional distinctive clinical signs. The genetic heterogeneity is evidenced by the myriad of associated genes and underlying genetic defects identified. In this study, we describe a new spinocerebellar ataxia subtype in nine members of a Spanish five-generation family from Menorca with affected individuals variably presenting with ataxia, nystagmus, dysarthria, polyneuropathy, pyramidal signs, cerebellar atrophy and distinctive cerebral demyelination. Affected individuals presented with horizontal and vertical gaze-evoked nystagmus and hyperreflexia as initial clinical signs, and a variable age of onset ranging from 12 to 60 years. Neurophysiological studies showed moderate axonal sensory polyneuropathy with altered sympathetic skin response predominantly in the lower limbs. We identified the c.1877C>T (p.Ser626Leu) pathogenic variant within the \textit{SAMD9L} gene as the disease causative genetic defect with a significant log-odds score ($Z_{\text{max}} = 3.43; \theta = 0.00; P = 3.53 \times 10^{-5}$). We demonstrate the mitochondrial location of human \textit{SAMD9L} protein, and its decreased levels in patients’ fibroblasts in addition to mitochondrial perturbations. Furthermore, mutant \textit{SAMD9L} in zebrafish impaired mobility and vestibular/sensory functions. This study describes a novel spinocerebellar ataxia subtype caused by \textit{SAMD9L} mutation, SCA49, which triggers mitochondrial alterations pointing to a role of \textit{SAMD9L} in neurological motor and sensory functions.
Introduction

The autosomal dominant spinocerebellar ataxias (SCAs) are a highly heterogeneous group of rare inherited movement disorders characterized by progressive cerebellar ataxia, often variably associated with pyramidal signs, parkinsonism or abnormal movements, lower motor neuron signs, peripheral neuropathy, ophthalmoplegia, pigmentary retinopathy, seizures or dementia. The disease onset presents typically in adulthood, albeit some clinical signs appear before ataxia is clinically apparent. Over 48 well-defined SCA subtypes have been described evidencing the high clinical and genetic heterogeneity. Common primary disease signs include cerebellar ataxia, dysarthria and oculomotor abnormalities and, when present, secondary signs including sensory deficits, parkinsonism, dysphagia, cognitive dysfunction or deafness which often are distinctive in each specific SCA subtype. Variable degeneration of the cerebellum, basal ganglia, cerebral cortex, optic nerve, pontomedullary systems, spinal tracts or peripheral nerves appears to underlie the heterogeneity of clinical signs and symptoms. Treatments are scarce and mostly symptomatic. At least 40 loci presenting with different pathogenic types of genetic mutations have been described associated with the rare Mendelian forms of SCAs, which can trigger cellular toxicity by gain-, loss-of-function effects or a combination of them, either at the protein or RNA levels in vulnerable neurons (reviewed in Klockgether et al.2). Identifying the genetic deficits has led to understand the
underlying cellular and molecular pathogenic pathways implicated, revealing recurrent alterations of transcription, protein aggregation and clearance, autophagy, ion channel physiology, calcium homeostasis or mitochondrial defects as the main pathological mechanisms underlying the different SCAs phenotypes.10–13 Dysregulation of mitochondrial function is a common pathogenic trigger of neurodegeneration, also identified in spinocerebellar and spastic ataxias, ataxia syndrome, multiple system atrophy and polyneuropathy, many of them presenting with variable degrees of ataxia, neuropathy and hypo- or hyperreflexia.14,15

Mutations within the SAMD9L gene or its parologue SAMD9, both genes located side-by-side on Chromosome 7q21, have been previously associated with a myriad of intra- and inter-familiar pleiotropic phenotypes.16 Germline frameshift SAMD9L mutations were first described in children presenting with inflammation of subcutaneous fat,17 and missense gain-of-function mutations in SAMD9L in ataxia-pancytopenia syndrome (ATXPC), characterized by a variable expression of a progressive neurological phenotype, pancytopenia and hypocellular bone marrow.18,19 On the other hand, heterozygous missense mutations within the parologue SAMD9 have been associated with MIRAGE syndrome, a severe early-onset condition characterized by myelodysplasia, infections, restricted growth, adrenal hypoplasia, genital phenotypes and enteropathy.20 The common unifying phenotype caused by SAMD9 or SAMD9L mutations is the early-onset myelodysplasia syndrome with partial or complete Monosomy 7, mainly described in paediatric cohorts without syndromic manifestation.21,22 While the penetrance of SAMD9L mutations in haematological malignancies is incomplete (estimated 70%),16 their pathogenic contribution to neurological phenotypes is much lower and unclear.21

In this study, we identify a SAMD9L mutation defining a novel SCA phenotype in kindred (M-SCA) from Menorca. Affected patients variably present with ataxia, nystagmus, dystarthritis, polyneuropathy, pyramidal signs and cerebral demyelination. In patients’ fibroblast and a zebrafish model, we demonstrate the mitochondrial perturbations underlying the molecular pathology in this new SCA pointing to the role of SAMD9L in neurological motor and sensory functions.

**Patients and methods**

This study was conducted according to the ethical principles for medical research involving human subjects according to the Declaration of Helsinki. Informed consents were obtained for all individuals participating in the study, which were approved by the ethical committee of the University Hospital Germans Trias i Pujol in Badalona, Spain.

**Clinical studies**

Clinical information from medical histories was obtained for 37 members from the M-SCA family including 11 affected individuals, while the clinical examination was performed in nine of them with a detailed neurological exam. The scale for the assessment and rating of ataxia (SARA) scores23 were assessed for six patients (III:7, IV:2, IV:6, IV:8, IV:14 and V:1). Affected patients were further investigated using an enhanced clinical protocol, including electrocardiograms, echocardiograms, audiometric tests, nerve conduction studies, MRI of the brain and evoked potentials, during follow-up visits.

**Electrophysiological studies**

Six patients from the M-SCA family (III:7, IV:2, IV:6, IV:8, IV:14 and V:1) underwent nerve conduction, EMG and somatosensory evoked potentials (SEPs) studies. In three patients, we could also assess autonomic nervous system function. All neurophysiological studies were performed by means of TruTrace® EMG system (DEYMED Diagnostic) and conducted according to standard methodology.24 A detailed protocol is included in the Supplementary material. Reference control values were obtained from Kimura as described.25

**Cerebellar volume quantification**

To quantify cerebellar volume, we used CERES algorithm26 to calculate the percentage of the relative volume to the total intracranial volume on T1-weighted 1.5 T MRI images from five patients (Patients IV:2, IV:6, IV:8, IV:14 and V:1). Normal boundaries were used from 30 age- and gender-matched controls (range: 24–75 years) randomly selected from the open-access IXI dataset as reported.27

**Genetic studies**

Genomic DNA samples were isolated from blood leucocytes using automated DNA purification (Chemagen, PerkinElmer). DNA samples were obtained from seven affected and six unaffected relatives. Additionally, 80 genomic DNA samples from the Menorcan general population were analysed to study the SAMD9L variant frequency. Genomic mitochondrial DNAs (mtDNAs) were sequenced using the Illumina MiSeq platform (Illumina, Inc.), and mtDNA copy number was determined by qPCR. Mitochondria DNA lesions were quantified using an adapted long-run qPCR method for DNA-damage quantification.28 Detailed protocols are described in Supplementary material (Materials and Methods section).

**Genome-wide linkage analysis**

Eleven family members, five healthy and six affected, were included in a genome-wide linkage study performed in collaboration with the Centro Nacional de Genotipado using the Illumina Infinium HumanOmni5 Chip (Illumina, Inc.). Genotypes were assigned using the BeadStudio genotyping module software (Illumina, Inc.), and initial non-parametric and parametric linkage were analysed using Merlin 1.1.2.29 Two-point and multipoint genetic linkage analyses were performed with MLINK and LINKMAP (version 5.10), from the Quiklink compilation (version 16) of LINKAGE.30,31
We subsequently used two additional samples (V:2 and V:3) and informative single nucleotide polymorphisms (SNPs) within the candidate region and their allele frequencies as the genetic markers. SLINK simulation program was used to calculate the P-value associated with the multipoint logarithm of the odds ratio (LOD) score that was obtained in 1000 replicates of the pedigree. Both complete-penetrance and age-dependent penetrance models were considered (Supplementary material, Materials and Methods section).

Whole-exome sequencing

Whole-exome sequencing (WES) was performed with genomic DNA obtained from peripheral blood of Patients IV:2 and IV:14 at the National Centre for Genomic Analysis (CNAG) in Barcelona, Spain. Exome capture was performed using Nimblegen SeqCapEZ Exome V.3 (Roche) for 64 Mb according to the manufacturer’s protocol. Sequencing and bioinformatics analysis protocols are included in the Supplementary material.

Sodium dodecyl sulphate–polyacrylamide gel electrophoresis and immunoblotting

Proteins were extracted from fibroblast samples from Patients IV:14 and V:1 and two age-matched controls by homogenization following standard and cell fraction protocols summarized in the Supplementary material. Primary antibodies used were anti-β-actin (AC15; Sigma–Aldrich), anti-ATP5H (SAB4500107; Sigma–Aldrich), anti-DRP1 (sc-271583; Santa Cruz), anti-GAPDH (G9545; Sigma–Aldrich), anti-LAMP1 (H4A3; DSHB), anti-LC3 (NB100-2220; Novus Biological), anti-MFN1 (sc-166644; Santa Cruz), anti-SAMD9L (25173-1-AP; Proteintech), anti-SQSTM1 (2C11; Thermo Fisher Scientific) and anti-VDAC (D73D12; Cell Signaling). Infrared-dye conjugated secondary antibodies (2C11; Thermo Fisher Scienti) and anti-VDAC (D73D12; Cell Signaling). Infrared-dye conjugated secondary antibodies anti-mouse IRDye-800CW and anti-Rabbit IRDye 700CW (Li-Cor) were used. Signals were detected and analysed with Odyssey analyser software (Li-Cor).

Human SAMD9L expression

Total RNAs were obtained from fibroblast samples from Patients IV:14 and V:1 and two age-matched controls following standard protocols included in the Supplementary material. ENCODE SAMD9L RNA sequence reads were compared on BAM files from adult cerebellar Purkinje cells, adult granular and pyramidal cells, 6-year-old child cerebellum and cerebellar and spinal cord human embryos.

Computational protein structure prediction and protein–protein interaction analysis

Sequence alignments against NCBI Conserved Domains (CD)_v3.16, PDB_mmCIF70_4_Feb, Pfam-A_v32.0 and SMART_V6.0 domain and structural databases were performed using HHpred. Protein structure was generated with PROSITE (https://prosite.expasy.org/mydomains/). Microtubule-binding motifs were predicted using MAPanalyzer and PredictProtein, DisEMBL, InterPro and MobiDB-lite, were used for intrinsically disordered regions (IDRs) prediction. NetPhos 3.1 and NetworKIN servers were used to predict serine, threonine or tyrosine phosphorylation sites. hSAMD9L (NP_001290425.1) reference protein was used. Protein–protein functional network interaction analysis was performed using the STRING database.

Transmission electron microscopy

Fibroblasts from patient IV:14 and one age- and gender-matched control were grown and fixed in 2.5% glutaraldehyde, 2% paraformaldehyde in 0.1 M phosphate buffer (pH 7.4) for 2 h at 4°C and processed as described. Ultrathin sections were stained with uranyl acetate and lead citrate and examined in an EX-electron microscope (JEM1200; JEOL) at 80 kV. Pictures were acquired using from 12 to 100 K magnifications. Subcellular organelles from 20 fibroblasts sections were classified according to previous publications identifying autophagosomes surrounded by double membrane and autolysosomes filled with undigested lipids.

Zebrafish ataxia model

All applicable international, national and/or institutional guidelines for the care and use of animals were followed. Animal procedures were approved by the Ethics Committee on Animal Experimentation of the Germans Trias i Pujol Research Institute. Zebrafish studies were conducted in collaboration with Zeclinics (Supplementary material). Locomotion and response to visual and physical stimuli were traced and analysed by the EthoVision XT 12 software (Noldus) and the DanioVision device in 32 5-days post-fertilization (dpf) zebrafish larvae from non-injected, control injected with lynGFP, wild-type and mutant SAMD9L groups distributed in six-well plates. To detect changes in the larvae locomotion deviations from the stereotyped behaviour characterized by motility in the dark phase and reduced motility in the light phase, larvae underwent 50 min dark/light alternating environments of 5 min each. This experiment was replicated with 10 min of alternating dark/light cycles. Data points for distances travelled (mm) during each minute and the number of head turns was measured. Data outside the 1.5 × interquartile range were considered outliers.

Statistical analysis

Statistical analyses were performed using SPSS 21.0 (IBM Corp.). Statistical significance was defined as P < 0.05. The Spearman’s rank correlation coefficient for non-linear relationship was used to assess correlation between SARA scores and patients’ age. Statistical differences in fibroblast samples for ATP determination and transmission electron microscopy (TEM) studies were analysed using
the Shapiro–Wilk test to assess for normal distribution followed by the parametric t-test, one-way ANOVA or alternatively with the non-parametric Mann–Whitney U-test. For zebrafish studies, behavioural time series statistical data were analysed using two-ways repeated mixed-effects ANOVA followed by post hoc comparisons with the Tukey HSD for data groups with equal variances. Total distance travelled and number of turns were further analysed using one-way mixed-effects ANOVA followed by post hoc comparisons with the Tukey HSD for data groups with equal variances. Those values that did not meet normality or equal variances criteria were individually tested with non-parametric test Mann–Whitney U-test. SEM denotes for standard error of the mean.

Data availability

The data supporting the findings of this study are available on request from the corresponding author. The data are not publicly available due to the fact that the genetic information may compromise the privacy of research participants.

Results

Clinical description

Eleven affected patients from the M-SCA family originally from Menorca, Spain, were clinically diagnosed by a neurologist (P.C., N.G.R., L.I., R.A.-R.) and a detailed clinical examination was performed in nine of them (Fig. 1A). Horizontal and vertical gaze-evoked nystagmus (GEN) was present as the first clinical sign in four patients (IV:8, IV:14, V:1, V:3; Table 1) in the absence of ataxia. Hyperreflexia was also present at the first initial clinical examination. Ataxia onset ranged from 30 to 60 years old (Table 1). Initial nystagmus progressed with diplopia or oscillopsia in four patients (III:7, IV:6, IV:8 and IV:14; Table 1). Patient IV:14 also referred recurrent vertigo. Younger patients (V:1 and V:3) presented with horizontal and vertical GEN and hyperreflexia without ataxic signs. Patient V:1 was neurologically evaluated for the first time at the age of 12 years and disclosed horizontal and vertical GEN, hyperreflexia with bilateral Babinski’s sign, clonus and pes cavus. Due to family history of falls and unsteady gait, clinical examination was complemented with CT which showed cerebellar atrophy (Fig. 1B), mega-cisterna magna with discrete ventricular dilatation, suggestive of initial signs of brain demyelination (Fig. 1C). His father (IV:2) was presented with gait instability since the age of 50. Clinical examination after 15 years of disease progression revealed GEN, dysmetria, dysdiadochokinesia, severe dysarthria and lower limbs hyperreflexia with bilateral extensor plantar response. MRI at 65 years of age showed global and symmetric dilatation of the ventricular system, lateral sulcus and cortical grooves with an accused diffuse vermal and hemispheric cerebellar atrophy (Fig. 1D). Cerebral parenchyma showed diffuse demyelination involving periventricular and centrum semiovale white matter suggestive of hypoxic leucoencephalopathy (Fig. 1E).

MRI of the brain showed cerebellar atrophy in three additional affected patients (IV:6, IV:8 and IV:14) with brain demyelination. Volumetric assessment in the five patients revealed a 28.96% significant decrease in cerebellar volume relative to the total intracranial volume ($6.55 \pm 0.55\%$) compared with 30 age- and gender-matched controls (Supplementary Fig. 1) with cerebellar relative volume ($9.22 \pm 0.22\%; F(1,8) = 99.506; P < 0.0001; Table 2; Supplementary Fig. 2), accompanied with a 36.86% decrease in cerebellar grey matter relative to the total brain grey matter (affected patients $= 4.32\% \pm 0.47\%$; controls $= 6.84\% \pm 0.17\%; F(1,8) = 122.136; P < 0.0001$; Table 2, Supplementary Fig. 3). Furthermore, all affected patients presented different grades of diffuse demyelination of the cerebrum white matter. Five patients (III:7, IV:2, IV:6, IV:8 and IV:14) presented with moderate axonal sensory polyneuropathy predominantly in the lower limbs (Fig. 2A and B) compared with control values (Supplementary Fig. 4), albeit at later stages of disease progression patients also revealed reduction of sensory nerve amplitudes in upper limbs (Supplementary Fig. 4). EMG studies were normal in all seven patients evaluated. Tibial nerve SSEPs showed an abnormal latency of P37 response in patients diagnosed with sensory axonal polyneuropathy (Fig. 2C and D). In two patients (IV:2 and IV:6), sympathetic skin responses (SSRs) were abnormal in lower limbs (Fig. 2E), and one patient (IV:2) also showed an abnormal R–R interval (Fig. 2G).

Interestingly, younger patient (V:1) presenting with horizontal and vertical GEN, hyperreflexia and cerebellar atrophy without apparent ataxia, recorded normal electrophysiological values at 36 years of age. The progression of the disease is slow, although variable, and the SARA scores$^{23}$ in this family follow an exponential trend ($y = 0.92, P = 0.008$; Fig. 1F). The disease progressed with dysmetria and dysarthria, and severity varied among patients. Relevantly, Patients III:2 and III:7 became wheelchair-bound after 15 and 31 years from ataxia onset, respectively. None of these patients showed fasciculations, epileptic seizures, cognitive impairment or haematological alterations (Supplementary Fig. 5A). Partial or total haploinsufficiency on Chromosome 7 in their haematological lineages was discarded by NGS or Sanger sequencing analysis from blood white cells DNA in their haematological lineages (Supplementary Fig. 5B). In conclusion, for this specific SCA phenotype, we found GEN and hyperreflexia as characteristic initial signs before ataxia onset. All five patients with neuromaging data revealed diffuse cerebral demyelination (Fig. 1C and E; data not shown). No extrapyramidal signs were identified in any affected patient.

Genomic studies

The genome-wide linkage analysis from six affected patients and five healthy family members from the M-SCA family
Figure 1 Pedigree, MRI and SARA scale from M-SCA patients. (A) Pedigree of the five-generation family from the Balearic island of Menorca with 11 affected individuals. Black asterisks denote individuals included in genome-wide linkage studies; red asterisks denote individuals studied by WES. (B–E) Sagittal and transverse T1-weighted MRI scans of Patient V:1 brain revealing cerebellar atrophy (B), and initial signs of focal brain demyelination lesions noted by red arrows (C). Imaging of the brain of the father of patient V:1 (IV:2) also showed severe cerebellar vermis atrophy (D), and marked cerebral demyelination noted by red arrows (E) compared with age- and gender-matched controls (Supplementary Fig. 3). (F) SARA clinical scale of six affected patients showing variable severity and progression of the disease following an exponential pattern \( r_{(4)} = 0.92, P = 0.008 \).
using the Illumina Infinium HumanOmni5 Chip including over 4,000,000 markers, revealed significant genetic linkage to 7q21 with the highest LOD score ($Z_{\text{max}} = 3.01$; $P < 0.0001$) defining a 19.1 Mb candidate region flanked by the polymorphic markers rs12705836 and rs16869440 (Fig. 3A). No other significant genetic linkage regions other than SAMD9L on 7q21 were identified. WES of Patients IV:2 and IV:14 identified 445 shared heterozygous candidate variants compatible with an autosomal dominant inheritance. After filtering variants by their position within the gene, population frequency, and their predicted pathogenicity, a missense c.1877C $>$ T (p.Ser626Leu) variant within the fifth exon of the SAMD9L gene located on 7q21, was selected as the candidate pathogenic variant. This variant segregated with the disease and was absent in all six healthy individuals studied from the M-SCA family, and was not identified in 80 healthy unrelated individuals from Menorca’s general population. Further linkage analysis with two additional family members, and considering a complete-penetrance model, revealed a significant two-point LOD score between the locus trait and the identified variant proposed as the causative disease mutation ($Z_{\text{max}} = 3.43$; $\theta = 0.00$; $P < 3.53 \times 10^{-5}$; Table 3). The multipoint LOD score analysis assuming a complete-penetrance model with three SNPs (rs17393952, rs4455763 and rs10263800) in the 7q21 candidate region and the candidate variant, resulted in a maximum significant multipoint LOD score $Z_{\text{max}}$ of 3.46 ($P < 3.28 \times 10^{-5}$; Supplementary material, Results section). Furthermore, linkage analysis using an age-dependent penetrance model with five multiple liability classes also resulted in significant two-point ($Z_{\text{max}} = 3.10$; $\theta = 0.00$; $P < 7.89 \times 10^{-5}$) and multipoint ($Z_{\text{max}} = 3.12$; $\theta = 0.00$; $P < 7.51 \times 10^{-5}$) LOD scores using the same markers. The c.1877C $>$ T (p.Ser626Leu) (Fig. 3B) variant within the SAMD9L gene involves a serine to leucine amino acid change, and was predicted to be deleterious by SIFT, FATHMM, Mutation Assessor, Mutation Taster and Provean algorithms. The variant is highly conserved through evolution (Fig. 3C), it is classified in the ClinVar database as a variant of uncertain significance (rs1554341671; RCV000498009.2), without minor allele frequency, and identified in two related paediatric cases presenting with hypopcellular marrow, dyserythropoiesis and Monosomy 7 in the absence of ataxia or cerebellar syndrome.21

**SAMD9L protein structure, domain and protein–protein interaction**

To investigate hSAMD9L protein structure and domains, we analysed the hSAMD9L protein sequence with HHpred protein sequence profile predictor and identified protein regions significantly similar to SAM (Ser6-85His), AAA+ ATP-ase/Hydrolase (711Lys–910Lys) and Apaf-1 ADP bound (716Glu–1018Glu) domains (Fig. 3D; Supplementary Table 4) in consonance with previously reported domain structures in SAMD9 family members.47 Furthermore, the analysis revealed the significant similarity with SPAST,
SKD1/VPS4B, katanin and torsin-1A AAA-ATPase specific protein domains, implicated in microtubule and lysosomal trafficking (Supplementary Table 4). Microtubule-associated protein (MAP) analyser classified hSAMD9L protein as a MAP under a 90% of specificity threshold and identified a significantly enriched motif in MAPs proteins, S....LKS between amino acids 619–626 (STILKLKS) where the identified mutation c.1877C>T entails the p.Ser626Leu amino acid change (Fig. 3E; Supplementary Table 5).

Comprehensive protein analysis with structural predictors identified an intrinsic disorder (ID) region around the SAMD9L amino acid residue Ser626 (Supplementary Fig. 6). The SAMD9L amino acid change p.Ser626Leu alters the ID region predicted by PredictProtein (PROFbval and Ucon) and DisEMBL (hot loops) algorithms, and it abolishes the identified ID region according to the prediction by InterPro (MobiDB-lite) (Supplementary Table 6). Furthermore, PredictProtein (PROFAcc) identified the 624–629 protein region of SAMD9L as a solvent-exposed region (Supplementary Fig. 6) and NetPhos 3.1 and NetworKIN algorithms predicted the residue Ser626 within in SAMD9L to be potentially phosphorylated by protein kinase C (Supplementary Fig. 7), a kinase directly implicated in cerebellar neurodegeneration in the SCA 14 subtype.48

With the STRING algorithm, we studied the interactions among SAMD9L and KATNA1, SPAST, VPS4B proteins identified by the HHpred disorder ID region profile predictor algorithm (Supplementary Table 4), and among the previously reported Samd9l-interacting proteins in mice Eea1 and Rab549 in the context of the whole human genome (Supplementary Fig. 8). The SAMD9L and SPAST network interaction was identified via IFI44I, also known as MAP 44, and ATL1 proteins. The protein–protein interaction enrichment analysis showed significantly more interactions among these proteins than expected (P < 2.52 × 10−7). Co-expression and interaction analyses linked both IFI44I and EEA1 with SAMD9L and revealed an association of the SAMD9L protein network with MAPs and endosomal/lysosomal pathways. Interestingly, mutations in the ATL1, SPAST, Rab7A or KATNB1 genes have been previously associated with diseases presenting with hyperreflexia, sensory axonal neuropathy or sensory impairment among other signs, which overlap with clinical signs identified in this new ataxia subtype in our family (Supplementary Table 7).

**Expression and mitochondrial localization of human SAMD9L**

Immunostaining with anti-SAMD9L antibody in control cerebellar sections revealed a punctate staining mainly in Purkinje cell soma in the human cerebellar cortex and multipolar neurons of the cerebellar dentate nucleus (Fig. 4A), similar to mitochondrial ATP5B protein staining (Supplementary Fig. 9), indicative of mitochondrial localization. Antibody specificity control staining is included in Supplementary Fig. 9. Some basket cells also appeared slightly stained. SAMD9L staining was also examined in fibroblast cells from age-matched control (Fig. 4B) and one affected patient (Fig. 4C; Supplementary Fig. 10), revealing mitochondrial staining and co-localization to the mitochondrial marker mitoTracker Red CMXRs. No significant differences in the mitochondrial network were observed in patients’ fibroblasts compared with controls (Supplementary material, Results section). ENCODE RNAseq analysis confirmed relative expression of SAMD9L in adult Purkinje cells and almost no expression in cerebellar granule and pyramidal cells (Supplementary Fig. 11). RNA tissue expression also showed higher levels of SAMD9L in child cerebellum and in embryonic cerebellum and spinal cord, in consonance with stages of high mitochondrial activity and tissues affected in the pathogenesis of this new ataxia subtype.

Immunoblotting of lysed and cellular fractioned fibroblasts confirmed SAMD9L mitochondrial localization and showed decreased SAMD9L protein levels (Fig. 5A; Supplementary Figs 18 and 19). To corroborate these findings, total cellular extraction from lysed fibroblasts was immunoblotted and reduced levels of SAMD9L in fibroblasts from affected patients were found (Fig. 5B; Supplementary Fig. 20) without alteration of SAMD9L cDNA levels (Fig. 5C; Supplementary Fig. 21).

**Mitochondrial perturbations in patients’ fibroblasts**

Potential alterations of fibroblasts’ mtDNAs were analysed for mtDNA depletions and found an increased copy number of the D-LOOP mitochondrial genomic region compared with the MT-CO3 region, suggesting increased activity of

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**Table 2 MRI cerebellar volume and grey matter assessment**

| Patient ID | Age at MRI | Relative cerebellar volume (%) | Age- and gender-matched controls relative cerebellar volumes range (%) | Relative grey matter (%) | Age- and gender-matched controls relative cerebellar volumes range (%) |
|------------|------------|--------------------------------|---------------------------------------------------------------|--------------------------|---------------------------------------------------------------|
| IV:8       | 61         | 6.64                           | 7.86–10.65                                                   | 4.49                     | 5.65–7.95                                                     |
| IV:6       | 59         | 7.05                           | 7.93–10.72                                                   | 4.64                     | 5.70–7.99                                                     |
| IV:14      | 46         | 7.07                           | 7.93–10.59                                                   | 4.78                     | 5.82–8.00                                                     |
| IV:2       | 64         | 5.76                           | 7.51–10.17                                                   | 3.59                     | 5.50–7.68                                                     |
| V:1        | 36         | 6.26                           | 8.10–10.75                                                   | 4.12                     | 5.99–8.16                                                     |

MRI assessment in five patients revealed significant decrease in cerebellar volume and cerebellar grey matter compared with age- and gender-matched controls (Supplementary Table 3).
Figure 2 Moderate axonal sensory polyneuropathy predominantly in the lower limbs and abnormal SSR and R–R interval in affected patients III:7, IV:6 and IV:2, respectively. (A and B) Moderate axonal sensory polyneuropathy predominantly in the lower limbs in IV:2. At later stages of disease progression, patients revealed reduction of sensory nerve amplitudes also in upper limbs (Supplementary Fig. 4). Tibial nerve SSEPs showed an abnormal latency of P37 response in IV:2 (C) and III:7 (D). (E) Abnormal SSR in lower limbs displayed abnormality in two patients (IV:2 and IV:6). (G) Altered R–R interval in Patient IV:2.
mitochondrial replication indicative of mitochondrial biogenesis (Fig. 5D). In affected fibroblast cell lines, the ATP concentration was found increased compared with those from age-matched control fibroblasts (Fig. 5E). Despite these mitochondrial alterations suggestive of mitochondrial stress, no significant DNA alterations were identified in the mitochondrial genomes from fibroblasts’ patients, discarding further mtDNA damage (Supplementary Fig. 12).
To further investigate mitochondrial perturbations, we immunoblotted with anti-ATP5H, a critical component of the energy-producing apparatus in eukaryotic cells, and found that it was increased in patients’ fibroblasts despite normal levels of VDAC protein located in the mitochondrial outer membrane (Fig. 6A and B, Supplementary Figs 22–27).

Table 3 Two-point LOD score

| θ       | 0      | 0.05 | 0.1   | 0.15  | 0.2   | 0.25  | 0.3   | 0.35  | 0.4   | 0.5   |
|---------|--------|------|-------|-------|-------|-------|-------|-------|-------|-------|
| Z (θ)   | 3.43   | 3.13 | 2.819 | 2.48  | 2.13  | 1.76  | 1.37  | 0.97  | 0.57  | 0     |

The maximum LODs score was obtained with the c.1877C > T (p.Ser626Leu) variation within the SAMD9L gene on 7q21 ($Z_{max} = 3.43, \theta = 0.00; P = 3.53 \times 10^{-5}$).

To further investigate mitochondrial perturbations, we immunoblotted with anti-ATP5H, a critical component of the energy-producing apparatus in eukaryotic cells, and found that it was increased in patients’ fibroblasts despite normal levels of VDAC protein located in the mitochondrial outer membrane (Fig. 6A and B, Supplementary Figs 22–27).

**Figure 4** Mitochondrial localization of SAMD9L in human cerebellar Purkinje cells and multipolar neurons of the dentate nucleus and fibroblasts. (A) Immunostaining with anti-SAMD9L antibody in human control cerebellar sections revealed a punctate staining mainly in Purkinje cell soma (left) and in multipolar neurons (right) of the human dentate nucleus indicative of mitochondrial staining. Black arrows point to SAMD9L staining in Purkinje and multipolar neurons. (B and C) Immunofluorescence staining of fibroblasts from human control (B) and a M-SCA affected patient (C) showed co-localization of SAMD9L to mitoTracker Red CMXRos demonstrating mitochondrial co-localization. Magnification bars: 50 µm (A), 20 µm (B and C).
Figure 5 Reduced SAMD9L protein levels and mitochondrial alterations in M-SCA patients’ fibroblasts. (A) Immunoblotting of lysed and subcellular fractioned fibroblasts confirmed SAMD9L mitochondrial localization, and decreased SAMD9L protein levels in the mitochondrial (A) and total (B) cellular extracts from patients’ fibroblasts samples (N = 2) without alteration of SAMD9L cDNA levels compared by qRT-PCR (C) [N = 2 fibroblasts samples for each group of individuals (patients and controls); three technical replicates each for SAMD9L and three technical replicates each for GAPDH housekeeping gene expression]. (D) Increased copy number of the mtDNA D-LOOP mitochondrial genome region in patient’s fibroblasts samples is suggestive of an increased activity of mtDNA replication. (E) A trend of increased ATP was found in mitochondrial fractions of affected fibroblasts [N = 2 fibroblasts samples for each group of individuals (patients and controls); four technical replicates each] compared with age-matched controls [N = 2 fibroblasts samples for each group of individuals (patients and controls); four technical replicates each]. Each point represents relative ATP levels compared with controls after normalizing for protein concentration. Controls values were set to 1 (dotted line).
Because ATP5H plays a crucial role in regulating apoptosis, this evidence further supports the upregulation of ATP synthase identified. Additionally, DRP1 protein levels also appeared to increase. Remarkably, DRP1 is a dynamin-like GTPase involved in the mitochondrial division during mitochondrial biogenesis, which has also been associated with lysosomes in brain, and implicated in Purkinje cells mitochondrial transport. Related to lysosomes and autophagy, the lysosomal-associated membrane protein LAMP1, and the autophagy-related proteins LC3-II and p62/SQSTM1 also appeared overexpressed in patients’ fibroblasts (Fig. 6A and B). Overall, these data demonstrate SAMD9L mitochondrial localization, suggest an increase of mtDNA replication and biogenesis, and point to dysregulation of the lysosomal/autophagy pathway in this novel SCA subtype caused by the SAMD9L mutation.

To investigate subcellular structural changes, ultrastructural images from fibroblasts obtained from Patient IV:14 obtained by TEM, showed diffused mitochondrial crests (Fig. 6C), evidence of mitophagy (Fig. 6D) and dilated endoplasmic reticulum (Fig. 6E and F; Supplementary Fig. 13). Significant higher number of lysosomes in affected fibroblasts was also identified [F(1,18) = 6.135, P = 0.023; Fig. 6G; Supplementary Fig. 13]. No differences in the number of mitochondria were observed.

The c.1877C>T (p.Ser626Leu) mutation in SAMD9L impairs locomotion and neurosensory phenotypes in zebrafish

Four putative genomic regions containing the candidate human SAMD9L ortholog gene sequence were identified in zebrafish by using meta-analysis data from previously generated transgenic zebrafish models (http://zfim.org), and compared with zebrafish cerebellar RNA-seq data from the SRA database (access: SRX4184229; Supplementary Fig. 14). All four regions presented low predicted translated protein identity (<30%) with that of human SAMD9L and did not show conservation for the Ser626 amino acid. For this reason, transient expression of the human wild-type did not show conservation for the Ser626 amino acid. For this reason, transient expression of the human wild-type and mutated SAMD9L mRNAs was generated in zebrafish instead of knocking out or down the endogenous zebrafish ortholog candidates.

Accumulated survival at 24 and 96 h post-fertilization was not significantly impaired with any mRNA concentration tested, and the teratogenic rate did not show any concentration dependency (Supplementary Fig. 15). Mean distance travelled was significantly reduced in five dpf fish exposed to SAMD9L-S626L (N = 24) compared with the wild-type SAMD9L group (N = 26) under dark intervals of high locomotor activity [F(1,48) = 4.745, P = 0.034; Fig. 7A]. Furthermore, fish overexpressing wild-type SAMD9L showed a significantly increase in mean distance moved in the dark compared with control injected group (N = 28) [F(1,52) = 6.583, P = 0.013; Fig. 7A]. Additionally, the number of head turns during the five light–dark cycles was significantly increased in WT-SAMD9L compared with the control group (Mann–Whitney U = 216, P = 0.010; Fig. 7B), and significantly decreased in S626L-SAMD9L zebrafish larvae compared with the WT-SAMD9L (Mann–Whitney U = 192.5, P = 0.022; Fig. 7B) demonstrating vestibular and sensory impairment by the mutation in SAMD9L. This evidence of higher functional activity of the human wild-type SAMD9L protein was not caused by RNA overexpression (Fig. 7C). Immunoblotting of lysed cellular fractions corroborated recombinant human SAMD9L mitochondrial localization in zebrafish larvae (Fig. 7D; Supplementary Fig. 16). Furthermore, the levels of the mitochondrial fusion protein DRP1 were found increased in mutant SAMD9L compared with either wild-type or controls in zebrafish embryos (Fig. 7D). Increased DRP1 is suggestive of mitochondrial biogenesis.

Whole-mount immunofluorescence of zebrafish larvae showed SAMD9L mitochondrial staining in the zebrafish spinal cord and peripheral nerves (Supplementary Fig. 17) and in the hindbrain (Fig. 7E), co-localizing with the ATP5B mitochondrial marker (Fig. 7F) and parvalbumin (Fig. 7E) in WT-SAMD9L and S626L-SAMD9L zebrafish larvae neurons. Parvalbumin showed positive immunostaining in inhibitory and excitatory interneurons and motoneurons in the zebrafish spinal cord.

Discussion

In the present study, we describe a new SCA subtype in nine affected individuals of a five-generation family of Menorcan ancestry, characterized by ataxia, horizontal and vertical GEN, dysarthria, polyneuropathy, pyramidal signs and cerebellar demyelination as distinctive features. We identified the c.1877C > T (p.Ser626Leu) mutation within the fifth exon of the gene encoding for the sterile alpha motif domain containing 9 like protein SAMD9L as the genetic cause underlying this new SCA. The functional studies in patients’ fibroblasts demonstrated for the first time the mitochondrial localization of human SAMD9L protein and mitochondrial perturbations underlying the molecular pathology in this new ataxia subtype.

The clinical phenotype of this new SCA subtype is characterized by early horizontal and vertical GEN and hypertonia, which precede the sensory polyneuropathy and ataxia symptoms. Whereas GEN in all directions indicates cerebellar dysfunction having multiple causes in cerebellar ataxias, purely vertical GEN is caused by midbrain lesions and purely horizontal GEN is due to pontomedullary affection.

Albeit GEN is a common feature sign in many SCAs subtypes, including SCA1, SCA3, SCA4, SCA5, SCA6, SCA10, SCA11, SCA12, SCA13, SCA14, SCA15, SCA16, SCA18, SCA19, SCA20, rarely in SCA23, SCA25, SCA27, SCA28, SCA31, frequently in SCA36 (in 30% of the cases), SCA38 and SCA42 (only...
Figure 6 Mitochondrial and lysosome/autophagy alterations in fibroblasts from ataxia patients. (A and B) The ATP5H and DRP1 mitochondrial associated proteins levels were found increased without changes in VDAC and MFN1 suggestive of mitochondrial biogenesis. Furthermore, lysosomal LAMP1 together with SQSTM1 and LC3-II protein levels were found increased suggestive of mitophagy. LAMP1, MFN1, SQSTM1, ATP5H AND LC3 proteins levels were normalized to actin B (ACTIN) and DRP1 proteins levels to GAPDH (Supplementary Figs 22–27). Controls values were set to 1 (dotted line). TEM showed diffused crests in the mitochondrial matrix (C) together with some mitophagy (D) and dilated endoplasmic reticulum (E and F) compared with age-matched control (Supplementary Fig. 13). (G) One-way ANOVA confirmed significantly increased number of lysosomes on patients' fibroblasts samples (N = 10) compared with controls [N = 10; F(1,18) = 6.135, P = 0.023] (Supplementary Fig. 13). Black arrowheads point to mitochondria; white arrowheads point to the dilated endoplasmic reticulum. Asterisks indicate autophagosomes. Magnification bars: 0.5 µm (C, D and F), 1 µm (E). Asterisk denotes significance at P < 0.05. Vertical bars denote SEM.
Figure 7 Mutant Ser626Leu SAMD9L triggers locomotive and neurosensory impairment and co-localizes with ATP5B or parvalbumin in zebrafish neurons. (A) One-way ANOVA with the mean distance travelled (mm) during five dark periods of high activity revealed significantly increase in the WT-SAMD9L zebrafish ($N = 26$) compared with the lynGFP control group ($N = 28$) ($F(1,52) = 6.583$, (continued)
Multiple demyelinating lesions in the periventricular subcortical white matter have only been described in SCA9 patients from a large multigenerational American family. Consequently, SCA1, SCA3, SCA6, SCA11, SCA14, SCA15/16 and SCA28 should all be considered in the differential clinical diagnosis, but when MRI data also reveal early brain demyelinating lesions, SAMD9L mutations should be first tested. Besides brain demyelinating lesions, GEN and hyperreflexia also appeared early at the pre-symptomatic stages preceding ataxia symptoms in this novel SCA subtype.

Herein, we provide evidence that the missense mutation c.1877C>T (p.Ser626Leu) within the fifth exon of the SAMD9L gene decreases SAMD9L protein levels along with increased ATP5H, DRP1 and ATP evidencing mitochondrial dysregulation in patients’ fibroblasts. Furthermore, we describe for the first time the mitochondrial localization of human SAMD9L protein in human fibroblasts and zebrafish.

To date, 35 different mutations have been identified with in SAMD9L gene on Chromosome 7q21 in 93 patients from 44 unrelated families variably associated with haematological malignancies including cytopenias or myelodysplastic syndrome (MDS), ATM and ATPX with neurologic manifestations or infantile autoinflammatory disease. All those patients with SAMD9L-related haematological malignancies presented, in addition to the SAMD9L mutation, either uniparental monosomy of Chromosome 7, partial haploinsufficiency del(7q) including SAMD9L, or somatic driver mutations within SETBP1, ASXL1, RUNX1, PTPN11, KRAS, CBL, EZH2, ETV6, BRAF or RAD21 genes. It seems plausible that for developing MDS, SAMD9L mutations need to occur along with Monosomy 7, either partial or total del(7q) haploinsufficiency, or with mutations in those driver genes. In consonance, our SAMD9L ataxia patients did not reveal abnormal blood cell counts, MDS or cytopenia in their longitudinal medical records, or loss of c.1877C>T SAMD9L heterozygosity detected by NGS or Sanger sequencing from blood white-cells DNA, thus ruling out SAMD9L partial or total haploinsufficiency in their haematological lineages. This may explain the absence of haematological malignancy in our patients. Thus, ours is the first family reported to date with a SCA phenotype.
caused by a SAMD9L mutation without the presence of any haematological pathology. This supports the hypothesis proposed that primary SAMD9L mutations do not lead per se to tumour cell proliferation, but that a secondary genomic event consisting of either Chromosome 7 monosomy or somatic mutations in driver genes would be needed to predispose to MDS and haematological malignancy.\(^{19,107}\)

The p.Ser626Leu mutation identified in SAMD9L in our patients localizes to a predicted protein IDR. Over 20% of human disease mutations occur in IDRs,\(^ {115}\) and Serine residue is the third most common amino acid present in IDRs.\(^ {116}\) Furthermore, IDRs are strongly associated with mitochondrial localization and function.\(^ {117}\) IDRs are polypeptide segments typically containing a higher proportion of polar or charged amino acids and do not contain sufficient hydrophobic amino acids to mediate co-operative folding.\(^ {118}\) Mutations within IDR domains, such as those within the IDR of the ITTP1 protein associated with SCA15/29, appear to trigger pathogenicity by creating dileucine motifs affecting protein interactions.\(^ {119}\) Our IDR c.1877C>T SAMD9L mutation promotes a serine (hydrophobic and polar) to leucine (hydrophilic and non-polar) amino acid change at the 626 aa position. This may trigger a similar effect like that in SCA15/29.

Several lines of evidence support mitochondrial perturbations triggered by the SAMD9L mutation in our ataxia patients. We demonstrate for the first time the subcellular localization of human SAMD9L protein to mitochondria in zebrafish and human cerebellar Purkinje cells and fibroblast cell lines. Likewise, the analysis of human ENCODE RNA data corroborated the expression of SAMD9L predominantly in Purkinje cells, cerebellum and spinal cord, which are the main tissues affected in this new ataxia subtype. To investigate how the c.1877C>T (p.Ser626Leu) SAMD9L mutation triggers mitochondrial dysregulation, we used in silico algorithm which showed that the 626ser in SAMD9L protein locates in the predicted S...LKS microtubule-binding motif, which is highly conserved along primate species and is absent in the hSAMD9 protein paralog. Remarkably, homology and structural predictor algorithms predicted an AAA-ATPase domain within SAMD9L protein similar to those in spastin, VPS4B (also known as SKD1), katanin and torsin-1A, which are all associated with microtubules or have been involved in lysosomal trafficking.\(^ {120–122}\) Mutations within the \(SPAST\) gene have been previously associated with autosomal dominant SPG4\(^ {123}\) presenting with hyperreflexia and pyramidal signs, with some patients also manifesting cerebellar ataxia.\(^ {103}\) Furthermore, some spastin mutations including R115C, N184T, L195V and N386K have been identified disrupting lysosomal function,\(^ {104}\) and to alter interaction with microtubules entailing to the abnormal cellular distribution of mitochondria, suggesting organelle transport alteration on the microtubule cytoskeleton, including transport to distal axons.\(^ {124,125}\) Ultrastructural alterations were identified in the patient’s fibroblasts including markedly dilated RER, diffuse mitochondrial crests and the presence of mitophagy vacuoles. This evidence would support the mutation in preventing proper protein folding leading to abnormal mitochondria clearance by mitophagy and compensatory mitochondrial biogenesis. The suggestive increased levels of p62/SQSTM1, DRP1, LAM1 and LC3-II together with the increased copy number of the mtDNA D-LOOP mitochondrial genomic region in patient’s fibroblasts support this hypothesis.

SAMD9L shows sequence similarity to VPS4B, a protein implicated in lysosomal/endoosomal membrane trafficking, regulating the maturation of autophagosomes to degradative autolysosomes, a process also implicating Rab7.\(^ {126,127}\) Rab7 belongs to Rab GTPases proteins which are key regulators of multivesicular body maturation from early endosomes, as well as the fusion of multivesicular bodies mediated by lysosomal degradation.\(^ {128,129}\) Another Rab GTPase, Rab5, mediates the signalling for endosome internalization entailing neurite outgrowth and dendritic branching,\(^ {130}\) regulates endosomes motility by stimulating their stable association with microtubules,\(^ {131}\) co-localizing with SAMD9L in mouse lung fibroblasts and KL cells.\(^ {49}\) Rab GTPases and associated membrane trafficking are implicated in different neurodegenerative diseases as Charcot–Marie–Tooth, amyotrophic lateral sclerosis (ALS), Alzheimer’s disease, Parkinson’s disease and Huntington’s disease,\(^ {129}\) and dysregulation of the lysosomal pathway in the pathogenesis of SCAs including Machado–Joseph disease,\(^ {132}\) SCA6,\(^ {133}\) SCA7\(^ {134}\) and SCA21.\(^ {135}\) To date, only mutations in the AFG3L2 gene coding for mitochondrial protein ATP-dependent metallopro tease are associated with a dominantly inherited ataxia, the SCA subtype 28 (SCA28),\(^ {136}\) albeit mitochondrial dysfunction is a common pathogenic mechanism underlying at least SCA1, SCA2, SCA3, SCA7, SCA10 and SCA12.\(^ {137–142}\)

To demonstrate in vivo pathogenicity of the c.1877C>T (p.Ser626Leu) SAMD9L mutation, we evaluated the molecular implication of SAMD9L and the mutation on locomotion and neurosensory functions in zebrafish. Similar effects on zebrafish phenotype by overexpressing wild-type proteins, such as in our study, were previously described for other genes involved in neurodegenerative diseases such as SPG and neuropathy, \(ATL1\) gene; \(ALS, SOD1\) and \(TDP43\) genes\(^ {143–145}\); as well as with genes associated with ataxia including \(RNF170\) and \(VLDLR.\(^ {146,147}\) Based on our evidence, we propose a dominant negative or haploinsufficiency effect triggered by the mutation within SAMD9L→S626L in zebrafish when compared with the wild-type protein when both are similarly overexpressed for comparison purposes. In fact, autosomal dominant mutations in residues located in molecular interfaces, such as our mutation within SAMD9L, predominantly trigger negative effects by affecting protein–protein interactions.\(^ {148}\) We demonstrate that the SAMD9L mRNA mutation in zebrafish impaired mobility and vestibular/sensory functions compared with the wild-type, pointing to the role of SAMD9L in neurological motor and sensory functions, which were both found altered in our patients.

In conclusion, in this study, we describe a novel SCA subtype, SCA49, caused by a c.1877C>T (p.Ser626Leu)
mutation within the gene encoding for the sterile alpha motif domain containing 9 like protein SAMD9L. Based on the evidence provided, we propose SAMD9L as a protein involved in mitochondrial function underlying neurological motor and sensory functions in this new SCA subtype.

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Competing interests

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Supplementary material

Supplementary material is available at Brain Communications online.

References

1. Matilla-Dueñas A, Goold R, Giunti P. Molecular pathogenesis of spinocerebellar ataxias. Brain. 2006;129(6):1357–1370.
2. Klockgether T, Mariotti C, Paulson HL. Spinocerebellar ataxia. Nat Rev Dis Prim. 2019;5(1):24.
3. Diallo A, Jacobi H, Tezenas du Montcel S, Klockgether T. Natural history of most common spinocerebellar ataxia: A systematic review and meta-analysis. J Neurol. 2021;268(8):2749–2756.
4. Parodi L, Coarelli G, Stevanin G, Brice A, Durr A. Hereditary ataxias and paraparesias: Clinical and genetic update. Curr Opin Neurol. 2018;31(4):462–471.
5. Jacobi H, du Montcel ST, Bauer P, et al. Long-term disease progression in spinocerebellar ataxia types 1, 2, 3, and 6: A longitudinal cohort study. Lancet Neurol. 2015;14(11):1101–1108.
6. de Silva R, Greenfield J, Cook A, et al. Guidelines on the diagnosis and management of the progressive ataxias. Orphanet J Rare Dis. 2019;14(1):51.
7. Meira AT, Arruda WO, Ono SE, et al. Neuroradiological findings in the spinocerebellar ataxias. Tremor Other Hyperkinet Mov. 2019;9:1–8.
8. Rüü U, Schols L, Paulson H, et al. Clinical features, neurogenetics and neuropathology of the polyglutamine spinocerebellar ataxias type 1, 2, 3, 6 and 7. Prog Neurobiol. 2013;104:38–66.
9. Matilla-Dueñas A, Infante J, Serrano-Munuera C, Ivánovic-Barbeiro Y, Álvarez R, Sánchez I. Novel therapeutic challenges in cerebellar diseases. In: Manto M Schmahmann JD Rossi F Ivánovic-Barbeito Y, Alvarez R, Sánchez I. Novel therapeutic challenges in cerebellar diseases. In: Manto M Schmahmann JD Rossi F. Handbook of the cerebellum and cerebellar disorders. Springer International Publishing; 2020:1–33.
10. Huang M, Verbeek DS. Why do so many genetic insults lead to Purkinje Cell degeneration and spinocerebellar ataxia? Neurosci Lett. 2019;688:49–57.
11. Paulson HL, Shakkottai VG, Clark HB, Orr HT. Polyglutamine spinocerebellar ataxias—from genes to potential treatments. Nat Rev Neurosci. 2017;18(10):613–626.
12. Jones L, Houlden H, Tabrizi SJ. DNA repair in the trinucleotide repeat disorders. Lancet Neurol. 2017;16(1):88–96.
13. Matilla-Dueñas A, Ashizawa T, Brice A, et al. Consensus paper: Pathological mechanisms underlying neurodegeneration in spinocerebellar ataxias. Cerebellum. 2014;13(2):269–302.
14. Bea L. Mitochondrial dysfunction in neurodegenerative diseases. Biochim Biophys Acta. 1998;1366(1–2):211–223.
15. Higuchi Y, Okunushi R, Hara T, et al. Mutations in COA7 cause spinocerebellar ataxia with axonal neuropathy. Brain. 2018;141(6):1622–1636.
16. Sahoo SS, Kozyra EJ, Wlodarski MW. Germline predisposition in myeloid neoplasms: Unique clinical and cellular features of GATA2 deficiency and SAMD9/SAMD9L syndromes. Best Pract Res Clin Haematol. 2020;33(3):101197.
17. de Jesus AA, Hou Y, Brooks S, et al. Distinct interferon signatures and cytokine patterns define additional systemic autoinflammatory diseases. J Clin Invest. 2020;130(4):1669–1682.
18. Chen D-H, Below JE, Shimamura A, et al. Ataxia-pancytopenia syndrome is caused by missense mutations in SAMD9L. Am J Hum Genet. 2016;98(6):1146–1158.
19. Tesi B, Davidson J, Voss M, et al. Gain-of-function SAMD9L mutations cause a syndrome of cytopenia, immunodeficiency, MDS, and neurological symptoms. Blood. 2017;129(16):2266–2279.
20. Narumi S, Amano N, Ishii T, et al. SAMD9 mutations cause a novel multisystem disorder, MIRAGE syndrome, and are associated with loss of chromosome 7. Nat Genet. 2016;48(7):792–797.
21. Schwartz JR, Ma J, Lamprecht T, et al. The genomic landscape of pediatric myelodysplastic syndromes. Nat Commun. 2017;8(1):1537.
22. Wong JC, Bryant V, Lamprecht T, et al. Germline SAMD9 and SAMD9L mutations are associated with extensive genetic evolution and diverse hematologic outcomes. JCI Insight. 2018;3(14):e121086.
23. Schmitz-Hübsch T, du Montcel ST, Baliko L, et al. Scale for the assessment and rating of ataxia: Development of a new clinical scale. Neurology. 2006;66(11):1717–1720.
24. Jablecki CK, Andary MT, Di BM, et al. American Association of Electrodiagnostic Medicine guidelines for outcome studies in electrodiagnostic medicine. Muscle Nerve. 1996;19(12):1626–1635.
25. Kimura J. Electrodiagnosis in diseases of nerve and muscle. Oxford University Press; 2013.
26. Romero JE, Coupé P, Giraud R, et al. CERES: A new cerebellum lobe segmentation method. Neuroimage. 2017;147:916–924.
27. Manjon JV, Coupé P, volBrain: An online MRI brain volumetry system. Front Neuroinform. 2016;10:30.
28. Lehrs S, Hildebrand DG, Merz B, et al. LORD-Q: A long-run real-time PCR-based DNA-damage quantification method for nuclear and mitochondrial genome analysis. Nucleic Acids Res. 2014;42(6):e41.
29. Abecasis GR, Cheny SS, Cookson WO, Cardon LR. Merlin—rapid analysis of dense genetic maps using sparse gene flow trees. Nat Genet. 2002;30(3):89–97.
30. Cortingham RW, Idury RM, Schaffer AA. Faster sequential genetic linkage computations. Am J Hum Genet. 1993;53(1):252–263.
31. Schäffer AA, Gupta SK, Shriram K, Cortingham RW. Avoiding re-computation in linkage analysis. Hum Hered. 1994;44(4):225–237.
32. Ott J. Computer-simulation methods in human linkage analysis. Proc Natl Acad Sci U S A. 1989;86(11):4175–4178.
33. ENCODE Project Consortium. An integrated encyclopedia of DNA elements in the human genome. Nature. 2012;489(7414):57–74.
34. Manjón JV, Coupé P. volBrain: An online MRI brain volumetry tool. Nucleic Acids Res. 2015;43(2):W373–W343.
35. Hildebrand A, Remmert M, Biegert A, Soding J. Fast and accurate automatic structure prediction with HHpred. Proteins. 2009;77(Suppl 9):128–132.
36. Hulst N, Barouch A, Buillard V, et al. The 20 years of PROSITE. Nucleic Acids Res. 2008;36:D245–D249.
37. Zhou Y, Yang S, Mao T, Zhang Z. MAPAnalyzer: A novel online tool for analyzing mitocutubule-associated proteins. Database. 2015;2015:bav108.
38. Yachdav G, Kloppmann E, Kajari L, et al. PredictProtein—an open resource for online prediction of protein structural and functional features. Nucleic Acids Res. 2014;42(Web Server issue):W337–W343.
39. Linding R, Jensen LJ, Diella F, Bork P, Gibson TJ, Russell RB. Protein disorder prediction: Implications for structural proteomes. Structure. 2003;11(11):1453–1459.
40. Mitchell AL, Artwood TK, Bab inert PC, et al. InterPro in 2019: Improving coverage, classification and access to protein sequence annotations. Nucleic Acids Res. 2019;47(D1):D351–D360.
41. Necci M, Piovesan D, Dostatyń Z, Tosatto SCe. MobiDBLite: Fast and highly specific consensus prediction of intrinsic disorder in proteins. Bioinformatics. 2017;33(9):1402–1404.
42. Blom N, Gammeltoft S, Brunak S. Sequence and structure-based prediction of eukaryotic protein phosphorylation sites. J Mol Biol. 1999;294(5):1351–1362.
43. Horn H, Schoof EM, Kim J, et al. KinomeXplorer: An integrated platform for kinome biology studies. Nat Methods. 2014;11(6):590–604.
44. Szklarczyk D, Gable AL, Lyon D, et al. STRING v11: Protein–protein association networks with increased coverage, supporting functional discovery in genome-wide experimental datasets. Nucleic Acids Res. 2019;47(D1):D607–D613.
45. Sánchez I, Hassinger L, Paskevich PA, Shine HD, Nixson RA. Oligodendroglioma regulate the regional expansion of axon caliber and local accumulation of neurofilaments during development independently of myelin formation. J Neurosci. 1996;16(16):5095–5105.
46. Eskelinen EL. To be or not to be? Examples of incorrect identification of autophagic compartments in conventional transmission electron microscopy of mammalian cells. Autophagy. 2008;4(2):257–260.
47. Meijer AJ, Codogno P. Autophagy: Regulation and role in disease. Crit Rev Clin Lab Sci. 2009;46(4):210–240.
48. Mekhedov SL, Makarova KS, Koonin EV. The complex domain architecture of SAMD9 family proteins, predicted STAND-like NTPass, suggests new links to inflammation and apoptosis. Biol Direct. 2017;12(1):13.
49. Chen D-H, Jrkancad Z, Verlinden CLMJ, et al. Missense mutations in the regulatory domain of PKC gamma: A new mechanism for dominant nonprogressive cerebellar ataxia. Am J Hum Genet. 2003;72(4):839–849.
50. Nissanka N, Moraes CT. Mitochondrial DNA damage and reactive oxygen species in neurodegenerative disease. FEBS Lett. 2018;592(5):728–742.
51. Song HK, Kim JH, Lee YH, et al. Mitochondrial reprogramming via ATP5H loss promotes multimodal cancer therapy resistance. J Clin Invest. 2018;128(9):4098–4114.
52. Iroki K, Adachi Y, Yamada T, et al. A brain-enriched Drp1 isoform associates with lysosomes, late endosomes, and the plasma membrane. J Biol Chem. 2018;293(30):11309–11322.
53. Fukumitsu K, Hatsu K, Yoshimura A, Heuser J, Fujishima K, Kengaku M. Mitochondrial fission protein Drp1 regulates mitochondrial transport and dendritic arborization in cerebellar Purkinje cells. Mol Cell Neurosci. 2016;71:56–65.
54. Leinonen R, Sugawara H, Shumway M. The sequence read archive. Nucleic Acids Res. 2011;39(Database):D19–D21.
55. Berg EM, Bertuzzi M, Ampatzis K. Complementary expression of calcium binding proteins delineates the functional organization of the locomotor network. Brain Struct Funct. 2018;223(5):2181–2196.
56. Strupp M, Hüfner K, Sandmann R, et al. Central oculomotor disturbances and nystagmus: A window into the brainstem and cerebellum. Dtsch Aerztebl Int. 2011;108(12):197–204.
57. Rivaud-Pechoux S, Durr A, Gaymard B, et al. Eye movement abnormalities correlate with genotype in autosomal dominant cerebellar ataxia type I. Ann Neurol. 1998;43(3):297–302.
58. Bürk K, Fetter M, Abele M, et al. Autosomal dominant cerebellar ataxia type I: Oculomotor abnormalities in families with SCA1, SCA2, and SCA3. J Neurol. 1999;246(9):789–797.
59. Jacobi H, Reetz K, du Montcel ST, et al. Biological and clinical characteristics of individuals at risk for spinocerebellar ataxia types 1, 2, 3, and 6 in the longitudinal RISCA study: Analysis of baseline data. Lancet Neurol. 2013;12(7):650–658.
60. Bürk K, Zühlke C, König IR, et al. Spinocerebellar ataxia type 5: Clinical and molecular genetic features of a German kindred. Neurology. 2004;62(2):327–329.
61. Gomez CM, Thompson RM, Gammack JT, et al. Spinocerebellar ataxia type 6: Gaze-evoked and vertical nystagmus, Purkinje cell degeneration, and variable age of onset. Ann Neurol. 1997;42(6):933–950.
62. Teive HAG, Munhoz RP, Raskin S, et al. Spinocerebellar ataxia type 10: Frequency of epilepsy in a large sample of Brazilian patients. Mov Disord. 2003;18(25):1875–1878.
63. Johnson J, Wood N, Guotti P, Houliden H. Clinical and genetic analysis of spinocerebellar ataxia type 11. Cerebellum. 2008;7(2):159–164.
64. Herman-Bert A, Stevanin G, Netter JC, et al. Mapping of spinocerebellar ataxia 13 to chromosome 19q13.3–q13.4 in a family with autosomal dominant cerebellar ataxia and mental retardation. Am J Hum Genet. 2000;67(1):229–235.
65. Rosini F, Pretegiani E, Battisti C, Dotti MT, Federico A, Rafa A. Eye movement changes in autosomal dominant spinocerebellar ataxias. Neurol Sci. 2020;41(7):1719–1734.
66. Storey E, Gardner RJM. Spinocerebellar ataxia type 15. Handb Clin Neurol. 2012;103:561–565.
67. Bekanac Z, Fernandez M, Matsushita M, et al. Autosomal dominant sensory/motor neuropathy with ataxia (SMNA): Linkage to chromosome 7q22–q32. Am J Med Genet. 2002;114(4):450–457.
68. Chung M-Y, Lu Y, Cheng N, Soong B. A novel autosomal dominant spinocerebellar ataxia (SCA22) linked to chromosome 1p21–q23. Brain. 2003;126(Pr 6):1293–1299.
69. Knight MA, Gardner RJMM, Bahlo M, et al. Dominantly inherited ataxia and dysphonia with dentate calcification: Spinocerebellar ataxia type 20. Brain. 2004;127(Pr 5):1172–1181.
70. Jeziorska J, Stevanin G, Watanabe H, et al. Identification and characterization of novel PDYN mutations in dominant cerebellar ataxia cases. J Neurol. 2013;260(7):1807–1812.

71. Stevanin G, Bouslam N, Thibois S, et al. Spinocerebellar ataxia with sensory neuropathy (SCA25) maps to chromosome 2p. Ann Neurol. 2004;55(1):97–104.

72. van Swieten JC, Brusse E, de Graaf BM, et al. Mutation in the fibroblast growth factor 14 gene is associated with autosomal dominant cerebellar ataxia [corrected]. Am J Hum Genet. 2003;72(1):199–199.

73. Cagnoli C, Mariotti C, Taroni F, et al. SCA28, a novel form of autosomal dominant cerebellar ataxia on chromosome 18q11.22–q11.2. Brain. 2006;129(Pt 1):235–242.

74. Nagaoka U, Ishikawa K, et al. A gene on SCA4 locus causes dominantly inherited pure cerebellar ataxia. Neurology. 2000;54(10):1971–1975.

75. Ikeda Y, Ohta Y, Kobayashi H, et al. A recurrent mutation in CCDC88C causes spinocerebellar ataxia type 22. Ann Neurol. 2014;95(2):209–217.

76. Couteilier M, Blanesac I, Monteil A, et al. A recurrent mutation in CACNA1G alters Cav3.1T-type calcium-channel conduction and causes autosomal-dominant cerebellar ataxia. Am J Hum Genet. 2015;97(5):726–737.

77. Raposo M, Vasconcelos J, Bettencourt C, Kay T, Coutinho P, Lima M. Nystagmus as an early ocular alteration in Machado-Joseph disease (MJD/SCA3). BMC Neuro. 2014;14(1):17.

78. Christova P, Anderson JH, Gomez CM. Impaired eye movements in presymptomatic spinocerebellar ataxia type 6. Arch Neurol. 2008;65(4):530–536.

79. Bürk K, Abele M, Fetter M, et al. Autosomal dominant cerebellar ataxia type I. Clinical features and MRI in families with SCA1, SCA2 and SCA3. Brain. 1996;119(5):1497–1505.

80. Velázquez-Pérez LC, Rodríguez-Labrada R, Fernandez-Ruiz J. Spinocerebellar ataxia type 2: Clinico genetic aspects, mechanistic insights, and management approaches. Front Neurol. 2017;8:472.

81. Teive H, Munhoz R, Arruda W, et al. Spinocerebellar ataxias—genotype-phenotype correlations in 104 Brazilian families. Clinics. 2012;67(5):443–449.

82. Infante J, Combarros O, Volpini V, Corral J, Llorca J, Berciano J. Autosomal dominant cerebellar ataxias in Spain: Molecular and clinical correlations, prevalence estimation and survival analysis. Acta Neurol Scand. 2003;111(5):391–399.

83. Day JW, Surt LJ, Moseley ML, Durand AC, Ranum LP. Spinocerebellar ataxia type 8: Clinical features in a large family. Neurology. 2000;55(5):649–657.

84. Worth PF, Giunti P, Gardner-Thorpe C, Dixon PH, Davis MB, et al. A landscape of germ line mutations in spinocerebellar ataxia with motor neuron involvement (Asidan). Neurology. 2012;79(4):333–341.

85. Di Gregorio E, Borroni B, Giorgio E, et al. ELOVL5 mutations cause spinocerebellar ataxia 38. Am J Hum Genet. 2014;95(2):209–217.

86. Inoue Y, Ohta Y, Kobayashi H, et al. A novel missense mutation in ELOVL5 identified as a novel causative gene of spinocerebellar ataxias using exome sequencing. Brain. 2010;133(PT 12):3510–3518.

87. Kobayashi H, Abe K, Matsuura T, et al. Novel SCA14 mutation in a Dutch autosomal spinocerebellar ataxia family. Neurology. 2003;61(12):1599–1603.

88. Lee YC, Durr A, Majczynsko K, et al. Mutations in KCND3 cause spinocerebellar ataxia type 22. Ann Neurol. 2012;72(6):859–869.

89. Verbeek DS, van de Warrenburg BP, Wesseling P, Pearson PL, Kremer HP, Sinke RJ. Mapping of the SCA23 locus involved in autosomal dominant cerebellar ataxia to chromosome region 20p13–12.3. Brain. 2004;127(Pt 11):2351–2357.

90. Storey E, Bahlo M, Fahey M, Sisson O, Lueck CJ, Gardner RJM. A novel dominantly inherited pure cerebellar ataxia, SCA 30. J Neurol Neurosurg Psychiatry. 2009;80(4):408–411.

91. Ozaki K, Doi H, Mitsui J, et al. Novel missense mutation in GCCCTG hexanucleotide repeat in NOP56 causes SCA36, a type of spinocerebellar ataxia accompanied by motor neuron involvement. Am J Hum Genet. 2011;89(1):121–130.

92. Tsoi H, Yu ACS, Chen ZS, et al. A novel missense mutation in CCDC88C activates the JNK pathway and causes a dominant form of spinocerebellar ataxia. J Med Genet. 2014;51(9):590–595.

93. Liang L, Chen T, Wu Y. The electrophysiology of spinocerebellar ataxias. Neurophysiol Clin Neurophysiol. 2016;46(1):27–34.

94. Indelicato E, Fanciulli A, Ndayisaba JP, et al. Autonomic function testing in spinocerebellar ataxia type 2. Clin Auton Res. 2018;28(3):341–346.

95. Montes-Brown J, Sánchez-Cruz G, García AM, Baez ME, Velázquez-Pérez L. Heart rate variability in type 2 spinocerebellar ataxia. Acta Neurol Scand. 2010;122(5):329–335.

96. Yeh TH, Lu CS, Chou YHW, et al. Autonomic dysfunction in Machado-Joseph disease. Arch Neurol. 2005;62(4):630–636.

97. González-Salazar C, Takazaki KAG, Martinez ARM, et al. Autonomic dysfunction in hereditary spastic paraplegia type 4. Eur J Neurol. 2019;26(4):687–693.

98. Nielsen JE, Johnsen B, Koefoed P, et al. Hereditary spastic paraplegia with cerebellar ataxia: A complex phenotype associated with a new SPG4 gene mutation. Eur J Neurol. 2004;11(12):817–824.

99. Allison R, Edgar JR, Reid E. Spastin MT domain disease-associated mutations disrupt lysosomal function. Front Neurosci. 2019;13:1179.

100. Higgins JJ, Ph LT, Ide SE, Nee LE, Polymepououlos MH. Evidence for a new spinocerebellar ataxia locus. Mov Disord. 1997;12(3):412–417.

101. Bluteau O, Sebert M, Leblanc T, et al. A landscape of germ line mutations in a cohort of inherited bone marrow failure patients. Blood. 2018;131(7):717–732.

102. Nagata Y, Narumi S, Guan Y, et al. Germline loss-of-function SAMD9 and SAMD9L alterations in adult myelodysplastic syndromes. Blood. 2018;132(21):2309–2313.

103. Pastor VB, Sahoo SS, Boklan J, et al. Constitutional SAMD9L mutations cause familial myelodysplastic syndrome and transient monosomy 7. Haematologica. 2018;103(3):427–437.

104. Ahmed IA, Farooqi MS, Vander Lugt MT, et al. Outcomes of hematopoietic cell transplantation in patients with germline SAMD9/SAMD9L mutations. Biol Blood Marrow Transplant. 2019;25(11):2186–2196.

105. Gercenko S, Komulainen-Ebrahim J, Nordborg K, et al. Ataxia-pancytopenia syndrome with SAMD9L mutations. Neuro Genet. 2017;5(1):e183.

106. Cheah JJC, Brown AL, Schreiber AW, et al. A novel germline SAMD9L mutation in a family with ataxia-pancytopenia syndrome and pediatric acute lymphoblastic leukemia. Haematologica. 2019;104(7):e318–e321.

107. Thunström S, Axellson M. Leukoencephalopathy, demyelinating peripheral neuropathy and dural ectasia explained by a not formerly
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described de novo mutation in the SAMD9L gene, ends 27 years of investigations—a case report. BMC Neuro. 2019;19(1):89.

113. Vaughan D, Bogdanova-Mihaylova P, Costello DJ, et al. Ataxia pancytopenia syndrome due to SAMD9L mutation presenting as demyelinating neuropathy. J Peripher Nerv Syst. 2020;25(4):433–437.

114. King-Robson J, Marshall J, Smith F, Willoughby L, Mansour S, Sztriha L. Ataxia-pancytopenia syndrome due to a de Novo SAMD9L mutation. Neurol Genet. 2021;7(3):e580.

115. Vacic V, Markwick PRL, Oldfield CJ, et al. Disease-associated mutations disrupt functionally important regions of intrinsic protein disorder. PLoS Comput Biol. 2012;8(10):e1002709.

116. Uversky VN. The intrinsic disorder alphabet. III. Dual personality of serine. Intrinsically Disord Proteins. 2015;3(1):e1027032.

117. Zarin T, Strome B, Nguyen Ba AN, Alberti S, Forman-Kay JD, Moses AM. Proteome-wide signatures of function in highly diverged intrinsically disordered regions. Elife. 2019;8:e46883.

118. Babu MM. The contribution of intrinsically disordered regions to protein function, cellular complexity, and human disease. Biochem Soc Trans. 2016;44(5):1185–1200.

119. Babu MM. The contribution of intrinsically disordered regions to protein function, cellular complexity, and human disease. Biochem Soc Trans. 2016;44(5):1185–1200.

120. White SR, Lauring B. AAA+ ATPases: Achieving diversity of function with conserved machinery. Traffic. 2007;8(12):1653–1677.

121. Matamoros AJ, Baas PW. Microtubules in health and degenerative disease of the nervous system. Brain Res Bull. 2016;126(Pr 3):217–225.

122. Hasegawa T, Konno M, Baba T, et al. Mitochondrial dysfunction and oxidative stress contribute to the pathogenesis of spinocerebellar ataxia type 12 (SCA12). J Biol Chem. 2011;286(24):21742–21754.

123. White MC, Gao R, Xu W, et al. Inactivation of hnRNP K by expanded intrinsic AUUCU repeat induces apoptosis via translocation of PKCδ to mitochondria in spinocerebellar ataxia 10. PLoS Genet. 2010;6(6):e1000984.

124. Sánchez I, Balagüé E, Matilla-Dueñas A. Ataxin-1 regulates the cerebellar bioenergetics proteome through the GSK3β-mTOR pathway which is altered in Spinocerebellar ataxia type 1 (SCA1). Hum Mol Genet. 2016;25(18):4021–4040.

125. Ward JM, Stoyas CA, Switoski PM, et al. Metabolic and organellar morphology defects in mice and human patients define spinocerebellar ataxia type 7 as a mitochondrial disease. Cell Rep. 2019;26(5):1189–1202.e6.

126. Beyer A, Scheuring S, Müller S, Mincheva A, Lichter P, Köhrer K. Deregulation of autophagy in postmortem brains of Machado-Joseph disease patients. Neuropathology. 2018;38(2):113–124.

127. Watase K, Unno T, Koike M, et al. Lysosomal contribution to the Purkinje cell degeneration in mouse models of spinocerebellar ataxia type 6. J Neurochem. 2013;125:194–200.

128. Alves S, Cormier-Dequaire F, Marinello M, et al. The autophagy/lysosome pathway is impaired in SCA7 patients and SCA7 knock-in mice. Acta Neuropathol. 2014;128(5):705–722.

129. Di Bella D, Lazzaro F, Busso A, et al. Mutations in the mitochondrial protease gene AFG3L2 cause dominant hereditary ataxia SCA28. Nat Genet. 2010;42(4):313–321.

130. Liu J, Lamb D, Chou MM, Liu Y-J, Li G. Nerve growth factor-mediated neurite outgrowth via regulation of Rab5. Mol Biol Cell. 2007;18(4):1375–1384.

131. Nielsen E, Severin F, Backer JM, Hyman AA, Zerial M. Rab5 regulates motility of early endosomes on microtubules. Nat Cell Biol. 1999;1(6):376–382.

132. Stittler A, Muriel M-P, Marinello M, Brice A, den Dunnen W, Alves S. Deregulation of autophagy in postmortem brains of Machado-Joseph disease patients. Neuropathology. 2018;38(2):113–124.