





RESEARCH ARTICLE

Spinocerebellar ataxia type 14: refining clinicogenetic diagnosis in a rare adult-onset disorder

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Abstract

Objectives: Genetic variant classification is a challenge in rare adult-onset disorders as in SCA-PRKCG (prior spinocerebellar ataxia type 14) with mostly private conventional mutations and nonspecific phenotype. We here propose a refined approach for clinicogenetic diagnosis by including protein modeling and provide for confirmed SCA-PRKCG a comprehensive phenotype

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Introduction

Spinocerebellar ataxias (SCAs) denote rare autosomal-dominant progressive ataxias and the most frequently diagnosed genotypes harbor trinucleotide-repeat expansions.^{1,2} In 2000, the first of now more than 20 conventional mutation SCA genotypes was identified in the protein kinase C gamma (*PRKCG*) gene^{3,4} (MIM 176980) and termed SCA 14⁵ or more recently SCA-PRKCG.⁶ Its prevalence estimates are continually rising and range from <1% to <6% in ataxia cohorts after repeat expansion SCAs have been excluded.^{7–14}

The neuron-specific gamma isoform of PRKC is most abundantly expressed in cerebellar Purkinje cells.¹⁵ It regulates their dendritic growth and calcium permeability and the elimination of climbing fiber synapses.^{16,17} It is yet unclear how different variants affect protein localization, aggregation, and kinase activity.^{18,19} Histopathological reports in variants at residue H101 describe selective Purkinje cell loss in the cerebellar cortex without atrophy in neocortex or deep cerebellar nuclei.^{20,21}

SCA-PRKCG diagnosis relies on genetic testing, however, the classification of variants according to current

description from a German multi-center cohort, including standardized 3D MR imaging. **Methods:** This cross-sectional study prospectively obtained neurological, neuropsychological, and brain imaging data in 33 PRKCG variant carriers. Protein modeling was added as a classification criterion in variants of uncertain significance (VUS). **Results:** Our sample included 25 cases confirmed as SCA-PRKCG (14 variants, thereof seven novel variants) and eight carriers of variants assigned as VUS (four variants) or benign/likely benign (two variants). Phenotype in SCA-PRKCG included slowly progressive ataxia (onset at 4–50 years), preceded in some by early-onset nonprogressive symptoms. Ataxia was often combined with action myoclonus, dystonia, or mild cognitive-affective disturbance. Inspection of brain MRI revealed nonprogressive cerebellar atrophy. As a novel finding, a previously not described T2 hyperintense dentate nucleus was seen in all SCA-PRKCG cases but in none of the controls. **Interpretation:** In this largest cohort to date, SCA-PRKCG was characterized as a slowly progressive cerebellar syndrome with some clinical and imaging features suggestive of a developmental disorder. The observed non-ataxia movement disorders and cognitive-affective disturbance may well be attributed to cerebellar pathology. Protein modeling emerged as a valuable diagnostic tool for variant classification and the newly described T2 hyperintense dentate sign could serve as a supportive diagnostic marker of SCA-PRKCG.

guidelines^{22,23} is challenging. Some of the proposed criteria do not apply to this rare disorder: There is no established model of pathogenicity (criterion PS3) and mostly private mutations render population frequencies/novelty less informative (criterion PS4). Segregation analysis (supporting criterion PP1) is valuable but difficult to pursue outside the research context. Thus, novel *PRKCG* variants are often classified as of uncertain significance (VUS) and must be interpreted against clinical findings²⁴. Conversely, informative phenotype description critically depends on correct genetic case ascertainment. Previous clinical descriptions (comprehensive list in Table S1) suggest a rather unspecific mildly progressive cerebellar ataxia of variable age of onset, in some cases with additional symptoms usually considered of extra-cerebellar etiology.

To improve the clinicogenetic diagnosis of SCA-PRKCG, this observational study investigates the SCA-PRKCG phenotype in a multicenter cohort of PRKCG variant carriers with prospective and standardized data that is available to the clinician in an ataxia patient work-up: clinical exam, brain structural MRI, and neuropsychological and neuro-ophthalmologic testing.²⁵ We add protein modeling as a supporting PP3-criterion in the

classification of pathogenic genetic variants and we explore phenotypic differences to nonconfirmed cases.

Instrumental gait analysis,²⁶ MR spectroscopy,²⁷ and detailed visual testing²⁸ from subcohorts are reported elsewhere.

Methods

The study included symptomatic subjects carrying a *PRKCG* variant considered of either pathogenic or uncertain significance, with referrals from five German university ataxia clinics. Exclusion criteria were any other disease involving the central nervous system, history of severe head trauma, severe psychiatric comorbidity, or contraindication for MRI investigation. A matched group of control subjects was included for the analysis of neuropsychology and brain imaging. Controls were assessed with identical protocol and had no history of neurologic or psychiatric disease, head trauma, no abnormal findings in neurological examination or contraindication for MRI investigation.

Subjects were investigated at one or both of the coordinating and neuroimaging centers (Berlin and Jülich). The study was approved by their respective Institutional Review Boards. Written informed consent was obtained from all participants.

All *PRKCG* variants were re-evaluated by a geneticist (P. B.), first, according to current guidelines put forward by the American College of Medical Genetics and Genomics.²² Minor allele frequency above 1% derived from published databases was set as stand-alone evidence for “benign” variants. Second, a refined approach was applied that included results of protein modeling as a supporting criterion. This modeling evaluated the protein-specific functional impact of a given variant (A. G.). Multitemplate homology modeling using the SwissModel web-server²⁹ was generated that covered the full *PRKCG* protein except for residues 0–35 (see Data S1). Within this model, two zinc-binding cavities are formed by residues C49, C52, C77, and H74 (1st zinc-binding site) and C85, C35, C69, and H36 (2nd zinc-binding site).

The onset of ataxia was defined as onset of permanent gait ataxia. The clinical assessment comprised a structured medical history (including questions to capture history of seizures, myoclonus, dystonia, tremor, spasticity, cognitive or affective disturbance, pain, impairment of mobility and hand function), clinical examination, and application of clinical ratings of ataxia (SARA³⁰, range 0–40) and non-ataxia symptoms (INAS³¹, range 0–16). Comprehensive neuropsychological tests were applied (description and reference in Data S1) and validated screening for affective disturbance³² (HADS) or cognitive impairment³³ (Dem-Tect) performed using published cut-offs.

The afferent visual pathway was assessed by functional testing (visual acuity) and retinal imaging (optical coherence tomography).

Brain MRI included 3D T1- and T2-weighted sequences obtained at 3T (Magnetom Trio system, Siemens Healthineers, Germany).

Electrophysiology results and previously obtained routine brain MRI for longitudinal assessment were made available by patients and not part of the prospective protocol.

Further detail on methods is provided in Data S1.

Data processing and statistical analysis

PRKCG variants were checked against published reports and ordered by location to detect possible feature clusters (Tables 1 and 2, Table S1). Missing information was handled per item as indicated. Neuropsychological test results in confirmed SCA-*PRKCG* were compared to results obtained in healthy controls matched for age, sex, education levels according to the International Standard of Education³⁴ and handedness according to Edinburgh Handedness Inventory.³⁵ Between-group comparisons used t-tests or Wilcoxon rank-sum test as indicated in Table 4. In the case of between-group difference in test results, correlations with ataxia ratings (SARA) and depression score (HADS-D) were performed and, if significant, additional effects of age explored via partial correlations. Spearman or Pearson test was used as indicated in Table 5.

Results of visual pathway assessment (H. Z., T. O.) and brain MRI (M. Sch., S. G.) were each independently inspected and interpreted by two experienced raters. Interpretation of imaging results included a comparison to healthy controls with groups matched for age and sex. The results of electrophysiological testing were reviewed by examiners of the respective centers.

Results

Genetic findings

We investigated 33 subjects (22 families) with 20 *PRKCG* variants, thereof 11 novel variants (Fig. 1).

All participants were of Caucasian ethnicity. The genetic re-evaluation according to current guidelines suggested (likely) pathogenicity in only 6/20 variants and (likely) benign variants in 2/20, whereas VUS was assigned in 12/20 (this included nine novel variants and three variants with suggested pathogenicity in previous reports (p. C77S, p. H116P, p. I173S, see Table S1). Of note, five novel variants were in residues previously published as disease-causing (p. A24S, p. G123A, p. G131S, p. C150Y, p. M256I).

Table 1. List of 20 PRKCG variants ordered by residue along with genetic classification by current (ACMG) guidelines and comprehensive classification decision which included results of protein modeling as supporting criterion, and CADD scores (not available for insertions or deletions as indicated).

PRKCG domain	PRKCG variants (all heterozygous)	n Subjects/families	Interpretation of protein modeling	Classification by current (ACMG) guidelines	Classification including protein modeling results	CADD PHRED score	CADD score	
N-terminal	c.68G > A, p. G23E	3/2	Not covered	VUS	Likely pathogenic ¹	26.8	3.88	
Regulatory domain C1	c.70G > T, p. A24S	1/1	Not covered	VUS	Likely pathogenic ²	26.0	3.74	
	c.146G > A, p. C49Y	1/1	1st Zinc-binding site probably disrupted	VUS	Likely pathogenic ³	27.7	4.00	
	c.197G > A, p. C66Y	5/2		Pathogenic	Pathogenic ⁴	27.0	3.91	
	c.207C > T, p. Cys69Cys	2/1	Benign	Benign	Benign ⁵	14.8	1.24	
	c.229T > A, p. C77S	1/1	1st Zinc-binding site probably disrupted	VUS	Likely pathogenic ³	25.2	3.57	
	c.244-252delACCTCGAG, p. T82_E84del	2/1	2nd zinc-binding site may be structurally affected	VUS	Likely pathogenic ⁶	N/A	N/A	
	c.338_340delTCT, p. F113_C114delInsC	2/1		Likely pathogenic	Likely pathogenic ⁷	N/A	N/A	
	c.347A > C, p. H116P	1/1	Close to 2nd zinc-binding site, may probably disrupt zinc binding	VUS	Likely pathogenic ⁸	27.0	3.91	
	c.353G > A, p. E118D	2/1		Pathogenic	Pathogenic ⁴	27.6	4.00	
	c.367G > A, p. G123R	1/1		Likely pathogenic	Likely pathogenic ⁷	29.3	4.19	
Regulatory domain C2	c.368G > C, p. G123A	1/1	Change in local environment that may affect protein structure	VUS	Likely pathogenic ²	26.3	3.81	
	c.391T > C, p. C131R	2/1		Pathogenic	Pathogenic ⁴	28.6	4.10	
	c.392G > C, p. C131S	2/1		Pathogenic	Pathogenic ⁴	25.9	3.73	
	c.449G > A, p. C150Y	1/1	2nd Zinc-binding site probably disrupted	VUS	Likely pathogenic ²	29.7	4.23	
	c.518T > G, p. I173S	1/1	Change to polar residue in conserved hydrophobic region may affect structure	VUS	VUS ⁹	25.2	3.55	
	c.638G > A, p. R213Q	2/1	Benign	Likely benign	Likely benign ⁹	23.2	2.84	
	c.768G > C, p. M256I	1/1	Near putative calcium-binding site, but no change predicted in chemical properties	VUS	VUS ⁹	22.4	2.44	
	Kinase domain	c.1901G > A, p. R634H	1/1	Benign	VUS	VUS ⁹	24.8	3.44
		c.2032C > G, p. P678A	1/1	Benign	VUS	VUS ⁹	23.2	2.85

Novel variants are written in bold. ACMG, American College of Medical Genetics and Genomics; CADD, combined annotation dependent depletion; PRKCG, protein kinase C gamma; VUS, variant of uncertain significance.

¹Based on ACMG variant classification (VUS), typical phenotype in two independent families within this study.

²Based on ACMG variant classification (VUS), typical phenotype plus another SCA-PRKCG patient with PRKCG missense variant at same residue.

³Based on ACMG variant classification (VUS), typical phenotype plus abnormal PRKCG protein modeling.

⁴Based on ACMG variant classification (pathogenic).

⁵Based on ACMG variant classification (benign).

⁶Based on ACMG variant classification (VUS), typical phenotype plus abnormal PRKCG protein modeling.

⁷Based on ACMG variant classification (likely pathogenic).

⁸Based on ACMG variant classification (VUS), typical phenotype in two families plus abnormal PRKCG protein modeling.

⁹Based on ACMG variant classification (VUS), PRKCG protein modeling suggests functional consequence.

¹⁰Based on ACMG variant classification (likely benign).

¹¹Based on ACMG variant classification (VUS); no further supportive evidence.

*asterisk relates to explanation given in table caption, that CADD scores are not available for insertions or deletions.

Table 2. Individual findings of selected outcomes in all 33 carriers of PRKCG variants, including four subjects with (likely) benign variants and four

PRKCG variant		Disease onset		Clinical rating		Nonataxia movement disorder			Possible pyramidal	
Domain	Variant classification including protein modeling results	Age at onset	disease duration (y)	SARA	INAS count	Myoclonus	Dystonia	Tremor	Increased tone/plantar extensor	Hyperreflexia
N-terminal	Likely pathogenic	30	23	10	2	no	yes	no	no	yes
	Likely pathogenic	26	16	7.25	2	yes	yes	no	no	no
	Likely pathogenic	35	20	8	2	no	no	no	no	yes
	Likely pathogenic	30	21	7	1	yes	no	no	no	no
Regulatory domain C1	Likely pathogenic	37	17	15.5	2	no	yes	no	no	no
	Pathogenic	13	20	7	1	yes	no	no	no	yes
	Pathogenic	48	14	25	2	yes	no	no	no	no
	Pathogenic	50	15	6	3	no	no	no	no	no
	Pathogenic	33	4	8	0	no	no	no	no	no
	Pathogenic	48	4	15	2	no	no	no	no	yes
	Benign	40	27	15.5	5	no	no	no	no	no
	Benign	38	9	2	3	no	yes	no	no	no
	Likely pathogenic	20	34	11.5	5	no	no	no	no	no
	Likely pathogenic	36	34	12	2	no	no	no	no	no
	Likely pathogenic	43	19	10	3	no	no	no	no	yes
	Likely pathogenic	47	11	12	2	yes	yes	no	no	no
	Likely pathogenic	20	11	4.5	0	no	no	yes	no	no
	Likely pathogenic	4	41	13	0	no	no	no	no	no
	Pathogenic	45	11	12	2	no	yes	no	no	no
	Pathogenic	50	3	5	0	yes	no	no	no	no
	Likely pathogenic	31	35	12.25	2	no	no	no	no	no
	Likely pathogenic	37	34	11	5	yes	no	no	no	no
	Pathogenic	11	46	11	2	yes	yes	yes	no	no
	Pathogenic	29	2	7	2	no	no	no	no	no
Pathogenic	41	8	12.25	2	yes	yes	yes	no	no	
Pathogenic	26	3	3	0	yes	no	no	no	no	
Likely pathogenic	20	29	5	1	no	yes	no	no	no	
Regulatory domain C2	VUS	44	9	7	0	no	no	yes	no	no
	Likely benign	no ataxia	n.a.	2	6	yes	no	yes	no	no
	Likely benign		n.a.	0	3	no	no	no	no	no
	VUS	49	8	7.5	3	no	yes	no	yes	no
Kinase domain	VUS	46	6	12.5	1	no	no	no	no	no
C-terminal	VUS	47	4	7	2	no	no	yes	no	no

Subjects are ordered by location of variant (same order as Table 1).

PRKCG protein kinase C gamma; VUS variant of uncertain significance.

n.a. not assessed.

yes/no refers to symptom, sign or abnormal finding present:

MRI cerebellar atrophy rated by inspection as (0) none, (1) mild, (2) moderate and (3) severe.

*MRI results: only report of routine MRI available.

When protein modeling and evaluation of clinical findings from this and previous reports were considered in the second step of variant classification, a genetic diagnosis of SCA-PRKCG was assigned to 14/20 variants (25 subjects/16 families), including seven novel likely pathogenic variants (Table 1, Fig. 1). All these variants were located in the N-terminal or C1 regulatory domain. Results of structural modeling clearly supported a

pathogenic relevance in five variants classified as VUS by current guidelines, as they were likely to impose critical changes at zinc-binding sites. In two other variants, possibly deleterious conformational changes were assigned due to changes in local structural properties (p. G123A) or change from hydrophobic to polar residue (p. I173S). In three other VUS and two benign/likely benign variants, no relevant effects were predicted on protein structure or

Table 2. Continued

Possible peripheral Areflexia (a) or mild pallhypesthesia (p)	Cognitive/psychiatric screening				Nerve conduction studies abnormal			Brain MRI findings		
	Muscle atrophy	DemTect	HADS depression	HADS anxiety	Peripheral nerve	Tibial nerve somatosensory- evoked potentials	Central motor conduction time	Cerebellar atrophy	Brainstem atrophy	T2 hyperintense dentate nucleus
no	no	14	1	3	no	no	no	2	no	yes
no	no	18	7	3	no	no	no	1	no	yes
no	no	11	1	3	yes	no	n.a.	2	no	yes
no	no	17	4	6	no	no	n.a.	1	no	yes
a	no	13	7	4	no	yes	n.a.	1	no	yes
no	no	18	9	4	no	no	no	2	no	yes
no	no	17	9	12	yes	yes	no	2	no	yes
a, p	no	13	4	5	yes	yes	n.a.	1	no	yes
no	no	18	9	6	n.a.	n.a.	n.a.	2	no	yes
no	no	7	12	4	n.a.	n.a.	n.a.	1	no	yes
a, p	yes	12	2	3	n.a.	n.a.	n.a.	1	no	yes
a, p	no	13	0	1	n.a.	n.a.	n.a.	1	no	yes
p	yes	14	7	5	no	no	n.a.	2	no	yes
no	yes	14	7	4	no	yes	no	2	no	yes
no	no	12	6	1	n.a.	n.a.	n.a.	2	no	yes
no	no	13	8	10	yes	no	no	2	no	yes
no	no	14	3	n.a.	no	no	no	2	no	yes
no	no	12	4	11	yes	yes	no	1	no	yes
no	yes	14	9	8	yes	n.a.	n.a.	3	no	yes
no	no	15	11	12	n.a.	n.a.	n.a.	2	no	yes
no	no	15	n.a.	n.a.	n.a.	n.a.	n.a.	2	no	yes
p	yes	n.a.	n.a.	n.a.	no	n.a.	n.a.	2	no	yes
no	no	15	12	2	n.a.	n.a.	n.a.	2	no	yes
no	no	10	3	6	yes	no	n.a.	1	no	yes
p	no	12	4	4	no	n.a.	n.a.	2	no	yes
no	no	18	9	6	n.a.	n.a.	n.a.	1	no	yes
no	no	14	10	10	yes	no	n.a.	2	no	yes
no	no	n.a.	n.a.	n.a.	n.a.	n.a.	n.a.	3	no	yes
a	yes	n.a.	n.a.	n.a.	n.a.	n.a.	n.a.	0	no	n.a.
p	yes	n.a.	n.a.	n.a.	n.a.	n.a.	n.a.	0	no	n.a.
no	no	13	n.a.	n.a.	no	yes	no	n.a.*	n.a.*	n.a.*
no	no	13	6	3	yes	yes	no	2	yes	no
no	no	9	6	4	n.a.	n.a.	n.a.	2	yes	no

function, whereas the two N-terminal variants were not covered by the model. Four remaining VUS were located in the C2 regulatory, kinase, or C-terminal domain. Two variants were classified as (likely) benign despite one (p. C69C) located within the mutational hotspot/1st zinc-binding site.

Family history was negative or not informative in only 3/25 SCA-PRKCG subjects – thereof one singular index

case – but in 3/4 VUS carriers and 2/4 carriers of benign/likely benign PRKCG variants.

Phenotype in SCA-PRKCG

An excerpt of individual findings in 33 subjects (whether confirmed SCA-PRKCG or benign/VUS) is presented in Table 2, whereas Table 3 summarizes the SCA-PRKCG

phenotype based on 25 subjects confirmed as SCA-PRKCG as annotated in Table 2.

The confirmed SCA-PRKCG had a mean age of 38 (SD 13.4) years and a disease duration of 19 (SD 12.5) years. Patients featured mild to moderate ataxia (SARA < 20) in all but one patient (score 25) presenting with additional myoclonus. Three patients reported permanent use of walking aids and none were wheel-chair dependent. INAS count indicated up to five nonataxia signs per patient (none in five patients). Myoclonus involved trunk or hands in most cases and stimulus sensitivity was observed in one patient. Mild focal dystonia was often reported as action-induced or task-specific. Although hyperreflexia was noted in five and a sensation of leg stiffness was reported by two subjects, no spasticity or extensor plantar response was observed. Five subjects reported persistent bone or muscle pain located in the legs or back that increased with exertion without other identifiable cause. Fasciculations or mild to moderate muscle atrophy affected proximal or distal muscle groups with mild to moderate weakness in three subjects.

There was clinical suspicion or subjective complaint of mild cognitive dysfunction in almost half of the patients, whereas DemTect indicated mild cognitive impairment in only five subjects. Screening tests indicated dementia in one subject but coincident with relevant depressive symptoms.

Neuropsychological test results indicated lower performance compared to healthy controls in mainly two domains: attentional functions and executive function (Table 4). Results of SCA-PRKCG in respective tests were unrelated to depressive symptoms (HADS-D) while an association with ataxia severity (SARA) was seen for visuospatial mental rotation and selective attention (Table 5). When effects of age were taken into account using partial correlations, the associations with ataxia scores were no longer significant (Fig. S1).

Assessments of the visual pathway did not indicate pathology of the optic nerve (see Ihl *et al.*²⁸ for detail).

Electroneurographic signs of mild axonal or mixed neuropathy of single nerves were seen in some subjects but did not qualify for a diagnosis of polyneuropathy. Of note, findings were normal in three of four patients who featured reduced vibration sense. Central motor conduction time was normal in all eight subjects with reports available.

Symptom onset and progression

Due to cross-sectional study design, the information in this section relies on patient report. The onset of gait ataxia varied between 4 and 50 years of age (mean/SD 38/13). In two very mildly affected subjects (SARA score 3

and 7), one subjectively unaware of ataxia, limb ataxia of the legs was more prominent than gait/stance ataxia. Disease manifestation coincided with giving birth to the 2nd child in one subject.

Several subjects reported possible early manifestations: minor difficulty with locomotor coordination since childhood (four patients, combination with early learning deficits in one), childhood-onset, nonprogressive slurring of speech (two patients), and reading–writing difficulties (one patient).

The onset of dysarthria was mostly close to or even coincident with the onset of gait ataxia, whereas (mild) dysphagia started later in the disease course. The onset of impaired hand coordination was on average >10 years after the onset of gait ataxia. Early mild writing difficulties before the onset of gait ataxia in one subject were likely attributable to task-specific dystonia. The onset of myoclonus remained unresolved as it often went unrecognized by patients themselves.

Progression of ataxia was slow (SARA annual progression rate 0.99 ± 1.01 pt/year, estimated as SARA scores by disease duration). In the subject with the most severe ataxia (SARA 25), valproate 900mg/day almost completely resolved the action-induced truncal myoclonus with subsequent sustained SARA improvement by five points. Results of earlier neuropsychological testing, available in one patient, indicated only a mild decrease in tests of attention and semantic verbal fluency over a period of 8 years but no impairment in semantic or verbal episodic memory.

Brain imaging

There was no atrophy of cerebrum, brainstem, or cervical spinal cord, but cerebellar atrophy was seen in all SCA-PRKCG subjects (Table 2), particularly pronounced in anterior lobe and upper vermis and including middle or superior cerebellar peduncles in three and two subjects, respectively.

A peculiar symmetrical hyperintensity of the dentate nucleus on T2-weighted images was unequivocally seen in all SCA-PRKCG subjects but none of the healthy controls. It extended from the dentate nucleus toward the superior cerebellar peduncle, whereas in healthy subjects the dentate nucleus was generally hypointense, presenting only a central clear-cut hyperintense spot in some cases resembling dilated perivascular space. The T2-hyperintense signal of the dentate nucleus had a hypointense correlate in T1-weighted images (Fig. 2). The detection of the T2-hyperintense dentate sign was improved by (para)coronal angulation of images along the superior cerebellar peduncles.

Both, the cerebellar atrophy and T2-hyperintense dentate sign, were clearly observed also in two subjects with

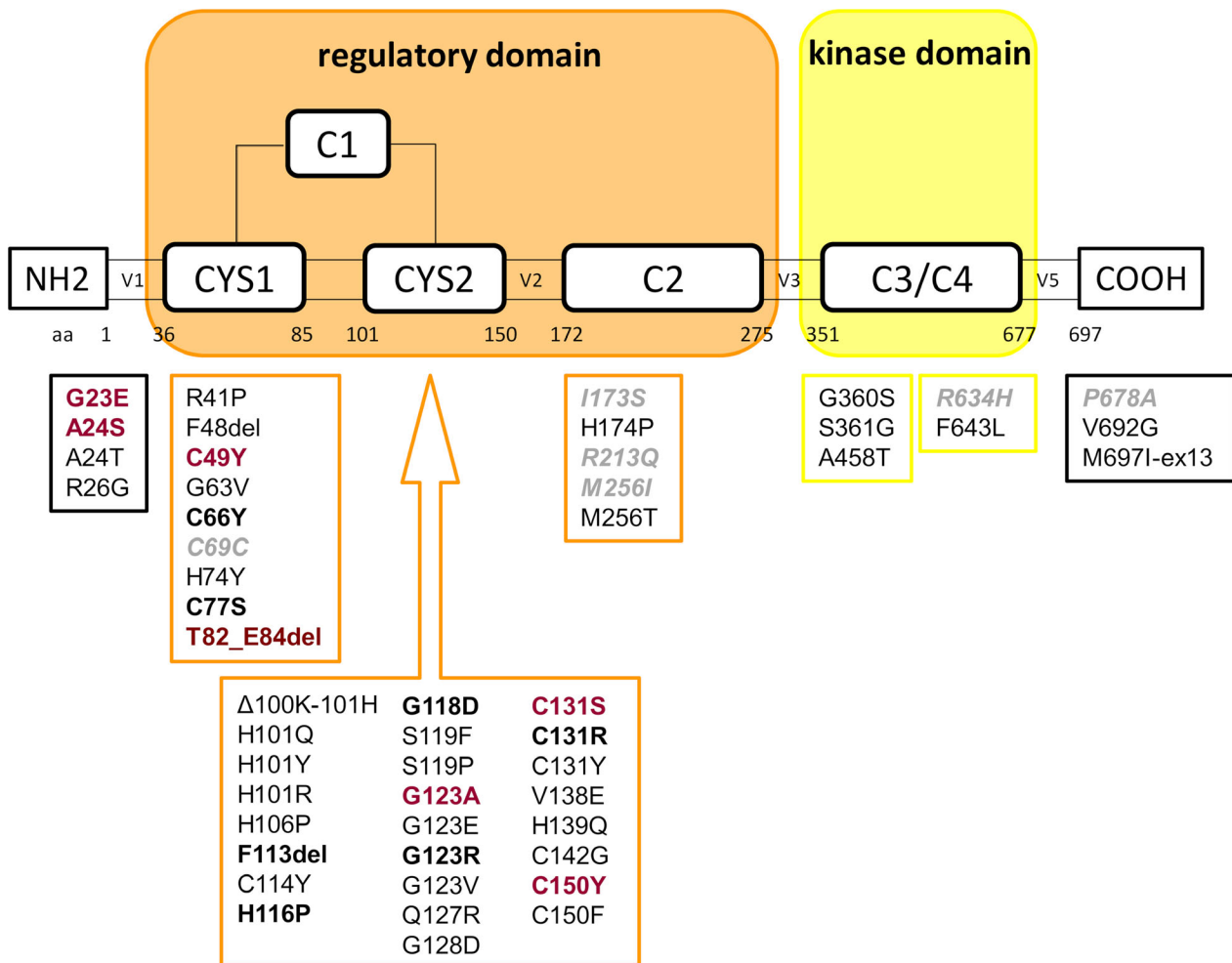


Figure 1. Overview of all PRKCG variants published to date. Variants included in this case series are marked in bold, novel (likely) pathogenic variants are marked in red. Variants of uncertain significance or likely benign variants of our cohort are marked in gray italic.

clinically incipient manifestation. Cerebellar atrophy even preceded clinical manifestation by 8 years in one of the three subjects with preceding routine clinical MRI available. By inspection, no obvious progression of atrophy was seen over periods of 8 to 17 years (Fig. 3). Volume loss could not be quantified since prior routine MRI was not obtained in 3D and slicing did not allow a statement on dentate signal alterations.

Clinical and imaging findings in VUS/(likely) benign cases

The four subjects classified as VUS had nominally older age (mean 46.5 years) and shorter disease duration (mean 6.8 years) than those confirmed as SCA-PRKCG. Nonataxia movement disorder was seen in three of four carriers of VUS and disturbed memory was reported by two

(Table 2). Signs of spasticity were reported in one subject (p. M256I) despite normal central motor conduction times and slowed saccades and horizontal ophthalmoparesis were seen in another subject (p. P678A).

In one parent-offspring pair of a likely benign variant (p. R213Q), no signs of ataxia were observed but myoclonus, resting tremor, mild muscle atrophy, and weakness in the index case. The other family carrying a benign variant (p. C69C) presented with slowly progressive ataxia, areflexia, mild muscle atrophy (1), focal dystonia (1), and moderate to severe sensory disturbance.

Structural brain MRI in carriers of VUS showed extracerebellar pathology in three of four cases with brainstem atrophy (p. R634H and p. P678A), whole-brain atrophy (p. M256I and p. P678A), or hyperintense middle cerebellar peduncle (p. R634H). Such features were not observed in any SCA-PRKCG subject. Furthermore, no T2

Table 3. Summary of clinical findings in 25 cases of confirmed SCA-PRKCG given as proportion (%) of sample with specific findings, ordered by possible structural attribution.

Structure	System	Sign	Observed or reported (% of sample)	<i>n</i> = 25 unless stated otherwise
Cerebellum	Cerebellar ataxia (SARA ratings > 0)	Gait ataxia	25 (100)	
		Stance ataxia	21 (84)	
		Dysarthria	23 (92)	
		Limb ataxia	25 (100)	
	Cerebellar oculomotor signs	Saccadic pursuit	25 (100)	
		Saccadic dysmetria	24 (96)	
		Gaze-evoked nystagmus	15 (60)	
	Non-ataxia movement disorder, observed or reported	Myoclonus	10 (40)	
		Dystonia	8 (32)	
		Tremor	3 (12)	
	Other symptoms or signs of suspected cerebellar attribution	Diplopia	11 (44)	
		Dysphagia	12 (48)	
		Mild cognitive impairment by clinical suspicion or subjective complaint	11 (44)	
Cognitive screening test positive		6 (25)	(<i>n</i> = 24)	
Brainstem	Brainstem oculomotor signs	Ophthalmoparesis	0	
		Slowing of saccades	0	
		Reduced visual acuity (monocular)	0	(<i>n</i> = 13)
Retina/optic nerve	Symptoms or signs of retinal/optic nerve involvement	Optical coherence tomography pRNFL reduction	0	(<i>n</i> = 13)
		Hyperreflexia	5 (20)	
Spinal tract	Symptoms or signs of pyramidal involvement	Spasticity	0	
		Plantar extensor	0	
		Electrophysiology: CMCT abnormal	0	(<i>n</i> = 8)
		Fasciculations	5 (20)	
	Symptoms or signs of spinal or peripheral involvement	Muscle atrophy	4 (16)	
		Pareses	3 (12)	
		Reduced vibration sense (ankle)	4 (16)	(<i>n</i> = 25)
		Electrophysiology: mild neuropathy	8 (44)	(<i>n</i> = 18)
		Electrophysiology: SSEP abnormal	5 (33)	(<i>n</i> = 15)
		Depression/anxiety screening test positive	11 (48)	(<i>n</i> = 23)
Undefined	Symptoms of unclear attribution	Depression/anxiety clinically relevant	5 (22)	(<i>n</i> = 23)
		Cramps or sensation of muscle stiffness	10 (40)	
		Pain in legs or lower back	5 (20)	
		unexplained otherwise		

CMCT, central motor conduction time; PRKCG, protein kinase C gamma; pRNFL, peripapillary retinal nerve fiber layer; SARA, scale for the assessment and rating of ataxia; SSEP, somatosensory-evoked potentials; VUS, variant of uncertain significance.

hyperintense dentate sign was seen in two cases (only report of routine MRI was available for p. M256I carrier). In both carriers of variant p. R213Q, brain MRI was unremarkable without cerebellar atrophy. However, both carriers of variant p. C69C and a singular carrier of VUS (p. I173S) had imaging findings compatible with SCA-PRKCG, including the hyperintense T2 dentate sign.

Discussion

As a main result, we describe a refined variant classification for the clinico-genetic diagnosis of SCA-PRKCG and summarize the SCA-PRKCG phenotype from prospective

investigation of clinical, neuropsychological, and imaging findings in the largest cohort to date. The novel brain MRI finding of T2 hyperintense/T1 hypointense dentate nuclei was shared by all confirmed SCA-PRKCG and may serve as a supportive marker for PRKCG variant classification.

Clinical findings support a variable combination of three motor symptoms: (1) mild to moderate cerebellar ataxia, (2) multifocal action myoclonus, and (3) task-specific or cervical dystonia (including dystonic tremor). The age of ataxia onset had a remarkably wide range apparently unrelated to phenotype or progression. The onset of ataxia related to childbirth in one of our subjects

was also described in other PRKCG variants^{11,36} and in SCA-ATXN10³⁷ but possible mechanisms of aggravation remain speculative. Long-standing mild or nonprogressive symptoms of walking, speech, or learning dysfunction were reported by 8/25 subjects and also noted in previous reports,^{7,10,38,39} suggestive of an early developmental component.

Action myoclonus in SCA-PRKCG may aggravate ataxia or be even mistaken for ataxia in commonly used motor coordination tests and obviously interfered with SARA rating in one of our subjects. Such interference is known from early-onset ataxias.⁴⁰ Further, history taking for myoclonus required specific enquiry, for example, for “muscle jerks at rest or action like you would sometimes experience falling asleep,” as most subjects did not complain of jerks spontaneously, even if clinically observed.

Action as a trigger argues for a cortical origin,⁴¹ further supported by the previous notion of negative myoclonus,⁴² response to valproate⁵ and this first description of stimulus-sensitivity in one patient. Dystonia was mild or intermittent in our study though disabling predominant myoclonus-dystonia has been described in SCA-PRKCG.⁴² The observed head and upper limb tremor was difficult to classify; classification in previous reports included tremulous dystonia, rhythmic myoclonus, or segmental myorhythmia (Table S1).

The nonataxia movement disorders observed in SCA-PRKCG have been referred to as extracerebellar signs in previous reports and are not (yet) considered part of the clinical cerebellar syndrome.⁴³ However, converging arguments attribute them to cerebellar pathology. For myoclonus, the coincidence with symptoms of ataxia has long

Table 4. Results of neuropsychological testing performed in 23 confirmed SCA-PRKCG (13 females; age 49 ± 11 years) and 23 age- and sex-matched controls (13 females; age 49 ± 11 years) along with statistics for group comparison (*t*-test or *Wilcoxon rank-sum (WRS) test).

Domain	Specific skill	Test acronym	<i>n</i>		Mean/median	SD/SE	<i>T/U*</i>	<i>P</i> -value
			SCA-PRKCG/control					
Attention	Selective attention	TAP-Flexibility	22		783.5	247.8	2.7	0.011
			22		605.7	191.6		
	Inhibition	TAP-Go/NoGo	22		550.1	58.7	2.7	0.010
Executive functioning	Processing speed	TAP-Alertness*	23		506.3	48.2		0.007*
			22		298.5	12.1	135*	
	Affinity of interference	FWIT*	23		246	13.6		0.207*
			23		30.7	5	207*	
			23		27.6	2.1		
Interhemispheric motor inhibition	COMO*	23		4.6	0.7	122.5*	0.001*	
		23		0	0.7			
Visuospatial mental rotation	LPS 50 + subtest 7	23		11.8	4.3	-3.1	0.004	
		22		18.3	9			
		23		11.8	4.3	-3.1		
Language	Vocabulary	MWT-B*	23		28	1	213*	0.254*
			23		29	0.8		
	Phonemic verbal fluency	RWT phon. *	23		21	1.2	242*	0.802*
			22		19.5	1.1		
	Semantic verbal fluency	RWT sem.	23		24.6	5.7	-0.6	0.553
22				25.6	5.3			
Memory	Figural memory	ROCFT learning*	23		18.5	1.5	192*	0.111*
			23		23	1.3		
			23		18	1.5	211.5*	
	Visual spatial working memory	CBT*	23		22	1.5		0.244*
			23		10	0.3	207.5*	
	Verbal episodic memory	VLMT learning*	23		10	0.4		0.196*
			23		59	1.8	260*	
			23		57	1.6		
Verbal working memory	VLMT delayed*	23		13	0.5	226*	0.391*	
		23		12	0.5			
		23		11	0.3	185*		
Verbal working memory	Digit-span test*	23		12	0.5		0.077*	
		23		12	0.5			
Perception	Emotional perception	FEFA	22		42.5	3.3	-0.6	0.524
			22		43.1	2.8		

Groups did not differ regarding education according to the International Standard of Education or handedness according to Edinburgh Handedness Inventory. For test descriptions and references see Table S1.

Table 5. Correlation of neuropsychological test results – performed only for those tests that indicated group differences, see Table 4 – to ataxia severity, depressive symptoms, and age, using Spearman's rho or Pearson's *r* as indicated with results.

Domain	Test acronym	Zero-order correlations SCA-PRKCG/control			Partial correlations SCA-PRKCG Ataxia (SARA) controlled for age
		Ataxia (SARA)	Depression symptoms (HADS-D)	Age	
Attention	TAP Flexibility	$\rho = 0.43$ $P = 0.044$	$\rho = 0.01$ $P = 0.950$	$r = 0.41$ $P = 0.060$	$\rho = 0.24$ $P = 0.298$
		–	$\rho = 0.11$ $P = 0.639$	$r = 0.58$ $P = 0.004$	
	TAP Go/NoGo	$\rho = 0.33$ $P = 0.133$	$\rho = -0.15$ $P = 0.509$	$r = 0.64$ $P = 0.001$	–
Executive functioning	TAP Alertness	$\rho = 0.20$ $P = 0.380$	$\rho = 0.18$ $P = 0.416$	$r = 0.64$ $P = 0.001$	–
		–	$\rho = 0.38$ $P = 0.013$	$\rho = 0.52$ $P = 0.013$	–
	COMO	$\rho = 0.33$ $P = 0.129$	$\rho = 0.08$ $P = 0.710$	$\rho = 0.38$ $P = 0.071$	–
LPS 50 + substest 7		$\rho = 0.33$ $P = 0.129$	$\rho = 0.03$ $P = 0.909$	$\rho = 0.34$ $P = 0.110$	–
		–	$\rho = 0.04$ $P = 0.849$	$\rho = 0.34$ $P = 0.110$	–
	LPS 50 + substest 7	$\rho = 0.55$ $P = 0.008$	$\rho = 0.21$ $P = 0.358$	$r = -0.46$ $P = 0.026$	$\rho = -0.21$ $P = 0.351$
	–	$\rho = 0.11$ $P = 0.619$	$r = -0.46$ $P = 0.032$		

been described⁴⁴ For dystonia, an ataxia to dystonia continuum was suggested from animal models⁴⁵ related to the irregularity of excitatory outflow from deep cerebellar nuclei impacting on cerebral cortical functions.^{46,47} Consistently, a previous SCA-PRKCG series³⁶ revealed changes in intracortical inhibition similar to those reported in cortical myoclonus or DYT-TOR1A/DYT1 carriers and clinical signs of reduced interhemispheric motor inhibition (contralateral movement test, Table 4) were observed in our study. In sum, nonataxia movement disorders in SCA-PRKCG may be related to a distinct cerebellar pathology that SCA-PRKCG possibly shares with other movement disorders: a pure Purkinje cell dysfunction/loss in coincidence with structurally intact but disinhibited deep cerebellar nuclei. This pattern is in line with a recent histopathological report of SCA-PRKCG (p. H101Q).²¹

Aside from hyperreflexia, there were no other signs of pyramidal affection and motor-evoked potentials were normal as in all previous reports (one previous report of abnormal central motor conduction times⁴⁸ was found unremarkable later-on, personal communication D. T.). The etiology of muscle atrophy/pareses, fasciculations, mild sensory symptoms, and pain remains unclear. Electrophysiological findings here and in other studies do not support large-fiber neuropathy as a feature of SCA-PRKCG (severe axonal neuropathy has hitherto been described in only one singular index case (p. A458T)¹⁴). PRKCG expression in dorsal horn and nucleus gracilis⁴⁹ may be of relevance and requires further investigations of spinal structures in SCA-PRKCG.

The results of neuropsychological testing were compatible with previously described cognitive features of cerebellar pathology.⁵⁰ Longitudinal data of neuropsychological testing available in one of our subjects indicated mild progression of cognitive dysfunction, in line with few

previous descriptions.^{7,51,52} Few reports of overt dementia were all in SCA-PRKCG with long-standing disease (Table S1) or probable comorbidity. In one report, marked cognitive decline coincided with hearing loss, diabetes, and epilepsy, suggestive of other pathology.⁵³ A role of (physiologically weak) neocortical expression of mutant PRKCG is not excluded, but dementia of rather subcortical type, normal structural MRI, MR spectroscopy,²⁷ and histopathology of cerebral cortex^{20,21} argue against it.

Standardized structural brain MRI confirmed pure cerebellar atrophy, predominantly of vermis and anterior lobe. This may precede clinical manifestation and disclose carrier status in premanifest stages as in other SCAs.⁵⁴ Atrophy was nonprogressive in serial MRI of three cases spanning up to 17 years and may thus be interpreted as a maldevelopmental or early degenerative change that occurs independent of the manifestation or progression of ataxia. Of note, cerebellar (cortical) atrophy in absence of ataxia has been reported in other movement disorders.^{55,56} The clinical manifestation of SCA-PRKCG may thus be more related to dysfunctional cerebellar signaling than to cerebellar structural change, whereas the early developmental or even congenital cerebellar atrophy/hypoplasia may explain early nonprogressive subtle clinical signs. Both hypotheses await further exploration in longitudinal, histopathological, and functional studies.

The finding of symmetrically T2 hyperintense/T1 hypointense dentate nuclei was consistently seen in all 25 SCA-PRKCG cases irrespective of time since onset. We were unable to relate this finding to previous reports, as these displayed only sagittal view images. This sign was not seen in any of our healthy controls in whom a physiological decrease of T2 signal in the dentate nucleus is expected throughout the lifespan.⁵⁷ Brain T2 hyperintensity with corresponding T1 hypointensity has been proposed to indicate myelin degradation.⁵⁸ In SCA-PRKCG,

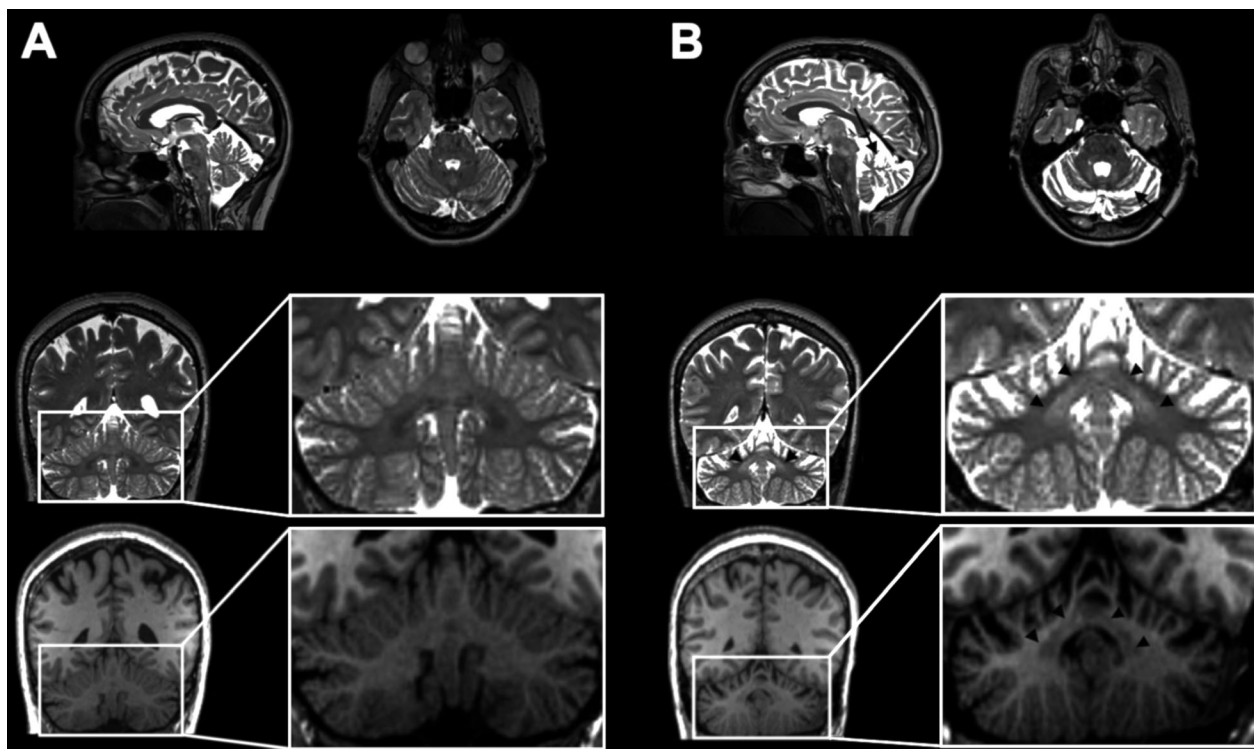


Figure 2. Example of MRI findings, specifically T2 signal of the dentate nuclei, in (A) a healthy subject aged 54 years and (B) a subject with SCA-PRKCG aged 53 years with 23 years since disease onset. As seen in all confirmed SCA-PRKCG of our cohort, there was a peculiar homogeneous T2 hyperintensity of the dentate nucleus (arrow head coronar image) that was not seen in any of the age- and gender-matched control subjects. This hyperintensity had a hypointense correlate in T1-weighted images. Further, as in all SCA-PRKCG, the patient featured cerebellar atrophy most pronounced in upper vermis (arrow sagittal image) and anterior lobe (arrow axial image).

the extension of this signal along the superior cerebellar peduncles suggests that this may also affect the dentato-thalamic efferents. The T2 hyperintense dentate sign was not observed in two VUS carriers who showed instead clinical and imaging features not seen in any confirmed SCA-PRKCG case. However, it was present in two related carriers of a benign variant (p. C69C) and in one VUS carrier (p. I173S). As all three shared a phenotype compatible with SCA-PRKCG this would rather support pathogenicity in the latter and should stimulate further (e.g., intronic) genetic investigation of PRKCG in the other family. Specificity and histopathological correlates of this novel sign are yet unknown. There have been reports of altered dentate signals in T2 weighted or FLAIR sequences in different movement disorders, usually as part of a more widespread pattern of imaging abnormalities.⁵⁹ Signal alterations confined to the dentate nucleus have recently been described in few genetic (ataxia) movement disorders^{60–62} but have not been systematically investigated and T2 hyperintensity of the dentate nucleus is currently not considered a characteristic imaging finding in neurodegenerative ataxia.^{63,64} Although it is possibly not specific, our results suggest the T2 hyperintense dentate

sign as a supporting criterion for PRKCG variant classification in cases with typical phenotype. Its absence, as well as the presence of atypical clinical findings (e.g., brainstem or pyramidal affection, early cognitive decline, retinal atrophy), or extracerebellar pathology on brain MRI may contribute to exclude PRKCG variants as causative. This should then stimulate further investigation into alternative causes or even genetic comorbidity.

The validity of this clinicogenetic description is strengthened by the use of standardized phenotype assessment applied in a prospective manner and a standardized refined procedure of variant classification. This is expected to reduce reporting bias for phenotypic features often seen with retrospective studies and to reduce misclassifications of pathogenicity. Some previous descriptions of SCA-PRKCG were published before the consensus guidelines on variant interpretation, the application of which, in fact, led to re-assignment as VUS in some (Table 1). All (likely) pathogenic variants in this study were within N-terminal or C1 regulatory domain. Conclusions on the (rarer) kinase domain mutations could thus only rely on literature review (Table S1) which did not convincingly reveal distinctive features.¹⁸ The

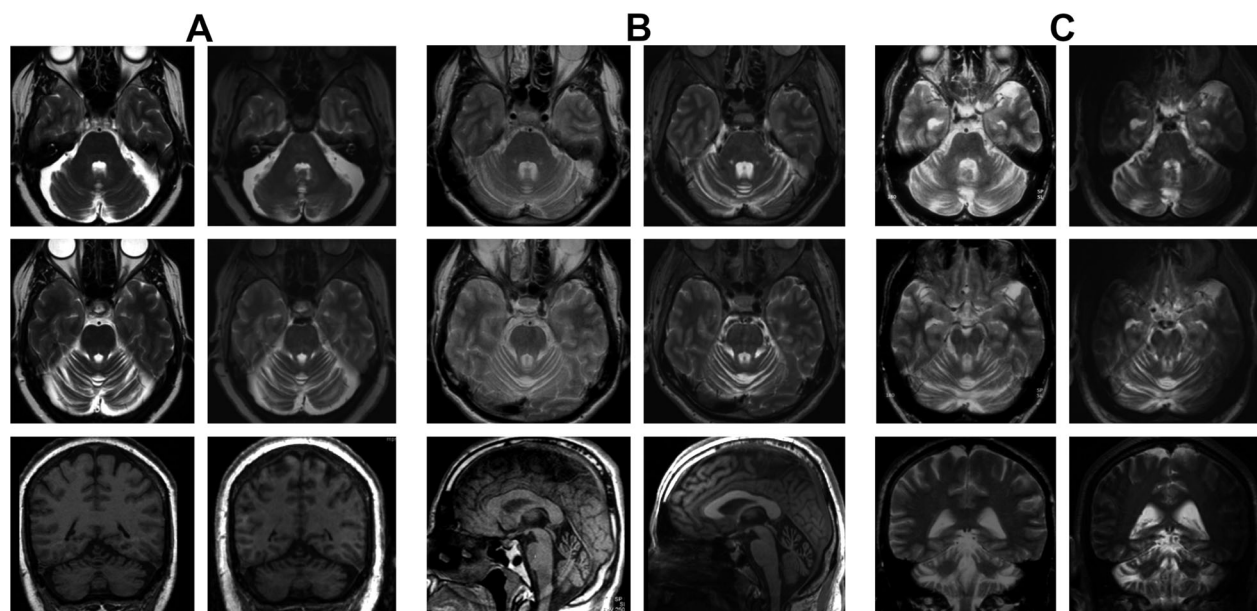


Figure 3. Evolution of MRI findings over eight (A), 12 (B) and 17 (C) years in three clinically manifest confirmed SCA-PRKCG subjects who made prior clinical MR imaging available (age at second scan 57 (A), 37 (B) and 56 (C) years). Most recent imaging is presented on the right hand columns, previous MRI presented on the left for each case. In case B, the first scan was obtained for other symptoms than ataxia, that is, in the premanifest stage.

comprehensive variant classification proposed here clearly increased diagnostic yield by inclusion of protein modeling results. Their interpretation weighs the structural and functional consequences of each variant on PRKCG function. Such a protein-specific approach can be considered more specific than generic pathogenicity prediction tools that are largely based on evolutionary conservation. Segregation analysis may have added certainty but we decided to systematically not consider such information, as its unavailability reflects the prevalent clinical reality. Clearly, most valid claims of pathogenicity would require functional study in a valid disease model, which has not yet been established for SCA-PRKCG. Thus, this refined classification approach may be generalizable to assign pathogenicity to missense variants in the case of other very rare, multi-allelic adult-onset disorders, in a gene with low tolerance to variability and in the absence of reliable biomarkers, functional models, or a highly specific phenotype. It should be noted, that the interpretation of both, genetic variants and protein modeling results, requires relevant expertise but would be feasible in the context of emerging research networks for rare diseases.

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Conflict of Interest

Dr. Schmitz-Hübsch reports honoraria from Biogen and Bayer AG outside the submitted work. Dr. Lux, Dr. Bauer, E. Schlapakow, Dr. Greschus, Dr. Scheel, Dr. Gärtner, Dr. Gras, Dr. Timmann, Dr. Synofzik, Dr. Giorgetti, Dr. Carloni, Dr. Shah, Dr. Schöls, Dr. Kopp, Dr. Bußenius, T. Oberwahrenbrock, Dr. Pfueller, Ms. Grosch, Dr. Amunts, Dr. Doss, M. Rönnefarth, and Dr. Minnerop have nothing to disclose. Dr. Brandt is cofounder and shareholder of medical technology companies Motognosis GmbH, Germany, and Nocturne GmbH, Germany, outside the submitted work. Dr. Kirlangic reports a position at the Gegenbauer Services GmbH, and a patent DE102016214575 with Volkswagen Aktiengesellschaft, outside the submitted work. Dr. Zimmermann reports grants from Novartis, outside the submitted work. E. Kadas is cofounder of Nocturne GmbH, Germany,

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Author Contributions

T. S-H. contributed to the conception and design of the study, acquisition and analysis of data, and drafting the manuscript and tables. S. L. contributed to the conception and design of the study (especially neuropsychology and imaging), analysis of data, and drafting parts of the manuscript and figures. P. B. contributed to study design, provided genetic analysis and interpretation, and co-drafted the manuscript and tables. A. U. B. contributed to the conception and design of the study, acquisition, and analysis of data and revising the manuscript for intellectual content. E. S. contributed to the conception of the study, acquisition of data, and revising the manuscript. S. G. provided acquisition and interpretation of imaging data, co-drafted the manuscript and figures. M. Sch. contributed to the acquisition and analysis of imaging data and revised the manuscript. H. G. contributed to the acquisition and analysis of data and revised the manuscript. M. E. K. contributed to the acquisition of data (particularly motor function and brain imaging), and editing and revising the manuscript for intellectual content. V. G. contributed to the conception and design of the MRI protocol, supported the acquisition of data, and revised the manuscript. D. T. contributed to the interpretation of clinical data and MRI data, and revised the manuscript for intellectual content. M. Sy. contributed to the discussion of the study concept, contributed to the interpretation of clinical and genetic findings, and revised the manuscript for intellectual content. A. G. contributed to the conception and design of the study, calculation, and analysis of protein modeling, and drafting the manuscript. P. C. contributed to the discussion of the study concept and revised the manuscript for intellectual content. N. J. S. contributed to the conception and design of the MRI protocol; revision of the manuscript. L. S. contributed to the discussion of the study concept, contributed to the interpretation of clinical and genetic findings, and revised

the manuscript for intellectual content. U. K. contributed to the discussion of study concept, to data acquisition and analysis (especially neuropsychology) and revised manuscript for intellectual content. L. B. contributed to the data analysis of neuropsychological statistics and drafting part of the figures and tables. T. O. contributed to data acquisition and interpretation (specifically the assessment of afferent visual pathway), revised manuscript for intellectual content. H. Z. contributed to data acquisition and interpretation (specifically assessment of afferent visual pathway), revised manuscript for intellectual content. C. P. contributed to coordinate study visits, to data acquisition and interpretation, and revised the manuscript for intellectual content. E. M. K. contributed to interpretation (specifically assessment of afferent visual pathway), and revised the manuscript for intellectual content. M. R. contributed to the acquisition and analysis of patients' and clinical data as well as revising the manuscript. A. S. G. contributed to the acquisition and analysis of data as well as revising the manuscript. M. E. involved in the discussion of study concept and revision of the manuscript for intellectual content. K. A. contributed to the conception and design of the study and revision of the manuscript for intellectual content. F. P. contributed to the conception and design of the study and revision of the manuscript for intellectual content. S. D. contributed to the conception and design of the study, acquisition and analysis of data, and revising the manuscript draft for intellectual content. M. M. contributed to the conception and design of the study, acquisition and analysis of data, and drafting the manuscript and figures.

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Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Table S1. Content summary and full reference list of all available published clinicogenetic descriptions of SCA-PRKCG ordered by location of variant from lower to higher number of amino acid residue.

Data S1. Details on methods (genetic classification, protein modeling, assessment protocols and test references (Table S2)) and additional results (Fig. S1: effects of age on cognitive test results).

Table S2. Description and reference of neuropsychological tests and screening instruments applied in this study.