

SYNEI ataxia is a common recessive ataxia with major non-cerebellar features: a large multi-centre study

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Mutations in the synaptic nuclear envelope protein 1 (SYNE1) gene have been reported to cause a relatively pure, slowly progressive cerebellar recessive ataxia mostly identified in Quebec, Canada. Combining next-generation sequencing techniques and deep-phenotyping (clinics, magnetic resonance imaging, positron emission tomography, muscle histology), we here established the frequency, phenotypic spectrum and genetic spectrum of SYNE1 in a screening of 434 non-Canadian index patients from seven centres across Europe. Patients were screened by whole-exome sequencing or targeted panel sequencing, yielding 23 unrelated families with recessive truncating SYNE1 mutations (23/434 = 5.3%). In these families, 35 different mutations were identified, 34 of them not previously linked to human disease. While only 5/26 patients (19%) showed the classical SYNE1 phenotype of mildly progressive pure cerebellar ataxia, 21/26 (81%) exhibited additional complicating features, including motor neuron features in 15/26 (58%). In three patients, respiratory dysfunction was part of an early-onset multisystemic neuromuscular phenotype with mental retardation, leading to premature death at age 36 years in one of them. Positron emission tomography imaging confirmed hypometabolism in extra-cerebellar regions such as the brainstem. Muscle biopsy reliably showed severely reduced or absent SYNE1 staining, indicating its potential use as a non-genetic indicator for underlying SYNE1 mutations. Our findings, which present the largest systematic series of SYNE1 patients and mutations outside Canada, revise the view that SYNE1 ataxia causes mainly a relatively pure cerebellar recessive ataxia and that it is largely limited to Quebec. Instead, complex phenotypes with a wide range of extra-cerebellar neurological and nonneurological dysfunctions are frequent, including in particular motor neuron and brainstem dysfunction. The disease course in this multisystemic neurodegenerative disease can be fatal, including premature death due to respiratory dysfunction. With a relative frequency of ~5%, SYNE1 is one of the more common recessive ataxias worldwide.

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Abbreviations: ARCA = autosomal recessive cerebellar ataxia; EDMD4 = Emery-Dreifuss muscular dystrophy type 4; FDG = fluorodeoxyglucose

Introduction

SYNE1 (OMIM 608441) is one of the largest genes in the human genome, with the longest isoform comprising 146 exons that encode the 8797 amino-acid synaptic nuclear envelope protein 1 (Gros-Louis et al., 2007). This protein, also known as Nesprin 1 (Nuclear envelope spectrin 1), is part of the spectrin family of structural proteins that share a common function of linking the plasma membrane to the actin cytoskeleton (Gros-Louis et al., 2007). Truncating recessive mutations in SYNE1 have been reported to cause a slowly progressive, relatively pure cerebellar ataxia with only few extra-cerebellar symptoms (spinocerebellar ataxia, autosomal-recessive 8; SCAR8/autosomal-recessive cerebellar ataxia type 1, ARCA1) (Dupre et al., 1993/ 2012, 2007; Gros-Louis et al., 2007; Noreau et al., 2013; Fogel et al., 2014). So far, this ataxia has been mainly observed in Quebec, Canada: while it presents the third most common hereditary ataxia in Quebec [after (autosomal recessive spastic ataxia Charlevoix-Saguenay) and Friedreich ataxia] (Dupre et al., 1993/2012, Gros-Louis et al., 2007), only few families have been identified outside French-Canadian populations

so far (Izumi et al., 2013; Noreau et al., 2013; Hamza et al., 2015). This view on the phenotype and geographic distribution of SYNE1 ataxia, however, might be rather preliminary, given that systematic screenings outside this founder population were confined to highly selected individual cases (Izumi et al., 2013; Noreau et al., 2013). A systematic characterization of the clinico-genetic spectrum of a larger non-French-Canadian subject group is missing. We here hypothesized that (i) SYNE1 is a frequent cause of recessive ataxia also outside French-Canadian populations, given the large size of the SYNE1 gene; and (ii) dysfunction in extra-cerebellar systems is not the exception but the rule. To test these hypotheses, we aggregated the genetic and phenotypic findings of a screening of 434 ataxia patients compiled by seven different European centres, unravelling 23 novel index patients with truncating SYNE1 mutations, including 34 mutations not previously linked to human disease. This large collection of SYNE1 patients demonstrates that SYNE1 ataxia is a common recessive ataxia also outside the French-Canadian founder population, and that it commonly presents with multisystemic neurodegenerative disease. This includes in particular motor neuron and brainstem features and even complex neuromuscular

syndromes, where respiratory dysfunction can lead to premature death.

Materials and methods

Patients

Four hundred and thirty-four index subjects with unexplained early-onset degenerative ataxia (age of onset <40 years) compatible with autosomal recessive inheritance (no ataxia in the parental generation) and negative for trinucleotide repeat expansions causing Friedreich's ataxia (FRDA) were compiled from seven different European ataxia centres (n = 116 Tübingen, Germany; n = 16 Essen, Germany; n = 13 Munich, Germany; n = 109 Milano, Italy; n = 139 Strasbourg and Paris, France; n = 41, Antwerp, Belgium). These patients originated from 36 different countries (Supplementary material).

Genetic screening using next-generation sequencing

Subjects were screened for *SYNE1* mutations by one of the following five next-generation sequencing methods: (i) as part of a high coverage HaloPlex gene panel kit (Agilent), which included >120 known ataxia genes (n = 179); (ii) as part of a targeted exon-capture sequencing strategy (Agilent SureSelect kit), which included 57 known ataxia genes (n = 139); (iii) as part of an amplicon-based customized panel (Illumina TrueSeq Custom Amplicon, TSCA) covering 76 known ataxia genes (n = 21); (iv) as part of a targeted exon-capture sequencing strategy (Illumina Nextera Rapid Capture Custom kit), which included 107 known ataxia genes (n = 54); or (v) as part of whole-exome sequencing using the SureSelect Human All Exon 50Mb kit (Agilent) (n = 41). Further details of these five methods can be found in the Supplementary material.

Variants were filtered for (i) non-synonymous homozygous or compound heterozygous truncating mutations in *SYNE1* (frameshift insertions or deletions, splice mutations, and stopgain mutations); that were (ii) absent or extremely rare (minor allele frequency <0.5%) in the public databases GEM.app (5992 exomes from 4279 families) (Gonzalez *et al.*, 2013, 2015), dbSNP137, NHLBI ESP6500, 1000Genomes project, and ExAc (60706 exomes; Exome Aggregation Consortium; http://exac.broadinstitute.org, accessed 06/2015) (for further details on the next-generation sequencing bioinformatics and filter criteria, see Supplementary material). Ninety to ninety-five per cent of the coding exons of *SYNE1* have been covered with > 20 reads by all five test formats.

In addition, to screen for *SYNE1* exon deletions/duplications, we performed a preliminary copy number variation analysis on our panel–sequencing datasets from 229 patients of the ataxia screening cohort (Supplementary material).

Inclusion of SYNE1 missense variants

Due to the high number of private missense variants in the large SYNE1 gene (often located in the numerous, modestly conserved spectrin repeats of the gene) and given the fact that only truncating SYNE1 variants have been established as a

cause of SYNE1 ataxia, we did not include cases carrying only SYNE1 missense variants. Following a conservative approach towards SYNE1 missense variants, subjects carrying a missense variant were only included if the respective missense variant (i) segregated in trans with a truncating SYNE1 variant; (ii) was absent or rare in public exome databases (same criteria as above); (iii) located in highly conserved positions of the N-terminal actin-binding domain (codon 1-289); and (iv) predicted to be damaging by at least two out of three in silico algorithms [Mutation Taster (Schwarz et al., 2010; Wang et al., 2010); SIFT (Sim et al., 2012); and PolyPhen-2 (Adzhubei et al., 2010)]. These criteria lead to the inclusion of one SYNE1 missense mutation (p.F220S) in the final cohort. The mutation p.D5140G (Family 7), which presented as a missense mutation on the genomic level, was shown to result in aberrant splicing of SYNE1 transcript, thus representing de facto a null mutation.

Burden analysis of SYNEI missense variants

To further explore the significance of SYNE1 missense variants in ataxia, we tested whether rare missense SYNE1 variants predicted to be damaging were more frequent in cases with ataxia than in controls. Cases comprised of a consecutive series of n = 96 whole exome datasets from index patients with early-onset ataxia (same inclusion criteria as above). Controls comprised of a consecutive series of n = 250 whole exome datasets from index subjects with early-onset Alzheimer's disease, a condition that is not part of the extended phenotypic spectrum of SYNE1 demonstrated here (Supplementary material).

Messenger RNA analysis

Messenger RNA analysis was performed for selected mutations (mutations of Families 6 and 7) to confirm the effect of the cryptic splice mutation identified in this study (Family 7), and to exemplarily confirm the loss-of-function mechanism of truncating SYNE1 mutations through nonsense-mediated decay of mutant mRNA (Family 6). Total RNA was extracted from lymphocytes using a Maxwell® Extractor (Promega) with the Maxwell® 16 LEV simply RNA Cells kit. Complementary DNA synthesis was carried out using Transcriptor FirstStrand cDNA synthesis kit (Roche) with random primers. PCR was carried out in 20 µl with an annealing temperature of 58 °C and specific primer pairs designed to amplify cDNA fragments encompassing the different mutations of Families 6 and 7. The amplified fragments were directly sequenced on an automated sequencer (ABI Prism 3100 Genetic Analyzer, Applied Biosystems) using the BigDye® Deoxy Terminator Cycle Sequencing Kit (Applied Biosystems) according to the manufacturer's recommendations.

Histological analysis and immunohistochemistry of muscle tissue

Biopsies of the quadriceps muscle were available from three *SYNE1* subjects and compared to a healthy age-matched control. The samples were analysed by classical histological, histochemical, histoenzymatic and immunohistochemical

techniques, as described previously (Martin *et al.*, 1997). The antibodies used were mouse monoclonal anti-emerin (NCL-emerin, 1/100, acetone fixation, Novocastra); mouse monoclonal anti-lamin A/C (NCL-LAM-A/C, 1/1000, aceton/methanol fixation, Novocastra) and rabbit polyclonal anti-nesprin-1 (PRB-439P, 1/2000, unfixated, Covance). The avidin-biotin complex technique for the monoclonal antibodies and the peroxidase-antiperoxidase technique were used for the polyclonal antibody.

Clinical and electrophysiological assessments

All index patients carrying two pathogenic *SYNE1* alleles (either in homozygous or compound heterozygous state) as well as their affected siblings received a systematic clinical assessment for disturbances in multiple neurological systems (Table 2) by a movement disorder specialist. Nerve conduction studies and EMG investigations were performed in all patients available for these investigations. Patients were classified into two phenotypic categories: (i) pure cerebellar ataxia (only ataxia features without any evidence of non-ataxia features); or (ii) cerebellar ataxia plus (ataxia plus clinical evidence for damage of at least one additional neurological system, such as e.g. upper and/or lower motor neuron disease). Disease severity was rated by the Scale for the Assessment and Rating of Ataxia (SARA) (Schmitz-Hubsch *et al.*, 2006).

MRI and PET imaging

Routine brain MRI including T₁-, T₂-, diffusion-weighted images and fluid attenuated inversion recovery T2 (FLAIR) images were performed for at least one patient per family and were reviewed by both neuroradiologists and a neurologist. ¹⁸F-fluorodeoxyglucose (FDG)-PET imaging was performed for two exemplary subjects with ataxia-motor neuron disease on a Siemens ECAT HR + scanner (CTI) with a 128 × 128 voxel matrix (voxel size 2 mm), an axial field of view of 15.5 mm and a full-width at half-maximal resolution of 5 mm, according to previously published tracer-specific protocols (Forster et al., 2010). Resting-state PET recordings were obtained in the interval 30-50 min after intravenous application of 200 MBq 18-Fluoro-2-deoxyglucose (18F-FDG) in fasting state (>6h), a brief attenuation scan was obtained with integrated ⁶⁸Ge rod sources (for details of image analysis, see Supplementary material).

Results

A large cohort of novel SYNEI index families and mutations from European and non-European populations

We identified 22 index patients carrying two truncating SYNE1 alleles and one index patient carrying one truncating plus one missense SYNE1 allele, thus yielding a total of 23 index patients out of 434 early-onset ataxia patients

(5.3%). Five index patients were from multiplex families, the remaining 18 index patients were simplex cases. Six index patients had consanguineous parents. In the 23 index patients, we observed a total of 35 different mutations, consisting of seven frameshift, 20 nonsense mutations, six nucleotide changes affecting constitutive splice sites, one nucleotide change generating a cryptic exonic splice site, and one missense mutation in the actin binding domain (Table 1; for a discussion of the cryptic exonic splice site and the missense mutation, see below). The mutations were spread across the giant SYNE1 gene, including the acting binding domain (also called calponin homology domain containing actin binding site) and the spectrin repeat domains (Fig. 1A). Five mutations (mutations 1-5) also affect the coding sequence of the SYNE1 isoform Nesprin 1α (nucleotide reference: http://www.ncbi.nlm.nih. gov/nucleotide/119120815), a region where no mutations linked to ataxia have been previously found (Razafsky and Hodzic, 2015). Thirty-four of thirty-five mutations have not reported in association with human disease before, thus more than doubling the total amount of all SYNE1 truncating mutations that have been published so far (for overview of all novel and published mutations, see Fig. 1B). Only the variant c.3736G > T, p.E1246* (mutation 29, Table 1) has been reported before (Hamza et al., 2015). Two mutations were listed in dbSNP/1000 Genomes database, but also not yet linked with human disease. All 35 mutations were absent or had extremely low minor allele frequency (<0.001%) in GEM.app (5992 exomes from 4279 families), dbSNP137, 1000 Genomes database, NHLBI ESP (6500 exomes), ExAc (60706 exomes) (Table 1). Moreover, all 35 mutations yielded a scaled CADD score (combined annotation dependent depletion) of ≥ 19 (Table 1), ranking the predicted pathogenicity of each mutation among the top 1% of all 8.6 billion single nucleotide variants in the GRCh37/hg19 (Kircher et al., 2014). All mutations were confirmed by Sanger sequencing. In all families where DNA from the siblings was available (10/ 23), the variants co-segregated with disease in affected siblings, and unaffected siblings carried at most one pathogenic allele. For all families where DNA of at least one parent was available (12/23 families), we were able to show that the respective parent carried only one of the respective two SYNE1 variants, indicating a biallelic localization of the variants in the index child. In four of the remaining 11 families, consanguinity was also suggestive of a biallelic location of the observed homozygous mutations. No other significant variants in known ataxia genes were identified in any of the 23 index subjects, as demonstrated by screening through whole exome sequencing and targeted panel sequencing, respectively.

No obvious *SYNE1* multi-exon deletions/duplications were observed by our preliminary next generation sequencing-based copy number variation analysis (Supplementary material). This result, however, has to be interpreted with caution as no consensus currently exists on a method to perform copy number variation analysis on next generation sequencing data.

Table | SYNE! mutations identified in this study

	8#				cnange					Genomes rs# dbsnp			
	1	CA plus	chr6:152456368C>T	c.25516-1G>A	-KA-	Splicing	Het	7.62	29.1	0	0	0	Absent
	#I8	pCA	chr6:152473139C>T	c.24054G > A	p.W8018*	Stopgain SNV	Het	7.625	19	0	0	0	Absent
	#	CA plus	chr6:152473168G > A	c.24025C>T	p.Q8009*	Stopgain SNV	Het	5.634	09	0	0	0	Absent
	#5, #23	CA plus	chr6:152476161G>A	c.23782C>T	p.R7928*	Stopgain SNV	Het	1.316	29	0 rs570 916267	0	0	Absent
2	#7	PCA	chr6:152477125insCAC TCGGCATCAGTGGC	c.23684_23685insACG CCTGTGCCACT	p.C7895*	Frameshift duplication	Het	Ϋ́ Ϋ́	24	0	0	0	Absent
	¢	4 V	ACAGGCGT	GATGCCGAGTG	*7414	stopgain Ctopgain CNIV	į	0 153	α	c	c	c	Abcont
	7 0	spld V	chrc.1525.12077 C / 1	T/027512.5	P.17.178	Stopgain SNIV	j ;	0000	2 4	o c	o c	o c	Absont
	, ;	spid <	CII 0:1323428888	C.21318C / 1	p:(/)106	Stopgalli SIAV	ם בר	0.000	8 =	o c	o c	o c	Absent
	71#	CA pius	Chr6:152531729G > A	C.20935C > 1	p.K63/7	Stopgain SINV	Пет	1.3/1	+ ,	-	0 0	> 0	Absent
	#14	CA plus	chr6:1525550356 > A	C. 20380C > 1	p.C6/94	Stopgain SINV	Het	7.4/6	9 6	> (o (> (Absent
	#23	CA plus	chr6:152557241C>G	c.G20183 + IG>C	-NA-	Splicing	Het	7.808	33	0	0	0	Absent
	=	CA plus	chr6:152558041G>A	c.19897C>T	p.Q6633*	Stopgain SNV	Homo	7.646	29	0	0	0	Absent
	01#	CA plus	chr6:152558084C > G	c.19855-1G > C	-NA-	Splicing	Ношо	7.304	28.1	0	0	0	Absent
l3 [‡]	61#	CA plus	chr6:152605163G>A	c.I7944C>T	p.R5982*	Stopgain SNV	Homo	1.582	20	0	0	0	Absent
	#14	CA plus	chr6:152615251insT	c.17480dup	p.Y5827*	Frameshift	Het	Y-	38	0	0	0	Absent
						insertion stopgain							
± 12	81#	PCA	chr6:152639275- 152639281	c.16294_16300delA AACTGA	p.K5432Lfs*25	Frameshift deletion	Het	Ϋ́ Ϋ́	40	0	0	0	Absent
	#3	PCA	chr6:152642966G > A	c.15760C>T	p.R5254*	Stopgain SNV	Homo	5.373	53	0	0	0	Absent
[‡] ∠1	#	CA plus	chr6:152644651-	c.15665_15666deITC	p.L5222Hfs*21	Frameshift deletion	Het	₹ Z-	35	0	0	0	Absent
8	47	A	52644652delGA chr6- 52646244T > C	C 15419A > G	D5140Gfc1*	(Cryptic) splicing	Het	3 931	19.05	C	87 × 10 ⁻⁶	0 9	Ahsent
	#2	CA plus	chr6:152651038G > A	c.14569C>T	p.O4857*	Stopgain SNV	Het	7.928	51	0	0	0	Absent
	#2	CA plus	chr6:152651345insA	c 14261 dun	p. C. 4755Profe * 15	Frameshift insertion		A Z	. %		o C		Absent
2 -	. 4	CA plus	chr6:152651352-	c. 14255delG	p. 64757Ffc*10	Frameshift deletion		ξ Ł	36	0 0	o C	· c	Absent
	2	35	152651352delC		2.0		-		2	,	>	>	
22 #	#	PCA	chr6:152652521-	c.13086delC	p.H4362Qfs*2	Frameshift deletion	Homo	Ł Z	24.8	0	0	0	Absent
	#15	CA plus	chr6:152657975C>T	c.12315 + 1G>A	₹ Z-	Splicing	Homo	7.499	26.2	0	0	0	Absent
	6#	CA plus	chr6:152665320G > A	c.11908C>T	p.R3970*	Stopgain SNV	Het	6.313	51	0	0	0	Absent
	#2	CA plus	chr6:152706854insAT	c.8627_8628insAT	p.M2876lfs*19	Frameshift insertion	Het	Ϋ́	34	0	0	0	Absent
	8#	CA plus	chr6:152711589T > C	c.8026-2A > G	-k V-	Splicing	Het	7.542	25.6	0	0	0	Absent
	91#	PCA	chr6:152722345C>T	c.6978G > A	p.W2326*	Stopgain SNV	Homo	5.118	20.3	0	0	0	Absent
	#21	CA plus	chr6:152737559G > A	c.6034C > T	p.R2012*	Stopgain SNV	Het	0.595	32	0.0002	8.2×10^{-6}	0 ,	Absent
	#17	CA plus	chr6:152765668C > A	c.3736G>T	p.E1246*	Stopgain SNV	Homo	5.495	39	0	0	0	Absent
30 #	#20	CA plus	chr6:152776698T > A	c.2776A > T	p.K926*	Stopgain SNV	Het	3.905	36	0	0	0	Absent
	#22	CA plus	chr6:152826408G > A	c.727C>T	p.R243*	Stopgain SNV	Homo	0.771	35	0	0	0	Absent
32 ≠	#I3	CA plus	chr6:152826453G > A	c.682C>T	p.R228*	Stopgain SNV	Homo	4.534	36	0	8.2×10^{-6}	0 ,	Absent
	#20	CA plus	chr6:152826476A > G	c.659T > C	p.F220S	Missense	Het	9.339	28.8	0	0	0	Absent
34 #	9#	CA plus	chr6:152832174A>T	c.395T > A	p.L132*	Stopgain SNV	Het	9.14	38	0	1.6×10^{-5}	2 0	Absent
	#2	CA plus	chr6:152841593C>T	c.309 + IG>A	-NA-	Splicing	Het	7.187	27.8	0	0	0	Absent

Genomic positions of the variants according to genome build hg19. DNA changes according to NM_033071.3. Variant type and protein changes according to GVS function based on NP_14962. pCA = pure cerebellar ataxia; CA plus = cerebellar ataxia; CA plus = PhyloP conservation score based on base-wise conservation across 100 vertebrates; CADD score = scaled Combined Annotation Dependent Depletion score, integrating many diverse annotations into a single measure (C score) for each variant. The predicted pathogenicity of each variant is scored and ranked relative to all ~8.6 billion single nucleotide variants of the GRCh37/hg19 reference. A scaled CADD score of 20 indicates variants at the top 1%, a CADD score of 30 indicates variants at the top 0.1%, etc. (Kircher et al., 2014). MAF = minor allele frequency; EXAC= Exome Aggregation Consortium; EVS = Exome Variant Server 6500 exomes all from the NHLBI GO Exome Sequencing Project; HGMD = Human Gene Mutation Database.

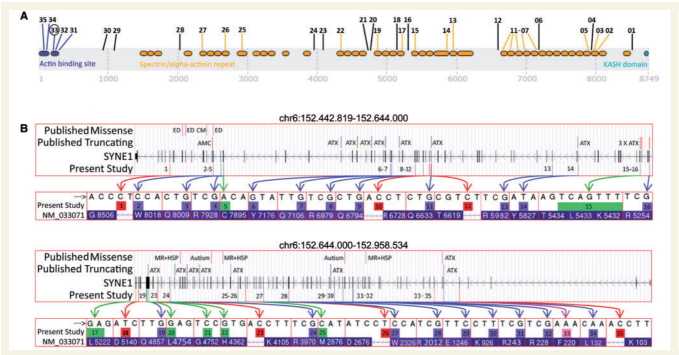


Figure 1 SYNE1 mutations. (A) Graphical overview of the mutations found in this study in relation to the SYNE1 domains. Numbers indicate the mutation IDs of the mutations identified in this study (Table 1). Their position indicates the position of the respective mutations in the SYNE1 gene. Blue = N-terminal actin binding domain [calponin homology domains containing actin binding sites (IPR001715)] and mutations affecting this domain; orange = spectrin/alpha-actinin repeat domains (IPR018159) and mutations affecting these domains; turquoise = KASH domain (C-terminal klarsicht domain) (IPR012315); black = mutations not affecting any of these domains. Mutation 33 (black circle) is the only missense mutation in the present study. (B) Overview of the variant types and their location of all published and novel SYNE1 mutations. The presentation of the giant SYNE1 gene is split in a first part (chr6:152.442.819–152.644.000; top) and a second part (chr6:152.644.000–152.958.534; bottom). It presents the variant types of all SYNE1 mutations found in the present study (bottom row of each panel) and other studies (Human Gene Mutation Database) (top row of each panel), their location and their annotation with the associated clinical phenotypes. ATX = ataxia;

AMC = arthrogryposis multiplex congenita; ED = Emery-Dreifuss muscular dystrophy; CM = cardiomyopathy; MR = mental retardation;

HSP = hereditary spastic paraplegia. Note that, except for ATX and AMC, all other phenotypes have been associated only with missense mutations, not truncating mutations in SYNE1. Green coloured arrows and boxes = indel mutations; blue coloured arrows and boxes = stop mutations; red coloured arrows and boxes = splice site mutations; purple coloured arrow and box = missense mutation. HG19 genome build. Transcript: NM_033071 > NP_149062.

Pathogenicity of selected base exchanges

Cryptic splice mutation

While six of seven splice mutations affected constitutive splice sites (Table 1), the base exchange c.15419A > G (mutation 18, Family 7)—which appeared as a p.D5140G missense change on the genomic level—was predicted by *in silico* analysis (NNSplice; Reese *et al.*, 1997) to act as a cryptic exonic splice mutation. Specifically, it was predicted to create a new donor site 26 bp upstream of the constitutive 5′ donor splice site of intron 80. To confirm this prediction, we performed reverse transcription and sequencing of *SYNE1* mRNA, revealing a heterozygous deletion of the last 26 nucleotides of exon 80 (r.15419_15444del26) (Fig. 2A), which would result in a protein truncated at residue 5140 (p.D5140Gfs1*). This variant occurred *in trans* with the frameshift duplication/ stop gain mutation p.C7895*.

Missense mutation

The missense variant c.659T > C, p.F220S (observed in Family 20): (i) segregated in trans with the nonsense variant c.2776A > T, p.K926*; (ii) was absent in all public databases described above; (iii) predicted to be damaging by three of three in silico algorithms [MutationTaster (Schwarz et al., 2010; Wang et al., 2010); SIFT (Sim et al., 2012), and PolyPhen-2 (Adzhubei et al., 2010)]; (iv) ranked among the top 1% of all 8.6 billion single nucleotide variants in the GRCh37/hg19 (CADD score: 28.8) (Kircher et al., 2014); and (v) located at a highly conserved position in the actin-binding domain of SYNE1 (Supplementary material). This domain is conserved even in non-metazoan species (paralogues in ichthyospora, fungi and slime mould, E-values in the range of 5×10^{-40} to 10^{-50}), whereas the spectrin domains of SYNE1 provide no significant alignment with paralogues of the same non-metazoan species (E-values > 10). The p.F220S mutation affects one of the six invariant amino

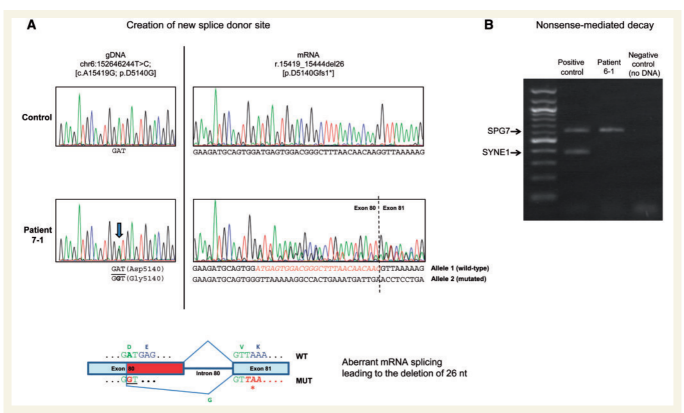


Figure 2 Exemplary mechanisms of SYNE1 mutations. (A) An exonic base exchange leads to activation of a cryptic splice site. (A) SYNE1 transcript analysis reveals that the c.15419A > G (p.D5140G) mutation (identified in Patient 7-1) creates a new splice donor site within exon 80 (GT, underlined in bottom panel) resulting in a 26-bp deletion (p.D5140Gfs1*). Top left: Sequence analysis of PCR products of amplified genomic DNA showing the A-to-G variant (minus strand) that changes codon 5140 from GAT (Asp) to GGT (Gly). Top right: Sequence analysis of amplification products following reverse transcriptase-PCR of SYNE1 transcript (exons 78–82) from patient's cells showing the presence of the 26-nt deletion (r.15419_15444del26). Bottom: Schematic diagram showing the aberrant splicing mechanism which leads to the premature truncation of the protein through the deletion of 26 nucleotides in exon 80 (in red). Premature TAA termination codon in exon 81 is shown in bold red italics. (B) Truncating SYNE1 mutations can lead to nonsense-mediated decay of SYNE1 mRNA. SYNE1 mRNA carrying the p.L132* and the p.G4752Efs*10 null mutations undergoes nonsense-mediated decay. PCR amplification from retro-transcribed mRNA was carried out with primers specific for SYNE1 (exons 4–9) and SPG7 (exons 6–10) transcripts. Note the complete absence of the SYNE1 mutant transcript in Patient 6-1 as compared to the control.

acid positions of the second calponin-homology domain in nesprin, spectrin and alpha-actinin homologues (Supplementary material). The biallelic truncating and the missense *SYNE1* variants were also found in the affected sibling of the index patient in Family 20, indicating segregation with disease.

Burden analysis for missense variants in SYNEI

Rare missense *SYNE1* variants were observed in 14/192 alleles (7.3%) from ataxia cases and in 28/500 alleles (5.6%) from disease controls (early-onset Alzheimer dementia) (Supplementary material). Also missense variants in the N-terminal actin-binding domain of *SYNE1* (codon 1–289) were not more frequent in ataxia cases than in controls (Supplementary material). This finding suggests that rare missense *SYNE1* variants seem to present a ubiquitous finding unrelated to specific disease conditions or phenotypes.

In general, these results from the cryptic splice and the missense mutation as well as from the missense variant burden analysis indicate that careful interpretation of single *SYNE1* nucleotide variants is necessary to evaluate possible effects detrimental to *SYNE1* function. Such a critical one-by-one debate of *SYNE1* single nucleotide variants is of particular relevance in the future, given the large size of *SYNE1* with its substantial variability.

Absence of SYNE1 transcript adds support for a loss-of-function mechanism

Truncation of proteins can lead to either toxic gain of function (exerted by the residual protein) or to loss-of-function. To exemplarily investigate the mechanism of action for truncating *SYNE1* mutations, we investigated the pathomechanism of the changes p.L132* and p.G4752Efs*10

(observed in Family 6). No *SYNE1* transcript was amplified by reverse transcriptase-polymerase chain reaction in patient's cells (Fig. 2B), suggesting that both the early nonsense mutation p.L132* and the later frameshift mutation p.G4752Efs*10 cause nonsense-mediated decay of mutant mRNA. The absence of the transcript adds support for a loss-of-function mechanism underlying the detrimental effect of truncating SYNE1 mutations in human SYNE1 disease, as hypothesized earlier (Gros-Louis *et al.*, 2007; Attali *et al.*, 2009).

Severely reduced to absent SYNEI staining in muscle tissue

Muscle biopsy [available for three different SYNE1 index patients (Patients 13-1, 10-1 and 6-1) originating from three different countries] showed neurogenic changes in all patients, thus corresponding to the clinical/EMG finding of frequent lower motor damage in SYNE1 disease (see below). Upon routine light microscopy and also immunohistochemistry with relevant antibodies, the nuclei had a normal peripheral distribution in the three patients comparable to the control individual (Fig. 3). More precisely, no myopathic and in particular no dystrophic muscle features were seen in any of the three patients, thus contrasting previous findings on dystrophic changes in patients with Emery-Dreifuss muscular dystrophy type 4 (EDMD4; MIM 612998) that were reported in association with heterozygous 'dominant' SYNE1 (and SYNE2) mutations (Zhang et al., 2007a).

In line with the suggested loss of protein in SYNE1 patients with truncating mutations, staining of SYNE1 was severely reduced in two patients and absent in one patient (Fig. 3). To control for the possibility that this lack of SYNE1 staining might be an unspecific result or artefact in the SYNE1 samples, control stainings were performed. Control immunolabelling showed normal results for emerin at the inner nuclear membrane and of lamin A/C also in the SYNE1 muscle samples (Fig. 3). No mislocalization of emerin or lamin A/C was seen, thus contrasting previous findings of a mislocalization of these proteins reported in patients with heterozygous SYNE1 mutations (Zhang et al., 2007a).

A broad range of additional non-cerebellar features, including severe and complex early-onset syndromes

We aggregated clinical data from 26 affected subjects belonging to 23 families (median age of disease onset: 22 years, range: 6–40 years; last examination in each individual performed after a median of 25 years of disease duration). Disease started with gait coordination disturbances in 24/26 subjects; in two subjects it began with focal upper limb dystonia (writer's cramp) (Patient 20-1) and lower limb weakness (Patient 23-1), respectively. Five of 26

patients (19%) showed the classical *SYNE1* phenotype of pure cerebellar ataxia that was mildly progressive and started in adulthood (range: 19–36 years) (pure cerebellar ataxia). However, the large majority of patients (21/26, 81%) exhibited at least one additional complicating feature from a large range of non-cerebellar features, both in neurological and non-neurological domains (cerebellar ataxia plus) (Table 2 and Fig. 4A and B).

The most frequent complicating finding was motor neuron dysfunction, observed in 15/26 (58%) of the total cohort (Fig. 4A). This comprised a combination of upper motor neuron dysfunction (bilateral positive extensor plantar reflex and/or spasticity) and lower motor neuron dysfunction (muscle atrophy combined with reduced reflexes; fasciculations clinically or on EMG; or chronic and/or acute neurogenic changes on EMG) in 5/26 patients (19%), only upper motor dysfunction in 8/26 patients (31%), and only lower mutation in 2/26 (8%) patients. Other frequent complicating features across the cohort included: scoliosis (5/26, 19%), sometimes combined with kyphosis, indicative of skeletal abnormalities in SYNE1; slowing of saccades (4/26, 15%), indicative of brainstem dysfunction in SYNE1; and reduced vibration sense of the lower limbs (3/26, 12%), indicative of peripheral nerve and/or dorsal column dysfunction in SYNE1 (for a relative frequency of all main features, see Fig. 4A). A broad range of non-cerebellar oculomotor deficits was seen in single patients across the cohort: square wave jerks, ophthalmoparesis, and strabism including esotropia (Table 2 and Fig. 4A). One patient exhibited mild macroglossia in addition to esotropia, that could not be related to any other disease other than SYNE1 (for image see Supplementary material).

Three cerebellar ataxia plus patients (3/26, 12%; Patients 13-1, 13-2 and 23-1; no consanguinity in either of the two families) showed a particularly severe and complex multisystemic phenotype, comprising very early onset (6–10 years of age) ataxia, spasticity, weakness and muscle wasting of all four limbs, mental retardation (IQ 49-60), dysphagia, pes cavus, and a broad range of variable skeletal and soft-tissue abnormalities such as sacral cysts, pseudarthrosis clavicula, hyperlaxity of joints, achilles tendon contractures, kyphosis, scoliosis, cataract, hypertelorism. All three developed respiratory dysfunction in late adolescence, necessitating BiPAP (bi-level positive airway pressure) ventilation in two of three subjects and leading to premature death at age 36 years in one of them. No obvious second hit in any other disease gene was detected in these patients (Supplementary material). Thus, the complex phenotype in these three subjects does not seem to be explained by a second hit in another gene or to present a qualitatively different condition; rather it seems to illustrate one end on the continuous spectrum of SYNE1 disease, which literally encompasses all of the aforementioned SYNE1associated domains and includes also mental retardation and respiratory distress (Fig. 4C).

One of the three subjects also showed developmental abnormalities of the visceral organs, such as malrotation

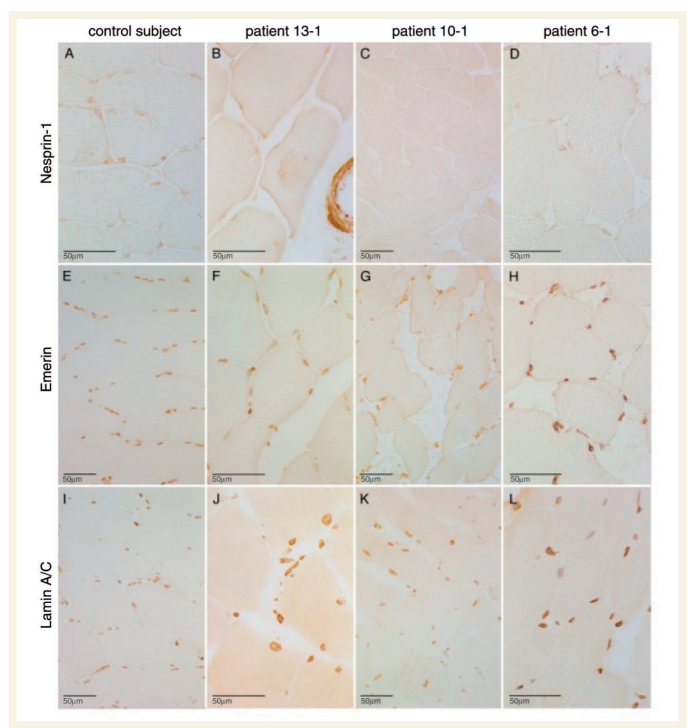


Figure 3 Severely reduced to absent SYNE1 staining in muscle tissue of SYNE1 patients. Immunohistological findings in control tissue (A, E and I), as well as in the quadriceps muscle biopsies of three different SYNE1 patients from three different countries: Patient 13-1, Belgian (B, F and J); Patient 10-1, German (C, G and K); and Patient 6-1, Italian (D, H and L). All three patients show a severely reduced to absent staining of the nuclear envelope after immunolabelling of nesprin-1 (B–D), whereas staining was normal in the control (A) (peroxidase-anti-peroxidase technique). To control for an unspecific lack of staining in the SYNE1 patients, further control stainings were performed. Immunolabelling of emerin and lamin A/C at the inner nuclear membrane was normal in all three patients as well as the control (E–H and I–L, respectively) (avidin-biotin complex technique).

of the colon and unilateral positioning of both kidneys at right pelvis. However, future studies in SYNE1 patients are warranted to confirm whether such visceral features are indeed part of its phenotypic spectrum.

Electrophysiological findings

Motor evoked potentials, available for eight patients, were prolonged or not evoked in six of eight patients (75%), all

Table 2 Clinical, imaging and electrophysiological features of SYNEI patients

Abnormal MEPS	n.d.	n.d.		7	<u>.</u>	n.d.	Abnormal LL		n.d.	n.d.	Abnormal UL + LL		Abnormal UL + LL		Abnormal LL					70	į				n.d.	n.d.		Abnormal	n.d.		n.d.		n.d.			Abnormal	UL + LL		
	-	n.d.		-	-		n.d.			n.d.			Chronic neurogenic		Acute /	0				Acute +	u	neurogenic)		n.d.			n.d.	n.d.		n.d.		n.d.			Chronic /	neurogenic		
Cerebellar Peripheral EMG atrophy (MRI) neuropathy changes (NCS) (muscle)									n.d.	n.d.			1																n.d.		n.d.		n.d.	+; mild	sensory	+, motor	axonal		
Cerebellar atrophy (MRI)	+	+	+	+	-	+	+		+	+	+		+		+					+					+	+		+	+		+		+	+		+			
itures							inence		es	Mild macroglossia; esotropia;	inence		Strabism divergens, fasciculation face, fibrillation tongue, reduced vibration sense,	depression, CK elevation	Respiratory distress, mental	on colon,	pseudarthrosis clavicula,	scoliosis, CK elevation,	pes cavus, strabism,	cataract, myoclonus	Spiracol Justi Cos (BIPAP all day). mental	retardation (IO = 60),	hypertelorism, CK elevation,	tendon contractures					le initiation,	horizontal ophtalmoparesis,	tacial myokymia Reduced vibration sense,	urge incontinence, bulging eyes,	upper limb dystonia rge incontinence	sphagia		Cognitive deficit ($IQ = 49$), slow	saccades, CK elevation,		
Additional clinical fea			1				Urge incontinence	;	Slow saccades	Mild macro	Urge incontinence		Strabism div face, fibri reduced	depressio	Respiratory	malrotation colon,	pseudarth	scoliosis,	pes cavus	cataract, myocloni	(BIPAP al	retardatio	hypertelo	tendon o				Pes cavus	Slow saccade initiation,	horizonta	facial myokymia Reduced vibration	urge inco	upper limb dyste Urge incontinence	Scoliosis, dysphagia		Cognitive d	saccades,	pes cavus	
Dysfunction Additional lower motor clinical features neuron									+				+		+					+								+			+; proximal			,		+			
Dysfunction Dysfundupper Iower n							Signs +;	spasticity LL		Signs +;	Signs +; spasticity	LL and UL	Signs +		Signs +;	LL and UL				Gans +	spasticity LL	(1)			Signs +;	spasticity LL; Signs +;	spastic gait	Signs +;	Signs +;	spasticity LL			Signs +	Signs +		Signs +;	spastic gait		
SARA Cerebellar Dysfunction score ataxia upper motor neur	+	+	+	+	-	+	+		+	+	+		+		+					+					+	+		+	+		+		+	+		+			
	<u>3</u>	12	52	α	0 (6	8	;	25	15,5	22		16,5	,	33					33	3				4	4		ω	28		24		20	8		22			
of Age at Iast examination (years)	32	9	84	ç	70	73	29	i	75	46	39		25	,	36					ä	3				42	4		29	49		0/		20	92		21			
Age of onset	22	25	36	7.0	/7	6	40	;		28	21		<u>®</u>	,	9					4	•				30	78		12	0		24		91	20		0			
Origin/ gender	Turkish/m	Turkish/f	Italian/f	Eronch/f	יו פוכוי	German/f	German/f		Italian/m	German/m	German/f		Turkish/f		Belgian/f					Relaisn/m	90				Moroccan/f 30	Moroccan/f 28		German/m 15	Italian/f		French/f		ltalian/f	Italian/m		Italian/m			
Patient ID Phenotype Genotype (cDNA) (family- category individual)	c.13086delC; c.13086delC	c.15760C > T; c.15760C > T	c.23684_23685insACGCC TGTGCCACTGATGCCGAGTG;	c.15419A > G	C \ D0//00: (C \ D0//00:	c.24054G>A; c.16294_16300delA AACTGA	c.24025C > T; c.15665_15666delTC		c.23782C > T; c.8627_8628insAT	c.21316C>T; c.11908C>T	c.19855-1G > C; c.19855-1G > C		c.19897C>T; c.19897C>T	H () 1000	c.21528C > A; c.20935C > 1					C 21528C > A : C 20935C > T					c.682C>T; c.682C>T	c.682C>T; c.682C>T		c.20380C > T; c.17480dup	c.17944C>T; c.17944C>T		c.2776A>T; c.659T > C		c.14261dup; c.6034C>T	c.727C>T; c.727C>T		c.23782C > T; c.G20183 + IG > C			
ID Phenotype category al)	Pure CA	Pure CA	Pure CA	Q. G.		Pure CA	CA plus	;	CA plus	CA plus	CA plus		CA plus		CA plus					sula 40					CA plus	CA plus		CA plus	CA plus		CA plus		CA plus	CA plus		CA plus			
Patient ID (family- individual)	Ξ	3-1	7-1	14.1	5	<u>.</u>	4-1		-5-	1-6	<u>-0</u>		Ξ		12-1					12-2					13-1	13-2		<u>-</u>	1-61		20-1		21-1	22-1		23-1			

Table 2 Continued

Slow vertical saccades	4 + + + + + + + + + + + + + + + + + + +
- Scoliosis	22 + + +
	22 + -
- Scoliosis; kyphosis	+
- Square wave jerks, horizontal	
ophtalmoparesis, mild microretrognathism,	
unilateral ptosis Scoliosis, seizures at age 2 year, n.d.	+
square wave jerks - Pes cavus	21 + -

Patient ID = family number: m = male; f = female; pure CA = pure cerebellar ataxia; CA plus = cerebellar ataxia plus at least one additional system damage; + = present; - = absent; CK = creatine kinase; SARA = scale for the tendon reflexes; UL = upper limb, LL = lower limb; NCS = nerve conduction studies; n.d = not done. plantar response positive and/or hyperreflexia of muscle rating of ataxia; upper motor

of them showing also clinical signs of upper motor neuron damage (Table 2). Nerve conduction studies, available for 19 patients, were normal in 17/19 patients (89%), while one patient showed a peripheral motor neuropathy and one patient a mild sensory axonal neuropathy, demonstrating that peripheral neuropathy is a rather infrequent feature in SYNE1 disease.

Imaging findings

Cerebral MRI, available from 24 patients, consistently showed marked cerebellar atrophy in all 24/24 (100%) patients (for exemplary illustration, see Fig. 5). ¹⁸F-FDG-PET imaging performed in two exemplary subjects with ataxia-motor neuron disease revealed a marked homogenous ¹⁸F-FDG decrease in both cerebellar hemispheres in both subjects and, on visual inspection, also of the pontine brainstem, which reached statistical significance in Patient 10-1 with a Z-score of 3 and a trend in Patient 9-1 with a Z-score of 1.5 (semiguantitative PET, normalization to global mean activity and comparison to an age-adjusted database) (Fig. 5). These imaging findings provide proofof-concept functional imaging evidence that dysfunctions in SYNE1 patients extend beyond the cerebellar domain to include in particular the pons, thus complementing our clinical findings of extra-cerebellar features in SYNE1.

Discussion

SYNEI: a recurrent recessive ataxia worldwide, with mutations spanning across the entire gene

So far, autosomal-recessive cerebellar ataxia (ARCA) due to SYNE1 mutations has been described mainly in Quebec, Canada, with few families having been identified outside French-Canadian populations (Izumi et al., 2013; Noreau et al., 2013). Reporting by far the largest series of SYNE1 patients outside Quebec, we here show that SYNE1 deficiency is in fact a relatively common cause of recessive ataxia worldwide, yielding an estimated frequency of at least 5.3% after exclusion of Friedreich's ataxia and the most common repeat expansion SCAs. This is probably a rather conservative estimate, given the fact that our approach used only a very preliminary NGS-based approach to screen for copy number variants (CNVs) and that it was very restrictive on including missense variants. Future studies might be able to identify damaging CNVs in the large SYNE1 gene, which would further increase the relative frequency of subjects with truncating SYNE1 mutations.

Our genetic findings more than double the number of truncating *SYNE1* mutations identified so far: while <15 different truncating *SYNE1* mutations have been described, we here identified 33 novel truncating (plus one novel missense) *SYNE1* mutations, which are scattered throughout

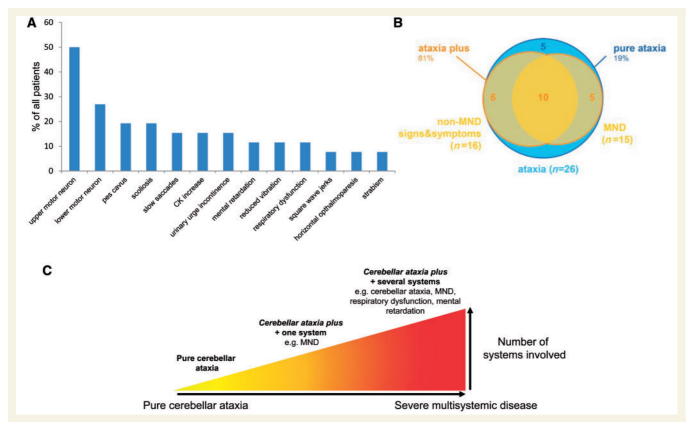


Figure 4 The phenotypic spectrum of SYNE1 disease. (A) Frequency of non-cerebellar features in SYNE1. Relative frequencies of non-cerebellar features observed in the n = 26 SYNE1 patients investigated in this study. Note that several patients showed ≥ 2 non-cerebellar disease features. (B) The relative share of main SYNE1 phenotypes. This figure illustrates that only a minor share of SYNE1 patients shows pure cerebellar ataxia (blue), whereas the large majority show cerebellar ataxia plus phenotypes (yellow), in particular cerebellar ataxia plus motor neuron disease (MND). The latter combination is thereby not a distinct ataxia plus-phenotype, but is commonly associated also with other non-motor neuron disease signs (see overlap group comprising of n = 10 subjects). (C) The continuous spectrum of SYNE1 disease. The clinical spectrum of neurological and non-neurological SYNE1 features presented in A unfolds along a continuous spectrum of disease. Variable combinations of these features can be found in different SYNE1 patients reaching from pure cerebellar ataxia (left) via cerebellar ataxia plus damage of one or two additional systems [e.g. motor neuron disease (MND)] to cerebellar ataxia as part of severe multisystemic neurodegenerative syndrome (right). In addition to ataxia and motor neuron disease, the latter can include also mental retardation, respiratory distress and visceral organs. The phenotype of mild, slowly progressive pure cerebellar ataxia, which has been considered the most prominent phenotype of SYNE1 so far (Gros-Louis et al., 2007; Noreau et al., 2013), thus represents only one far edge of this broad disease spectrum. CK = creatine kinase.

all parts of this gigantic gene. Almost all mutations are private, identified in single families without obvious mutational hot spot regions. Five mutations affected the coding sequence of the SYNE1 isoform Nesprin 1α (see mutations 1–5, Table 1). As no mutations linked to ataxia had been previously found in this domain (Razafsky and Hodzic, 2015), it was suggested that a protein encoded by these exons may underlie a distinct pathology in humans (Razafsky and Hodzic, 2015). However, here we show that mutations in this sequence can lead to the same phenotypic features as mutations in other parts of *SYNE1*.

Taken together, these findings have also major implications for future clinic-genetic testing strategies in unresolved ARCA patients around the world: they demonstrate that also in non-Canadian populations current genetic testing approaches have to include the large *SYNE1* gene by using approaches that tightly capture all of its exons.

SYNEI spectrum: from late-onset pure cerebellar ataxia to complex neuromuscular disease with mental retardation

Recessive ataxia due to *SYNE1* mutations is commonly perceived to present as a slowly progressive, relatively pure cerebellar ataxia with only mild and infrequent extra-cerebellar symptoms (e.g. brisk reflexes in 33%; Dupre *et al.*, 2007), starting in adult age (Dupre *et al.*, 1993/2012; Gros-Louis *et al.*, 2007; Noreau *et al.*, 2013; Fogel *et al.*, 2014). However, here we show that pure cerebellar ataxia represents rather a relatively infrequent phenotypic cluster along a broad multidimensional SYNE1-associated spectrum of neurodegenerative disorders. With 81% of all patients exhibiting variable additional non-cerebellar features both in neurological and

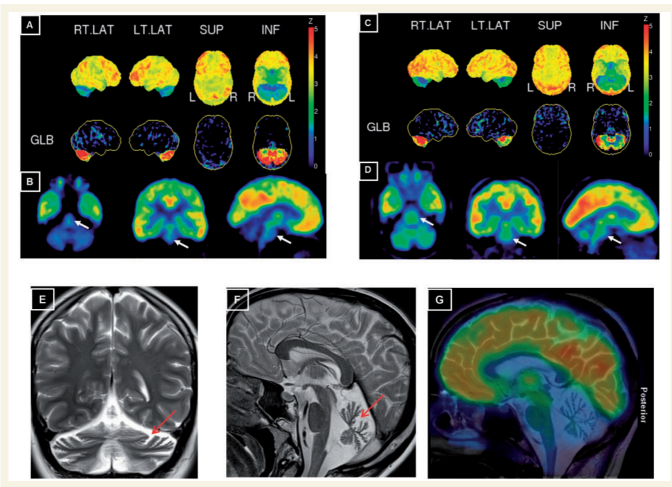


Figure 5 Cerebral ¹⁸F- FDG PET and MRI in exemplary SYNEI patients. (A and C) Surface projections of the ¹⁸F-FDG PET scans of Patient 10-1 at age 37 years (A) and Patient 9-1 at age 45 years (C). The upper rows in A and C show the surface projection map of cerebral glucose metabolism normalized to the maximum of the acquisition. The lower rows show the deviation of an age-adjusted normal database after normalization to a global mean calculation (GLB) of metabolic activity. On both projections, a marked homogenous bilateral reduction of cerebellar FDG metabolism can be seen in both subjects. (B and D) Tomographic projections after automated fitting. For both Patient 10-1 (B) and Patient 9-1 (D) a reduction of FDG-metabolism is also clearly visible in the brainstem (white arrows), which reached statistical significance in Patient 10 with a Z-score of 3 and a trend in Patient 9-1 with a Z-score of 1.5 (semiquantitative PET, deviation after normalization to the global mean). (E and F) T₂-weighted MRI scans in Patient 18-1 at age 21 years shows marked vermian atrophy (F) and cortical hemisphere atrophy (E) of the cerebellum (red arrows). (G) Fusion image of ¹⁸F-FDG PET scan of Patient 10-1 onto a central paramedian sagittal T₂-weighted MRI scan shows considerable hypometabolism in the cerebellum and in the pons, whereas FDG metabolism is normal in the cortical cerebral regions.

non-neurological domains, a multisystemic phenotype seems to be the rule rather than the exception in SYNE1 disease. Apart from cerebellar ataxia, this spectrum includes upper motor neuron disease, lower motor neuron disease, brainstem dysfunction (e.g. saccadic slowing), and a variable range of musculoskeletal abnormalities (e.g. kyphosis, scoliosis, pes cavus, contractures). Variable combinations of these features can be found in different SYNE1 patients along a continuous spectrum of disease. This spectrum ranges from (i) pure cerebellar ataxia; via (ii) cerebellar ataxia plus damage of one or two additional systems (such as motor neuron disease, thus giving rise to spastic ataxia and complicated hereditary spastic paraplegia phenotypes); to (iii) cerebellar ataxia as part of severe multisystemic neurodegenerative syndromes. The latter

encompass literally all of the aforementioned SYNE1-associated domains and include also mental retardation, respiratory distress and possibly visceral organ involvement (Table 2; for graphic illustration of the SYNE1 disease continuum, see Fig. 4B and C).

The complex early-onset neurodegenerative syndrome does not seem to present a mere coincidental finding (that might be unrelated to the *SYNE1* mutations), but rather reflects a phenotypic cluster systematically related to *SYNE1* mutations. This is indicated by the fact that we identified three patients from two different families with this severe phenotypic combination. In fact, a fourth patient with early-onset complex *SYNE1* ataxia-motor neuron disease and respiratory dysfunction has recently been described as a single case in a Japanese family (Izumi *et al.*, 2013),

thus increasing the evidence to three different families. We now show that this phenotypic cluster is not a qualitatively distinct syndrome, but rather a combination of *SYNE1*-assosciated features that can each be found in isolation or as part of less complex phenotypic clusters in other *SYNE1* patients along a continuous spectrum of disease. Although we did not find obvious mutation—phenotype correlations, different *SYNE1* mutations might differentially affect the multiple SYNE1 splicing isoforms which vary greatly in size and tissue-specific expression patterns (Zhang *et al.*, 2010; Razafsky and Hodzic, 2015), thus possibly explaining parts of the phenotypic variability observed here. In fact, *SYNE1* mutant mice show variable phenotypes depending on the splicing isoform that was targeted (for overview see Zhang *et al.*, 2010).

Our findings have important implications for clinico-genetic counselling. They revise current clinical notions about the benign disease course of *SYNE1*-disease, including the notion that life expectancy was normal (Dupre *et al.*, 1993/2012; Gros-Louis *et al.*, 2007). They show that, at least in some patients, *SYNE1*-disease can start already as a developmental disease with mental retardation and early-onset ataxia in the first decade of life, and that it can lead to premature death due to respiratory dysfunction in the midadult age (age 30–40 years).

Bridging the gap between SYNEI ataxia and arthrogryposis syndromes

Truncating recessive SYNE1 mutations have been described as a cause of an arthrogryposis multiplex congenita syndrome in a single Palestinian family presenting with infantile-onset hypotonia, bilateral club foot, progressive motor decline in the first decade, scoliosis and restrictive lung disease, yet without ataxia or motor neuron disease (Attali et al., 2009) (for location of these mutations, see Fig. 1B). This musculoskeletal infantile-onset phenotypic cluster was proposed as a 'distinct human disease phenotype' of SYNE1, separate from the neurological phenotype (Attali et al., 2009). As several of our ataxic SYNE1 patients, however, also show scoliosis/kyphosis, restrictive lung disease, foot deformities, and other neuromuscular abnormalities as part of their early-onset multisystemic disease, and also carry truncating mutations in the same or neighbouring gene region (Fig. 1B), we here suggest that such arthrogryposis syndromes do not represent qualitatively distinct phenotypes, but rather clusters of variably combined neurologic and non-neurological features along the continuum of

It remains more questionable whether this SYNE1 disease continuum also includes the dominant form EDMD4 (Zhang *et al.*, 2007*a*). Although a phenotypic overlap might be theoretically conceivable, as indicated by our clinical findings, this syndrome has been linked only to heterozygous missense mutations in *SYNE1* (Zhang *et al.*, 2007*a*) (Fig. 1B), but not to biallelic truncating mutations. As

advanced genomic techniques now allow us to appreciate the immense variability of the SYNE1 gene and to determine the frequency of rare SYNE1 missense variants in controls (e.g. 28/500 = 5.6% rare missense alleles in our control dataset with early-onset Alzheimer dementia; Supplementary material), the pathogenicity of such variants previously related to EDMD4 needs to be critically evaluated against current standards. Furthermore, findings on muscle biopsies between subjects with EDMD4 and the here reported cases with biallelic truncating mutations differ substantially; while EDMD4 subjects showed dystrophic myopathic changes and in particular mislocalization of emerin or lamin A/C (Zhang et al., 2007a), none of these features was seen in the muscle cells of any of our subjects (Fig. 3). Although this might be explained by a different SYNE1-related pathomechanism, this patho-morphological discrepancy, in addition to the genetic concerns mentioned above, indicates a possible over-interpretation of heterozygous SYNE1 variants in the context of EDMD4 and warrants a critical re-evaluation of these historic findings.

A widespread functional role of SYNEI in various tissues

Our findings on widespread extra-cerebellar dysfunctions in human SYNE1 disease complement and recapitulate some of the key features of SYNE1/Nesprin 1 knock-out mouse models. Key features in mice include kyphoscoliosis, respiratory failure, and shortened survival (Zhang *et al.*, 2007*b*, 2010; Puckelwartz *et al.*, 2009). We now show systematically that these features are also part of the human disease spectrum, thus bridging the cross-species gap.

Moreover, our findings on extra-cerebellar dysfunctions might provide new insights into the pathophysiology underlying SYNE1-associated neurodegeneration. Recent work has shown that a specific SYNE1 isoform devoid of the KASH domain (KLNes1g) is specifically abundant in the cerebellum (in particular in the granule cell layer), where it might be involved in vesicular trafficking and/or in dendritic membranes' structural organization (Razafsky and Hodzic, 2015). Our clinical, electrophysiological and imaging observations of multisystemic damage in SYNE1 patients, however, indicate that intact SYNE1 transcripts seem to be functionally important not only for cerebellar, but also for motor and brainstem neurons. They thus stimulate future molecular research aiming to identify the specific functional role of SYNE1 in these neuron types. For example, the proposed interaction of SYNE1 transcripts with vesicular trafficking proteins such as KIF5C, suggested for cerebellar neurons (Razafsky and Hodzic, 2015), might also apply to motor neurons. Aberrant vesicular trafficking dynamics is a common process underlying motor neuron degeneration (as in hereditary spastic paraplegias; Crosby and Proukakis, 2002), and mutations in KIF5C have been

shown to include upper motor neuron damage (Poirier et al., 2013).

In sum, our clinical, electrophysiological and imaging findings suggest a widespread functional role of SYNE1 in many different tissues in humans, which is in line with its ubiquitous expression (Apel et al., 2000; Gros-Louis et al., 2007; Zhang et al., 2010). This ubiquitous expression of SYNE1—and, in turn, its absence in case of protein truncation—might help to find diagnostic biomarkers for the disease. Our results suggest that severely reduced to absent SYNE1 staining in muscle tissue, which was observed in all three available samples, may represent a diagnostic marker that indicates underlying truncating SYNE1 mutations. If confirmed in larger sample cohorts, SYNE1 staining of muscle tissue might thus become a helpful auxiliary tool in the histochemical work-up of muscle biopsies of future patients with unresolved neurodegenerative and neuromuscular diseases. It is likely that many more SYNE1 patients with both relatively pure ataxia as well as complex neuromuscular phenotypes will be identified worldwide in the very next years.

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Conflict of interest

Dr Klopstock has performed consultancies for Actelion Pharmaceuticals Ltd, GenSight Biologics, Gerson Lehrman Group, USA, and FinTech Global Capital, Japan. He has been serving as a Section Editor for BMC Medical Genetics from 2011-2015. Dr Anheim received received honoraria, consulting fees and travel reimbursements from Actelion Pharmaceuticals Ltd, Abbvie, Novartis, Teva and Lundbeck.

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Supplementary material

Supplementary material is available at Brain online.

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