

SIL1 mutations and clinical spectrum in patients with Marinesco-Sjögren syndrome

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Marinesco-Sjögren syndrome is a rare autosomal recessive multisystem disorder featuring cerebellar ataxia, early-onset cataracts, chronic myopathy, variable intellectual disability and delayed motor development. More recently, mutations in the SIL1 gene, which encodes an endoplasmic reticulum resident co-chaperone, were identified as the main cause of Marinesco-Sjögren syndrome. Here we describe the results of SIL1 mutation analysis in 62 patients presenting with early-onset ataxia, cataracts and myopathy or combinations of at least two of these. We obtained a mutation detection rate of 60% (15/25) among patients with the characteristic Marinesco-Sjögren syndrome triad (ataxia, cataracts, myopathy) whereas the detection rate in the group of patients with more variable phenotypic presentation was below 3% (1/37). We report 16 unrelated families with a total of 19 different SIL1 mutations. Among these mutations are 15 previously unreported changes, including single- and multi-exon deletions. Based on data from our screening cohort and data compiled from the literature we found that SIL1 mutations are invariably associated with the combination of a cerebellar syndrome and chronic myopathy. Cataracts were observed in all patients beyond the age of 7 years, but might be missing in infants. Six patients with SIL1 mutations had no intellectual disability, extending the known wide range of cognitive capabilities in Marinesco-Sjögren syndrome to include normal intelligence. Modestly constant features were somatic growth retardation, skeletal abnormalities and pyramidal tract signs. Examination of mutant SIL1 expression in cultured patient lymphoblasts suggested that SIL1 mutations result in severely reduced SIL1 protein levels irrespective of the type and position of mutations. Our data broaden the SIL1 mutation spectrum and confirm that SIL1 is the major Marinesco-Sjögren syndrome gene. SIL1 patients usually present with the characteristic triad but cataracts might be missing in young children. As cognitive impairment is not obligatory, patients without intellectual disability but a Marinesco-Sjögren syndrome-compatible phenotype should receive SIL1 mutation analysis. Despite allelic heterogeneity and many families with private mutations, the phenotype related to SIL1 mutations is relatively homogenous. Based on SIL1 expression studies we speculate that this may arise from a uniform effect of different mutations on protein expression.

Keywords: Marinesco-Sjögren syndrome; ataxia; cataract; myopathy; SIL1 mutation

Introduction

Marinesco-Sjögren syndrome (MIM 248800) is a long-recognized autosomal recessively inherited, infantile-onset multisystem disorder that affects brain, eyes and skeletal muscles. The clinical triad of bilateral cataracts, ataxia and intellectual disability was noted in the first half of the last century (Moravcsik, 1904; Marinesco et al., 1931; Sjögren, 1947) and later confirmed in a series of additional reports. Pathoanatomical and brain imaging studies revealed cerebellar atrophy as the cause of the cerebellar syndrome (Todorov, 1965; Georgy et al., 1998). From histopathological and neurophysiological investigations chronic myopathy emerged as an additional feature (Chaco, 1969; Herva et al., 1987) and electron microscopy revealed particular electron-dense membranous structures around degenerating myonuclei (Herva et al., 1987; Sewry et al., 1988). Hypergonadotropic hypogonadism, skeletal abnormalities and short stature are additional features that have been reported with variable frequency (Berg and Skre, 1976; Brogdon et al., 1996). Although most patients are severely handicapped, life span in Marinesco-Sjögren syndrome is at least not drastically reduced as the oldest reported patients are in their 70s (Anttonen et al., 2005). More recently, using positional cloning strategies, we and others have shown that Marinesco-Sjögren syndrome is caused by homozygous or compound heterozygous point mutations in the SIL1 gene on chromosome 5q31.2 (Anttonen et al., 2005; Senderek et al., 2005). Following the two original reports, 14 additional families with SIL1 mutations have been described in the medical literature (Karim et al., 2006; Annesi et al., 2007; Anttonen et al., 2008; Eriguchi et al., 2008; Riazuddin et al., 2009; Takahata et al., 2010; Terracciano et al., 2012). A subgroup of patients with Marinesco-Sjögren syndrome was found to be negative for SIL1 point mutations, suggesting alternative mutation mechanisms or locus heterogeneity (Senderek et al., 2005; Anttonen et al., 2008).

SIL1, the human homolog of yeast Sil1p (suppressor of the △ire1 △lhs1 double mutant number 1), is an evolutionary conserved, ubiquitously expressed 461 amino-acid N-linked glycoprotein (Chung et al., 2002). SIL1 acts as a co-chaperone and nucleotide exchange factor for the HSP70 (heat-shock protein 70) ATPase BiP (immunoglobulin binding protein, also referred to as 78-kDa glucose-regulated protein GRP78) (Haas and Wabl, 1983; Munro and Pelham, 1986). BiP is a stress-inducible molecular chaperone and controls a plethora of essential processes in the endoplasmic reticulum including translocation of nascent proteins into the endoplasmic reticulum, accurate subsequent folding of the newly synthesized proteins in the endoplasmic reticulum lumen,

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elimination of proteins that fail to mature properly, response to cell stress, and calcium homeostasis (Hendershot, 2004). As at least some functions of BiP require its ATPase activity, it seems plausible to assume that the nucleotide exchange factor SIL1 is involved in the regulation of endoplasmic reticulum-associated processes through control of the BiP ATPase cycle and that impaired BiP activation is the relevant pathomechanism in Marinesco-Sjögren syndrome.

The prominent cerebellar atrophy seen in patients with Marinesco-Sjögren syndrome is caused by loss of Purkinje and granule cells (Todorov, 1965; Mahloudji et al., 1972; Skre and Berg, 1977). The same histopathological findings are present in the spontaneous mouse mutant woozy, which lacks a functional Sil1 gene resulting in adult-onset ataxia (Zhao et al., 2005). However, woozy mice seem to differ from humans with Marinesco-Sjögren syndrome as no symptoms like cataracts or myopathy have been described in this animal model. SIL1 is expressed in all tissues and organs in vertebrates with highest levels in secretory tissues such as liver, placenta and kidney (Chung et al., 2002). It remains unknown why loss of SIL1 function in man and mice does not cause lethality and why certain tissues and cell types such as cerebellum (in mice and humans), eye and skeletal muscle (in humans) are more vulnerable to loss of SIL1 function than other tissues. As in yeast (Tyson and Stirling, 2000), the presence of the additional nucleotide exchange factor HYOU1 (also known as ORP150 and GRP170) in various organs of mammals may be able to compensate the loss of SIL1 function. This hypothesis has recently been supported by the finding that the cerebellar atrophy in the woozy mouse can be compensated by over-expression of HYOU1 (Zhao et al., 2010).

Here, we screened a cohort of unselected patients referred for molecular genetic testing of the *SIL1* gene. We extend the phenotypic and mutation spectrum in patients with Marinesco-Sjögren syndrome and suggest inclusion criteria for *SIL1* mutation screening based on compilation of clinical data of all reported patients with *SIL1* mutations.

Materials and methods

Patients

All patients included in this study were referred to our molecular genetic diagnostic laboratory for SIL1 mutation analysis between 2005 and 2011. The general inclusion criteria for this study required definite or probable clinical evidence of early-onset cataracts, cerebellar atrophy/ ataxia and chronic myopathy (n = 25). We also enrolled a cohort of patients presenting with two out of the three cardinal features plus at least one of the following additional features: short stature, intellectual disability, delayed motor milestones, muscular hypotonia, hypogonadism, peripheral neuropathy, skeletal deformities (e.g. scoliosis, pes cavus) (n = 37). All biological materials (blood samples, DNA samples, cell lines, archived specimens of diagnostic muscle biopsies), medical and neurophysiological reports, clinical photographs, ultrasound images, and brain MRI scans were obtained under appropriate informed consent of the patients or their legal guardians. Muscle biopsies were processed as described previously (Weis and Schröder, 1988, 1989).

SIL1 gene sequence analysis

Primer sets for PCR amplification of human *SIL1* coding exons 2–10 (NM_022464.4) with 40 to 50 bp of flanking intronic sequences have been described previously (Senderek *et al.*, 2005). The resulting PCR products were subjected to fluorescence-based cycle sequencing using the BigDye® Terminator Cycle Sequencing Ready Reaction Kit, version 3.1 (Applied Biosystems). Samples were run and analysed on an ABI PRISM 3100 Genetic Analyzer (Applied Biosystems). Sequencing electropherograms were assessed by visual inspection in order to identify variants.

Identification of *SIL1* exon-deletion mutations

In Patient MSS87.1 we identified a seemingly homozygous mutation in exon 9 whereas we observed heterozygosity for two variants in exon 3 and intron 4. Based on these findings we assumed that Patient MSS87.1 might harbour a heterozygous intragenic deletion encompassing at least exon 9. High resolution genome wide single-nucleotide polymorphism array analysis was performed by using an Affymetrix 250K Nsp SNP array (Affymetrix), in accordance with the supplier's instructions, and copy-number data were analysed with the Affymetrix Genotyping Console 3.0.2 software. For confirmation and fine mapping of a copy number deviation, we developed quantitative real-time PCR assays for 12 fragments located in introns 7 and 9 and 3' of the SIL1 gene (Fig. 1B). As reference locus, exon 3 of the factor VIII gene (Wilke et al., 2000) was used. Quantitative real-time PCR was carried out on an ABI PRISM 7000 sequence detection system (Applied Biosystems) using the quantitative real-time PCR core kit for SYBR Green I according to the manufacturer's protocol (Eurogentec). Data evaluation was performed using the ABI PRISM 7000 sequence detection software as described previously (Wilke et al., 2000) and the ratios of test and reference fragments were calculated to determine copy numbers of the test fragments. Primers from the fragments found to be situated just outside the deletion (i.e. fragments yielding results in favour of two copies) were used in an attempt to amplify the breakpoint region by long-range PCR.

PCR amplification of SIL1 exon 5 consistently failed in the DNA sample of the index patient of Family MSS150.1 whereas amplification of exons 4 and 6 was normal. To obtain further evidence for a presumed intragenic deletion removing exon 5, total RNA was isolated from a fresh blood sample of the index patient, reverse transcribed and used as template for PCR with primers in SIL1 exons 1 and 10 (Fig. 1C). Sequencing of the PCR product was performed using ABI Prism BigDye® technology as described above. At the genomic level, the extent of the presumed deletion was defined using amplification of several short 100-200 bp genomic fragments placed every 2-3 kb within the 25 kb genomic region encompassing intron 4, exon 5 and intron 5. Primers from fragments found to lie just outside the deletion (i.e, these fragments yielded amplification in patient's DNA while fragments situated in the genomic region flanked by these markers gave no products) were combined to amplify a junction fragment characteristic for the deletion (Fig. 1D). The junction fragment was subjected to cycle sequencing in order to identify the precise positions of the deletion breakpoints.

Cell culture and western blotting

Epstein-Barr virus-immortalized B lymphoblast cell lines from control subjects and patients with Marinesco-Sjögren syndrome were established

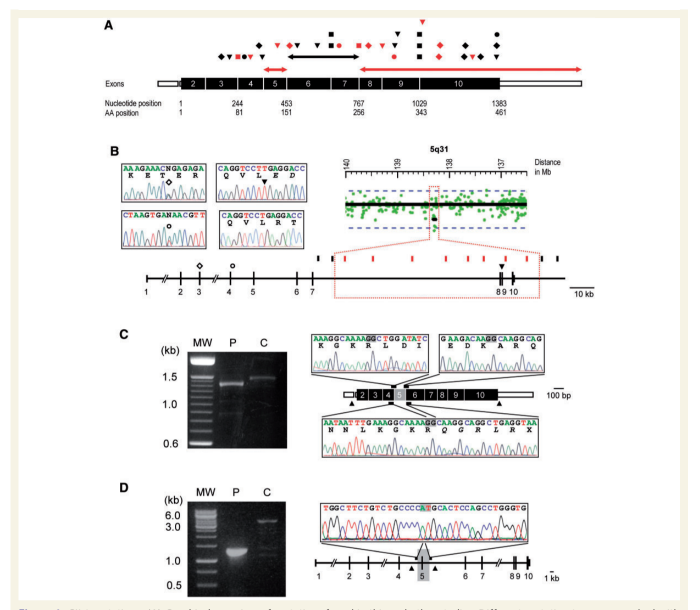


Figure 1 SIL1 mutations. (A) Graphical overview of mutations found in this and other studies. Different mutation types are marked with different symbols. Circles = missense mutations and small in-frame indels; rhombi = nonsense mutations; triangles = frameshift mutations; squares = splice site mutations; two-sided arrows = exon-deletion mutations. Previously unreported sequence variations identified in this study are coloured in red. Exon, nucleotide and amino acid numbering is according to NM_022464.4 and NP_071909.1. Introns are not drawn to scale. AA = amino acid. (B) Heterozygous multi-exon deletion in Patient MSS87.1. Sequencing electropherograms show heterozygous variants in exon 3 (rs3088052, open rhombus) and intron 4 (c.353 + 38T > A, open circle) as well as a seemingly homozygous frameshift mutation c.947dupT, p.R317fs in exon 9 (filled triangle, the electropherogram below represents the wild-type sequence). A high-density genome-wide single nucleotide polymorphism array (upper left) and copy number analysis of fragments located in the potential deletion region by quantitative real-time PCR (vertical bars above the schematic representation of the SIL1 gene, red: one copy, black: two copies) were suggestive of a partial, \sim 68–77 kb deletion involving exons 8–10 (boxed with dotted red lines). Long-range PCR with primers from the fragments found to be situated just outside the deletion failed to amplify a deletion specific junction fragment leaving the possibility of a more complex genomic rearrangement although routine-karyotyping had previously shown two normal chromosomes 5. (C and D) Homozygous exon 5 deletion in Patient MSS150.1. (C) Reverse transcription PCR using patient's messenger RNA (P) yielded a shorter 1370-bp fragment (C: control messenger RNA, 1470 bp). Positions of pimers are indicated by arrowheads underneath the schematic representation of the SIL1 messenger RNA. Sequencing electropherograms representing the normal exon 4/ exon 5 and normal exon 5/exon 6 junctions are shown above the schematic representation of the SIL1 messenger RNA; the exon 4/exon 6 junction resulting from the genomic deletion is shown below. Removal of the 100-bp exon 5 from the mature messenger RNA is predicted to result in frameshift and premature stop of translation (asterisk). (D) Sequencing of a PCR-amplified deletion-specific junction fragment at the genomic level. Primers placed in intron 4 and intron 5 of the SIL1 gene yielded a ~ 2.5 kb shorter product when using patient's (P) DNA (C: control DNA, 4029 bp). Positions of primers are indicated by arrowheads underneath the schematic representation of the SIL1 gene genomic region (not drawn to scale). The borders of the deletion which encompasses 2660 bp including exon 5 were determined by DNA sequencing with internal primers. The extent of the deletion is indicated as a grey box.

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according to standard protocols (Tohda et al., 1978). The immortalized lymphoblasts were cultured in RPMI-1640 growth medium, harvested by centrifugation (800 rpm, 7 min), washed twice in cold PBS and pelleted by centrifugation (14 000 rpm, 5 min, 4°C). Cell pellets were homogenized in lysis buffer (10 mM Tris-HCl, 5 mM EDTA, 150 mM NaCl, 1% TritonTM X-100) containing protease and phosphatase inhibitors. Post-nuclear supernatants were boiled in sample buffer (80 mM Tris pH 6.8, 10% glycerol, 2% SDS, 0.002% bromphenol blue), resolved by SDS-PAGE and electroblotted onto polyvinylidene fluoride membranes (Hybond-C; GE Healthcare). Immunoblots were developed by incubation with appropriate antibodies followed by horseradish peroxidasephosphatase chemiluminescence detection (ECL; GE Healthcare). The following antibodies were used: mouse monoclonal anti-SIL1 clone 1F9 (Origene Technologies; dilution 1:1000), mouse monoclonal anti-GAPDH (Genetex; dilution 1:1000), goat anti-rabbit immunoglobulin G antibody conjugated to horseradish peroxidase (Invitrogen; dilution 1:5000), goat anti-mouse immunoglobulin G conjugated to horseradish peroxidase (Invitrogen; dilution 1:5000).

Results

Mutation screening of the *SIL1* gene in a diagnostic sample of 62 unrelated, previously unreported index patients with Marinesco-Sjögren syndrome or Marinesco-Sjögren syndrome-like conditions revealed pathogenic sequence variants in 16 families of diverse ethnic backgrounds, representing roughly 25% of the total cohort. The mutation detection rate reached 60% (15/25) in the cohort of patients presenting with the clinical triad of cerebellar atrophy, myopathy and cataracts whereas the mutation detection rate was <3% (1/37) among Marinesco-Sjögren syndrome-like cases (two of the cardinal features and at least one of the following symptoms: short stature, intellectual disability, delayed motor milestones, hypotonia, hypogonadism, peripheral neuropathy, skeletal deformities).

We observed a total of 19 different SIL1 mutations consisting of a missense mutation, a two-amino acid in-frame deletion, seven frameshift mutations, four nonsense mutations, four nucleotide changes affecting splice sites and two genomic deletions (Fig. 1 and Table 1). Among these mutations, 15 had not been reported previously. None of these so far unrecognized changes was present in databases containing information on the 'normal' variability of the human genome (dbSNP version 135, www.1000genomes. org/; 1000 Genomes database, www.ncbi.nlm.nih.gov/projects/ SNP/; Exome Variant Server, evs.gs.washington.edu/EVS/). In addition, analysis of DNA samples obtained from healthy control individuals, adjusted to the type of mutation (i.e. at least 100 for each truncating and at least 400 for each missense mutation), did not yield any of the identified genotypes. All patients were identified with bi-allelic mutations, either in the homozygous or compound heterozygous state. When DNA from family members was available, we observed that the disease phenotype co-segregated with recessive inheritance of the SIL1 mutations. The parents carried mutations in the heterozygous state, and unaffected siblings carried either one heterozygous mutation or were homozygous for the wild-type alleles.

Detailed clinical, electrophysiological and muscle biopsy findings of 23 patients from the 16 families carrying pathogenic *SIL1* mutations are provided in Supplementary Table 1. All patients came to medical

attention with symptoms suggestive of Marinesco-Sjögren syndrome within the first 5 years of life. Initial presenting symptoms were nonetheless variable: some patients presented very soon after birth with severe hypotonia, whereas most showed delay in motor milestones or truncal ataxia when starting to sit or stand, after a normal neonatal period. The mean age at onset of cataracts was 3.7 years (±1.5 years). Congenital cataracts were only noted in one patient and the latest manifestation of cataracts was reported at the age of 7 years. In line with this observation, one girl did not show any lens opacities at the age of 4 years. Apart from this exception, all patients invariably presented with a cerebellar syndrome, symptoms and signs of a myopathy and bilateral cataracts. Modestly constant features were skeletal deformities, somatic growth retardation and pyramidal tract signs. Mental capacities of patients with SIL1 mutations were highly variable, ranging from normal (n = 6) over mild (n = 7) to moderate (n = 8) and severe intellectual disability (n = 2). Other clinical features were only occasionally recorded (e.g. epilepsy or microcephaly) or could only be reasonably assessed in a too small subgroup of patients to give meaningful figures (e.g. hypogonadism in postpubertal patients). Although motor development was often markedly delayed, patients generally became mobile with a walker or other orthopaedic devices (canes, crutches, braces). Marked atrophy of the cerebellum was observed in all patients who received brain MRI) (Fig. 2B-D). Serum creatine kinase (CK) levels were usually only slightly or moderately elevated (377 \pm 159 U/I). Through a review of the medical literature we recorded 65 additional patients with Marinesco-Sjögren syndrome with confirmed SIL1 mutations (30 families). We combined these data with information from our patient cohort to determine the relative frequency of signs and symptoms in SIL1-associated Marinesco-Sjögren syndrome (Supplementary Table 2 and Fig. 2A).

We reassessed muscle biopsy specimens that had been taken for diagnostic purposes from four patients with identified SIL1 mutations (MSS33.1, MSS87.1, MSS91.1, and MSS94.1). Histologically we observed different degrees of degeneration of skeletal muscle fibres (Fig. 3A): sarcoplasmic vacuoles, often associated with myonuclei, greater-than-normal variability in fibre size, rounded atrophic fibres, myofibre hypertrophy and splitting, increased numbers of internal nuclei, endomysial fibrosis and proliferation of fat tissue. At the ultrastructural level (Fig. 3B-D), autophagic vacuoles, which were often associated with degenerating myonuclei, were encountered frequently. Several degenerating myonuclei were surrounded by an electron-dense, membrane-like structure. This feature was constantly observed in all four muscle biopsies that were accessible to us. Muscle biopsy findings of 10 additional patients were available from medical records and were indicative of a chronic myopathic process. Five biopsy samples had also been examined by electron microscopy; however, no details concerning possible nuclear abnormalities were mentioned in the reports.

By immunoblotting, we found that SIL1 levels in lymphoblast lines of five patients with different *SIL1* mutations were substantially reduced compared with the levels in control subjects (Fig. 4). Importantly, reduced amounts of SIL1 were also seen in patients with seemingly 'milder' mutations [small in-frame deletion (MSS24.1), missense mutation (MSS32.1) and frameshift mutation in the last exon which is expected to escape nonsense-mediated messenger RNA decay (MSS64.1)].

Table 1 SIL1 mutations in patients with Marinesco-Sjögren syndrome identified in this study and earlier reports

Location	Nucleotide change	Effect on coding sequence	Ethnic origin	References
Exon 3	c.178G>T	p.E60X	Vietnam	Senderek et al., 2005
Exon 3	c.212dupA	p.H71fs	France	Anttonen et al., 2005
Intron 3	c.244 + 1G > A	splice error predicted	Turkey	This study
Exon 4	c.274C>T	p.R92W	Pakistan	Riazuddin et al., 2009
Exon 4	c.302_303delAG	p.E101fs	Pakistan	This study
Exon 4	c.331C>T	p.R111X	Iran, Turkey, Italy	Senderek et al., 2005 Anttonen et al., 2005 Annesi et al., 2007 Terracciano et al., 2012
Exon 4	c.347delG	p.G116fs	Vietnam	Senderek et al., 2005
Exon 5		•	India	
Exon 5	chr5:g.138376244_138378903del c.424delG	p.L119fs		This study
		p.A142fs	Germany	This study
Exons 6, 7	chr5:g.138311133_138369401delinsTGCA	p.A152fs	Japan	Takahata <i>et al.</i> , 2010
Exon 6	c.460C>T	p.Q154X	France	This study
Exon 6	c.506_509dupAAGA	p.D170fs	Finland, Sweden, Norway	Anttonen et al., 2005
Exon 6	c.603_607delGAAGA	p.E201fs	Japan	Takahata et al., 2010
Intron 6	c.645 + 1G > A	p.A152_Q215del	Turkey, Argentina	Senderek <i>et al.</i> , 2005 This study
Intron 6	c.645 + 2T > C	p.A152_Q215del	Sweden	Anttonen et al., 2005
Exon 7	c.691_696delGTGATC	p.V231_l232del	Turkey	This study
Intron 7	c.768-1G > A	p.S256fs	Italy	This study
Exons 8, 9, 10	chr5:g.(138271912_138277581) _(138345106_138349134)del	p.S256_R461del	Russia	This study
Exon 8	c.811C>T	p.Q271X	Turkey	This study
Exon 9	c.866dupT	p.L290fs	France	This study
Exon 9	c.934G > A	p.G312R	USA	This study
Exon 9	c.936dupG	p.L313fs	Japan, Argentina	Eriguchi <i>et al.</i> , 2008 Anttonen <i>et al.</i> , 2008 This study
Exon 9	c.947dupT	p.R317fs	Germany, Russia	Senderek <i>et al.</i> , 2005 This study
Intron 9	c.1029 + 1G > A	p.V289_K343del	Bosnia	Senderek et al., 2005
Intron 9	c.1030-9G > A	p.F345fs	Norway, USA, Pakistan	Anttonen <i>et al.</i> , 2008 This study
Intron 9	c.1030-18G>A	p.M344fs	Germany	Senderek et al., 2005
Exon 10	c.1035delC	p.F345fs	Germany	This study
Exon 10	c.1126C>T	p.Q376X	Pakistan	This study
Exon 10	c.1137C>A	p.C379X	Iraq	This study
Exon 10	c.1240C>T	p.Q414X	Pakistan	Riazuddin et al., 2009
Exon 10	c.1249C>T	p.Q417X	Mali	Senderek et al., 2005
Exon 10	c.1276_1282delCAGGCTG	p.Q426fs	Germany	This study
Exon 10	c.1312C>T	p.Q438X	Egypt	Karim <i>et al.</i> , 2006
Exon 10	c.1367delT	p.L456fs	Russia	Senderek et al., 2005
Exon 10	c.1367T > A	p.L456X	Turkey	Anttonen et al., 2008
Exon 10	c.1370T > C	p.L457P	Japan	Anttonen et al., 2008

Exon, nucleotide and amino acid numbering is according to NM_022464.4 and NP_071909.1. Genomic coordinates are based on GRCh37/hg19.

Discussion

In this study we report the results of *SIL1* mutation screening in patients who were referred to our laboratory for molecular genetic testing of Marinesco-Sjögren syndrome during the past 5 years. Our study confirms the previous findings of mutations in *SIL1* being the major cause of Marinesco-Sjögren syndrome. Our data extend the spectrum of Marinesco-Sjögren syndrome-causing *SIL1* mutations, increasing the number of different mutations to 36 and bringing the total number of published molecular-genetically

confirmed families to 46 (88 patients). Based on these data we describe the relative frequency of signs and symptoms of patients with Marinesco-Sjögren syndrome with identified *SIL1* mutations. This information may help to decide which patients should undergo *SIL1* mutation analysis.

We confirm that Marinesco-Sjögren syndrome caused by *SIL1* mutations is a panethnic condition. Some *SIL1* mutations have been found as recurrent events in more than one family and four mutations have been found repeatedly ($\geqslant 3 \times$) in independent families (p.R111X, p.D170fs, p.R317fs, p.F345fs, Table 1).

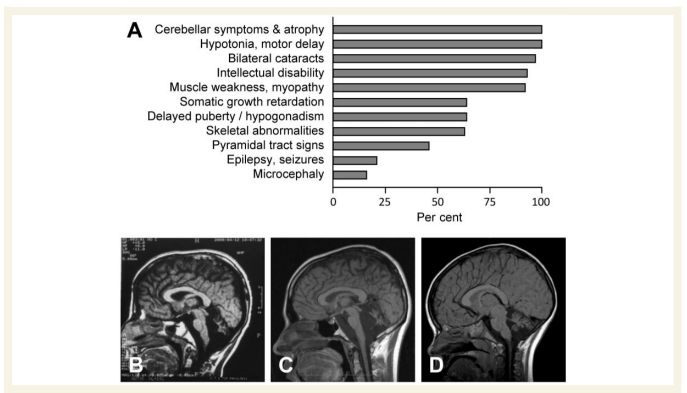


Figure 2 Clinical presentation of patients with Marinesco-Sjögren syndrome with identified SIL1 mutations. (A) Relative frequency of signs and symptoms based on data compiled from our patient cohort and reports from the literature (Anttonen et al., 2005; Senderek et al., 2005; Karim et al., 2006; Annesi et al., 2007; Anttonen et al., 2008; Eriguchi et al., 2008; Riazuddin et al., 2009; Takahata et al., 2010; Terracciano et al., 2012). For details see Supplementary Tables 1 and 2. (B-D) Sagittal MRI scans of three patients with Marinesco-Sjögren syndrome showing marked cerebellar atrophy. (B) Patient MSS25.2 at age 14 years, T₁-weighted image; (C) Patient MSS33.1 at age 24 years, T_1 -weighted image; (D) Patient MSS87.1 at age 3 years, T_1 -weighted image.

However, most mutations are private mutations identified in single families only making diagnostic recommendations for selected screening of exons or testing of selected mutations arguable. Most pathogenic variants identified in this study and reported earlier are micromutations affecting one or a few nucleotides. Moreover, we and others (Takahata et al., 2010) found that the SIL1 mutation spectrum also includes deletions of several kb of DNA. Although it is conceivable that some patients without an identified SIL1 mutation are compound heterozygous for nonoverlapping genomic deletions in the huge SIL1 gene (spanning ~0.25 Mb of genomic DNA), data from whole-genome single nucleotide polymorphism genotyping and copy number variation analysis available for 29 out of the 46 SIL1 mutation-negative patients (A. Roos and S. Spengler, unpublished data) showed no genetic imbalances of the SIL1 gene that is covered by 115 markers on the microarrays used (Affymetrix GeneChip Genome-Wide Human SNP 6.0-Array, Supplementary Fig. 1). Nevertheless, it is probably warranted to extend future SIL1 mutation screening to investigate the presence of pathogenic copy number aberrations. In addition to mutations missed by commonly applied diagnostic strategies there is evidence for locus heterogeneity as well. We excluded linkage or homozygosity to the SIL1 region on chromosome 5 in four families with a typical Marinesco-Sjögren syndrome phenotype confirming that at least one additional locus does exist (Families MSS96, MSS136, MSS144, MSS174). Others have

explored functional candidate genes for mutations in unclarified Marinesco-Sjögren syndrome cases without success (Anttonen et al., 2008).

Although Marinesco-Sjögren syndrome is caused by a wide spectrum of SIL1 mutations, patients present with a relatively homogeneous phenotype and show almost invariably the hallmark clinical features of cerebellar atrophy and ataxia, cataracts, intellectual disability and myopathy (Fig. 2). These somewhat contradictory observations might be reconciled by the notion that all SIL1 mutations are expected to eventually lead to loss of the protein or its critical functional domains. This has been predicted or experimentally confirmed for most nonsense and frameshift mutations, for two splice-site mutations resulting in in-frame deletions (Senderek et al., 2005) and for mutations altering the very C-terminus of the protein (Howes et al., 2012). In the present study we have shown that a SIL1 missense mutation (p.G312R), a two amino acid in-frame deletion (p.V231 I232del) and a frameshift mutation in the last exon, which is predicted to escape nonsense-mediated messenger RNA decay (p.Q426fs), also result in a prominent reduction of SIL1 protein levels in cultured lymphoblasts (Fig. 4). Although we cannot conclude from these data that reduced protein levels are a general consequence of seemingly 'milder' SIL1 mutations, such a mechanism might exist for at least a subset of mutations, and for some it has been experimentally shown that mutant proteins are particularly

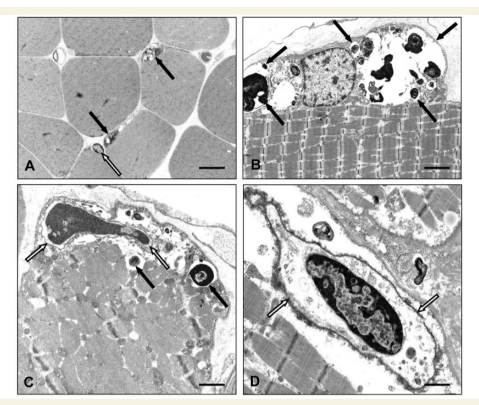


Figure 3 Histopathological and electron microscopic findings in skeletal muscle biopsies of patients with Marinesco-Sjögren syndrome. (A) Patient MSS33.1. Black arrows: abnormal nuclei and autophagic material. White arrow: normal myonucleus. Semi-thin section of glutaraldehyde-fixed, epon-embedded tissue; toluidine blue. Scale bar = 20 µm. (B) Patient MSS87.1. Arrows: autophagic vacuoles in the perinuclear sarcoplasm. Ultra-thin section of glutaraldehyde-fixed, epon-embedded tissue; electron microscopy. Scale bar = 3 µm. (C) Patient MSS87.1. Black arrows: autophagic vacuoles in the perinuclear sarcoplasm of an atrophic muscle fibre. White arrows: perinuclear osmiophilic membrane-like structure characteristic for Marinesco-Sjögren syndrome. The myonucleus shows degenerative condensation of chromatin. Electron microscopy. Scale bar = 2.5 µm. (D) Patient MSS33.1. Arrows: degenerating myonucleus ensheathed by a prominent osmiophilic membrane-like structure. Electron microscopy. Scale bar = 1.5 μm.

unstable and either form large aggregates in the endoplasmic reticulum or are rapidly degraded through the proteasome (Howes et al., 2012).

Our data confirm that the clinical triad of cataracts, cerebellar atrophy and myopathy strongly suggests the presence of SIL1 mutations (detection rate 60%) whereas patients presenting with variant phenotypes are unlikely to have SIL1 mutations (detection rate 2.7%). The only mutation-positive case in the latter cohort was a girl who had not developed cataracts until the age of 4 years (Patient MSS24.1). Two additional patients without cataracts (at age 1 and 4 years, respectively) have been reported (Senderek et al., 2005; Terracciano et al., 2012). As the latest reported onset of cataracts was in a 7-year-old girl in our cohort (Patient MSS142.1), these patients are likely to develop cataracts within the next few years. It has been noted previously that cataracts may develop unusually fast within a few weeks in patients with Marinesco-Sjögren syndrome (Ishikawa et al., 1993). Therefore it might be appropriate to include infants and preschool age children with cerebellar atrophy and myopathy without cataracts in SIL1 mutation screening while the full-blown phenotype including cataracts should be expected after the age of 10 years. Historical descriptions of patients with Marinesco-Sjögren syndrome stressed the presence of profound intellectual disability in most cases (Müller, 1962; Hayabara et al., 1975). More recent reports showed a large spectrum of mental impairment in Marinesco-Sjögren syndrome with most patients exhibiting only mild cognitive impairment and only few cases with severe intellectual deficits. In our cohort of patients with SIL1 mutations we identified six cases with normal intellectual capacities but an otherwise typical Marinesco-Sjögren syndrome phenotype (Fig. 5). Patients without intellectual disability should be considered for SIL1 mutation analysis if their clinical presentation was compatible with the diagnosis of Marinesco-Sjögren syndrome.

The differential diagnosis of Marinesco-Sjögren syndrome includes several conditions featuring combinations of brain, skeletal muscle and eye abnormalities. Congenital cataracts, facial dysmorphism, and neuropathy syndrome shares with Marinesco-Sjögren syndrome the features of cataracts and muscle weakness or hypotonia. The presence of cerebellar atrophy in Marinesco-Sjögren syndrome and demyelinating neuropathy in congenital cataracts, facial dysmorphism, and neuropathy syndrome distinguishes the two syndromes (Lagier-Tourenne et al., 2002). Congenital cataracts, facial dysmorphism, and neuropathy syndrome is caused by a founder mutation in the CTDP1 gene

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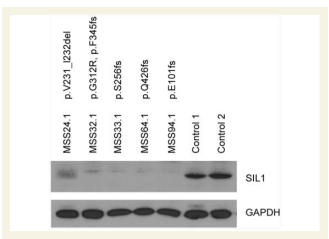


Figure 4 SIL1 mutations result in substantially decreased SIL1 protein levels. Immunoblot of SIL1 in immortalized lymphoblasts of patients with Marinesco-Sjögren syndrome. Cell lysates of Patients MSS24, MSS32, MSS33, MSS64 and MSS94 and two healthy control individuals were immunoblotted with an anti-SIL1 antibody (top) and an anti-GAPDH antibody (as loading control, bottom). SIL1 levels in patients' lymphoblasts were severely reduced compared to the levels in control subjects. Note that seemingly 'milder' mutations [small in-frame deletion (MSS24.1), missense mutation (MSS32.1) and frameshift mutation in the last exon (MSS64.1)] have similar effects as truncating mutations that are expected to lead to nonsense-mediated messenger RNA decay (MSS33.1 and MSS94.1).

(Varon et al., 2003) that seems to be restricted to the Gypsy population. In fact, two patients from our cohort of Marinesco-Sjögren syndrome-like cases, both of Gypsy origin and presenting with congenital cataracts and a history of hypotonia, muscle weakness and elevated creatine kinase levels initially suggesting a myopathic process were found to carry the CTDP1 founder mutation. Careful clinical and neurophysiological re-evaluation revealed a demyelinating neuropathy as the cause of the neuromuscular features (Patients MSS21.1 and MSS73.1, Supplementary Table 3). Cerebrotendinous xanthomatosis is an inherited disorder associated with the deposition of cholestanol in the brain and other tissues and is characterized by progressive cerebellar ataxia, cataracts, and tendineous or tuberous xanthomas (Van Bogaert et al., 1937). The diagnosis can be made by elevated serum cholestanol levels and identification of mutations in the CYP27A1 gene (Cali et al., 1991). Importantly, cerebrotendinous xanthomatosis is a treatable condition warranting proper diagnostic measures to be taken (Berginer et al., 1984). One patient with juvenile cataracts and neurological problems starting during adolescence included in our series (Patient MSS48.1, Supplementary Table 3) was found to have cerebrotendinous xanthomatosis. Other differential diagnoses include ataxia-microcephaly-cataract syndrome (Ziv et al., 1992), cataract-ataxia-deafness-retardation syndrome (Begeer et al., 1991) and VLDLR-associated cerebellar hypoplasia (Schurig et al., 1981; Boycott et al., 2005). Finally, Marinesco-Sjögren syndrome shares clinical features with mitochondrial disorders (Schapira, 2006) and disorders of defective N- and O-glycosylation of proteins (Eklund and Freeze, 2006;

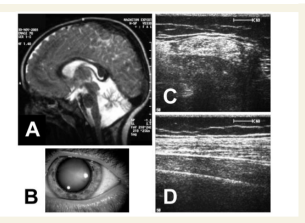


Figure 5 Patient MSS64.1 without intellectual disability but an otherwise characteristic Marinesco-Sjögren syndrome phenotype. (A) Brain MRI scan at age 20 months shows profound atrophy of the cerebellum (T_2 -weighted image). (B) Cataract of the right eye which developed within a few weeks at age 3.5 years. (C and D) Sonography of the M. quadriceps at the age of 6.75 years shows an increase in echogenicity, especially in the area of the M. rectus femoris with poorly identifiable structures of fascia and lack of echogenicity of the bone.

Muntoni *et al.*, 2011). One of the *SIL1* mutation-negative patients in our series (Patient MSS53.1, Supplementary Table 3) was diagnosed with Sengers syndrome, a form of mitochondrial DNA depletion disorder caused by an *AGK* gene mutation (Calvo *et al.*, 2012; Mayr *et al.*, 2012).

To our knowledge, there are no comprehensive screening studies that have explored a potential role of *SIL1* mutations in cohorts of patients with non-syndromic, isolated myopathy, cataracts or ataxia. However, we assume that *SIL1* has only a minor relevance in the screening of these patients. Data obtained using whole-exome sequencing and next generation sequencing ataxia panels (including all known genes for isolated or syndromic ataxias) did not reveal clearly pathogenic *SIL1* alleles in a cohort of >90 patients with early-onset ataxia that had already been screened negative for all common early-onset ataxia genes (i.e. *FXN*, *AOA2*, *AOA1*, *POLG*), thus highly enriched for mutations in rarer early onset ataxia genes (M. Synofzik and P. Bauer, unpublished data).

Muscle biopsies obtained from patients with Marinesco-Sjögren syndrome show various non-specific signs indicating degeneration of skeletal muscle fibres. At the ultrastructural level, degenerating myonuclei are occasionally surrounded by an electron-dense, membrane-like structure (Herva *et al.*, 1987; Sewry *et al.*, 1988). We have confirmed this finding in all muscle biopsies of patients with Marinesco-Sjögren syndrome that were available for electron microscopy in this study (n = 4; MSS33.1, MSS87.1, MSS91.1 and MSS94.1) and in our earlier report (n = 6) (Senderek *et al.*, 2005) suggesting that this finding is a characteristic feature related to SIL1 pathology. In general, muscle biopsies are considered invasive procedures that are best avoided if a diagnosis can be established using other methods. Although with the availability of *SIL1* sequencing there is no longer a diagnostic need

for muscle biopsies in patients with Marinesco-Sjögren syndrome with typical symptoms, this procedure still may be important in the diagnosis of Marinesco-Sjögren syndrome, especially in less clearcut cases. First of all a muscle biopsy can be helpful in differentiating myopathy from other causes of hypotonia and weakness. Secondly, the full Marinesco-Sjögren syndrome phenotype might not be seen in very young children as cataracts tend to develop later making the differentiation from other causes of a cerebellar syndrome and hypotonia more difficult. In addition, as parts of the EMG procedure (to establish a myopathy) are uncomfortable and not well tolerated by young children, results might be inconclusive. In this scenario, electron microscopy of a muscle biopsy unveiling vacuolar myopathy and the characteristic perinuclear membranelike structures can help to consider Marinesco-Sjögren syndrome as a differential diagnosis and can direct to SIL1 mutation analysis (Terracciano et al., 2012).

The current study contributes to our understanding of the clinical and genetic basis of Marinesco-Sjögren syndrome. A molecular diagnosis was obtained in ~60% of patients from a screening cohort of 25 unrelated index patients with a full-blown Marinesco-Sjögren syndrome phenotype leaving a considerable proportion of patients without genetic diagnosis to date. This further underscores the fact that other still unknown mutations must exist in known or unknown disease associated genes. As a co-chaperone for BiP, SIL1 is involved in protein synthesis and quality control in the endoplasmic reticulum. Thus, secretory pathway proteins not properly processed or not reaching their site of action in SIL1 deficient cells will represent plausible functional candidates. No such factors have been identified so far, but appropriate tools for proteomic analysis of cell compartments and cell surface proteins are now at hand. Combining such data with data obtained by high-throughput DNA sequencing projects in non-SIL1 Marinesco-Sjögren syndrome families will likely disclose the missing Marinesco-Sjögren syndrome gene(s) within the next few years and may also hold promise to identify potential therapeutic targets.

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Supplementary material

Supplementary material is available at Brain online.

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