

# A frameshift mutation in *LRSAM1* is responsible for a dominant hereditary polyneuropathy

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Despite the high number of genes identified in hereditary polyneuropathies/Charcot–Marie–Tooth (CMT) disease, the genetic defect in many families is still unknown. Here we report the identification of a new gene for autosomal dominant axonal neuropathy in a large three-generation family. Linkage analysis identified a 5 Mb region on 9q33–34 with a LOD score of 5.12. Sequence capture and next-generation sequencing of the region of interest identified five previously unreported non-synonymous heterozygous single nucleotide changes or indels, four of which were confirmed by Sanger sequencing. Two sequence variants co-segregated with the disease, and one, a 2 bp insertion in the last exon of *LRSAM1*, was also absent in 676 ethnicity-matched control chromosomes. This frameshift mutation (p.Leu708Argfx28) is located in the C-terminal RING finger motif of the encoded protein. Ubiquitin ligase activity in transfected cells with constructs carrying the patient mutation was affected as measured by a higher level of abundance of TSG101, the only reported target of *LRSAM1*. Injections of morpholino oligonucleotides in zebrafish embryos directed against the ATG or last splice site of zebrafish *Lrsam1* disturbed neurodevelopment, showing a less organized neural structure and, in addition, affected tail formation and movement. *LRSAM1* is highly expressed in adult spinal cord motoneurons as well as in fetal spinal cord and muscle tissue. Recently, a homozygous mutation in *LRSAM1* was proposed as a strong candidate for the disease in a family with recessive axonal polyneuropathy. Our data strongly support the hypothesis that *LRSAM1* mutations can cause both dominant and recessive forms of CMT.

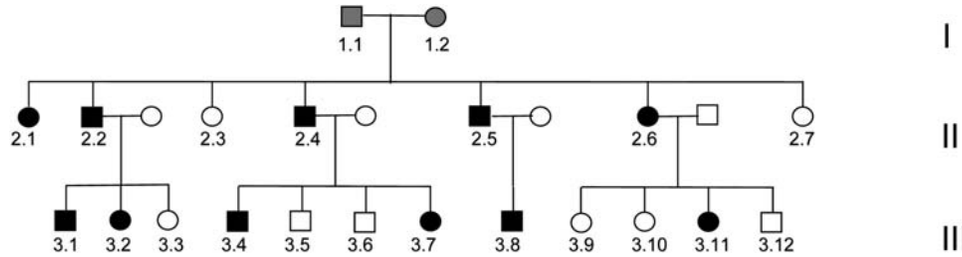
## INTRODUCTION

Hereditary motor and sensory neuropathies (HMSN), also known as Charcot–Marie–Tooth (CMT) disease, consist of a group of common inherited peripheral neuropathies that are heterogeneous at the clinical and genetic levels. Clinically, they can be divided into demyelinating and axonal forms of the disease, primarily based on electrophysiological data. Demyelinating neuropathies show a marked decrease in

median motor nerve conduction velocities (NCVs). The most frequent type of CMT with an autosomal dominant mode of inheritance, CMT1A, is most commonly caused by duplication of *PMP22*, although mutations and copy number variants in the vicinity of *PMP22* are also described (1–6).

CMT2 is an axonal type of CMT with reduced compound motor axon potentials but initially no reduction in NCVs. However, the clinical features of patients of both types can be quite similar. X-linked forms of CMT also exist as well

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**Figure 1.** Pedigree of family with dominant axonal neuropathy. Filled symbols indicate disease, and open symbols represent unaffected persons. Grey symbols: not known if affected.

as subtypes with intermediate forms of NCVs and/or recessive modes of inheritance.

Many loci have been linked to CMT/HMSN and to date 44 genes have been implicated in the disease; mutations in 15 different genes have been identified to cause CMT2. Except for the *PMP22* duplication that is responsible for CMT1A, mutations in other CMT genes occur at relatively low frequencies. Despite the high number of loci, the defect in many families remains unknown thereby precluding any genetic testing.

Most genes are associated with either demyelinating or axonal forms of subtypes of the disease, although *GJB1* and *MPZ* have been associated with both (7–9). Severity of the disease varies between different individuals, even when an identical mutation is present in members of the same family. So far, only few clinical characteristics have been linked to a specific gene or mutation, thus presenting a diagnostic tool to determine which genes might be screened first. Here we report on a three-generation family with axonal neuropathy (CMT2) and a clear dominant mode of inheritance. Linkage analysis followed by next-generation sequencing of the linkage region on chromosome 9q33–34 and functional analysis led to the identification of the causative gene for this dominant type of CMT. We propose CMT2o for dominant LRSAM1-associated CMT2.

## RESULTS

### The clinical phenotype is associated with the chromosomal region 9q33–34

The index patient of Dutch descent presented at the age of 32 with progressive distal muscle weakness and mild sensory disturbances of both feet. Over decades, his symptoms were slowly progressive now also including weakness of hand muscles. He had relatively mild foot deformities with pes cavus and hammer toes and a marked atrophy of his lower legs with absent tendon reflexes. The vibration sense was mildly reduced, showing a proximal to distal gradient. His family history revealed various other family members with the same features (Fig. 1; Table 1). Nerve conduction studies and needle EMG showed a relatively severe axonal neuropathy with absent sural nerve sensory nerve action potentials, mildly reduced motor NCVs and severe distal neurogenic features on needle examination. Over two decades ago, a sural nerve biopsy was performed that showed severe axonal degeneration confirming the diagnosis

of CMT2. The clinical phenotype can be summarized as a relatively mild, very slowly progressive axonal neuropathy with age of onset in the second or third decade of life. The pedigree of the family clearly showed an autosomal dominant inheritance of the disease.

Mutations in the coding regions of *MFN2*, *MPZ*, *GJB1* and *GARS*, which are known to be associated with CMT2, were excluded. In view of the large number of CMT2-associated genes and the availability of many affected individuals in this family, we decided to locate the genetic defect by genome-wide linkage analysis using an Affymetrix 250K single nucleotide polymorphism (SNP) array. Due to the large number of family members, the family was split into two smaller pedigrees for fast multipoint parametric analysis using the ALLEGRO (10) program with the assumption of autosomal dominant inheritance and full penetrance. Analysis showed a single peak on chromosome 9 with a maximum LOD score of 4.52 (Fig. 2). When confining the analysis to markers on chromosome 9 using SimWalk2 (11), multipoint LOD scores could be calculated for the family as a whole resulting in a maximum LOD score of 5.12. Therefore, the single critical interval on 9q33–34, previously unknown as a CMT locus, of approximately 7 cm must harbor the causative mutation for the disease. Consequent construction of haplotypes with ALLEGRO revealed a specific haplotype that showed perfect co-segregation with the disease (Fig. 3). The borders of this haplotype shared between all affected individuals were defined by two limiting SNPs (rs10081624 and rs2039463) which are 4.4 Mb apart.

### Identification of heterozygous changes associated with the clinical phenotype

We designed a custom sequence capture array covering the entire linkage region (chr9: 127624486–132419012 hg18/NCBI36). Genomic DNA of an affected individual (subject 2.2) was enriched by sequence capture for this region, sequenced on a FLX Titanium sequencer (Roche) and mapped to the human consensus genome (hg18) with Newbler software. The captured region contains 78 annotated genes, 10 loci encoding predicted proteins and 2 microRNAs. Over 94% of all exons were sequenced with coverage of more than 10 times. Analysis yielded 5827 reliable nucleotide changes which is within the estimated range of nucleotide changes to be expected for a 5 Mb region. Since the mode of inheritance is autosomal dominant, the expected disease-causing mutation would be a heterozygous change. Within

Table 1. Characteristics of family members

Clinical features Code in pedigree	Affected family members										
	1.2	2.1	2.2	2.5	2.6	2.4	3.1	3.8	3.2	3.4	3.11
Gender	F	F	M	M	F	M	M	M	F	M	F
Age at onset	40	Unknown	30	30	37	40	27	20	28	No complaints	No complaints
Age	died at the age of 72	70	68	64	62	63	34	32	32	26	25
Distal muscle wasting/weakness (LL)	-	Unknown	+	+	+	+	+	+	+	+	-
Distal muscle wasting/weakness (UL)	-	Unknown	+(mild)	+(mild)	+	+	+	+	+	+	-
Upper limb reflexes	-	Unknown	N	-	Reduced	Reduced	Reduced	Reduced	N	N	Reduced
Knee reflexes	-	Unknown	-	-	-	-	Reduced	Reduced	N	N	Reduced
Ankle reflexes	-	Unknown	-	-	-	-	-	-	N	N	Reduced
Vibration sense	Unknown	Unknown	Reduced	Reduced	Reduced	Reduced	Reduced	Reduced	Reduced	ND	N
Pin-prick	Mildly reduced	Unknown	Mildly reduced	Mildly reduced	Reduced	Reduced	Reduced	Reduced	N	ND	N
Hearing loss	Unknown	Unknown	-	Mildly perceptible	-	-	-	-	-	-	-
Additional features	None	None	Parkinson	None	None	Parkinson	None	None	None	None	None
Nerve conduction studies	ND	+	+	+	+	+	+	+	+	+	+
Sural nerve response	-	-	-	-	-	-	-	-	-	-	-
Peroneal nerve response	-	-	-	-	Strongly reduced	-	N	ND	Reduced	N	ND
Needle EMG	-	ND	Neuro	Neuro	Neuro	Neuro	Neuro	ND	Neuro	Neuro	Neuro

-, absent; +, present; F, female; M, male; N, normal; ND, not done; Neuro, neurogenic.

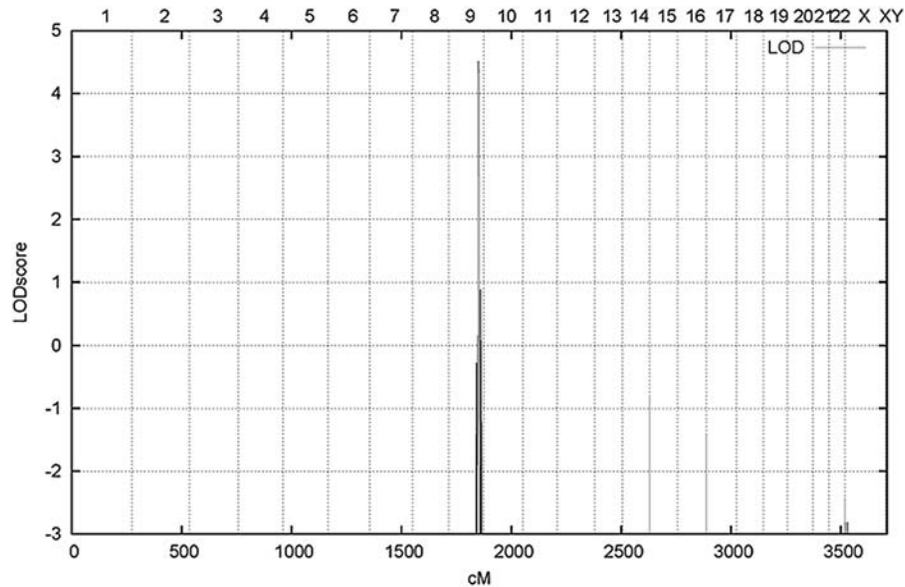
annotated genes, we found 330 nucleotide variations, 51 of these were heterozygous changes within exons or within 20 nucleotides of the exon-intron boundaries. The majority of sequence variants were already known as polymorphisms in the public databases. Five heterozygous changes were not known as polymorphisms and were located in the coding region of the corresponding genes, two of which were frameshift mutations, the other three being missense mutations (Table 2; Supplementary Material, Table S2). The sequence variants in *GOLGA2*, *SLC27A4*, *WDR34* and *LRSAMI* were all confirmed by Sanger sequencing, whereas the frameshift in *ST6GALNAC6* represented a pyrosequencing artifact due to the presence of homopolymer stretches.

**LRSAMI is a candidate gene for CMT2**

For segregation analysis, the presence of the four validated changes was determined by direct sequencing in seven additional family members (five affected and three non-affected persons). The missense mutation in *WDR34* and the frameshift mutation in *LRSAMI* both segregated with the disease. The other two missense mutations in *GOLGA2* and *SLC27A4* were variations located on the non-disease-associated allele. They were both found in two of three healthy individuals and, in addition, were absent in four of five affected individuals and should therefore be regarded as rare polymorphisms not related to the disease. Prediction programs (Polyphen, SIFT) indicated that the change in *WDR34* was not expected to be harmful to the protein. Moreover, we identified two heterozygotes in 352 ethnicity-matched controls. In contrast, the 2 bp insertion in *LRSAMI* was absent in 676 control chromosomes. Screening of all available family members, 11 affected and 8 non-affected individuals, showed complete co-segregation with the disease. The 2 bp insertion leads to a replacement of a leucine by an arginine residue near the predicted C-terminal RING finger domain and a concomitant frameshift with a stop codon 28 codons downstream (p.Leu708Argfx28). This would make the mutant protein slightly longer than the wild-type one, since normal translation would terminate only 16 amino acids downstream of this position. Thus, *LRSAMI* is a very strong candidate gene for the neuropathy in this family. In addition, it was published recently that another mutation in *LRSAMI* in a Canadian family was thought to be the cause for a recessive form of CMT2 (12). To determine if the found mutation of the Dutch family would possibly occur more frequently in Dutch CMT2 pedigrees, 11 patients as representatives of 11 additional CMT2 pedigrees in which no genetic aberration had been identified as well as 32 unrelated CMT1 patients without known mutations were screened for this specific insertion but we could not identify other Dutch families carrying the identical mutation.

**LRSAMI is highly expressed in the fetal and adult nervous system**

We examined *LRSAMI* expression by using locked nucleic acid (LNA) probes in RNA *in situ* experiments in neural tissue on sections of a 6-8-week-old human fetus and adult spinal cord. Motoneurons in the human adult spinal cord



**Figure 2.** Linkage analysis identifies 9q33–34 as the disease gene-containing region: LOD scores are plotted against the position (cM) on the chromosomes from 1–22, XX, XY and aligned from p-ter to q-ter.

showed high *LRSAMI* expression (Fig. 4) and a clear signal was also seen in some intermediate neurons (data not shown). Expression in a human fetus was found to be high in the fetal spinal cord, sensory ganglia and all developing muscles but was also observed in some other tissues such as the esophageal endothelium, epithelium, Meckel's cartilage and at a location that probably contains osteoblasts in the developing pharyngeal arches. Although expression is not restricted to neural tissue, a high *LRSAMI* expression is present in all tissues that are affected by the disease (Fig. 4).

#### Interference with *Lrsam1* in zebrafish embryos results in a neurodevelopmental phenotype

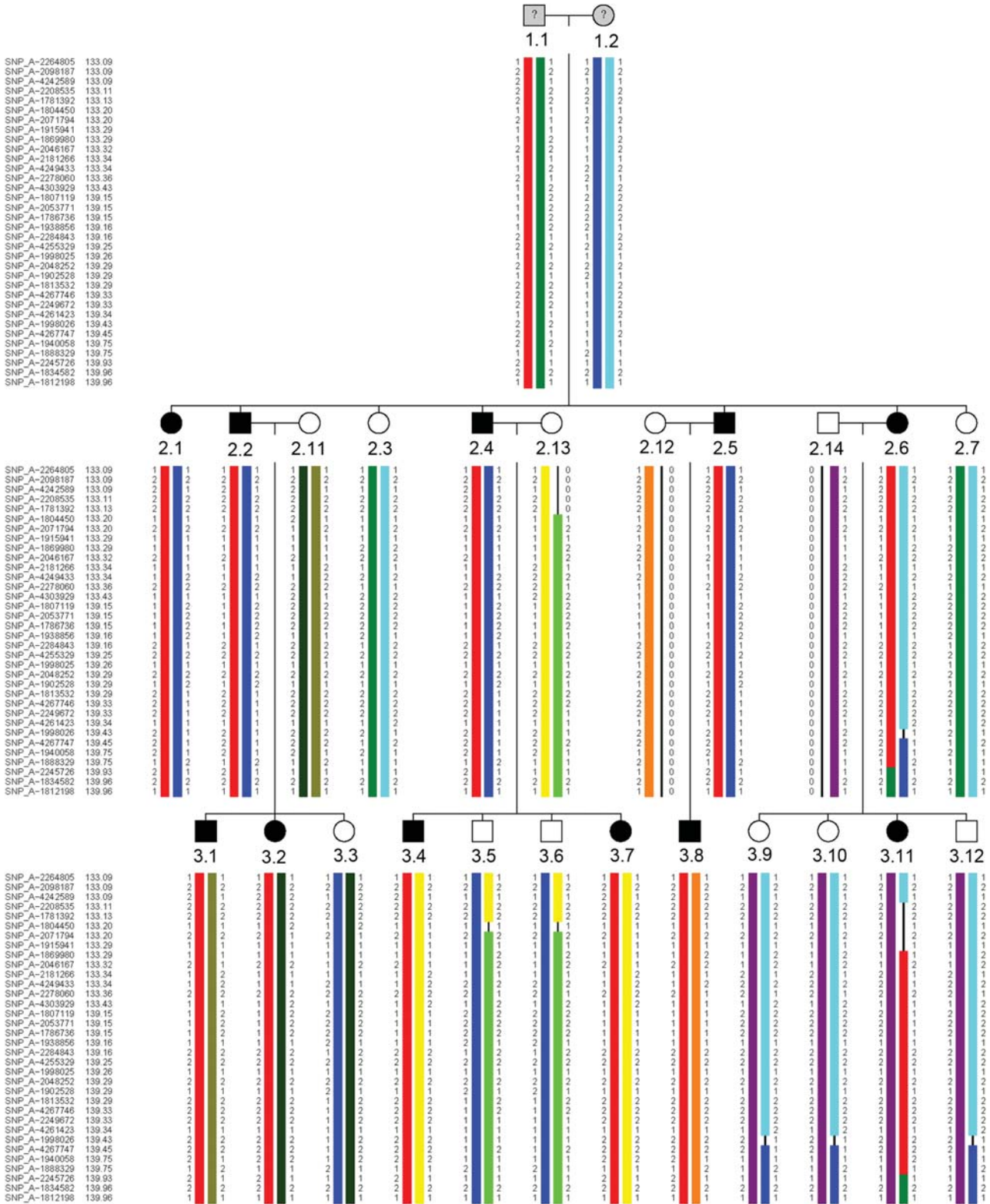
To study the role of *LRSAM1* in neurodevelopment, we injected zebrafish embryos with morpholino oligonucleotides (MOs) to target the *Lrsam1* gene (ENSDARG0000060683). We used MOs directed against the ATG to obtain a knock-down phenotype. We designed a splice site MO targeting the last exon, thereby mimicking the situation as found in our patients as closely as possible. MOs directed against splice sites usually lead to exon skipping but we expected that cryptic splice acceptor sites present at a short distance from the original site would be activated, resulting in a protein that is slightly different at the C-terminal end but not prone to nonsense-mediated decay similar to the situation in the patients. RT-PCR followed by sequence analysis on zebrafish embryos indeed showed that an alternative acceptor splice site at 39 nucleotides upstream of the normal acceptor site was used (data not shown) leading to a frameshift followed by a stopcodon after 13 codons, replacing the last 43 codons of the wild-type transcript.

Several injection experiments were performed to obtain the optimal concentration for the MO used. After 6 h post-fertilization (hpf), a delay in development was readily observed for the ATG morphants and to a lesser extent for

the splice MO-injected embryos, whereas it was hardly present if at all for the control MO-injected embryos (Fig. 5). After 24 hpf, death had occurred in a relatively high percentage and most of the surviving ATG and splice morphants were clearly affected. Only a low percentage of death and abnormalities was seen for the wild-type non-injected or control injected embryos. The phenotype at 2–3 days post-fertilization (dpf) varied from near-normal with a slightly smaller head, a slightly shorter body axis, slightly less pigmentation and bent tail tips to completely curled up and smaller embryos with bent tails with little pigmentation, smaller eyes, abnormal brain development and a less organized structure of the somites. The morphology of all MO-injected embryos at 54 hpf is given in Figure 5, typically affecting about 70–80% of the embryos with about one-third severely affected. The data for six independent experiments are given in Table 3. Despite the observed variation in the severity of the phenotype, the injections reproducibly gave similar results with a phenotype that was comparable between ATG and splice morphants. When allowed to develop for a longer time (experiment 1), the majority of morphants would not hatch. When excluding the most severely affected morphants which obviously would also show abnormal swimming behavior, we observed quite a high percentage with swimming abnormalities which was expected for the ones with curly tails but was also seen in the morphants that looked (near) normal. While wobbly swimming behaviour may partially be related to a developmental delay, some embryos showed no normal response at all when repeatedly touched using a tip of a pipette.

#### The abnormal phenotype in zebrafish is associated with abnormal motoneuron development

As CMT2 is an axonal neuropathy, the observed phenotype in zebrafish morphants would probably originate in neural



**Figure 3.** Haplotypes of affected and non-affected family members. Haplotypes were inferred for the grandparents. Different haplotypes were given a different color. All affected persons share the red haplotype.

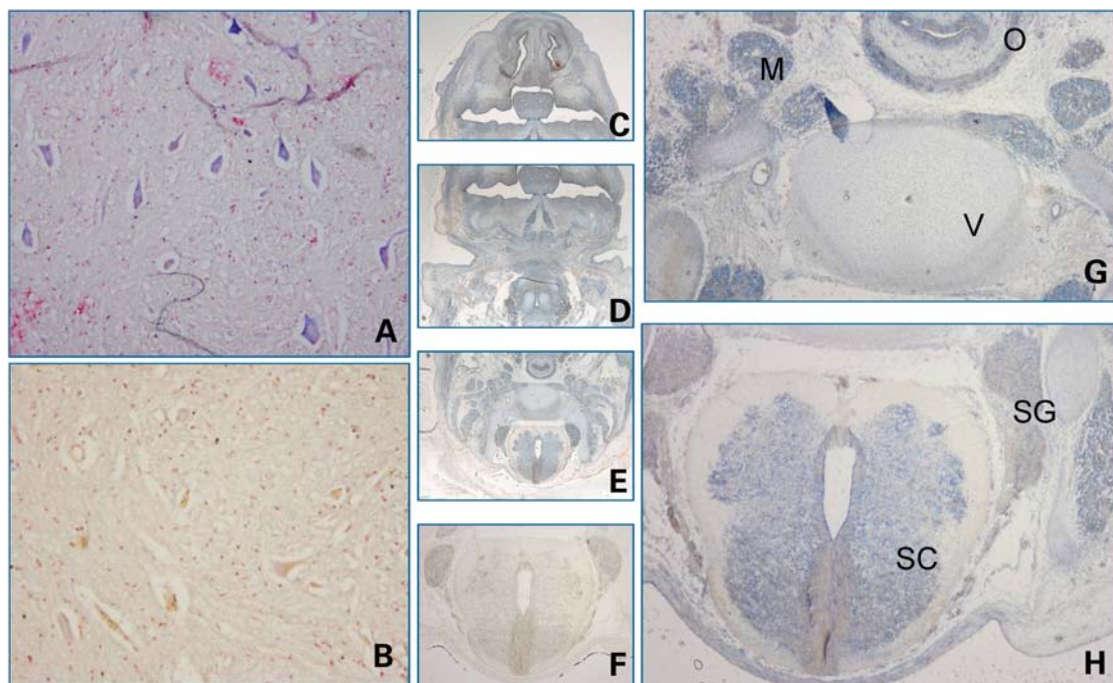
abnormalities. To explore this possibility, we used an antibody, *znpl1*, that detects caudal and middle motoneurons and Rohon Beard primary neurons. Immunohistochemical staining

of embryos with a slightly to moderately affected phenotype of 30 hpf to 4 dpf showed several abnormalities. Severely affected embryos displayed a very weak staining due to a

**Table 2.** Found heterozygous changes

GENE	Pos. chr9 (hg18)	Change	NT	Amino acid	Exon	Coverage	% mutant reads
<i>GOLGA2</i>	130059259	G>A	c.2881C>T	p.Pro961Ser	26	34 reads	47
<i>SLC27A4</i>	130155189	A>G	c.1052A>G	p.Asn351Ser	8	62 reads	56
<i>WDR34</i>	130435956	G>A	c.1499G>A	p.Ala500Val	9	50 reads	50
<i>ST6GALNAC6</i>	129692883–884	CG>-	c.557_558delGC	p.Pro186Gln fsX38	5	63 reads	27
<i>LRSAMI</i>	129304950–951	->GC	c.2121_2122insGC	p.Leu708Arg fsX28	26	20 reads	70

The position (pos.) on the chromosome is given, as well as the consequence for the gene involved at the nucleotide level (NT) and the protein level.



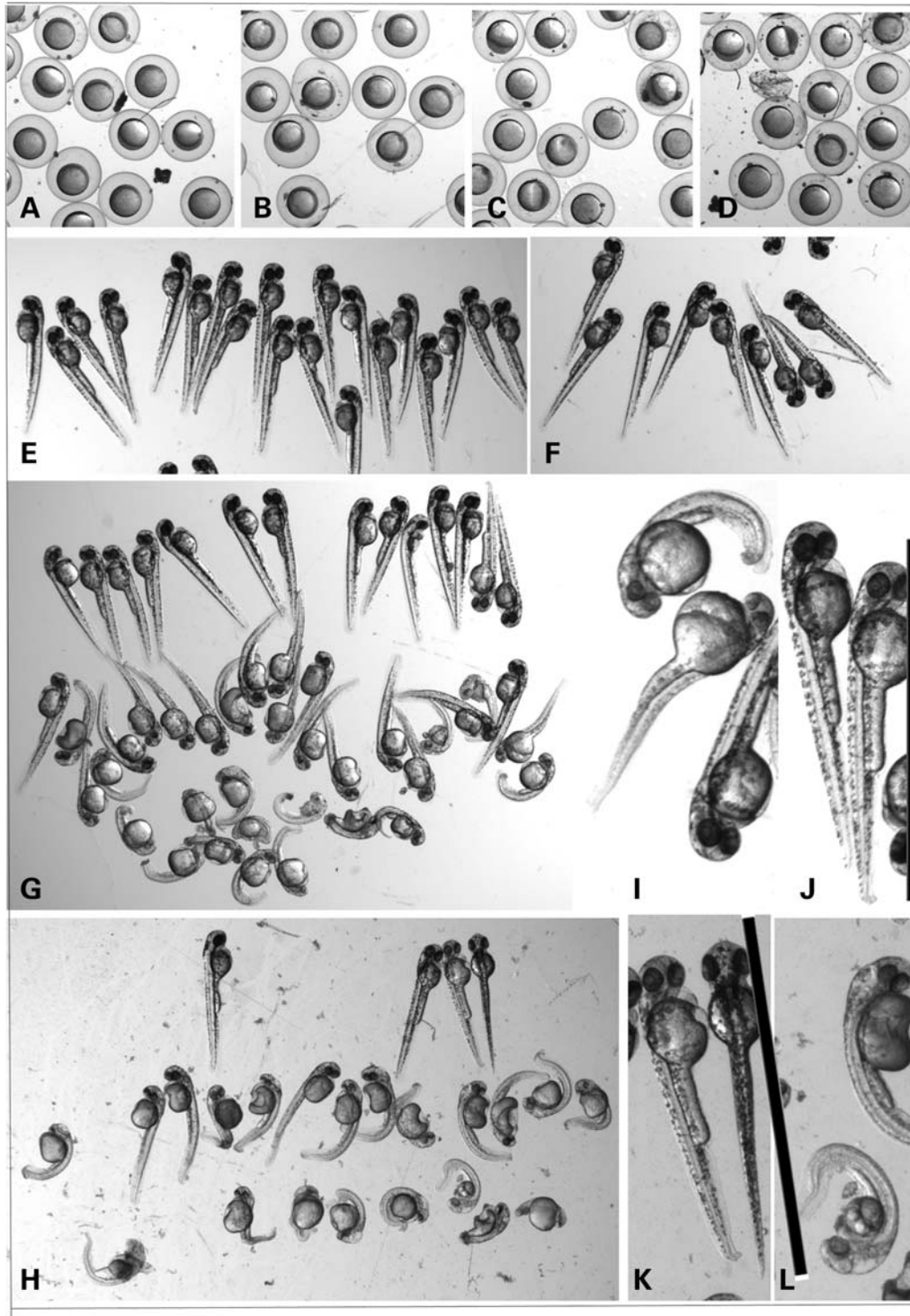
**Figure 4.** *LRSAMI* shows a high expression in neural and muscle tissue. RNA *in situ* hybridization using LNA probes on (A) adult spinal cord (10 × 10 magnification), (B) negative control on spinal cord; (C–H): cross-sections of human fetus 6–8 weeks, approximately at a level between the heart and the brain. (C–E) Overlapping pictures at 1.6 × magnification, (F) negative control on the fetal spinal cord and (G and H) 4 × 10 magnification of specific parts of (E). Detection of RNA was visualized by the blue color, and counterstaining was performed with fast Nuclear red. M = muscle, V = developing vertebra, O = oesophagus, SC = spinal cord, SG = sensory ganglia

completely disorganized structure. In less affected embryos, staining seems reduced for both ATG and splice morphants and, in addition, showed a more irregular structure in contrast to the wild-type staining (Fig. 6). Motoneurons normally migrate halfway over the midline of the somite and then develop more extensive arborizations. This structure is less organized in the morphants. When examining older embryos at 4 dpf, a time point where the effects of the MOs are likely to have worn off, staining intensity increases to a level comparable to the controls, but the neural structure remains less well organized (Fig. 6, bottom panels). The observed neuronal aberrations probably explain at least a large part of the phenotype also affecting movement and swimming behavior.

#### The mutation in *LRSAMI* affects its function

*LRSAMI* encodes a multidomain protein and has been shown to possess E3-ubiquitin ligase activity owing to the C-terminal

RING finger domain (13). E3-ubiquitin ligases constitute a large family of enzymes that can modify their target proteins through covalent attachment of ubiquitin molecules, which will promote their proteosomal degradation. The product of tumor suppressor gene 101, TSG101, is the only *LRSAMI* target reported to date. Since the CMT2-associated mutation is located at the border of the C-terminal RING finger domain, it could perturb its E3 ligase activity. To test this hypothesis, HEK293T cells were co-transfected with *LRSAMI* and *TSG101* expression constructs. The introduction of a RING-inactivating mutation in *LRSAMI* which completely abolishes its E3-ligase activity (p.C675A) (13) resulted in an increased abundance of exogenously introduced TSG101, despite equal transfection efficiencies (Fig. 7). Similarly, the introduction of the patient *LRSAMI* mutation increased the abundance of the tagged TSG101 to a level even higher than that observed with the C675A mutant, thus supporting the notion that E3-ligase function of the CMT2-associated mutated *LRSAMI* is affected.



**Figure 5.** Phenotype of MO-injected zebrafish embryos at 6 and 54 hpf. (A) Wt embryos at 6 hpf and embryos injected with a control MO (B), ATG MO (C) or splice MO (D). (E–H) Wt embryos at 54 hpf (e) and (F–H) embryos injected with a control MO (f), ATG MO (g) or splice MO (h) at 54 hpf. Magnification used was 0.8 $\times$ . (I–L) A further magnification of the same ATG (I and J) and splice (K and L) MO-injected embryos where the black bar indicates the size of a wild-type embryo.

## DISCUSSION

We identified a mutation in *LRSAMI*, c.2121\_2122insGC, p.Leu708ArgfsX28, causative for CMT in a large three-generation

family with axonal neuropathy. The 2 bp insertion in the last exon of this gene results in the exchange of the last leucine by an arginine at the border of the C-terminal RING finger motif (amino acids 674–713). In addition, it leads to a

**Table 3.** Injection experiments of zebrafish embryos.

Exp.	MO	Conc. ( $\mu\text{M}$ )	#	%† 24 hpf	%† final	morphology and hatching	swimming
2	ATG	750a	47	44	65	2 NN, 9 ABN (5 SA)	1/2 NN and all ABN swim abnormal
1	ATG	600	31	48	58	6/13 non-hatched and SA	ND
3	ATG	550	66	41	59	6 NN, 15 ABN (5 SA)	2 N swimmers rest ABN
4	ATG	550	60	20	20	13 NN, 35 ABN (10 SA)	4 N swimmers
2	ATG	500	62	6.5	17	30 NN, 6 ABN (2 SA)	14 ABN swimmers
5	ATG	450	75	13	17	All N except 7 ABN 3 SA)	10 ABN swimmers, rest N
6	ATG	450	82	15	16	All N except 10 weakly affected	13 abnormal swimmers, rest normal
1	ATG	300	32	15	15	All N, 2/27 w abnormal tails	ND
1	splice	800	41	100	100	All dead	ND
1	splice	400	46	39	60	1 N, 1 curly tail, rest ABN	ND
2	splice	300	51	63	67	1 N, 11 ABN (5SA, 2 weak)	1 N one swims abnormal
3	splice	275	67	50	51	6 NN, 21 ABN (12 SA)	2 N swimmers, rest ABN
4	splice	275	51	47	49	4 NN 21 ABN (8 SA)	3 N swimmers, rest ABN
2	splice	250	44	38	65	1 N, 12 ABN	ABN ones swim ABN
5	splice	200	138	30	33	30 NN, 54 ABN (15 SA)	2–3 swim N, rest wobbly to ABN
6	splice	200	86	14	18	44 NN, 24 ABN (3 SA)	16 ABN swimmers, 3 N, rest wobbly
1	como	800	39	0	0	All N	ND
3	como	550	39	5.1	9.1	All N except one with curly tip of tail	ABN one swims ABN
4	como	550	25	0	0	All N except 1	5 ABN swimmers
2	como	500	40	0	3.7	All N except 2 with pigtails	2 ABN ones move a bit do not swim
5	como	450	42	19	2.1	All N except 3 (1 curly, 1 curly tail tip)	ABN one swims ABN
6	como	450	23	22	26	All N except 1 with slightly curly tailtip	ABN one swims ABN
1	–(wt)		33	60	6.0	All N	ND
2	–(wt)		22	4.5	4.5	All N	All swim N
2	–(wt)		40	2.5	4.1	All N	All swim N
3	–(wt)		75	1	4.6	All N except 1 with curly tail tip	Curly one swims ABN
4	–(wt)		41	2.5	2.5	All N except 1 with curly tail	4 ABN swimmers
5	–(wt)		114	11	12	All N except 3 curly (2 body, 1 tip)	3–4 ABN swimmers
6	–(wt)		45	6.6	7.1	All N except 1	ABN one swims ABN

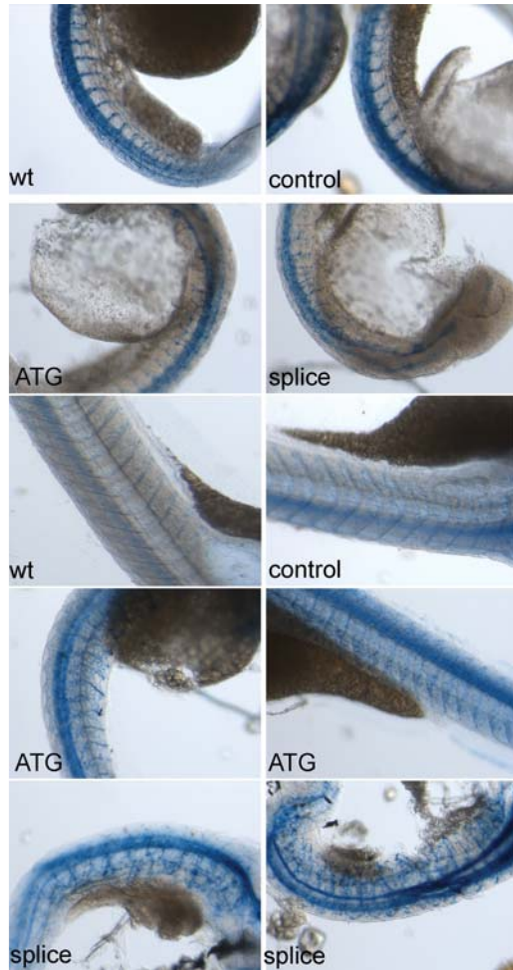
Embryos in experiment (exp.) 1 were not dechorionated, but scored for the percentage of hatching and sacrificed at 4 dpf, embryos in exp. 2–4 were dechorionated at 48 hpf and sacrificed at 54 hpf, embryos in exp. 5 and 6 were dechorionated at 72 hpf if necessary and sacrificed at 78 hpf. For staining purposes, six embryos were removed at 30 hpf (splice), 6–16 (ATG), 6–12 embryos (como) and 10–16 (wt) (exp. 2 and 3). Removed embryos and embryos damaged by dechorionation were excluded from final counts. 750a: Volumes of 1 nl were injected except for exp. 2 in which the estimated injection size was 1.15 nl which equals 1 nl of a solution of 750  $\mu\text{M}$ . ABN, abnormal; NN, near normal; A, clearly affected; SA, severely affected. †, Death.

frameshift predicted to result in a protein that is 12 amino acids longer than the normal protein. Since the mutation is located in the last exon, the mutant mRNA would not be prone to degradation by nonsense-mediated decay, which may explain the dominant nature of the phenotype. As expected, both mRNAs were found to be present at comparable levels in patient fibroblasts, as determined by RT-PCR followed by the sequence analysis of cDNA fragments containing the mutation site (Supplementary Material, Fig. S1). In contrast, the mutation identified in a Canadian family as the probable cause for CMT2 was a homozygous mutation in the consensus splice acceptor AG dinucleotide of the penultimate exon into an AA dinucleotide which leads to degradation by nonsense-mediated decay of the mRNA and the absence of the protein which was confirmed in EBV-transformed B-lymphocytes of one of their patients. The described clinical phenotype of the Canadian family was that of a relatively mild autosomal recessive CMT2 quite similar to the phenotype found in our family, albeit the mode of inheritance is different. This phenomenon has also been reported for other CMT-associated genes. Mutations in *HSP27/HSPB1* can cause both dominant and recessive forms of CMT2 (14). Likewise, most mutations in *GDAP* that have been reported (15,16) are associated with autosomal recessive

CMT disease (CMT4A) but some cause a dominant form of the disease (17).

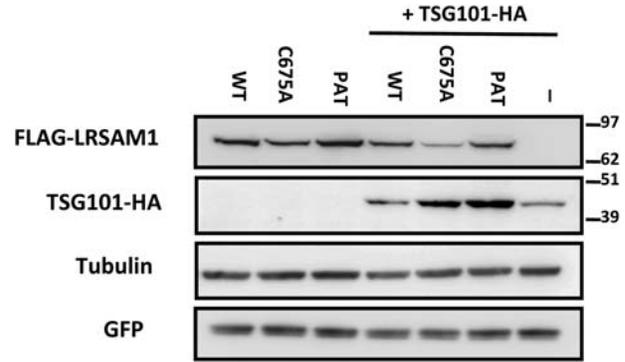
The phenotype we observed in zebrafish was also similar when comparing two different antisense morpholinos directed against *Lrsam1*, one blocking the transcription start site resembling a null mutation and the other one blocking splicing at the 3' end of the messenger, thereby mimicking the patient mutation described here. When screening the sequence of the used morpholinos against other zebrafish transcripts, no other mRNA was remotely similar when allowing up to four mismatches. With the concentrations used, it seems unlikely that the morphological effects and consequences on zebrafish and their ability to respond and move with underlying deficient and less organized motoneurons would be due to a block of other unrelated messengers, coincidentally in both cases leading to the same neurodevelopmental phenotype. Recently, another CMT2 model was described in zebrafish using mitofusin morpholinos (18) with a phenotype quite strikingly similar to *Lrsam1* morphants.

*LRSAMI* (leucine-rich repeat and sterile alpha motif-containing protein 1) was reported independently by two groups who named the gene *TAL*, for TSG101-associated protein, and *RIFLE*, which stands for RING finger and leucine repeat-rich protein, respectively (13,19). The protein



**Figure 6.** Znp-1 staining in MO-injected zebrafish embryos. Embryos were non-injected or injected as indicated with control MO, ATG MO or splice MO. The upper four pictures show 10 $\times$  magnifications of zebrafish at 30 hpf (injection exp. 2), the bottom six panels show 10 $\times$  magnifications of zebrafish at 4 dpf (injection exp.1).

was initially implicated in PC12 cells in E-cadherin-mediated cell adhesion and WNT signaling. Subsequent studies showed that it controls sorting of ubiquitinated cargo and viral budding, which is attributed to ubiquitinylation of TSG101, a component of the ESCRT-I (endosomal-sorting complexes required for transport) sorting machinery (20). Ubiquitinylation by LRSAM1 or Mahogunin RING finger 1 (MGRN1), another RING finger E3-ubiquitin ligase (21), is an important post-translational determinant for expression of TSG101. The found CMT2-associated mutation that is located in the C-terminal RING domain of LRSAM1 required for ubiquitinylation indeed affects TSG101 abundance in co-transfected cells akin to what we observed when using the RING finger dead *LRSAM1* mutant C675A (13) which acts as a non-functional allele in this assay. Although it is currently unknown if the readout for LRSAM1 function used here is the (most) relevant aspect ultimately leading to the disease, we can conclude from Figure 7 that the introduction of a high expression of wild-type *LRSAM1* does not significantly change the level of exogenously introduced TSG101 when comparing it with the



**Figure 7.** Mutation of the RING domain of LRSAM1 affects the abundance of its target TSG101. HEK293T cells were transfected with the indicated expression plasmids for FLAG-*LRSAM1* (450 ng) in combination with *TSG101*-HA (450 ng) and GFP (100 ng). Total cell lysates were analyzed by immunoblotting as indicated. WT, wild-type; C675, RING mutant; PAT, patient mutation; -, no construct.

sample in which no exogenous *LRSAM1* was introduced, whereas mutant *LRSAM1*, similar to the C675A loss-of-function mutant, does affect these levels. We propose that the found mutation is a loss-of-function mutant that exerts a downstream dominant effect, and therefore should be regarded as a dominant negative mutation. However, we cannot exclude that mutant LRSAM1 could have acquired some additional characteristic of a dominant nature which we cannot measure using this assay. Based on these results and the expression of both mutant and wild-type mRNAs in patient fibroblasts, haploinsufficiency can be excluded as the mechanism leading to the disease as is the case for the carriers of the Canadian recessive mutation, who have the wild-type allele in addition to a null allele and are not affected. Due to the presence of a sterile alpha motif domain that attributes to oligomerization processes (22), LRSAM1 may act as a homodimer, like other RING finger E3-ubiquitin ligases, or be part of a multimeric ligase complex, but be defective in evoking the normal response, probably by a change in ubiquitinylation signaling. Consequent downstream effects such as dysregulation of expression levels of TSG101, (wild-type) LRSAM1 or other targets or changes in dimerization capacity and/or binding affinity may explain the dominant outcome of the mutation with clinical features similar to having no LRSAM1 protein at all.

Recently, LRSAM1 was also reported as a novel candidate modifier for Huntington's disease, with a role in the regulation of localization and clearance of the protein (23). The upregulation of LRSAM1 led to the amelioration of the Htt phenotype in mice and prevented the loss of NeuN-positive cell bodies in primary striatum cultures, concomitant with a reduction in nuclear aggregates. Conversely, the loss of normal LRSAM1 expression or function in the case of a homozygous recessive mutation or a dominant negative mutation would result in the loss of this neuroprotective effect. Except for TSG101 and Huntington protein, more proteins may be regulated by LRSAM1 activity and affect CNS neurons as well. In this respect, it is remarkable that two of our CMT2 patients with the *LRSAM1* mutation also developed Parkinson's disease. Another report described LRSAM1 as a potential

component of the antibacterial autophagic response. This would be regulated by its leucine-rich repeats which are typically involved in the detection of pathogen-associated molecular patterns by the innate immune system (24) that is known to be relevant for neural regeneration (25). Clearly, *LRSAM1* may affect several pathways and the mechanism how this will lead to a neuropathy will be the subject for further study.

Based on the number of expressed sequence tags and other information in publicly available databases, *LRSAM1* appears to be widely expressed, although it was also reported to be higher in tissues with a high energy demand such as the brain, heart, skeletal muscle, liver and kidney. Although we did find a high expression in the tissues that are affected in CMT, it was not restricted to these tissues. Many CMT-associated genes are widely expressed and have been implied in the protein machinery, ranging from amino-acyl transferases to transport. As such, a mutation in *LRSAM1* with a potential role in protein clearance would fit into this picture. It is currently unclear whether the large distances in and energy requirement of neural cells make these cells more vulnerable to these mutations or whether the proteins involved in CMT have an additional neural-specific function, as for the case of *LRSAM1* the outcome of the phenotype in the zebrafish model could suggest. Our data indicate that *LRSAM1* function is relevant for neural development and functioning, and although not neural-specific, it is highly expressed in tissues relevant for CMT both at the fetal level and at the adult level. The identified mutation in the C-terminal part of the protein affects its function to such an extent that the consequent effect is of a dominant nature. In summary, we can conclude that the *LRSAM1* mutation we identified in this family is causative for a dominant form of CMT2 enabling further genetic testing for next generations in this family. The absence of *LRSAM1* in zebrafish which would represent a null mutation results in a phenotype comparable to the mimicked patient mutation in this model. This provides evidence that the homozygous recessive *LRSAM1* mutation that was reported is indeed the causative mutation. We have shown that *LRSAM1* mutations can cause both dominant and recessive forms of CMT2 and have also provided a new CMT2 model for inherited neuropathies.

## MATERIALS AND METHODS

### Patients and DNA

Patients were all seen by an experienced neurologist and subjected to detailed examination. The index person of the family was a Dutch male presenting with the disease at 32 years of age (subject 2.5). There was no medical history of note. Detailed clinical information for all affected family members is given in Table 1. In addition, two brothers of the index patient developed Parkinson's disease at the age of 64 (subject 2.2) and 52 (subject 2.4), respectively. Both showed slowly progressive extrapyramidal features with an asymmetrical onset, a relatively good response to levodopa and a slowly progressive course. Extrapyramidal features were not observed in the younger family members. Clinical data and blood samples were collected as part of the diagnostic procedure for hereditary neuropathies for which all patients gave their consent. Genomic DNA was isolated from blood by proteinase K/SDS treatment followed

by phenol/chloroform extractions and ethanol precipitation. Tissue samples for expression analysis were obtained from the department of pathology.

### Linkage analysis

DNA samples from 11 affected and 8 unaffected members of the family were genotyped using the Affymetrix GeneChip® Human Mapping 250K Sty Array as described before (26). The gender of samples was verified by counting heterozygous SNPs on the X chromosome. Relationship errors were evaluated with the help of the program graphical representation of relationships (27). Linkage analysis was performed assuming autosomal dominant inheritance, full penetrance and a disease gene frequency of 0.0001. Multipoint LOD scores were calculated using the programs ALLEGRO (10) and SimWalk2 (11). The latter program is much slower but allows for the analysis of large families without splitting them. Haplotypes were constructed with ALLEGRO and presented graphically with HaploPainter (28). All data handling was performed using the graphical interface ALOHOMORA (29).

### Next-generation and Sanger sequencing

Sequence capture was performed using a custom design 385K capture array (chr9: 127624486–132419012 hg18/NCBI36) according to the protocol supplied by the manufacturer (Nimblegen, Roche). Basically, a representing library was made of the patient's DNA sheared to an average length of 500–800 bp by cavitation (Covaris). After hybridization, DNA enriched for target sequences was eluted, amplified, quantified (Qubit) and analyzed on a BioAnalyzer DNA 7500 chip (Agilent) followed by pyrosequencing on an FLX titanium sequencer. Analysis was performed with the Newbler program (v2.3, Roche). About  $1 \times 10^6$  reads were obtained from  $1.4 \times 10^6$  key passed sequences. Of the  $6.5 \times 10^5$  filter passed reads, 96% could be mapped, and 50% of the mapped reads were uniquely mapped to the target region and were scanned for differences with the human consensus sequence using the Roche software. All high-quality differences were analyzed for their heterozygous or homozygous state, position within coding exons or distance of at most 20 nucleotides from exon–intron boundaries, known occurrence as polymorphisms and potential effect on the protein.

Direct Sanger sequencing was performed on amplified exons containing the heterozygous nucleotide changes. Amplification was performed with exon-specific M13-tagged primers using 20 ng of genomic DNA, 4 mM MgCl<sub>2</sub>, Hotfire polymerase (Solys Biodyne) and a touchdown PCR program. Primer sequences are provided in Supplementary Material, Table S1. The resulting PCR products were treated with shrimp alkaline phosphatase and exonuclease I prior to sequencing using the ABI Big Dye Terminator cycle sequencing kit and an ABI3730xl sequencer (Applied Biosystems).

### LNA *in situ* hybridizations

Fluorescein-labeled LNA/2-*O*-Me probes (locked nucleic acid/2'-*O*-methyl RNA) were synthesized by Ribotask ApS (Odense, Denmark) and hybridized according to Budde *et al.* (26) followed

by anti-fluorescein-labeled alkaline phosphatase-coupled Fab fragment (Roche, 1:200) incubation as a secondary step. After several washes with a decreasing salt concentration to  $0.2\times$  SSC, a final wash was done at ambient temperature and the SK5300 alkaline phosphatase detection kit (Vector Laboratories, Burlingame, CA, USA) was used for visualization. Pretreatment conditions were slightly different for adult and fetal sections. Fetal tissue sections were pretreated using 20  $\mu\text{g}/\text{ml}$  protK, followed by post-fixation in 4% paraformaldehyde (PFA), and denaturation of proteins in adult sections was achieved by a 10 min incubation in a microwave in 0.1 M citric acid/0.1 M trisodium citrate (pH = 6). Probe concentrations were 0.25–1  $\mu\text{M}$ . Secondary antibody fragment dilutions were 1:200 and 1:1000, respectively. Counterstaining was performed using fast Nuclear red. Pictures were taken on an Olympus BX41 microscope using CellD software. As a control, samples were taken where the probe was omitted from the hybridization mixture. The sequence of the LNA probe used was: 5'-FAM-UcuGgcUccUgaUcuGccG-3' where capital letters represent LNA, normal letters are 2-*O*-methyl oligos.

### Morpholino experiments in zebrafish

All zebrafish (*Danio rerio*) procedures complied with standard animal care guidelines according to Dutch Animal Ethics Committee guidelines. Breeding and maintenance of all zebrafish stocks were performed as described by others (30). Wild-type TL zebrafish outbred strains were crossed to generate embryos for MO injections. MO antisense 25mers were designed against the sequences surrounding the ATG and the last splice acceptor site by the manufacturing company (GeneTools; ATG MO: GAACAAAACATCGTGAGGCTTAGC; splice MO: TGAGACTACACAAAGCAGAACAAGA). A range of 1.7–6.4 ng of MO (1 nl of a 200–750  $\mu\text{M}$  MO solution) was injected into the yolk sac of one to two cell-stage embryos using fine glass needles and a microinjector (World Precision Instruments). Standard control MOs (GeneTools) and non-injected wild-types were taken as controls (31). The developing embryos were allowed to recover in E3 medium and incubated at 28°C. Morphology was monitored over time and images were taken using a Leica MZFLIII equipped with a Retiga 2000R Fast1394 Q imaging camera and the QCapture Pro program. Zebrafish were dechorionated in all experiments at 2–3 dpf, at least 6 h before final pictures were taken except for one experiment where the percentage of hatching was examined. Swimming behavior was examined by the response upon repeatedly touching with a pipette tip. Surviving zebrafish were fixed in 4% PFA after which they were kept in PBS prior to immunohistochemistry (IHC) or stored at –20°C in methanol.

### Wholemout IHC zebrafish

Zebrafish older than 30 hpf were bleached for 3–4 min using a bleaching solution containing 5% formamide,  $0.5\times$  SSC and 9%  $\text{H}_2\text{O}_2$ . After several washes in PBST, the embryos were permeabilized with 2.5 mg/ml trypsin in PBST for 3–4 min on ice, followed by three PBST washes and incubation with the primary antibody in a 1:400–1:2000 dilution in blocking

buffer at 4°C for 16–48 h under gentle agitation. Similarly, secondary antibody detections were performed overnight at 4°C in a 1:1000–2000 dilution (Southern Biotechnology, AP-coupled goat anti-mouse IgG) followed by extensive washing prior to detection using the SK5300 (Vector Laboratories) alkaline phosphatase detection kit. Images were captured using a Leica M125 stereomicroscope equipped with a DFC425 camera and LASv3.7 software.

### Transfections and western blot analysis

HEK 293T cells were obtained from the American Tissue Culture Collection and maintained in DMEM (Invitrogen) supplemented with 10% FBS at 37°C and 5%  $\text{CO}_2$ . Cells were transfected with Lipofectamine 2000 (Invitrogen). Full-length wild-type *LRSAMI* cDNA was amplified by PCR from IMAGE clone ID 5852; cat#OHS1770-9382089 (Thermo Scientific) using oligonucleotides flanking the coding part of *LRSAMI* and recombined into the gateway entry plasmid pDONR221 (Invitrogen). Site-directed mutagenesis was used to introduce the C675A mutation in the RING domain of LRSAM1 with the QuickChange site mutagenesis kit (Stratagene). Full-length *LRSAMI* carrying the c.2121\_2122duplGC mutation was cloned with a two-step PCR strategy. Briefly, genomic DNA from fibroblasts of an individual carrying this mutation was used to amplify exon 26 of *LRSAMI*. Subsequently, the obtained PCR fragment was used to amplify the full-length *LRSAMI* cDNA in combination with the forward primer that was used to clone full-length wt *LRSAMI*. The pCAGGS-hTSG101-HA vector was a kind gift from Dr R. Harty (University of Pennsylvania) (32). To generate mammalian expression plasmids for *LRSAMI*, we used LR recombination between *LRSAMI*-containing pDONR221 and pDEST-FLAG. Transfection efficiency was monitored by co-transfection of an expression plasmid for GFP and was consistently >90%.

Total cell lysates were prepared in RIPA buffer (150 mM NaCl, 1% NP-40, 0.1% sodium deoxycholate, 0.1% SDS, 100 mM Tris–HCl, pH 7.4) supplemented with protease inhibitors (Roche Molecular Biochemicals). Lysates were cleared by centrifugation at 4°C for 10 min at 10 000g. Protein concentration was determined using the Bradford assay (Biorad) with BSA as reference. Samples (10–40  $\mu\text{g}$ ) were separated on NuPAGE Bis–Tris gels (Invitrogen) and transferred to nitrocellulose. Membranes were probed with the following antibodies: HA (Covance, clone HA.11, 1:4000–10000), tubulin (Calbiochem, clone CP06, 1:10000), FLAG-HRP (Sigma, clone M2, 1:8000), GFP (affinity purified rabbit polyclonal anti-GFP was a gift from Dr Mireille Riedinger, UCLA, 1:10000). Secondary HRP-conjugated antibodies (Zymed) were used and visualized with chemiluminescence on a Fuji LAS4000 (GE Healthcare). The immunoblots are representative of at least three independent experiments.

### WEB RESOURCES

CMT mutation databases: <http://www.molgen.ua.ac.be>, <http://www.neuromuscular.wustl.edu>.

## SUPPLEMENTARY MATERIAL

Supplementary Material is available at *HMG* online.

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