Novel *SPG11* mutations in Asian kindreds and disruption of spatacsin function in the zebrafish

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**Abstract** Autosomal recessive hereditary spastic paraplegia with thin corpus callosum (HSP-TCC) maps to the *SPG11* locus in the majority of cases. Mutations in the *KIAA1840* gene, encoding spatacsin, have been shown to underlie *SPG11*-linked HSP-TCC. The aim of this study was to perform candidate gene analysis in HSP-TCC subjects from Asian families and to characterize disruption of spatacsin function during zebrafish development. Homozygosity mapping and direct sequencing were used to assess the *ACCPN*, *SPG11*, and *SPG21* loci in four inbred kindreds originating from the Indian subcontinent. Four novel homozygous *SPG11* mutations (c.442+1G>A, c.2146C>T, c.3602_3603delAT, and c.4846C>T) were identified, predicting a loss of spatacsin function in each case. To investigate the role of spatacsin during development, we additionally ascertained the complete zebrafish *spg11* ortholog by reverse transcriptase PCR and 5′ RACE. Analysis of transcript expression through whole-mount in situ hybridization demonstrated ubiquitous distribution, with highest levels detected in the brain. Morpholino antisense oligonucleotide injection was used to knock down spatacsin function in zebrafish embryos. Examination of *spg11* morphant embryos revealed a range of developmental defects and CNS abnormalities, and analysis of axon...
pathway formation demonstrated an overall perturbation of neuronal differentiation. These data confirm loss of spatacsin as the cause of SPG11-linked HSP-TCC in Asian kindreds, expanding the mutation spectrum recognized in this disorder. This study represents the first investigation in zebrafish addressing the function of a causative gene in autosomal recessive HSP and identifies a critical role for spatacsin during early neural development in vivo.

**Keywords** Hereditary spastic paraplegia · SPG11 · Molecular genetics · Zebrafish studies

**Introduction**

Hereditary spastic paraplegia (HSP) encompasses a group of inherited neurological disorders characterized by degeneration of the corticospinal tract motor neurons and resulting in progressive lower limb spasticity. Over 35 HSP (or SPG) loci have been reported [1] and 18 disease genes identified, accounting for autosomal dominant [2–10], autosomal recessive [11–17], and X-linked [18, 19] subtypes of HSP. Importantly, many of the protein products of these genes share a role in intracellular transport and axonal trafficking [1, 20], suggesting that a common pathogenic mechanism may underlie a number of these disorders.

Hereditary spastic paraplegia with thin corpus callosum (HSP-TCC) is a distinct “complicated” form of autosomal recessive (AR) HSP associated with thinning of the corpus callosum. Despite phenotypic overlap with other ARHSPs, including Mast syndrome (SPG21) [14], agenesis of the corpus callosum with peripheral neuropathy (ACCPN) [21], and a form of HSP-TCC with epilepsy [22], the majority of HSP-TCC cases appear to be linked to the SPG11 locus on chromosome 15q15.1-q21.1 [23–27]. The SPG11 gene has been cloned and mutations identified in 11 of 12 European and North African families analyzed [15]. Since then, independent confirmation that spatacsin loss-of-function mutations cause HSP-TCC has been reported in patients of European, North African, Turkish, Middle Eastern, South American, and Asian origin. In this paper, we present further verification of the SPG11 gene as a major genetic determinant in HSP-TCC by describing novel mutations in four kindreds of Asian descent.

To gain further insight into the physiological and cellular bases of HSP-TCC pathology, we have also used morpholino-based gene knockdown in the zebrafish, Danio rerio, to investigate spg11 function during development. Heterozygous mutations in the human SPG4 and SPG8 genes, encoding spastin and strumpellin, respectively, cause autosomal dominant HSP with predominantly adult onset of disease [2, 10]. Previous investigations have shown that knockdown of spg4 and spg8 during zebrafish development is characterized by aberrant motor neuron axon outgrowth and widespread CNS developmental defects [10, 28]. Here, we report initial studies to address the function of a gene underlying autosomal recessive HSP.

**Materials and methods**

**Patients**

We assessed six affected and seven unaffected family members from four HSP-TCC kindreds of Indian (families 1, 3, and 4) and Pakistani (family 2) descent. Parental consanguinity was documented in all families. The diagnosis of HSP-TCC was established according to published criteria [22]. On examination, all affected individuals demonstrated significant lower limb spasticity, brisk deep tendon reflexes, and bilateral extensor plantar responses. Onset of symptoms occurred between 16 and 20 years of age, with the exception of the two affected siblings in family 3 presenting with subnormal intelligence and difficulty in walking since preschool age. Affected female probands from families 2 and 4 also had mild learning difficulties. Brain MRI revealed substantial thinning and hypoplasia of the corpus callosum in all affected cases, together with generalized moderate cerebral atrophy.

The study was approved by the University Hospitals of Leicester and Guy’s & St Thomas’ NHS Foundation Trust local research ethics committees, and all participants provided written informed consent. Control samples were of mixed Asian origin, ascertained in relation to a program of autozygosity mapping with inbred kindreds [29].

**Genetic analysis**

DNA was extracted from peripheral blood leukocytes by standard techniques. Microsatellite markers flanking the ACCPN, SPG11, and SPG21 loci were identified from the NCBI UniSTS database and amplified by PCR. Genotypes were assigned from PCR products using an ABI3730xl automated sequencer and GeneMapper v3.7 software (Applied Biosystems, Foster City, CA, USA). To confirm identity-by-descent, additional polymorphic CA-repeat regions were detected at 43.416 and 43.775 Mb and genotyped as above (primer details available on request). Marker positions were assigned according to the NCBI sequence map (Build 37.1).

All coding exons and intron–exon boundaries of the SPG21 and SPG11 genes were screened by direct DNA sequencing. Primers were designed using Primer3 software [30] or were as previously described [15]. PCR products were purified with ExoSAP-IT (GE Healthcare, Munich, Germany), sequenced using BigDye Terminator v3.1
chemistry (Applied Biosystems), and analyzed on an ABI3730xl automated sequencer. Sequence traces were aligned to reference using Sequecher v.4.5 software (Gene Codes Corporation, Ann Arbor, MI, USA).

To directly assess the consequence of the intron 2 splice site mutation upon transcript translation, total RNA was extracted from peripheral blood of subjects III:2 and IV:3 with the QIAamp RNA blood mini kit (Qiagen, Crawley, UK). First-strand complementary DNA (cDNA) synthesis was performed using the Reverse-iT 1st strand synthesis kit (ABgene, Epsom, UK). Gene-specific primers SPG11ex1-intF (5′ CAGCCTCCAAGTGCTTCTTTCTTTT 3′) and SPG11ex4intR (5′ GTCTCTGCTGCTGTCATTA 3′) were used for reverse transcriptase PCR (RT-PCR) and sequencing of messenger RNA (mRNA).

Zebrafish maintenance

Zebrafish embryos were obtained from natural matings, maintained at 28.5°C, and staged as previously described [31]. Pigmentation was inhibited by treatment with 0.2 nM phenylthiourea at 24 h post-fertilization (hpf).

Bioinformatics and cloning of the zebrafish ortholog

Searching the zebrafish genome assembly (Zv7 release v48) for KIAA1840 orthologs identified an Ensembl gene (ENSDARG0000045968) on chromosome 25, within contig Zv7_scaffold224041, at position 317,067–343,586 kb. This gene encodes several overlapping predicted transcripts sharing homology with the 3′ region of human SPG11 cDNA (exons 25–40). To delineate the rest of the zebrafish gene, human cDNA sequence was queried against the zebrafish genome and EST databases using the TBLASTN algorithm.

A predicted partial mRNA (XM_001346277), located within Zv7_NA70511.2, was identified exhibiting homology with human SPG11 exons 2–9. Using primer pairs JH37 (5′ TTCTTTGTGGGAGGATGTGAG 3′) and JH42 (5′ CGAACATACTCCCTTCCAGATT 3′), we generated partial cDNA fragments of 1,522 and 1,097 bp. Both PCR products were cloned into the pCR3 Ready amplification of cDNA ends

5′ Ready amplification of cDNA ends

To determine exon 1 of the zebrafish spg11 gene, 5′ ready amplification of cDNA ends (5′ RACE) was performed using the GeneRacer RACE ready cDNA kit (Invitrogen) according to the manufacturer’s guidelines. The gene-specific reverse primer JH40b (5′ AACACACACGTGTTC ACCAACACAC 3′) was used for RT-PCR amplification with the forward GeneRacer 5′ Primer, and a nested PCR was performed using zSPG11ex2-3intR (5′ ATGTCCGG CGTGTGTGTGTGTGATGA 3′) and GeneRacer 5′ Nested Primer. PCR products were gel excised and sequenced directly with primer zSPG11ex2-3intR. The ATG start site was identified by translating all three reading frames of the mRNA sequence using publicly accessible software [34] and correlating zebrafish and human sequences.

Whole-mount in situ hybridization and immunohistochemistry

In situ hybridization was performed as previously described [35]. Antisense and sense digoxigenin-labeled RNA probes were generated by in vitro transcription from plasmids pCR5′-spg11 and pCR3′-spg11. To address the role of spg11 during CNS development, axon pathway formation was analyzed by anti-acetylated α-tubulin immunohistochemical labeling at 52 hpf. Alkaline phosphate conjugated antibodies (6-11B-1) were used for detection (Roche Applied Science, Mannheim, Germany and Sigma-Aldrich, St. Louis, MO, USA). Embryos were mounted in 90% glycerol and photographed using a Nikon DS-5Mc cooled camera system with NIS-Elements Basic Research (BR) v3.0 software (Nikon Instruments Inc., Amstelveen, The Netherlands).

Morpholino knockdown of zebrafish spg11

The antisense splice-blocking morpholino oligonucleotides spg11E2I2 (5′ GTGATGTTCAGTCTCCTTTACCGGC 3′) and spg11E4I4 (5′ TGTTGGCCGTGTGCACCTGAA 3′), designed to target the spg11 exon2–intron2 and exon4–intron4 boundaries, respectively, were synthesized by Gene Tools, LLC (Philomath, OR, USA). The 5-mismatch morpholino spg11E2I2mm (5′ GTCATTTGAGTCTCTTTACCGGC 3′) was used as a specificity control. Lyophilized oligonucleotide resuspended in sterile water was used at a working concentration of 2 ng/µl containing 0.1% phenol red and rhodamine for visualization and sorting of injected embryos. Titrations were performed by injecting 0.5–2 ng into a single blastomere of 1–2 cell stage embryos, and an optimal dose minimizing mortality and toxic effects was determined. Total RNA was isolated from embryos to detect aberrant splicing events using RT-PCR with primers JH49 (5′
TCTGAGCAGGAGTGATGCTG 3′) and JH40 (5′ AACA-
CACACGTGTCTACCAAC 3′) across exon 2 and zSPG11ex3intF (5′ CGTCTG TGAGGCTGTTGTG 3′) and zSPG11ex6intR (5′ TTCGTCTGTCGTCGTC 3′) across exon 4. PCR products were isolated by gel purification and sequenced.

**Results**

Genotyping and mutation detection

Microsatellite genotyping in families 1 and 2 revealed all affected subjects were heterozygous for ACCPN markers, but showed homozygosity at the SPG11 (Fig. 1) and SPG21 loci (data not shown). Sequencing of SPG21 excluded this gene as the cause of HSP-TCC in both families by the identification of heterozygous SNPs in affected subjects. Genotyping data for families 3 and 4 were consistent with exclusion of the ACCPN and SPG21 loci, having heterozygous genotypes in affected cases, and were homozygous for SPG11 (Fig. 1). Following additional microsatellite genotyping across the SPG11 locus, the minimal disease gene interval was defined to a 3.96-Mb region, flanked by markers D15S779 and D15S659 (Figs. 1 and 2a).

Sequence analysis of the SPG11 gene revealed a splice donor site mutation in intron 2 (c.442+1G>A), co-segregating with the disease phenotype in family 1 (Fig. 2b, c). Affected subjects were homozygous and all unaffected family members were heterozygous for this mutation, consistent with the genotyping data generated previously. Using primers in exons 1 and 4, RT-PCR in a heterozygous parent and an affected subject revealed a wild-type fragment of 575 bp and a smaller mutant band of 390 bp, the expected size upon excision of exon 2. Sequencing of the mRNA transcript confirmed skipping of exon 2 in the mutant allele, resulting in premature termination and a truncated protein (p.H86QfsX15; Fig. 2d).

Mutation screening of the SPG11 gene in the remaining HSP-TCC kindreds identified three further novel mutations (Fig. 2c). The sequence variants detected in families 2 and 3 also predict generation of truncated transcripts due to homozygous c.4846C>T and c.2146C>T nonsense mutations in exons 28 and 11, incorporating stop codons in place of the native glutamines at positions 1616 and 716, respectively. In family 4, a homozygous 2-bp deletion was detected in exon 21 (c.3602_3603delAT), predicting a frameshift leading to premature termination of the polypeptide chain four amino acids downstream (p.Y1201LfsX4). None of the identified mutations were observed in a panel of 180 control chromosomes of similar ethnicity.

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**Fig. 1** Pedigree structures of the four families with genotyping data across the SPG11 locus. Pedigree identifiers indicate subjects from which DNA was available. All affected subjects are homozygous for microsatellite markers across this region. Critical recombinations in families 1 and 3 define the disease gene interval between markers D15S779 and D15S659.
Isolation and characterization of the zebrafish spg11 gene

Reference sequence for ~1.5 kb at each terminus of the spg11 ortholog was obtained through bioinformatics and 5′ RACE, facilitating amplification of the central portion by RT-PCR. Complete transcript sequence data have been deposited with the GenBank data library under accession numbers FJ217177 (full-length cDNA, 7,140 bp) and FJ217178 (alternative cDNA, 3,747 bp). Evaluation of sequence homology between human and zebrafish spastasin revealed 41% identity and 60% similarity across the length of the proteins (Fig. 3), with particularly high levels of nucleotide sequence identity across human SPG11 exons 35–36. Using a multiple sequence alignment across vertebrate species (data not shown), a highly conserved putative WD repeat signature within this region was subsequently detected with QuasiMotifFinder (Fig. 3).

Expression of spg11 during zebrafish development

Developmental RT-PCR analysis revealed that spg11 was expressed both maternally and zygotically, persisting until at least day 3 of development. Using whole-mount in situ hybridization, spg11 transcripts were ubiquitously distributed throughout the embryo at 26 hpf, with increased expression levels detected in the brain and low levels in the notochord. Corresponding sense probes produced no significant hybridization signal (data not shown).
Morpholino knockdown and immunohistochemistry

RT-PCR analysis of spg11E2I2 morphant embryos detected three transcripts: the wild-type 468-bp fragment, an abundant 413-bp product, corresponding to an mRNA lacking the terminal 55 bp of exon 2 and a less abundant 551-bp fragment caused by retention of intron 2. Both mis-splicing events are predicted to severely truncate the encoded proteins through introduction of a premature stop codon in exon 3 (Fig. 4a), providing a good model of the donor splice site mutation detected in family 1. Morpholino titration to determine an effective dose for phenotypic effects.

Fig. 3 Multiple sequence alignment of human and zebrafish spatacsin proteins. Human reference sequence was obtained from the Ensembl database (ENSP00000261866). Identical amino acid residues are highlighted in dark gray and residues sharing more than 90% similarity in light gray. Putative functional domains (i–iv), indicated by horizontal lines, correspond to the glycosyl hydrolase family 1, leucine zipper, coiled coil, and Myb domains, described previously [15]. The highly conserved WD repeat signature predicted by QuasiMotifFinder is boxed (Prosite ref. PS00678)
analysis revealed a dose-dependent blocking of splicing (Fig. 4b). Embryos injected with 0.5 ng demonstrated phenotypic recovery by 52 hpf, while 2 ng was toxic. The optimal dose for further analysis of *spg11* morphants was therefore established at 1 ng.

A range of developmental defects was apparent in *spg11*E2I2 morphants by 26 hpf. Morphologically, the embryos had a curly-tail down phenotype (data not shown), defective brain ventricle formation, and small eyes (Fig. 4c–e). At 52 hpf, *spg11* morphants had developed mild hydroceph-
aly, the mid-hindbrain boundary was reduced, and the eyes were still smaller (Fig. 4f, g). In the trunk, the typical chevron somite pattern appeared flatter in morphant embryos in comparison to uninjected controls (8% mild, 36% moderate, 55% severely affected embryos at 1-ng morpholino dose; Fig. 4h, i). Morphant embryos exhibited a generalized perturbation of neuronal differentiation, with reduced levels of staining visible in spg11 morphants compared to controls (Fig. 4j–s). Although retinal ganglion cell differentiation was reduced in the eye, the optic axons did form and exit the retina correctly before projecting to the tectum (Fig. 4j, k).

Analysis of cranial motor neurons revealed that the facial and branchiomotor neurons of spg11 morphant embryos were present, but appeared shorter and less differentiated than those of control embryos (Fig. 4l, m). The posterior commissure also formed normally by 52 hpf; however, the habenular commissure failed to project to the dorsal midline (Fig. 4n, o). In the hindbrain, neuronal differentiation was severely reduced and glial “curtain” cells were absent. In addition, an ectopic cluster of disorganized neurons was often apparent within spg11 morphant hindbrains (Fig. 4p, q). In the trunk, spinal motor neuron axons projected correctly but fascicules appeared thinner, suggesting a reduced number of neurons. Furthermore, there was some evidence of impaired orientation and axons appeared shortened, implying a failure to reach their ventral target (Fig. 4r, s). Embryos injected with an equivalent dose of the spg11E2I2 mismatch oligonucleotide appeared comparable to uninjected controls, indicating that the observed phenotype is due to knockdown of the spg11 transcript (Electronic supplementary materials).

To confirm the specificity of reduction of spg11 activity, zebrafish embryos were injected with a second spg11-specific morpholo targeted to the intron 4 splice donor site, mutation of which has also been previously reported in the human SPG11 gene [36]. RT-PCR analysis of embryos injected with the spg11E4I4 morpholino demonstrated efficient splice blocking, detecting an alternatively spliced transcript corresponding to retention of intron 4.

Marked CNS abnormalities were also apparent in spg11E4I4 morphants, and in comparison to spg11E2I2-injected embryos, a more severe truncation of the spinal motor neuron axons was observed at higher concentrations (Electronic supplementary materials). In addition to the CNS features, embryos also exhibited a range of developmental defects, including an enlarged heart cavity, which is a relatively common feature in injected fish, a curly tail and deformities of the fin, and were generally smaller than the uninjected controls. As observed with the spg11E2I2 morpholino, the maximum dose tested was toxic. Statistical analysis of the distribution of morphological abnormalities by one-way ANOVA revealed a significant difference between target and control morpholinos at the 1-ng dose (p<0.0001; Table 1).

### Discussion

The genetic basis of the hereditary spastic paraplegias is rapidly evolving with 18 disease genes now reported, accounting for approximately 50% of the known loci. In this study, we report the genetic analysis of four autosomal recessive HSP-TCC families, each originating from the Indian subcontinent. Following localization to chromosome 15 by autozygosity mapping, direct sequencing of the SPG11 gene identified novel pathogenic homozygous mutations (c.442+1G>A, c.2146C>T, c.3602_3603delAT, Table 1).

#### Table 1 Phenotypic characteristics of injected embryos at 52 hpf in comparison to uninjected controls

| Target region | Dose (ng) | No. of embryos (n) | Phenotypic characteristics observed |
|---------------|-----------|--------------------|-----------------------------------|
|               |           |                    | Normal (%) | Curly Tail (%) | Fin (%) | Hydrocephaly (%) | Severe deformity (%) |
| spg11E2I2     | 0.5       | 151                | 57         | 7             | 0       | 7                 | 5                     |
|               | 1         | 158                | 23         | 22            | 19      | 17                | 29                    |
|               | 1.5       | 201                | 18         | 22            | 30      | 15                | 24                    |
| spg11E2I2mm   | 0.5       | 80                 | 71         | 1             | 0       | 6                 | 1                     |
|               | 1         | 211                | 71         | 4             | 0       | 5                 | 6                     |
|               | 1.5       | 395                | 69         | 2             | 2       | 9                 | 10                    |
| spg11E4I4     | 0.5       | 87                 | 51         | 14            | 5       | 10                | 6                     |
|               | 1         | 173                | 16         | 36            | 12      | 5                 | 19                    |
|               | 1.5       | 227                | 10         | 20            | 23      | 15                | 25                    |
| Uninjected    | n/a       | 81                 | 100        | 0             | 0       | 0                 | 0                     |

Increased morphological features were observed with increasing concentrations of spg11E2I2, spg11E2I2 mismatch (mm) control, and spg11E4I4 morpholinos. The differences between the spg11E2I2 target and mismatch control groups at the 1-ng target dose are statistically significant, as assessed by one-way ANOVA test (p<0.0001). Embryos with enlarged heart cavities were scored, but this feature was not specific to a loss of spg11 activity.
and c.4846C>T in all kindreds. To date, spatacsin mutations have predominantly been reported in European (47%), North African (14%), and Chinese (11%) populations, as detailed in Electronic supplementary materials. However, only one study has described SPG11 mutations in patients of subcontinental descent [37], most likely representing an ascertainment bias but suggesting that SPG11 variation may be a rare cause of HSP in these patients.

All mutations identified in our patients lead to a truncated protein product, consistent with other reports that loss of spatacsin function is the pathogenic mechanism underlying SPG11-linked HSP. While a homozygous G>C substitution has previously been reported at the intron 2 splice donor site [38], the variant segregating in family 1 is the first example of a c.442+1G>A mutation and formally demonstrates that aberrant splicing results in exon 2 skipping. Families 2 and 4 harbor mutations in exons 28 and 21 of the SPG11 gene. Variants have previously been identified in more than half of all exons; however, <10% of HSP-TCC cases are accounted for by mutations in exons 18–29, encoding the central region of the spatacsin protein. Interestingly, previous bioinformatic analysis has indicated this to be a region of putative structural importance [15]. Although the function of spatacsin remains undetermined, the putative leucine zipper motif and Myb domain have implied a possible regulatory function during gene expression. Through resolution of sequence and amino acid homology between human and zebrafish, we detected a highly conserved WD repeat signature across vertebrate orthologs of spatacsin. Characteristically, WD proteins are composed of repeating units culminating in tryptophan (W) and aspartic acid (D) and share a functional role in the regulation of cellular processes such as cell division, gene transcription, transmembrane signaling, mRNA modification, and vesicle fusion [39]. It remains to be determined whether the restricted motif detected here reflects a similar function for spatacsin.

To investigate the role of spatacsin during development, we have performed whole-mount in situ hybridization studies in the zebrafish embryo. Previous studies of Spg11 mRNA expression in the rat brain have demonstrated that it is undetectable in the newborn but is present in the cerebellum from postnatal days 6–21 [15]. In contrast, RT-PCR in the zebrafish embryo has detected spg11 mRNA expression both maternally and zygotically, persisting until at least 72 hpf. Whole-mount in situ hybridization revealed that transcripts were ubiquitously distributed throughout the embryo at 26 hpf with high levels of expression in the CNS. This ubiquitous expression, similar to that reported for spg4 [28], is consistent with previous RT-PCR profiling of the SPG11 gene in human tissues [40] and suggests an important role for spatacsin during early developmental processes throughout the embryo.

To specifically assess the functional impact of the c.442+1G>A splice site mutation, we sought to recapitulate the phenotype in zebrafish by means of morpholino injection. We observed a range of developmental defects in spg11 morphant embryos, including CNS abnormalities. Overall, spg11 morphants exhibited perturbed neuronal differentiation, with markedly reduced levels of immunohistochemical staining in comparison to control embryos. Consistent with the features seen in spg4 and spg8 embryos [10, 28], spg11 morphants had a curly-tail phenotype and motor neuron axons appeared reduced with some evidence of impaired orientation, albeit less pronounced than in the autosomal dominant HSP counterparts. While axonal structures within the brain developed normally, subsequent neuronal projections appeared to be impeded, particularly within the forebrain and midbrain regions. The characteristic lower limb spasticity in HSP is caused by degeneration of the corticospinal tract motor neurons. The data presented in this study are in agreement with previous reports demonstrating a disruption of axon morphogenesis in loss of HSP-specific gene activity [10, 28], and while the function of the strumpellin protein remains unknown, this may suggest a similar role for spatacsin in microtubule dynamics. Further studies will be required to determine the neuronal-specific effects of spg11 expression during later development. In humans, autosomal recessive HSP-TCC typically manifests during infancy or puberty, and slow disease progression continues over a period of 10–20 years. In addition to lower limb spasticity and cognitive impairment, some individuals exhibit additional features including dysarthria, dysphagia, and ocular defects. Interestingly, spatacsin knockdown also appears to impair eye development in spg11 morphants.

In summary, this study provides confirmation that loss of spatacsin function is the primary molecular defect in the pathogenesis of SPG11-linked HSP-TCC and points to a critical role for spatacsin during early neural development. For clarity, we have also collated all SPG11 mutations reported to date, providing a comprehensive resource for neurologists and molecular geneticists in genetic screening for HSP-TCC.

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Ethical standards These experiments comply with the current laws of the United Kingdom.

Conflict of interest The authors declare that they have no conflict of interest.
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