REPORT

TBCE Mutations Cause Early-Onset Progressive Encephalopathy with Distal Spinal Muscular Atrophy

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Tubulinopathies constitute a family of neurodevelopmental/neurodegenerative disorders caused by mutations in several genes encoding tubulin isoforms. Loss-of-function mutations in *TBCE*, encoding one of the five tubulin-specific chaperones involved in tubulin folding and polymerization, cause two rare neurodevelopmental syndromes, hypoparathyroidism-retardation-dysmorphism and Kenny-Caffey syndrome. Although a missense mutation in *Tbce* has been associated with progressive distal motor neuronopathy in the *pmn/pmn* mice, no similar degenerative phenotype has been recognized in humans. We report on the identification of an early-onset and progressive neurodegenerative encephalopathy with distal spinal muscular atrophy resembling the phenotype of *pmn/pmn* mice and caused by biallelic *TBCE* mutations, with the c.464T>A (p.Ile155Asn) change occurring at the heterozygous/homozygous state in six affected subjects from four unrelated families originated from the same geographical area in Southern Italy. Western blot analysis of patient fibroblasts documented a reduced amount of TBCE, suggestive of rapid degradation of the mutant protein, similarly to what was observed in *pmn/pmn* fibroblasts. The impact of *TBCE* mutations on microtubule polymerization was determined using biochemical fractionation and analyzing the nucleation and growth of microtubules at the centrosome and extracentrosomal sites after treatment with nocodazole. Primary fibroblasts obtained from affected subjects displayed a reduced level of polymerized α -tubulin, similarly to tail fibroblasts of *pmn/pmn* mice. Moreover, markedly delayed microtubule re-polymerization and abnormal mitotic spindles with disorganized microtubule arrangement were also documented. Although loss of function of TBCE has been documented to impact multiple developmental processes, the present findings provide evidence that hypomorphic *TBCE* mutations primarily drive neurodegeneration.

Tubulinopathies are a family of severe neurodevelopmental disorders caused by mutations in multiple genes encoding tubulin isoforms (e.g., TUBA1A [MIM: 602529], TUBA8 [MIM: 605742], TUBB2B [MIM: 612850], TUBB3 [MIM: 602661], and TUBB5 [MIM: 191130]), with defective neuronal migration and cerebral cortex malformation representing their major features.¹ Inactivating mutations in TBCE (tubulin folding cofactor E [MIM: 604934]), encoding one of the five tubulin-specific chaperones involved in tubulin folding and polymerization, are known to cause two rare neurodevelopmental syndromes, hypoparathyroidism-retardation-dysmorphism syndrome (HRDS [MIM: 241410]), which is characterized by congenital hypoparathyroidism, intellectual disability, facial dysmorphism, and extreme growth failure, and Kenny-Caffey syndrome type 1 (KCS1 [MIM: 244460]), which overlaps HRDS but exhibits osteosclerosis and recurrent bacterial infections as additional features.² All reported children with HRDS and KCS1 but one originate from inbred families of Middle Eastern origin (Saudi and Israeli pedigrees) and carry a recurrent homozygous in-frame deletion (c.155_ 166delGCCACGAAGGGA [p.Ser52_Gly55del]) in TBCE (GenBank: NM_001079515.2) resulting in virtually absent protein level.^{2,3} Two additional mutations, both truncating (c.66_67delAG [p.Val23fs48*] and c.T1113A [p.Cys371*]), have been reported in a compound heterozygous state in a Belgian pedigree with two siblings manifesting features of HRDS.² More recently, microtubule-disrupting mutations in tubulin genes have also been shown to trigger neurodegeneration, as illustrated by TUBA4A (MIM: 191110) and TUBB4A (MIM: 602662) mutations underlying amyotrophic lateral sclerosis (MIM: 616208) and hypomyelinating leukodystrophy associated with progressive spastic dystonic tetraplegia (MIM: 612438), respectively.^{4,5} Similarly, a missense *Tbce* mutation predicting the p.Trp524Gly amino acid substitution has been associated with progressive distal motor neuronopathy in pmn/pmn mice,^{6,7} but no similar phenotype has so far

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been reported in humans. Here, we report on the identification of an early-onset neurodegenerative encephalopathy resembling the phenotype of *pmn/pmn* mice, caused by biallelic *TBCE* mutations, with the c.464T>A missense change (p.Ile155Asn) occurring at the heterozygous/homozygous state in all affected subjects. We also show that the mutation results in a reduced amount of TBCE and that defective TBCE function in primary fibroblasts is associated with a lowered amount of polymerized α -tubulin, markedly delayed microtubule re-polymerization, and abnormal mitotic spindle morphology. Consistent with what we had previously been observed in *pmn/pmn* mice, the present findings demonstrate that hypomorphic TBCE function primarily drives neurodegeneration.

Six individuals from four apparently unrelated families originating from the same geographical area (metropolitan area of Naples in Campania, Southern Italy, and Ischia Island, just off the coast) were included in the study. Clinical features are schematically listed in Table 1 and summarized in the Supplemental Case Reports. Clinical data and biological material collection and storage were attained from the participating families after written informed consent was obtained, and the study was approved by the Ethical Committees of the participating centers. Genomic DNA and total RNA was isolated from peripheral blood leukocytes, using standard protocols. We originally performed whole-exome sequencing (WES) on genomic DNA obtained from subjects 1544334 and 2518864 (Figure 1). Both subjects had been followed at the Ospedale Pediatrico Bambino Gesù, Rome, and shared a neurodegenerative disorder characterized by early-onset slowly progressive distal motor neuropathy resembling distal spinal muscular atrophy (SMA) with bilateral foot drop associated to spasticity and cerebellar ataxia. They did not exhibit dysmorphia or growth defects. Muscle biopsy confirmed neurogenic muscle atrophy, and brain magnetic resonance imaging (MRI) documented cerebellar atrophy (Figure 1). In both cases, we used a trio-based (proband and unaffected parents) strategy. Exome capture was carried out using SureSelect Human All Exon v.4 (Agilent), and WES data analysis was performed using an in-house implemented pipeline, as previously reported.⁸⁻¹⁰ Pairedend reads were aligned to the human genome (UCSC GRCh37/hg19) with the Burrows-Wheeler Aligner (BWA v.0.7.10). Presumed PCR duplicates were discarded using Picard tools' MarkDuplicates. GATK tools were used for realignment of sequences encompassing indels and for base quality recalibration.¹¹ SNPs and small indels were identified by means of the GATK's HaplotypeCaller used in gVCF mode, followed by family-level joint genotyping and phasing, according to GATK's latest best practices. Called variants were filtered to retain variants with quality > 100 and quality-by-depth score > 1.5. High-quality variants were then filtered against public databases (dbSNP142 and ExAC v.0.3) to retain novel and clinically associated variants, and annotated variants with unknown frequency or having MAF < 0.1% and occurring with a

quency data from approximately 600 population-matched WES. SnpEff toolbox (v.4.1)¹² was used to predict the functional impact of variants, which were filtered to retain only those located in exons with any effect on the coding sequence, and splice site regions (-3 to +8 with respect to)exon-intron junctions). Functional annotation of variants was performed with SnpEff v.4.1 and dbNSFP v.2.8,^{12,13} and their functional impact was analyzed by Combined Annotation Dependent Depletion (CADD) v.1.3 and dbNSFP Support Vector Machine (SVM) v.3.0 algorithms.^{14,15} For sequencing statistics, see Table S1. Data annotation predicted 13,541 and 13,484 high-quality variants having functional impact (i.e., non-synonymous and splice site changes) in subjects 1544334 and 2518864, respectively. Among them, 356 and 350 clinically associated, private, and low/unknown frequency variants were retained for further analyses. Because family 1 had documented consanguinity, filtering was first directed to identify genes shared by the probands having variants compatible with a recessive inheritance models. We identified only one gene, TBCE, satisfying the filtering criteria. Specifically, subject 1544334 (family 1) was homozygous for the c.464T>A missense change (p.Ile155Asn) and subject 2518864 was a compound heterozygote for the same nucleotide substitution and the truncating c.1076delC (p.Leu360Ter) change. Sanger sequencing confirmed both variants and segregation in each family (Figure S1). No gene with a putative de novo change was shared by the two probands, and none of the genes with putative de novo variants in each individual was functionally linked to processes relevant to neurodegeneration (Table S1), strongly pointing to TBCE as the gene implicated in the disorder. Mutation scan of the TBCE coding sequence was successively performed on two monozygotic twins (subjects 00997847 and 00997844) of a third family originating from the same geographical area. Both individuals exhibited MRI profiles and an overall clinical phenotype overlapping that of the two previous cases, and Sanger sequencing allowed us to identify homozygosity for the recurrent c.464T>A change (Figure S1). Finally, interrogation of the Telethon Institute of Genetics and Medicine WES variant database allowed us to identify an additional sibling pair homozygous for the c.464T>A missense substitution (Figure S1). In that family, WES had been performed to identify the genetic cause of an unsolved apparently recessive neurodegenerative disorder characterized by spastic tetraparesis, optic atrophy, and abnormal brain MRI (cerebellar and brainstem atrophy, corpus callosum hypotrophy, and white matter hyperintensity with iron deposition in the globus pallidus and substantia nigra) resembling a pattern of neurodegeneration with brain iron accumulation (NBIA [MIM: 234200]) (Figure 2). Clinical reassessment of the two affected subjects revealed clinical features matching the disorder characterizing the previously identified subjects homozygous/compound heterozygous for the TBCE c.464T>A change. Remarkably,

frequency < 2% in an in-house database including fre-

Table 1.	Clinical Features of Sub	piects with Biallelic Muta	tions in TBCE
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Affected Subjects	1544334	2518864	00997847	00997844	VN_X3359	VN_3360
Family	1	2	3	3	4	4
Gender	male	male	male	male	female	male
Year of birth	2007	2009	2	2000	1996	2001
Inheritance	simplex case	simplex case	simplex case		familial	familial
Consanguinity	yes	no	no		no	no
Pregnancy, delivery, neonatal period	normal, weight at birth 3,360 g at 39 th week of gestation	normal, weight at birth 3,490 g at 38 th week of gestation	normal, weight 3,030 g at 37 th week gestation	normal, weight 2,570 g at 37 th week gestation	normal, weight at birth 2,600 g at 38 th week of gestation	normal, weight at birth 3,250 g at 39 th week of gestation
Initial motor & cognitive development	neonatal period normal	abnormal	neonatal period normal		neonatal period normal	normal until 8 months
Age at presentation	around 4–5 months of age	at birth	around 4–5 months of age		14 months	8 months
Signs of presentation	hypotonia and developmental delay in the first year of life	hypotonia and developmental delay in the first year of life	hypotonia and developmental delay in the first year of life		spasticity and developmental delay in the first year of life	spasticity and developmental delay in the first year of life
Signs of regression	distal amyotrophy, ataxia, spasticity	distal amyotrophy, ataxia, spasticity	distal amyotrophy, ataxia, spasticity		distal amyotrophy, spastic tetraparesis, optic atrophy	distal amyotrophy, spastic paraparesis, optic atrophy
Outcome						
Age (years)	6	6	15	15	20	15
Motor function	able to sit	never able to sit	lost the ability to sit	lost the ability to sit	lost the ability to sit	lost the ability to sit
Cognitive level	mildly affected	moderately affected	moderately affected	moderately affected	severely affected	severely affected
Speech and language	able to speak, dysarthria	not able to speak	able to speak, severe dysarthria	able to speak, severe dysarthria	not able to speak	severe dysarthria
TBCE mutations ^a	c.463T>A (p.Ile155Asn), c.463T>A (p.Ile155Asn)	c.463T>A (p.11e155Asn), c.1076delC (p.Leu360Ter)	c.463T>A (p.Ile155Asn), c.463T>A (p.Ile155Asn)		c.463T>A (p.Ile155Asn), c.463T>A (p.Ile155Asn)	c.463T>A (p.Ile155Asn), c.463T>A (p.Ile155Asn)

^aBoth variants have been reported with low recurrence in public databases (c.464T>A, rs780472451, 0.0000082/1, ExAC; c.1076delC, rs750781063, 0.00002/3, ExAC). Mutations are annotated according to GenBank: NM_001079515.2 (NP_001072983.1).



Figure 1. *TBCE* Mutations Cause Early-Onset Encephalopathy with Distal Spinal Muscular Atrophy Clinical features of subjects 1544334 (A) and 2518864 (B–D); muscle biopsy of 1544334 stained with ATPase 4.3 showing fiber type grouping and scattered atrophic angulated fibers (E) and atrophic angulated fibers that are denervated because positively stained with non-specific esterase (F); brain MRIs of subjects 1544334 (G–I, at 2 years; J–L, at 3 years) and 2518864 (M–O, at 1 year; P–R, at 2 years). Note the hypoplasia of corpus callosum and the cerebellar atrophy with prominent folia (K, Q).

similar to the other case subjects, the family was from Pozzuoli and Mergellina, located in the Naples metropolitan area, along the coast near Ischia Island. Parents were not consanguineous. WES data statistics are reported in Table S1.

All affected subjects were homozygous or compound heterozygous for the c.464T>A missense change and were from the same geographical area, strongly suggesting a founder effect. The variant has recently been reported in the ExAC database (rs780472451) with an overall estimated frequency of 0.000008 (1/121,411, information on the ethnic origin of the carrier not available). Genotyping of 400 unrelated and apparently unaffected individuals from the Naples metropolitan area by ARMS-PCR (primers and PCR conditions available on request) failed in identifying any allele carrying the variant (0–0.0025, 95% confidence interval). Based on the informative variants identified by WES to flank the c.464T>A change, the genomic region encompassing the disease-causing variant shared by the three families was estimated to cover approximately 1.3 Mb (chr1: 235,357,303–236,687,580). To more precisely define the size of the shared fragment, eight microsatellite markers covering the *TBCE* locus were selected, and the four trios were genotyped using an ABI PRISM 3500 Sequencer and the GeneMapper 4.0 software (Applied Biosystems). Reconstruction of the haplotypes allowed us to refine the length of the shared genomic region, which spanned approximately 0.5 Mb (Table S2), confirming the identity by descent of the c.464T>A missense variant.

All individuals homozygous or compound heterozygous for the c.464T>A variant in *TBCE* shared a strikingly homogeneous phenotype with clinical signs of progressive neurodegeneration, associated with normal calcium/ phosphate metabolism and absence of repeated infections (Table 1 and Supplemental Case Reports), clearly differing from HRDS and KCS. By contrast, their clinical features were reminiscent of the *pmn/pmn* mouse,¹⁶ in which the progressive motor neuronopathy, hindlimb atrophy, and



Figure 2. TBCE Mutations Cause a Pattern of Neurodegeneration with Brain Iron Accumulation in the Second Decade of Life of Affected Individuals

(A–J) Brain MRIs of subjects VN_X5360 (A–E, performed at age 13 years) and VN_X3359 (F–J, performed at age 17 years). Axial FLAIR-weighted images shown in (A), (B), (F), and (G); coronal T2-weighted images shown in (C) and (D); T1-weighted sagittal images shown in (E) and (J); (A), (B), (F), (G), (H), and (I) are FLAIR-weighted images. Note T2 or FLAIR hypointense areas corresponding to globi pallidi, substantia nigra, and mesencephalic red nuclei in both siblings, indicating abnormal iron deposits.

(K–O) Brain MRIs of subject 00997847 performed at age 16 years. Axial T2-weighted image shown in (K); a T2 coronal weighted image shown in (N); two axial T2* relaxation-weighted images documenting the presence of iron shown in (L) and (M); and a sagittal T1-weighted image shown in (O). The hypointense areas detected in T2-weighted images correspond to the increased iron for age accumulated at the level of the globi pallidi and substantia nigra.

reduced brain size is caused by a missense substitution in the same gene (p.Trp524Gly) documented to partially impair TBCE stability and function.^{6,7}

TBCE is a 527-amino-acid-long protein characterized by an N-terminal CAP-Gly domain, a motif mediating protein binding to α-tubulin, followed by nine tandemly arranged leucine-rich repeat (LRR) motifs, and a C-terminal ubiquitin-like domain possibly implicated in α-tubulin degradation via the proteasome (Figure S1).^{17,18} Ile155 is a highly conserved residue across orthologs (Figure S1) and is located within the LRR domain. A homology model of the protein region spanning the nine LRRs (residues 123-348) was generated on the basis of the available LRR domain structure of internalin H (PDB: 1H6U), using the program Swiss-PdbViewer 4.1;¹⁹ loops were modeled with MODELER v.9.13.²⁰ The generated model fitted well with the established structure of these motifs²¹ and localized the affected residue in the hydrophobic core stabilizing the second LRR motif (Figure 3A). At the same position, the introduction of the hydrophilic asparagine

was predicted to perturb dramatically the structure of the motif and the overall folding of the LRR domain. Since LRR domains are functional modules that mediate protein-protein interaction, this amino acid substitution was expected to affect TBCE function or the overall TBCE structure and protein stability. To verify the latter hypothesis, the level of TBCE was evaluated by western blot analysis in skin fibroblasts obtained from affected subjects 1544334 and 2518864 and four healthy control subjects. Fibroblasts were lysed with RIPA buffer containing protease inhibitors cocktail (Roche), and 30 µg of total proteins were separated on a 4%-12% Bis-Tris gel (Invitrogen) and transferred to polyvinylidene difluoride membrane (Amersham). After blocking, membranes were incubated with anti-TBCE (Novus) (1:1,000) and anti-GAPDH (Sigma) (1:10,000) antibodies, the latter used to normalize samples loading. After washing, membranes were incubated with the secondary antibodies (Jackson ImmunoResearch), and immunoreactive bands were visualized using Lite Ablot Extend Long Lasting Chemiluminescent Substrate



Figure 3. Predicted Structural Impact of the p.IIe155Asn Substitution, and Consequence of TBCE Mutations on mRNA and Protein Levels

(A) Homology model of the TBCE leucine-rich repeat (LRR) domain (residues 123–348, GenBank: NP_001072983.1). Ile155 with surrounding hydrophobic residues contributes to the hydrophobic core required for proper folding of the domain. Introduction of Asn is predicted to disrupt the intramolecular binding network stabilizing the LRR structure.

(B) Western blot (WB) analysis documenting reduced TBCE levels in fibroblasts from affected subjects. Data are presented as means \pm SD, *p < 0.05 (Mann-Whitney test).

(C) Real-time qRT-PCR showing reduced expression level of *TBCE* mRNA in fibroblasts from subject 2518864. *p < 0.05 (Mann-Whitney test). Such reduced level of the transcript is caused by transcript decay of the allele carrying the truncating c.1076delC variant, as documented by Sanger sequencing of *TBCE* cDNA showing solely the transcript with the c.464T>A missense change.

(D) WB analysis demonstrating reduced levels of Tbce in tail fibroblasts from the *pmn/pmn* mouse (Tbce^{Trp524Gly}); recombinant protein (rec. TBCE) is also reported. Data are presented as means \pm SD, *p < 0.01 (Mann-Whitney test).

(Euroclone). Western blot analysis documented a significantly reduced amount of TBCE in fibroblasts from affected individuals compared to control cells (Figure 3B), suggestive of rapid degradation of the TBCE mutant. Of note, reduction in TBCE levels was more accentuated in fibroblasts from subject 2518864, who was compound heterozygote for the p.Ile155Asn and p.Leu360Ter changes. To compare the levels of TBCE mRNA among samples, total RNA from cell lysates was isolated using TRIzol solution (Invitrogen) and 1 µg RNA was reverse transcribed with the SuperScript First-Strand Synthesis system (Invitrogen), using random hexamers as primers. TBCE mRNA expression levels were determined by quantitative realtime PCR using an ABI PRISM 7000 Sequence Detection System (Applied Biosystems) and Power SYBR Green I dye chemistry. Assays were performed in triplicate, with GAPDH as internal controls, and documented a reduction to half of normal of TBCE mRNA in fibroblasts from subject 2518864 compared to what was observed in individual 1544334 and control cells (Figure 3C). Consistent with this finding, the frameshift variant was undetectable at

the mRNA level, as determined by cDNA Sanger sequencing (Figure 3C), indicating nonsense-mediated RNA decay. A similar reduction in Tbce levels was documented in fibroblasts (Figure 3D) of *pmn/pmn* mice. Overall these data indicated that TBCE levels in fibroblasts of subjects with isolated neurodegenerative disease is preserved, in part, in sharp contrast with the extremely reduced or completely absent level of the protein documented in HRDS and KCS.^{3,22}

TBCE is required for the folding of α -tubulin, α/β -tubulin dimerization, and subsequent heterodimer polymerization into microtubules, which are major components of the cytoskeleton.^{23,24} To explore the impact of the two *TBCE* mutations on microtubule polymerization, we first determined the levels of total, soluble, and polymerized α -tubulin in fibroblasts of affected and control subjects, using biochemical fractionation. Total, soluble, and polymerized tubulins from fibroblast lysates were separated by centrifugation (16,000 × g, 30 min, room temperature) as previously described.²⁵ Proteins were solubilized by heating in Laemmli buffer, separated on 10% SDS-PAGE,



Figure 4. Defective Microtubule Polymerization in Subjects with Homozygous/Heterozygous TBCE c.464T>A Missense Change and in pmn/pmn Mice

(A) Western blot (WB) analyses showing total (T), soluble (S), and polymerized (P) α -tubulin (above), detyrosinated tubulin (middle), and β -actin (below) in fibroblasts from affected individuals (1544334, 2518864) and in control cells (Ctrl1, Ctrl2). Diagrams (right) show reduced fractions of polymerized α -tubulin and detyrosinated tubulin in fibroblasts of affected subjects. In both comparisons, differences were statistically significant by Mann-Whitney test (p < 0.01).

(B) WB analyses showing reduced amount of polymerized α -tubulin (P) in fibroblasts from *pmn/pmn* mice compared to control fibroblasts (wt). Differences were statistically significant by Mann-Whitney test (p < 0.01).

(C) Immunofluorescence analysis documenting loss of microtubules containing detyrosinated tubulin (detyr-Tub) in fibroblasts of affected subjects. Note that detyrosinated microtubules (red) in control fibroblasts are closely associated with the GM130-stained Golgi apparatus (green), while the Golgi apparatus in fibroblasts carrying mutated *TBCE* alleles appears to have lost compaction. Nuclei are DAPI stained (blue). Scale bar represents 10 µm.

(D) Immunofluorescence analyses showing reduced re-polymerization of microtubules (labeled for α -tubulin, red) after nocodazole treatment in fibroblasts from subjects 1544334 and 2518864, as compared to control cells (Ctrl). Nuclei are labeled with DAPI (blue). Scale bar represents 20 μ m.

and transferred on membranes, which were immunoblotted with antibodies against α -tubulin (Sigma) (1:5,000), detyrosinated tubulin (a kind gift of Dr. A. Andrieux, INSERM, Grenoble, France) (1:10,000), and β -actin (TuJ1, Babco) (1:10,000). The level of total α -tubulin did not significantly differ among cells (Figure 4A). By contrast, the amount of polymerized α -tubulin was significantly reduced in fibroblasts from affected subjects compared to control cells (Figure 4A), in line with what was observed in tail fibroblasts of *pmn/pmn* mice (Figure 4B). Moreover, significantly reduced levels of detyrosinated α -tubulin, a marker of stable and long-lived microtubules in neurons, were also noted (Figure 4A), indicating again a reduced microtubule stability in fibroblasts with biallelic mutations in *TBCE*.

We hypothesized that defective TBCE function might alter microtubule nucleation and dynamics. To verify this hypothesis, synchronized fibroblasts from individuals with mutated *TBCE* alleles and unaffected controls were cultured in Dulbecco's Modified Eagle Medium supplemented with 10% FBS, seeded on coverslips, and allowed to attach for 24 hr. Cells were treated with the microtubule-disrupting drug nocodazole (Sigma) (10 µM, 35 min at 37°C), and microtubule regrowth was analyzed after drug washout by confocal microscopy analysis, in time-course experiments, after fibroblasts were fixed, and immunostained for a-tubulin (Sigma) and with DAPI (Life Technologies). Cultures were visualized with a confocal microscope Fluoview FV1000 (Olympus) and images were acquired with the software FV10-ASW v.2.0. In control fibroblasts, microtubules began to polymerize efficiently as early as 10 min after nocodazole washout, and extended polymerized structures were well formed at 30 min (Figure 4C). By contrast, microtubule re-polymerization appeared markedly delayed in fibroblasts from affected subjects; microtubules were also less abundant and strongly disorganized in both early and late stages of re-polymerization. Such altered dynamics appeared particularly pronounced in fibroblasts with compound heterozygosity for the p.Ile155Asn and p.Leu360Ter changes, consistent with the more profound reduction of TBCE levels.

TBCE is concentrated at the Golgi membrane in an Arf1regulated manner and facilitates both nucleation rates and polymerization speed of Golgi-derived microtubules.²⁶ Immunofluorescence of synchronized primary fibroblasts was performed to visualize tubulin polymerization using antibodies against detyrosinated tubulin (1:1,000), as a marker of microtubules, and the Golgi membrane, using an anti-GM130 antibody (Becton Dickinson) (1:300). A decreased level of detyrosinated tubulin was observed in fibroblasts from affected subjects (Figure 4D), in line with the previously collected data and the reported loss of Golgi-derived microtubules in motor neurons of *pmn/pmn* mice.²⁶ Of note, such reduced nucleation of microtubules was associated with a decreased compaction of the Golgi apparatus (Figure 4D).

Microtubules are a key cytoskeletal component of the mitotic apparatus, and their centrosomal nucleation coordinates formation and dynamics of the mitotic spindle during mitosis. To explore the occurrence of altered microtubule spindle organization, confocal microscopy analysis of synchronized fibroblasts from subjects with mutated TBCE alleles and unaffected individuals was performed. Fixed cells were stained with pericentrin (Abcam), γ -tubulin (Sigma), and α-tubulin (Abcam) antibodies, followed by the appropriate secondary antibodies (Invitrogen) and DAPI to visualize chromosomes. An abnormal mitotic morphology and disorganized mitotic microtubules were observed; spindles appeared asymmetric and unrefined and exhibited an irregular aster structure (Figure S2), documenting the impact of the hypomorphic defect in TBCE on mitotic spindle formation and organization.

The level of α/β -tubulin heterodimers as well as their polymerization and depolymerization at the microtubule ends are tightly controlled in cells to guarantee proper dynamics of this cytoskeletal component. Since microtubules control multiple neuronal processes, including cell division, migration, function, and survival,²³ neurons are particularly vulnerable to altered microtubule dynamics. In this report, we provide evidence that defective TBCE function underlies a recessive early-onset neurodegenerative disorder and is associated with perturbed microtubule dynamics characterized by decreased and delayed microtubule polymerization. TBCE is a tubulin-binding cofactor required for the formation of α/β -tubulin heterodimers, microtubule polymerization, and also tubulin scavenging. Virtually complete loss of TBCE function has previously been identified to underlie two clinically related developmental disorders, HRDS and KCS1.^{2,3} In contrast, hypomorphic Tbce function has specifically been reported to cause a neurodegenerative disorder in pmn/pmn mice due to retrograde dying back of motor axons and degeneration of motor neuron cell bodies.^{6,7,16,27} The present data provide evidence of a similar neurodegenerative phenotype in humans resulting from defective TBCE function.

Consistent with the presently observed reduced nucleation of microtubules at the Golgi apparatus and decreased compaction of the Golgi apparatus, previous observations in pmn/pmn mice and TBCE-depleted motor neurons have documented that defective TBCE function impairs microtubule polymerization at the Golgi apparatus, leading to defective trafficking of COPI vesicles and Golgi fragmentation, which interestingly represents one of the earliest pathological features of degenerating motor neurons in amyotrophic lateral sclerosis.^{28,29} In sharp contrast, HRDS and KCS are syndromic conditions mainly affecting multiple developmental processes and growth, with no sign of early-onset and progressive neurodegeneration, and in particular no sign of functional alterations involving upper and lower motor neurons. At the moment we have no explanation for this remarkable phenotypic difference. This picture has similarities with the phenotypic heterogeneity associated with mutations in ASAH1 (MIM: 613468), encoding N-acylsphingosine amidohydrolase 1, in which hypomorphic defects have been linked to childhood SMA associated with progressive myoclonic epilepsy (SMA-PME [MIM: 159950]), a rare autosomal-recessive phenotype restricted to the central nervous system generally starting with lower motor neuron disease,²⁹ while a different class of mutations dramatically affecting catalytic activity of the enzyme results in Farber disease (MIM: 228000), a lysosomal storage disease characterized by severe motor and intellectual disability, early-onset subcutaneous lipogranulomata, painful and progressively deformed joints, and hoarseness of the voice.

Another remarkable clinical aspect of TBCE-related neurodegeneration is the MRI pattern resembling NBIA that was documented in the four older affected sibs in their second decade (subjects VN_X3359, VN_X5360, 00997847, and 00997844). We suppose that this pattern might represent a feature evolving with age, possibly appearing during the second decade of life. This pattern is unexpected because iron accumulation has not been reported among the neuropathological features of the *pmn/pmn* mouse.¹⁶ Although additional observations are needed to confirm this unexpected association, this finding suggests a critical role of cytoskeletal components in the complex cellular processes implicated in iron homeostasis in the brain.³⁰

Overall, we describe a distal motor neuropathy, spastic ataxia, and juvenile-onset brain iron accumulation caused by biallelic mutations in *TBCE* overlapping the phenotype of the *pmn/pmn* mouse, a well-known model of motor neuron disease. Expanding beyond the previously documented impact of abolished TBCE function on developmental processes and growth, our findings indicate a stringent requirement of TBCE specifically for neuronal survival and function.

Supplemental Data

Supplemental Data include case reports, two figures, and two tables and can be found with this article online at http://dx.doi. org/10.1016/j.ajhg.2016.08.006.

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Web Resources

Burrows-Wheeler Aligner, http://bio-bwa.sourceforge.net/ CADD, http://cadd.gs.washington.edu/

Chimera, http://www.cgl.ucsf.edu/chimera

dbNSFP v.2.0, https://sites.google.com/site/jpopgen/dbNSFP

dbSNP, http://www.ncbi.nlm.nih.gov/projects/SNP/

ExAC Browser, http://exac.broadinstitute.org/

GATK Best Practices, https://www.broadinstitute.org/gatk/guide/ best-practices

GenBank, http://www.ncbi.nlm.nih.gov/genbank/

NCBI Gene, http://www.ncbi.nlm.nih.gov/gene

OMIM, http://www.omim.org/

Picard, http://broadinstitute.github.io/picard/

RCSB Protein Data Bank, http://www.rcsb.org/pdb/home/ home.do

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