

Spectrum Compact CE System

Sanger Sequencing and Fragment/STR Analysis



- benchtop 4-capillary electrophoresis instrument for 4-, 5- and 6-dye chemistries and up to 32 samples in a single run
- compatible with a broad range of commercial chemistries for DNA sequencing, forensic STR profiling, MLPA, MSI and many more
- flexible raw data analysis
- prefilled, plug-and-play consumables for maximal flexibility and efficiency
- controlled by the integrated touchscreen or by Remote Access Software
- on-site instrument and software installation, operational and application trainings

www.promega.com/spectrum-compact-system





Revised: 18 March 2021 Accepted: 8 April 2021

DOI: 10.1002/humu.24206

RESEARCH ARTICLE



Clinical, neuroimaging, and molecular spectrum of *TECPR2*-associated hereditary sensory and autonomic neuropathy with intellectual disability

Sonja Neuser ¹ Barbara Brechmann ^{2,3} Geli Heimer ^{4,5} Heimer ^{4,5} Keise Brösse ³
Susanna Schubert ¹ Lauren O'Grady ⁶ Michael Zech ^{7,8}
Siddharth Srivastava ² David A. Sweetser ⁶ <a>[Yasemin Dincer ^{9,10}
Volker Mall ^{9,11} Juliane Winkelmann ^{7,8,12,13} Christian Behrends ¹³ <a>D
Basil T. Darras ¹⁴ Robert J. Graham ¹⁵ Parul Jayakar ¹⁶ Barry Byrne ¹⁷
Bat El Bar-Aluma ^{4,5} Yael Haberman ^{4,5,18} Amir Szeinberg ^{4,5}
Hesham M. Aldhalaan ¹⁹ Mais Hashem ¹⁹ Amal Al Tenaiji ²⁰ Omar Ismayl ²⁰
Asma E. Al Nuaimi 20 Karima Maher 20 Shahnaz Ibrahim 21 Fatima Khan 21
Henry Houlden ²² Vijayalakshmi S. Ramakumaran ²³ Alistair T. Pagnamenta ²⁴ 💿
Jennifer E. Posey ²⁵ 💿 📔 James R. Lupski ^{25,26,27,28} 💿 📔 Wen-Hann Tan ²⁹ 💿 🛛
Gehad ElGhazali ²⁰ Isabella Herman ^{25,27,30} 💿 Tatiana Muñoz ³¹
Gabriela M. Repetto ³¹ 💿 Angelika Seitz ³² Mandy Krumbiegel ³³
Maria Cecilia Poli ^{26,31} 💿 Usha Kini ²³ 💿 Stephanie Efthymiou ²² 💿
Jens Meiler ^{34,35} 💿 Reza Maroofian ²² 💿 Fowzan S. Alkuraya ^{19,36} 💿
Rami Abou Jamra ¹ 💿 Bernt Popp ¹ 💿 Bruria Ben-Zeev ^{4,5}
Darius Ebrahimi-Fakhari ² 💿

¹Institute of Human Genetics, University of Leipzig Medical Center, Leipzig, Germany

²Department of Neurology, The F.M. Kirby Neurobiology Center, Boston Children's Hospital, Harvard Medical School, Boston, Massachusetts, USA

³Department of Pediatrics, Hospital for Children and Adolescents, Heidelberg University Hospital, Heidelberg, Germany

⁴Edmond and Lily Safra Children's Hospital, Sheba Medical Center, Ramat Gan, Israel

⁵Sackler Faculty of Medicine, Tel Aviv University, Tel Aviv, Israel

⁶Department of Pediatrics, Division of Medical Genetics and Metabolism, Massachusetts General Hospital, Boston, Massachusetts, USA

⁷Institute of Neurogenomics, Helmholtz Zentrum München, Munich, Germany

⁸Institute of Human Genetics, Klinikum rechts der Isar, Technical University of Munich, Munich, Germany

⁹Lehrstuhl für Sozialpädiatrie, Department of Pediatrics, Technische Universität München, Germany

¹⁰Zentrum für Humangenetik und Laboratoriumsdiagnostik (MVZ), Martinsried, Germany

¹¹kbo-Kinderzentrum München, Munich, Germany

¹²Lehrstuhl für Neurogenetik, Technische Universität München, Munich, Germany

Sonja Neuser, Barbara Brechmann, Gali Heimer, Bernt Popp, Bruria Ben-Zeev and Darius Ebrahimi-Fakhari are equally contributed to this study.

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

 $\ensuremath{\mathbb{C}}$ 2021 The Authors. Human Mutation published by Wiley Periodicals LLC

¹⁴Department of Neurology, Boston Children's Hospital, Harvard Medical School, Boston, Massachusetts, USA

¹⁵Department of Anesthesia, Critical Care and Pain Medicine, Boston Children's Hospital, Harvard Medical School, Boston, Massachusetts, USA

¹⁶Nicklaus Children's Hospital, Miami, Florida, USA

¹⁷Powell Gene Therapy Center, University of Florida, Gainesville, Florida, USA

¹⁸Cincinnati Children's Hospital Medical Center and the University of Cincinnati College of Medicine, Cincinnati, Ohio, USA

¹⁹Department of Translational Genomics, Center for Genomic Medicine, King Faisal Specialist Hospital and Research Center, Riyadh, Saudi Arabia

²⁰Sheikh Khalifa Medical City, Abu Dhabi, United Arab Emirates

²¹Department of Paediatrics and Child Health, Aga Khan University Hospital, Karachi, Pakistan

²²Department of Neuromuscular Disorders, Queen Square Institute of Neurology, University College London, London, UK

²³Oxford Centre for Genomic Medicine, Oxford, UK

²⁴NIHR Biomedical Research Centre, Wellcome Centre for Human Genetics, University of Oxford, Oxford, UK

²⁵Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, Texas, USA

²⁶Department of Pediatrics, Baylor College of Medicine, Houston, Texas, USA

²⁷Texas Children's Hospital, Houston, Texas, USA

²⁸Human Genome Sequencing Center, Baylor College of Medicine, Houston, Texas, USA

²⁹Division of Genetics and Genomics, Boston Children's Hospital, Boston, Massachusetts, USA

³⁰Department of Pediatrics, Section of Pediatric Neurology and Developmental Neuroscience, Baylor College of Medicine, Houston, Texas, USA

³¹Facultad de Medicina, Clinica Alemana Universidad del Desarrollo, Santiago, Chile

³²Department of Diagnostic and Interventional Radiology, Heidelberg University Hospital, Heidelberg, Germany

³³Institute of Human Genetics, Friedrich-Alexander-Universität (FAU), Erlangen, Germany

³⁴Department of Chemistry, Vanderbilt University, Nashville, Tennessee, USA

³⁵Institute for Drug Discovery, University of Leipzig Medical Center, Leipzig, Germany

³⁶Department of Anatomy and Cell Biology, College of Medicine, Alfaisal University, Riyadh, Saudi Arabia

Correspondence

Sonja Neuser, Institute of Human Genetics, University of Leipzig Medical Center, Philipp-Rosenthal-Straße 55, 04103 Leipzig, Germany.

Email: sonja.neuser@medizin.uni-leipzig.de and s.neuser.md@gmail.com

Abstract

Bi-allelic TECPR2 variants have been associated with a complex syndrome with features of both a neurodevelopmental and neurodegenerative disorder. Here, we provide a comprehensive clinical description and variant interpretation framework for this genetic locus. Through international collaboration, we identified 17 individuals from 15 families with bi-allelic TECPR2-variants. We systemically reviewed clinical and molecular data from this cohort and 11 cases previously reported. Phenotypes were standardized using Human Phenotype Ontology terms. A crosssectional analysis revealed global developmental delay/intellectual disability, muscular hypotonia, ataxia, hyporeflexia, respiratory infections, and central/nocturnal hypopnea as core manifestations. A review of brain magnetic resonance imaging scans demonstrated a thin corpus callosum in 52%. We evaluated 17 distinct variants. Missense variants in TECPR2 are predominantly located in the N- and C-terminal regions containing β -propeller repeats. Despite constituting nearly half of disease-associated TECPR2 variants, classifying missense variants as (likely) pathogenic according to ACMG criteria remains challenging. We estimate a pathogenic variant carrier frequency of 1/1221 in the general and 1/155 in the Jewish Ashkenazi populations. Based on clinical, neuroimaging, and genetic data, we provide recommendations for variant reporting, clinical assessment, and surveillance/ treatment of individuals with *TECPR2*-associated disorder. This sets the stage for future prospective natural history studies.

KEYWORDS

Human Phenotype Ontology, neurodevelopmental disorder, sensory autonomic neuropathy, spastic paraplegia, *TECPR2*

1 | INTRODUCTION

TECPR2 belongs to the tectonin β -propeller repeat-containing protein family and is implicated in the autophagy pathway (Oz-Levi et al., 2013; Stadel et al., 2015). Autophagy is critical to the development and function of the central nervous system. Loss-of-function variants in several genes of the autophagy pathway lead to both neurodevelopmental and neurodegenerative diseases (Ebrahimi-Fakhari et al., 2016; Menzies et al., 2017; Teinert et al., 2019).

In 2012, Oz-Levi et al. identified the homozygous *TECPR2* variant c.3416del, p.(Leu1139Argfs*75) in five individuals from three Jewish Bukharian families and classified the syndrome as a novel subtype of hereditary spastic paraplegia (HSP) (SPG49; MIM# 615000) (Oz-Levi et al., 2012). To date, 11 individuals with bi-allelic *TECPR2* variants have been reported (Covone et al., 2016; Heimer et al., 2016; Oz-Levi et al., 2012; Patwari et al., 2020; Zhu et al., 2015). All individuals showed muscular hypotonia and most had global developmental delay followed by intellectual disability. Only a subset of individuals displayed progressive spasticity as a characteristic HSP symptom. An autonomic and sensory neuropathy with respiratory, gastrointestinal and cardiovascular system involvement was present in a subset of individuals and central apnea was found to account for a large part of the morbidity (Heimer et al., 2016; Patwari et al., 2020).

Beside two founder variants (c.3416del, p.(Leu1139Argfs*75) in the Jewish Bukharian background and c.1319del, p.(Leu440Argfs*19) in the Jewish Ashkenazi background), likely derived as new variants under a Clan Genomics hypothesis (Lupski et al., 2011), two other truncating and three missense TECPR2 variants have been associated with the disease. Expression analyses in cell lines transfected with the p.(Leu1139Argfs*75) variant indicated escape from nonsense-mediated RNA-decay (NMD) but the degradation of the truncated protein (Oz-Levi et al., 2012). Functional data is largely missing for other described variants. This poses challenges for the interpretation of missense variants, for which normal expression of an altered protein is expected. All variants have been reported based on the clinical overlap but have yet to be scored through the five-tier variant classification system recommended by the American College of Medical Genetics and Genomics (ACMG) (Richards et al., 2015). The lack of functional data and reliable variant classification have prevented an estimation of carrier frequencies and disease incidence, genotype-phenotype correlation analyses and the ability to make a genetic diagnosis in novel cases.

Through international collaboration, we assembled a cohort of 28 individuals from 24 families of different ethnic backgrounds with

known/novel disease-associated *TECPR2*-variants. Based on a detailed review of the published cases and comparison with the herein described individuals, we provide a systematic quantitative clinical synopsis based on Human Phenotype Ontology (HPO) (Köhler et al., 2019). We provide recommendations for clinical management, including surveillance and symptomatic treatment. Annotation and classification of all disease-associated variants according to the current ACMG recommendations are provided (Richards et al., 2015). Using public databases, we estimate carrier frequencies and disease incidence. Based on this curated phenotype and genotype data set, we propose a framework for reporting and validating *TECPR2* variant alleles.

2 | MATERIALS AND METHODS

2.1 | Editorial policies and ethical considerations

This study adheres to the principles set out in the Declaration of Helsinki. The following Research Ethics Committees approved the study: Ethical Committee of the Medical Faculty, Leipzig University (P1), Institutional Review Board at Boston Children's Hospital (IRB-P00033016; P2, P4, and P5), Ethics Review Board of Technical University of Munich (P3), Institutional Review Board of King Faisal Specialist Hospital and Research Center (KFSRHC RAC# 2080006 and 2121053; P7, P8, and P13), Institutional Review Board at University College London (P14 and P15, SYNaPS cohort), East of England and South Cambridge Research Ethics Committee (REC: 14/EE/1112) for 100,00 Genomes Project Protocol (P16), Institutional Review Board at Baylor College of Medicine (H-29697) and Comité Etico Cientifico at Facultad de Medicina, and Clinica Alemana Universidad del Desarrollo (P17). Genetic testing for P6, P9, P10, P11, and P12 was performed in a diagnostic setting. The authors received and archived written consent of the legal guardians to publish genetic and clinical data (P1 - P17) as well as photographs, computed tomography (CT) scan, and magnetic resonance imaging (MRI) images (P1, P4, P6, P11, P13, P15, P16, P17).

2.2 Cohort

All 17 individuals described herein (P1-P17) were recruited through GeneMatcher (Sobreira et al., 2015) or personal communication, from

Human Mutation-WILEY

different institutions in Germany, Israel, the United States, Saudi Arabia, the United Arab Emirates, Great Britain, Pakistan, and Chile. Genotypic data from P3 and P13 were previously reported without a detailed clinical description (P3: reported as CB-DYS-125 in Zech et al., 2020; P13: reported as 09DG00835 (Shams Anazi et al., 2017)).

2.3 | Clinical spectrum

Molecular and clinical data were collected from the referring clinicians using a standardized questionnaire. All affected individuals were evaluated by a pediatric neurologist and/or geneticist. Reports of brain MRI scans were available from 15 individuals. Clinical terms were standardized using Human Phenotype Ontology (HPO) terminology (Köhler et al., 2019). Clinical features were grouped into six categories (phenotypical abnormalities of body and face, intellectual and social development, neurological system, respiratory system, gastrointestinal system, and diagnostic procedures). Detailed case descriptions for all included individuals are provided in Supporting Information Files S1 and S2 (sheet "clinical_table").

2.4 | Genetic analyses

Genomic DNA was extracted using standard methods from peripheral blood samples of probands/parents. For P1, P16, and P17 conventional karyotyping was performed and all individuals, except P4, P14, P15, P16, and P17, received a chromosomal microarray. *TECPR2* variants were identified by gene panel analysis (P13), exome (P14 and P15), trio exome (P1 to P6, P10, P17), quad exome (P7 and P8), trio genome (P16), or targeted Sanger sequencing (P9, P11, P12). All herein identified *TECPR2* variants have been submitted to ClinVar (Supporting Information File S3 sheet "TECPR2_variants").

2.5 | Review of published cases

A PubMed search identified five publications (Covone et al., 2016; Heimer et al., 2016; Oz-Levi et al., 2012; Patwari et al., 2020; Zhu et al., 2015) describing 11 individuals from nine families diagnosed with *TECPR2*-associated disease (searched on 2020-09-10). Phenotypic features were extracted from published reports using the same questionnaire applied to novel cases.

2.6 | Variant annotation and scoring

Variants were standardized to the *TECPR2* reference transcript NM_014844.4 (GRCh37/hg19) using Mutalyzer 2.0.32 (Wildeman et al., 2008) and annotated as described previously (Popp et al., 2017) with up-to-date versions of all tools (Cingolani, Patel, et al., 2012; Cingolani, Platts, et al., 2012; Freeman et al., 2018; Liu et al., 2013) and scores (Jian et al., 2014; Rentzsch et al., 2019; Xiong et al., 2015) (for

details see Supporting Information File S1). All diagnostic *TECPR2* variants were subsequently reclassified (Supporting Information File S3 sheet "TECPR2_variants") following ACMG guidelines (Richards et al., 2015).

2.7 | Estimation of carrier frequencies from public databases

We retrieved all *TECPR2* variants from gnomAD (Karczewski et al., 2020) and BRAVO (see Web Resources). These were annotated, scored, and filtered for classification as (likely) pathogenic as described before to calculate carrier frequencies (Hebebrand et al., 2019).

2.8 | Analysis of missense variant spectrum and modeling of TECPR2 protein structure

The distribution of *TECPR2* missense variants in the secondary protein structure was compared to missense variants reported as homozygous in public population databases and protein regions constrained for missense variation were analyzed as described (Hebebrand et al., 2019). For analysis of the tertiary structure, we used the GalaxyWEB pipeline (Heo et al., 2013; Ko, Park, Heo, et al., 2012; Ko, Park, & Seok, 2012) to divide TECPR2 protein sequence into modeling units, predict their structure, and refine the top model. Protein data bank (PDB) format structures (Popp & Neuser, 2020) were then used for visualization with a pipeline using the Pymol software (Meyer et al., 2016) and missense clustering analysis as described before (Hebebrand et al., 2019). For details, also see Supporting Information Notes S1.

2.9 | RNA expression analysis for the *TECPR2* variant c.2829del, p.(Asn944Thrfs*7) in P1

Messenger RNA (ribonucleic acid) from peripheral blood lymphocytes of P1 and both parents was used to generate complementary DNA (cDNA). Monoallelic expression was analyzed with reverse transcription polymerase chain reaction (RT-PCR), and Sanger sequencing and *TECPR2* expression were analyzed using quantitative PCR (qPCR) (see details in Supporting Information File S1).

3 | RESULTS

3.1 | TECPR2 Variant Spectrum

Genetic analyses, including conventional karyotyping, chromosomal microarray analysis, and multigene panels (except for P13) were unremarkable in all novel cases. Seventeen distinct variants in *TECPR2*, including nine truncating and eight missense variants, were identified. Of these, five truncating and five missense variants have not been reported previously (Figure 1a).

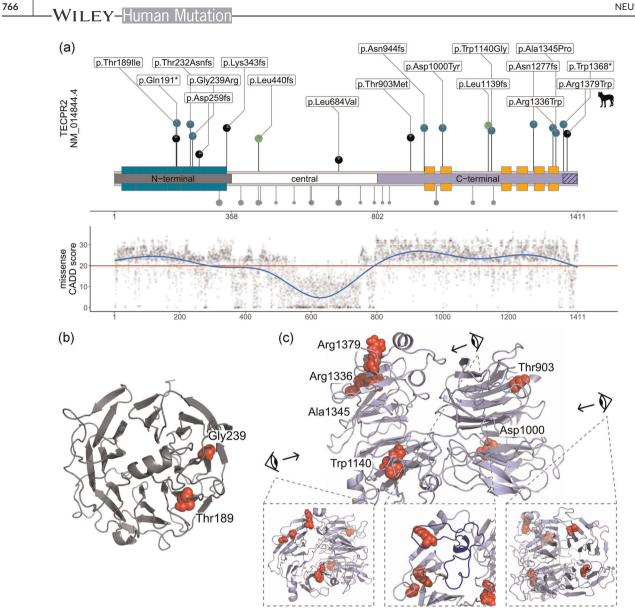


FIGURE 1 TECPR2 structure with variant distribution and computational scores. (a) Schematic of the TECPR2 protein with WD40 and TECPR repeat units (WD40: green, TECPR: orange; based on Uniprot O15040) and three modeling units ("N-terminal": gray, "central": white, "C-terminal": purple) identified by GalaxyDom. Disease-associated variants identified in the cohort are depicted toward the top. The length of the segments corresponds to each variant's CADD score. Blue dots represent novel identified variants, black dots represent variants reported in the literature, and green dots represent the founder variants. Gray dots downwards show homozygous variants from gnomAD, the dot size represents the logarithm of the allele count. In the panel below, a generalized additive model shows the values of CADD PHRED v1.6 for all possible missense variants in TECPR2 across the linear protein structure. The red horizontal line marks the recommended cut-off (20). (b) Homology model of the N-terminal domain (AA 1–357; gray) generated through the GalaxyTBM pipeline showing the 7-bladed β -propeller fold typical for WD40 repeat. The position of missense variants identified in the individual P3 (Gly239) from our study and "Family E II-1" (Thr189) from the literature review are presented as red spheres. Both missense variants affect conserved residues in β-propeller folds. (c) Lateral overview of the homology model of the C-terminal domain (AA 802–1411; blue) showing the two β -propeller folds in the TECPR repeat unit. The position of missense variants identified in the individuals P7 and P8 (Asp1000), P6 (Trp1140), P17 (Arg1336), and P3 (Ala1345) from our study and "Family H I-1" (Thr903) from the literature review and (Arg1379) from the Spanish water dogs (Supporting Information Notes S1 and Figure S1) are presented as red spheres. The blue highlighted part of the protein structure in the middle panel is truncated by the most downstream stop gained variant c.4103G>A, p.(Trp1368*) identified in P14 and contains the amino acid position described as pathogenic in Spanish water dogs

3.2 | Founder variants

The first reported founder variant (Oz-Levi et al., 2012) in the Jewish Bukharian population c.3416del, p.(Leu1139Argfs*75) was identified in

the homozygous allelic state in five individuals from the literature and in two cases in our cohort. Additionally, the variant was discovered in a compound heterozygous state with the Jewish Ashkenazi founder variant in one previously reported individual. Two previously reported individuals and four cases in our cohort were homozygous for the founder variant in the Jewish Ashkenazi population, c.1319del, p.(Leu440Argfs*19). This variant was also found in a compound heterozygous state with a missense variant (Heimer et al., 2016) and another truncating variant (in our cohort). The two founder variants are located in exons 8 and 16, respectively. GnomAD minor allele frequency (MAF) was 37/275,698 for c.1319del, p.(Leu440Argfs*19) and 2/247,472 for c.3416del, p.(Leu1139Argfs*75). There were no entries for the homozygous occurrence of these variants in the reference populations with data available.

3.3 | Other truncating variants

Among the cases derived from the literature, one individual carried compound heterozygous frameshift variants (c.774del, p.(As-p259Metfs*44); c.1028_1032del, p.(Lys343Argfs*2)). Novel identified truncating variants were c.571C>T, p.(Gln191*) (homozygous), c.694dup, p.(Thr232Asnfs*15) (homozygous), c.2829del, p.(Asn944Thrfs*7) (homozygous), c.3830del, p.(Asn1277Thrfs*43) (compound heterozygous with Ashkenazi founder variant) and c.4103G>A, p.(Trp1368*) (homozygous). The variants are located in exons 5, 6, 7, 12, 18, and 20. MAF in the heterozygous state was consistent with ultrarare variant alleles (Hansen et al., 2019) and between 0 and 2/251,490 (gnomAD).

3.4 | Expression analysis of the stop codon containing transcript in P1

Sanger sequencing of cDNA showed comparable detection of the normal allele and the allele with the c.2829del, p.(Asn944Thrfs*7) variant in both carrier parents of individual P1 (Figure 2a). Additionally, RT-PCR indicated normal expression in individual P1 who is homozygous for the variant (Figure 2b). Comparable expression of *TECPR2* in individual P1, his parents, and in-house controls was confirmed by qPCR (Figure S3).

3.5 | Missense variants

To date, only three disease-associated missense variants have been reported (c.566C>T, p.(Thr189IIe); c.2050C>G, p.(Leu684VaI); c.2708C>T, p.(Thr903Met)). Novel variants identified include three homozygous missense variants c.2998G>T, p.(Asp1000Tyr), c.3418T>G, p.(Trp1140Gly) and c.4006C>T, p.(Arg1336Trp) as well as two compound heterozygous missense variants c.715G>A, p.(Gly239Arg) and c.4033G>C, p.(Ala1345Pro). All variants are predicted to be deleterious by multiple in silico prediction programs except for the previously described variant c.2050C>G, p.(Leu684VaI) (CADD PHRED v1.6: 5.5; mean for all reported missense: 24.4). For a complete overview of in silico analyses please refer to Supporting Information File S3. Similar results were obtained for the MAF, which is between 0 and 21/282,852, again except for c.2050C>G, p.(Leu684VaI), which showed

an MAF of 11,974/282,150. In addition, this variant is found homozygous in gnomAD (440x).

Analysis of spatial distribution in the linear protein structure indicated that missense variants identified in the bi-allelic state in individuals with *TECPR2*-associated disease are predominantly located in the N-terminal (amino acid (AA) 1–357) and C-terminal (AA 802–1411) protein regions. These two regions display a higher restraint for missense variation as indicated by higher computational scores and depletion of homozygous missense variants (Figure 1a and S1).

This finding is further supported by the missense variant described in Spanish water dogs (Hahn et al., 2015), which is highly conserved (CADD PHRED score v1.6: 27.2) and located near to the c.4033G>C, p.(Ala1345Pro) variant (P3) in the C-terminal region; also the amino acid residue affected by this variant is truncated by the late stop variant c.4103G>A, p.(Trp1368*) identified in P14 (see Supporting Information Notes S1 and Figure S1).

Our spatial proximity analysis using predicted 3D protein structures failed to identify clusters of missense variants (Table S3) but showed that all affect highly conserved residues in the repeats forming the N-terminal 7-bladed WD40 β -propeller or the two predicted C-terminal β -propeller structures (Figures 1b,c and S2). While we chose the GalaxyTBM (Ko, Park, & Seok, 2012) model for visualization of the spatial missense distribution in Figure 1, the structural similarity of the model predicted de novo by the trRosetta algorithm (Yang et al., 2020) is remarkable (Figure S2 and Table S2). This convergence of structure prediction algorithms adds confidence to the derived models and will thus accelerate our understanding of missense variants in genetic disorders lacking experimentally derived protein structures.

3.6 | Carrier frequency for (likely) pathogenic *TECPR2* variants

Our results indicate that at least 1 in 1221 individuals (0.082%) in gnomAD and 1 in 1610 individuals (0.062%) in BRAVO is a carrier of a (likely) pathogenic variant in *TECPR2*. In gnomAD, we were able to estimate the carrier frequency for eight subpopulations, which ranged from 1 in 155 (0.650%; Jewish Ashkenazi) to 1 in 7654 (0.013%; South Asian). Using these frequencies, the expected incidence is at least 1 in 5,961,640 newborns (based on gnomAD) to 1 in 10,366,419 newborns (based on BRAVO). Of the analyzed populations (which did not include the Jewish Bukharian population) the highest incidence is expected in the Jewish Ashkenazi population with 1 in 95,864 newborns.

3.7 | Predicted tertiary TECPR2 protein structure

The three different protein modeling algorithms that we have used (Popp & Neuser, 2020) indicated similar results for the overall TECPR2 tertiary structure. The N-terminal domain (AA 1–357) containing seven WD-repeats is predicted to form a 7-bladed

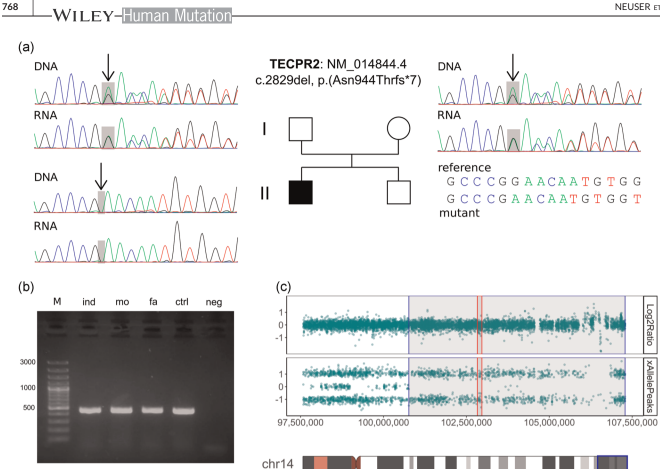


FIGURE 2 Exemplary Sanger sequences, RT-PCR, and CMA results for P1. (a) Chromatograms of DNA (Sanger sequencing) and RNA (RT-PCR on PAXgene stabilized blood) of P1 (down left) and his parents (up left and right). (b) Gel electrophoresis of cDNA-amplicon. (c) CMA data for individual P1 showing an unremarkable copy number of chromosome 14 (Log2Ratio top) and SNP allele peak distribution (AllelePeaks bottom) showing a 6.52 Mb run-of-homozygosity (blue) containing TECPR2 (red). cDNA, complementary DNA; CMA, chromosomal microarray; RT-PCR, reverse transcription polymerase chain reaction; SNP, single-nucleotide polymorphism

 β -propeller fold (WD40 domain) with high similarity in all models generated. The central region (AA 358-801) could either not be modeled completely due to a lack of template structures or resulted in unstructured and highly diverging models. The C-terminal domain (AA 802-1411), containing the six TECPR-repeats annotated from UniProt, was predicted to form a double β-propeller motif in most models with good structural similarity and five to seven blades per propeller. Overall, this indicates a structured C-terminal WD40domain and TECPR-repeat containing a structured double β -propeller motif in the C-terminus, linked by a 444 AA long unstructured peptide (Figures 1b,c and S2).

3.8 **Clinical spectrum**

In our cohort of newly diagnosed cases, 11 of 17 individuals were male. Age at last follow-up was between 16 months and 15 years with a mean of 65.2 ± 43.7 (SD) months. Consanguinity was reported in 7 out of the 15 families. Five families were of Jewish Ashkenazi descent, two families were of Jewish Bukharian descent. Except for P1, all individuals were born at term without significant pre- or perinatal complications. Three individuals were small for gestational age. Head circumference at birth was generally within normal limits. At last follow-up, only seven individuals displayed short stature with a height below -2 SD from age-matched controls, however, all 11 individuals with data available were below average height. Brachycephaly and microcephaly were observed in seven and four individuals, respectively, with three individuals presenting both. Distinct facial features were seen in 11 individuals though were not uniform. Shared characteristics included a short neck, synophrys and a triangular-shaped face, still a recognizable pattern, or facial gestalt, was not appreciated. Skeletal abnormalities, including significant lumbar kyphosis, a barrel-shaped chest, or hyperextension of the neck were present in five cases.

The ages at diagnosis in our cohort ranged between 13 months and 15 years with a mean of 55.6 ± 48.8 (SD) months. All affected individuals showed global developmental delay and later intellectual disability (DD/ ID) in the mild (n = 1), moderate (n = 7), and severe (n = 8) ranges. P2 had only mildly delayed gross motor skills at the last investigation, but her young age rendered a detailed assessment difficult. Six individuals with moderate or severe development delay were reported to have behavioral dysregulation with hyperactivity, restlessness, and aggressive behaviors. Two received a formal diagnosis of autism spectrum disorder. **4 D** Ten children (age range: 16 months to 8 years) had not started walking at the time of the last follow-up and 7 individuals walked independently (mean age: 40.5 ± 36.2 (SD) months). P3 was diagnosed with dystonic/

the time of the last follow-up and 7 individuals walked independently (mean age: 40.5 ± 36.2 (SD) months). P3 was diagnosed with dystonic/ dyskinetic cerebral palsy and started walking around the age of 10 years. Speech development was delayed in all children and speech remained limited to a few words with five individuals remaining completely nonverbal.

The most common neurological manifestations in our cohort included axial and appendicular hypotonia (17/17) accompanied by gait ataxia (11/11), hyporeflexia of the lower limbs (13/17), and dysarthria (6/8). Autonomic dysfunction, for example, temperature instability (3/14) and hyperhidrosis (2/14) were noticed in a subset of cases (5/15). Four individuals were reported to have impaired pain sensation (4/16). Febrile seizures were found in P1 as well as P10: P13, P14, and P15 were reported to have medically refractory epilepsy and peripheral neuropathy was diagnosed in P6. Hearing impairment (3/13) and visual impairment (5/12) were present in a subset. The constellation of central respiratory dysregulation, dysphagia, and neuromuscular-derived respiratory insufficiency was common, resulting in central nocturnal (8/13) and/or daytime (5/16) hypoventilation, dysphagia (9/17), and impaired clearance of secretions. This was complicated by recurrent respiratory infections (14/ 15), aspiration events (10/15), gastroesophageal reflux disease (9/ 15), necessitated noninvasive positive pressure ventilation (i.e., nocturnal BiPAP) (2/13), and utilization of gastrostomy tubes (6/11) in a subset. Airway malformation, such as laryngeal cleft or laryngomalacia, was identified in a subset (4/17). Five individuals (5/15)were reported to have chronic and significant constipation.

Clinical manifestations of previously reported individuals are summarized in Supporting Information File S2. One case (Covone et al., 2016) was excluded from further analysis since the variant c.2050C>G, p.(Leu684Val) was classified as likely benign according to ACMG criteria. In summary, manifestations shared by the majority of all 27 individuals include global development delay and intellectual disability (26/26, 100%), muscular hypotonia (27/27, 100%), hyporeflexia of the lower limbs (22/27, 83%), and gait ataxia (19/19, 100%). Peripheral neuropathy, dysarthria, and abnormal facial features were found in 9/12 (75%), 12/14 (86%), and 19/25 (76%) of individuals with sufficient data available (Table 1). Recurrent respiratory infections (23/25, 92%), gastroesophageal reflux in infancy (18/25, 72%), and nocturnal hypoventilation (12/17, 71%) affected most individuals.

3.9 | Brain imaging and EEG

A review of 16 brains' MRI studies from our cohort (Figures 3 and S4) and a review of reported cases in the literature defined a thin corpus callosum as a common feature (11/21, 52%). Additional findings in a subset of individuals included mild ventriculomegaly (often asymmetric colpocephaly), delayed myelination, and diffuse cerebral atrophy. EEG (electroencephalogram) was abnormal in four cases (4/15, 27%), but no specific pattern was reported.

4 | DISCUSSION

We here report a series of 17 individuals with bi-allelic TECPR2 variants from eight nonconsanguineous families and nine consanguineous families and combine the detailed clinical, imaging, and molecular characterization of these individuals with the 11 cases previously reported. Since the variant c.2050C>G, p.(Leu684Val) was classified as likely benign according to ACMG criteria, one previously reported case (Covone et al., 2016) was excluded. The proband's different clinical presentation without developmental delay, autonomic nervous system involvement or abnormal facial shape supports the variant assessment. Additionally, an inherited variant of unknown significance in SPG7 was reported as an additional genetic finding (Covone et al., 2016). The analysis of the remaining 27 individuals defines a core set of clinical and molecular features. These consist of global developmental delay and intellectual disability, axial and appendicular hypotonia, dysarthria, and an abnormal gait, often described as an ataxic gait. Peripheral neuropathy was found in two-thirds of all individuals in whom a detailed neurological assessment was available. Along with this, hyporeflexia was common and signs of autonomic dysfunction were prominent in the majority of cases. The latter included central hypoventilation, impaired temperature, and blood pressure regulation, repeat aspiration events, and evidence of abnormal gastrointestinal motility. These features imply the involvement of both the central and peripheral nervous systems and substantiate features of hereditary sensory and autonomic neuropathy (HSAN).

Whereas spasticity was recognized as a hallmark feature in the individuals initially reported (Oz-Levi et al., 2012), the overall prevalence of spasticity was limited to a subset in our analysis (24%). We recognize that this is a potentially age-dependent manifestation since increased tone was mainly reported in older individuals (P3 at age 15 years; Family B II-2 at age 20 years). P3 stands out because of the presence of dystonia, which was not present in previously published cases and possibly further broadens the spectrum of neurological symptoms. Of note, epilepsy was relatively infrequent in our cohort and consisted of two individuals who experienced febrile seizures, two previously reported siblings with infrequent generalized tonic-clonic seizures and three individuals with medically refractory seizures. Future studies will be necessary to reassess epilepsy as an associated feature. Overall, the wide neurological manifestations in individuals with TECPR2-associated disease along the age spectrum, point to an involvement of multiple areas of the central nervous system (i.e., cortico-spinal tracts, cerebral cortex, brain stem, possibly basal ganglia) as well as the peripheral nervous system.

A large part of the morbidity and mortality associated with *TECPR2* results from central hypoventilation requiring therapy with noninvasive positive pressure ventilation and occasionally active mechanical ventilatory support. Our findings are supported by a recently published, detailed analysis of the distinct breathing pattern from one affected individual (Patwari et al., 2020).

Based on our clinical experience and the reported disease manifestations, we suggest a framework for routine surveillance as detailed in Table 2. Symptomatic treatment should be tailored to 770

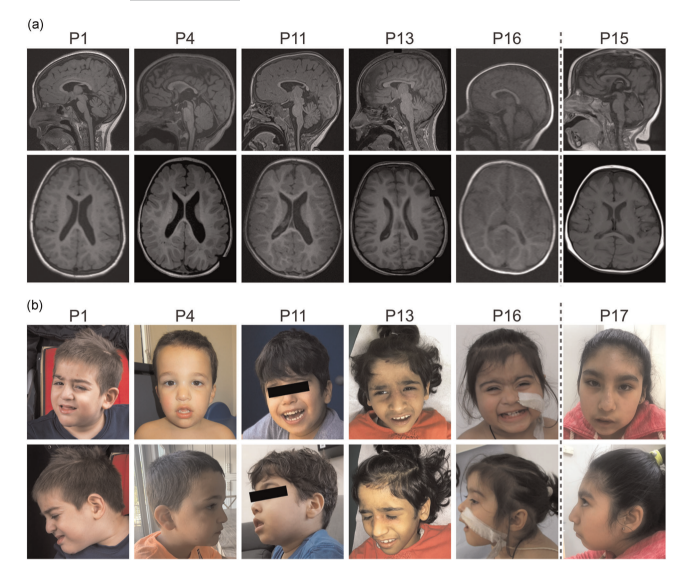


FIGURE 3 MRI and facial features of individuals with *TECPR2*-associated disease. (a) T1 axial and sagittal images of P1 at the age of four years show delayed myelination, mild ventriculomegaly, and periventricular gliosis. T1 axial and sagittal images of P4 at the age of one year show mild thinning of the posterior corpus callosum and/or mildly hypoplastic corpus callosum with mild lateral ventriculomegaly. T1 axial and sagittal images of P11 at the age of 2 years show thin corpus callosum, dysmorphic ventricles, and mild cerebral and cerebellar atrophy. T1 axial and sagittal images of P16 at the age of 2 years show thinning of the corpus callosum and cerebellar vermis mild atrophy. T1 axial and sagittal images of P16 at the age of 2 years show dysmorphic ventricles and a reduction in white matter volume. T1 axial and sagittal images of P15 at the age of 3 years show rounded posterior horns of the bilateral lateral ventricles, cerebral, and mild cerebellar atrophy. (b) Facial images of P1 (5 years 4 months), P4 (4 years front, 7 years lateral), P11 (4 years front, 3 years lateral), P13 (8 years), P16 (5 years), and P17 (10 years 11 months). Individuals with both facial and MRI are ordered vertically in (a) and (b); the dotted line indicates that for P15 only MRI images are shown while for P17 only facial images are shown. MRI, magnetic resonance imaging

each individual case and aims at preserving function and preventing long-term morbidity and mortality. Early developmental support should be maximized to harness the developmental potential.

Overall, our cross-sectional analysis suggests that there is evidence of disease progression from a predominantly neurodevelopmental disorder with global developmental delay and hypotonia in early childhood to progressive disease with corticospinal and corticobulbar dysfunction later in life. We know from personal communications about the disease course of previously reported patients (Heimer et al., 2016; Oz-Levi et al., 2012), who all lost the ability to walk.

Due to largely nonspecific initial clinical features, individuals with *TECPR2*-associated disease may initially receive a diagnosis of cerebral palsy. In addition to an often unremarkable perinatal history, clinical features that help distinguish *TECPR2*-disease from cerebral palsy include the findings of central apnea/hypoventilation, autonomic instability, hyporeflexia as well as other signs of peripheral neuropathy. Brain MRI in *TECPR2*-associated disease shows a

Group	Phenotype	ОДН	Novel cases (%)	ises (%)	Literature cases (%)	cases (%)	All cases (%)	es (%)
Phenotypical abnormalities of body and face	Abnormal facial shape	HP:0001999	69	(11/16)	89	(8/6)	76	(19/25)
	Short stature	HP:0004322	44	(7/16)	78	(2/6)	56	(14/25)
	Abnormality of skeletal morphology	HP:0011842	29	(5/17)	100	(2/2)	37	(7/19)
	Microcephaly	HP:0000252	47	(7/15)	75	(6/8)	57	(13/23)
	Brachycephaly	HP:0000248	25	(4/16)	56	(5/9)	36	(9/25)
Intellectual and social development	Behavioral abnormalities	HP:0000708	43	(6/14)	100	(4/4)	56	(10/18)
	Global developmental delay; Intellectual disability	HP:0001263; HP:0001249	100	(16/16)	100	(10/10)	100	(26/26)
	Moderate	HP:0011343; HP:0002342	44	(7/16)	14	(1/7)	35	(8/23)
	Severe	HP:0011344; HP:0010864	50	(8/16)	86	(6/7)	61	(14/23)
Neurological system	Muscular hypotonia	HP:0001252	100	(17/17)	100	(10/10)	100	(27/27)
	Hyporeflexia of lower limbs	HP:0002600	76	(13/17)	60	(9/10)	81	(22/27)
	Gait ataxia	HP:0002066	100	(11/11)	100	(8/8)	100	(19/19)
	Dysarthria	HP:0001260	75	(8/8)	100	(9/9)	86	(12/14)
	Peripheral neuropathy	HP:0009830	25	(1/4)	100	(8/8)	75	(9/12)
	Impaired pain sensation	HP:0007328	25	(4/16)	100	(4/4)	40	(8/20)
	Abnormality of the autonomic nervous system	HP:0002270	33	(5/15)	100	(10/10)	60	(15/25)
	Abnormal systemic blood pressure	HP:0030972	0	(0/13)	88	(2/8)	33	(7/21)
	Visual impairment	HP:0000505	44	(7/16)	75	(3/4)	50	(10/20)
	Hearing impairment	HP:0000365	24	(4/17)	100	(2/2)	32	(6/19)
Respiratory system	Recurrent respiratory infections	HP:0002205	93	(14/15)	60	(9/10)	92	(23/25)
	Aspiration	HP:0002835	67	(10/15)	50	(4/8)	61	(14/23)
	Nocturnal hypoventilation	HP:0002877	62	(8/13)	100	(4/4)	71	(12/17)
	Central hypoventilation	HP:0007110	31	(5/16)	100	(6/6)	56	(14/25)
GI system	Gastroesophageal reflux at infancy	HP:0002020	60	(9/15)	90	(9/10)	72	(18/25)
	Dysphagia	HP:0002015	53	(9/17)	11	(1/9)	38	(10/26)
Diagnostic procedures	Abnormal corpus callosum morphology	HP:0001273	53	(8/15)	50	(3/6)	52	(11/21)
	Abnormality of metabolism/homeostasis	HP:0001939	62	(8/13)	25	(1/4)	53	(9/17)

 TABLE 1
 Clinical manifestations of TECPR2-associated disease

Note: Phenotypes observed >70 % of all cases are presented in bold.

771

-WILEY-Human Mutation

Every 6 months

- Neurological examination, including a developmental assessment
- Evaluation by a pulmonologist
- Evaluation by a gastroenterologist and consultation with a dietician
- Consider orthopedic evaluation
- Consider venous blood gases
- Consider BERA test if there is clinical concern for hearing impairment

Every 12 months

- Polysomnography study
- Consider chest X-ray
- Consider sputum culture
- Consider spine X-ray
- Consider swallow study (unless fed by gastrostomy)
- Consider echocardiography for signs of pulmonary hypertension
- Consider blood pressure monitoring
- Consider arterial blood gases
- Fasting glucose, electrolytes, and liver function tests (also during intercurrent illnesses)
- Consider ENT evaluation if snoring or consistent tonsillar enlargement

Supportive therapy-recommendations

Routine treatments by a physiotherapist, occupational therapist, and speech therapist Routine chest physiotherapy and mechanical insufflator-exsufflator device Consider antacids, H2 blockers or PPI if GERD present Consider gastrostomy tube and fundoplication if severe GERD/aspiration present Consider adenoidectomy/tonsillectomy if obstructive sleep apnea present Consider continuous nighttime pulse oximetry depending on sleep study results Consider nighttime noninvasive ventilation depending on sleep study results Use sedatives with caution given reports of prolonged effects in this patient population

thinning of the posterior parts of the corpus callosum in about half of individuals. This finding can help guide diagnostic testing.

A diagnosis is achieved through molecular testing. With the identification of novel truncating and missense variants, we confirm and broaden the spectrum of disease-associated variants in *TECPR2*associated hereditary sensory and autonomic neuropathy with intellectual disability. All individuals in the cohort with distinct ethnic origins carried the respective founder variant. This observation affirms the expected genotypic trait in the Jewish Ashkenazi and Jewish Bukharian populations. However, the identification of other truncating variants provides evidence for the occurrence of *TECPR2*-associated disease in other ethnic groups. For all families with homozygous variants other than the founder variants, consanguinity of the parents was reported. This is exemplified for P1 where the run-of-homozygosity on chromosome 14 was not described in the CMA report because it was below the 10 Mb (mega base) filtering cutoff (Figure 2c). Similar results were reported for P2 and P16 (Supporting Information File S1).

Our analysis did not show clustering or specific distribution pattern of the truncating variants. RNA analysis of the novel frameshift variant c.2829del, p.(Asn944Thrfs*7), identified in P1, indicated escape from nonsense-mediated decay. This argues against NMD and is in line with previous results in cell lines showing no effect on mRNA levels for the Jewish Bukharian variant c.3416del, p.(Leu1139Argfs*75) (Oz-Levi et al., 2012), but instead the resulting truncated protein being targeted for proteasome-mediated degradation after translation.

In contrast, all disease-associated missense variants in this cohort affect conserved residues in repeats forming the blades of βpropeller structures at the C-terminal and the N-terminal ends of the protein (Figure 1). As we could not identify clustering in the tertiary structure, misfolding and subsequent degradation could cause loss of the protein carrying these missense substitutions. All five individuals from our cohort harboring missense variants showed moderate to severe DD/ID and are as severely affected as individuals with truncating variants. Therefore, our data do not indicate milder clinical manifestations in carriers of missense variants. This clinical observation further supports a similar pathomechanism, for example, degradation of truncated or misfolded proteins, for both truncating and missense variants. However, due to the currently limited knowledge about TECPR2 function and lack of well-established and readily available functional tests, in most cases, missense variants cannot be classified as (likely) pathogenic according to ACMG guidelines. Based on our computational analyses, we propose to consider the following criteria for the interpretation of TECPR2 missense variants: (1) variant position in the functional domains identified through our conservation and modeling analyses (PM1_Supporting; Figures 1a, S1, and S2), (2) deleterious effect predicted by in silico CADD score with cutoff >20 (PP3; Figure 1a), (3) the patient's phenotype matches the core features as well as TECPR2-specific symptoms of the HSAN-spectrum (Table 1) and exome-wide analyses do not reveal other clinically relevant findings (PP4), and (4) cosegregation of the identified variants with multiple

TABLE 2 Recommendations for surveillance and symptomatic treatment

Human Mutation-WILEY

affected family members (PP1). In this context, it should be noted that large deletions spanning the whole *TECPR2*-gene, in the homozygous state or *in trans* with another pathogenic variant, would reduce the gene dosage and be consequently classified as pathogenic. This also applies to intragenic duplications disrupting the *TECPR2* reading frame (cf. Supporting Information File S2 sheet "allClinVar"). However, large duplications encompassing all of *TECPR2* are not expected to disrupt the reading frame and should be evaluated according to the above recommendations for missense variants and followed up functionally (e.g., expression analyses on RNA level).

Our estimation of the carrier frequency is based on automated ACMG classification of variants and therefore includes only potentially truncating variants. Given that eight of the 17 unique variants are missense variants and considering a large number of uncharacterized TECPR2 missense variants (729 in gnomAD and 448 in BRAVO), we anticipate that the true carrier frequency for (likely) pathogenic TECPR2-variant might be double our current estimate of 0.082% in the general population. Notably, the estimated carrier frequency is 7.9× higher (0.650%) in individuals with Jewish Ashkenazi background and at least 16.2× higher (1.33%) with Jewish Bukharian background (Oz-Levi et al., 2012). A review of carrier screening tests for individuals of Jewish descent showed that TECPR2 is currently included in four offered tests (see Table S1). Overall, based on these high carrier frequencies, both founder variants should be included in commercial carrier screening tests to inform genetic counseling and diagnostics in Jewish couples at increased risk for children with TECPR2-associated disease.

TECPR2 encodes a protein that is implicated in the early steps of the autophagy pathway where it interacts with the Atg8 family proteins, including LC3, to promote autophagic vesicle formation (Behrends et al., 2010). Fibroblasts from affected individuals showed a decreased number of autophagosomes and reduced delivery of LC3 and p62 for lysosomal degradation; this suggests an impairment of autophagic flux (Oz-Levi et al., 2012). Providing insights into the mechanism of defective autophagy, a subsequent study showed that TECPR2 is involved in maintaining functional endoplasmic reticulum exit sites, which are implicated in the cargo from the endoplasmic reticulum to Golgi and may serve as scaffolds for the formation of autophagosomes (Stadel et al., 2015).

While the precise role of autophagy in *TECPR2*-associated disease remains to be established, there are several clinical features that are shared with other single-gene disorders of this pathway (Ebrahimi-Fakhari et al., 2016; Teinert et al., 2019). This includes the involvement of multiple brain areas, clinical signs that point to a progressive involvement of the long central nervous system tracts, such as the corticospinal tracts, as well as the imaging finding of a thinning of the corpus callosum. *TECPR2*-associated disease, however, stands out for its prominent involvement of brain stem function, autonomic dysregulation, and peripheral neuropathy.

In summary, our cross-sectional analysis provides a depiction of clinical and molecular features across the age spectrum. Functional analyses of the variant mechanisms are of great importance to confirm the intended effect by our in silico modeling approach. Future prospective longitudinal studies are required to better define the natural history and patterns of disease progression. Our present study provides a framework for assessing disease manifestations. Close follow-up and surveillance for neurological and nonneurological manifestations are recommended.

ACKNOWLEDGMENTS

We thank all involved families for participating in this study. We also thank the Exome Aggregation Consortium and therein involved groups for providing exome and genome variant data for comparison. Barbara Brechmann is supported by scholarships from the German National Academic Foundation and the Carl Duisberg Program by the Bayer Foundation. Bernt Popp is supported by the Deutsche Forschungsgemeinschaft (DFG) through grant PO2366/2-1. Darius Ebrahimi-Fakhari is supported by grants from the CureAP4 Foundation, CureSPG50 Foundation, Spastic Paraplegia Foundation, the Thrasher Research Fund, and a joint research agreement with Astellas Pharmaceutical Inc. and MitoBridge Inc. Trio-ES of P17 was supported by the US NIH National Human Genome Research Institute (NHGRI)/National Heart Lung and Blood Institute (NHLBI) UM1 HG006542 (James R. Lupski). Jennifer E. Posey is supported by NHGRI K08 HG008986. This study was made possible through access to the data and findings generated by the 100,000 Genomes Project. The 100,000 Genomes Project is managed by Genomics England Limited (a wholly owned company of the Department of Health and Social Care). The 100,000 Genomes Project is funded by the National Institute for Health Research and NHS England. The Wellcome Trust, Cancer Research UK and the Medical Research Council have also funded research infrastructure. The 100.000 Genomes Project uses data provided by patients and collected by the National Health Service as part of their care and support. Funding for this study was provided by the Baylor-Hopkins Center for Mendelian Genomics through National Human Genome Research Institute grant 5U54HG006542. Open access funding enabled and organized by Projekt DEAL.

CONFLICT OF INTERESTS

The authors declare that there are no conflict of interests.

AUTHOR CONTRIBUTIONS

Bernt Popp, Rami Abou Jamra, and Sonja Neuser conceived the initial study concept and coordinated collection of clinical and genetic data through matchmaking and personal communications. Sonja Neuser, Barbara Brechmann, and Gali Heimer reviewed literature data and standardized the clinical HPO terms. Angelika Seitz, Ines Brösse, Lauren O'Grady, David A. Sweetser, Barry Byrne, Wen-Hann Tan, Basil T. Darras, Siddharth Srivastava, Parul Jayakar, Robert J. Graham, Michael Zech, Yasemin Dincer, Volker Mall, Juliane Winkelmann, Gehad ElGhazali, Amal Al Tenaiji, Omar Ismayl, Asma E. Al Nuaimi, Karima Maher, Fowzan S. Alkuraya, Hesham M. Aldhalaan, Mais Hashem, Bat El Bar-Aluma, Yael Haberman, Amir Szeinberg, Shahnaz Ibrahim, Fatima Khan, Stephanie Efthymiou, Reza WILEY-Human Mutatior

Maroofian, Henry Houlden, Vijayalakshmi S. Ramakumaran, Usha Kini, Alistair T. Pagnamenta, Isabella Herman, Jennifer E. Posey, James R. Lupski, Maria Cecilia Poli, Tatiana Muñoz, Gabriela M. Repetto, Bruria Ben-Zeev, Gali Heimer, and Darius Ebrahimi-Fakhari provided clinical and genetic data and performed clinical assessments. Sonja Neuser, Ines Brösse, Lauren O'Grady, Siddharth Srivastava, Wen-Hann Tan, Gehad ElGhazali, Fowzan S. Alkuraya, Asma E. Al Nuaimi, Hesham M. Aldhalaan, Michael Zech, Bruria Ben-Zeev, Gali Heimer, Stephanie Efthymiou, and Vijayalakshmi S. Ramakumaran wrote the case reports. Ines Brösse provided RNA samples of P1 and his parents. Susanna Schubert conducted RT-PCR and qPCR analyses. Christian Behrends supported protein analysis. Jens Meiler supported computational protein modeling. Mandy Krumbiegel processed samples of P1 and his parents for establishing a cell line. Sonja Neuser and Bernt Popp created Figures 1-3 and the Supporting Information. Sonja Neuser, Barbara Brechmann, Gali Heimer, Bruria Ben-Zeev, Darius Ebrahimi-Fakhari, and Bernt Popp wrote and edited the manuscript. All authors reviewed and commented on the final draft of the manuscript.

DATA AVAILABILITY STATEMENT

All data generated or analyzed during this study can be found in the online version of this article at the publisher's website. The genomes from P16 and her parents are available within the Genomics England research environment, access to which can be obtained by applying to join a Genomics England Clinical Interpretation Partnership domain (www.genomicsengland.co.uk/join-a-gecip-domain).

WEB RESOURCES

gnomAD browser: http://gnomad.broadinstitute.org/ BRAVO/TOPmed: https://bravo.sph.umich.edu Mutalyzer: https://mutalyzer.nl ClinVar: https://www.ncbi.nlm.nih.gov/clinvar/ UCSC BLAT: https://genome.ucsc.edu/cgi-bin/hgBlat UCSC Browser: https://genome.ucsc.edu/cgi-bin/hgGateway SPIDEX/SPANR: http://tools.genes.toronto.edu dbscSNV: http://www.liulab.science/dbscsnv.html

ORCID

Sonja Neuser b http://orcid.org/0000-0002-2875-8791 Barbara Brechmann b https://orcid.org/0000-0003-1228-4349 Gali Heimer b http://orcid.org/0000-0001-6218-3170 David A. Sweetser b http://orcid.org/0000-0002-1621-3284 Christian Behrends b http://orcid.org/0000-0002-9184-7607 Alistair T. Pagnamenta b http://orcid.org/0000-0001-7334-0602 Jennifer E. Posey b http://orcid.org/0000-0003-4814-6765 James R. Lupski b http://orcid.org/0000-0001-9907-9246 Wen-Hann Tan b http://orcid.org/0000-0002-1593-6149 Isabella Herman b http://orcid.org/0000-0002-7359-6832 Gabriela M. Repetto b http://orcid.org/0000-0003-0120-5684 Maria Cecilia Poli b http://orcid.org/0000-0001-6817-0573 Usha Kini b http://orcid.org/0000-0001-9111-5433 Stephanie Efthymiou b http://orcid.org/0000-0003-4900-9877 Jens Meiler b http://orcid.org/0000-0001-8945-193X Reza Maroofian b https://orcid.org/0000-0001-6763-1542 Fowzan S. Alkuraya b https://orcid.org/0000-0003-4158-341X Rami Abou Jamra b http://orcid.org/0000-0002-1542-1399 Bernt Popp b http://orcid.org/0000-0002-3679-1081 Darius Ebrahimi-Fakhari b http://orcid.org/0000-0002-0026-4714

REFERENCES

- Anazi, S., Maddirevula, S., Salpietro, V., Asi, Y. T., Alsahli, S., Alhashem, A., Shamseldin, AlZahrani, F., Patel, N., Ibrahim, N., Abdulwahab, F. M., Hashem, M., Alhashmi, N., Al Murshedi, F., Al Kindy, A., Alshaer, A., Rumayyan, A., Al Tala, S., Kurdi, W., ... Alkuraya, F. S., & H. E. (2017). Expanding the genetic heterogeneity of intellectual disability. *Human Genetics*, 136(11), 1419–1429. https://doi.org/10.1007/ s00439-017-1843-2
- Behrends, C., Sowa, M. E., Gygi, S. P., & Harper, J. W. (2010). Network organization of the human autophagy system. *Nature*, 466(7302), 68–76. https://doi.org/10.1038/nature09204
- Cingolani, P., Patel, V. M., Coon, M., Nguyen, T., Land, S. J., Ruden, D. M., & Lu, X. (2012). Using *Drosophila melanogaster* as a model for genotoxic chemical mutational studies with a new program, SnpSift. *Frontiers in Genetics*, 3, 35. https://doi.org/10.3389/fgene.2012.00035
- Cingolani, P., Platts, A., Wang, L. L., Coon, M., Nguyen, T., Wang, L., Land, S. J., Lu, X., & Ruden, D. M. (2012). A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff: Snps in the genome of *Drosophila melanogaster* strain w1118; iso-2; iso-3. *Fly*, 6(2), 80–92. https://doi.org/10.4161/fly.19695
- Covone, A. E., Fiorillo, C., Acquaviva, M., Trucco, F., Morana, G., Ravazzolo, R., & Minetti, C. (2016). Wes in a family trio suggests involvement of TECPR2 in a complex form of progressive motor neuron disease: Wes in a family trio suggests involvement of TECPR2. *Clinical Genetics*, 90(2), 182–185. https://doi.org/10.1111/cge.12730
- Ebrahimi-Fakhari, D., Saffari, A., Wahlster, L., Lu, J., Byrne, S., Hoffmann, G. F., Jungbluth, H., & Sahin, M. (2016). Congenital disorders of autophagy: An emerging novel class of inborn errors of neuro-metabolism. *Brain*, 139(2), 317–337. https://doi.org/10.1093/ brain/awv371
- Freeman, P. J., Hart, R. K., Gretton, L. J., Brookes, A. J., & Dalgleish, R. (2018). Variantvalidator: Accurate validation, mapping, and formatting of sequence variation descriptions. *Human Mutation*, 39(1), 61–68. https://doi.org/10.1002/humu.23348
- Hahn, K., Rohdin, C., Jagannathan, V., Wohlsein, P., Baumgärtner, W., Seehusen, F., Spitzbarth, I., Grandon, R., Drögemüller, C., & Jäderlund, K. H. (2015). TECPR2 associated neuroaxonal dystrophy in Spanish water dogs. *PLOS One*, 10(11), e0141824. https://doi.org/ 10.1371/journal.pone.0141824
- Hansen, A. W., Murugan, M., Li, H., Khayat, M. M., Wang, L., Rosenfeld, J., Andrews, B. K., Jhangiani, S. N., Coban Akdemir, Z. H., Sedlazeck, F. J., Ashley-Koch, A. E., Liu, P., Muzny, D. M., Davis, E. E., Katsanis, N., Sabo, A., Posey, J. E., Yang, Y., Wangler, M. F., ... Wiener, J. (2019). A genocentric approach to discovery of mendelian disorders. *American Journal of Human Genetics*, 105(5), 974–986. https://doi.org/10.1016/j.ajhg.2019.09.027
- Hebebrand, M., Hüffmeier, U., Trollmann, R., Hehr, U., Uebe, S., Ekici, A. B., Kraus, C., Krumbiegel, M., Reis, A., Thiel, C. T., & Popp, B. (2019). The mutational and phenotypic spectrum of TUBA1A-associated tubulinopathy. Orphanet Journal of Rare Diseases, 14(1), 38. https:// doi.org/10.1186/s13023-019-1020-x

- Heimer, G., Oz-Levi, D., Eyal, E., Edvardson, S., Nissenkorn, A., Ruzzo, E. K., Szeinberg, A., Maayan, C., Mai-Zahav, M., Efrati, O., Pras, E., Reznik-Wolf, H., Lancet, D., Goldstein, D. B., Anikster, Y., Shalev, S. A., Elpeleg, O., & Ben Zeev, B. (2016). TECPR2 mutations cause a new subtype of familial dysautonomia like hereditary sensory autonomic neuropathy with intellectual disability. *European Journal of Paediatric Neurology*, 20(1), 69–79. https://doi.org/10.1016/j.ejpn.2015.10.003
- Heo, L., Park, H., & Seok, C. (2013). Galaxyrefine: Protein structure refinement driven by side-chain repacking. Nucleic Acids Research, 41(W1), W384–W388. https://doi.org/10.1093/nar/gkt458
- Jian, X., Boerwinkle, E., & Liu, X. (2014). In silico prediction of splicealtering single nucleotide variants in the human genome. Nucleic Acids Research, 42(22), 13534–13544. https://doi.org/10.1093/nar/ gku1206
- Karczewski, K. J., Francioli, L. C., Tiao, G., Cummings, B. B., Alföldi, J., Wang, Q., Collins, R. L., Laricchia, K. M., Ganna, A., Birnbaum, D. P., Gauthier, L. D., Brand, H., Solomonson, M., Watts, N. A., Rhodes, D., Singer-Berk, M., England, E. M., Seaby, E. G., Kosmicki, J. A., ... MacArthur, D. G. (2020). The mutational constraint spectrum quantified from variation in 141,456 humans. *Nature*, *581*(7809), 434–443. https://doi.org/10.1038/s41586-020-2308-7
- Ko, J., Park, H., Heo, L., & Seok, C. (2012). Galaxyweb server for protein structure prediction and refinement. *Nucleic Acids Research*, 40(W1), W294–W297. https://doi.org/10.1093/nar/gks493
- Ko, J., Park, H., & Seok, C. (2012). Galaxytbm: Template-based modeling by building a reliable core and refining unreliable local regions. BMC Bioinformatics, 13, 198. https://doi.org/10.1186/1471-2105-13-198
- Köhler, S., Carmody, L., Vasilevsky, N., Jacobsen, J. O. B., Danis, D., Gourdine, J. P., Gargano, M., Matentzoglu, N., McMurry, J. A., Osumi-Sutherland, D., Cipriani, V., Balhoff, J. P., Conlin, T., Blau, H., Baynam, G., Palmer, R., Gratian, D., Dawkins, H., Segal, M., ... Robinson, P. N. (2019). Expansion of the Human Phenotype Ontology (HPO) knowledge base and resources. *Nucleic Acids Research*, 47(D1), D1018–D1027. https:// doi.org/10.1093/nar/gky1105
- Liu, X., Jian, X., & Boerwinkle, E. (2013). Dbnsfp v2.0: A database of human non-synonymous SNVs and their functional predictions and annotations. *Human Mutation*, 34(9), E2393–E2402. https://doi.org/ 10.1002/humu.22376
- Lupski, J. R., Belmont, J. W., Boerwinkle, E., & Gibbs, R. A. (2011). Clan genomics and the complex architecture of human disease. *Cell*, 147(1), 32–43. https://doi.org/10.1016/j.cell.2011.09.008
- Menzies, F. M., Fleming, A., Caricasole, A., Bento, C. F., Andrews, S. P., Ashkenazi, A., Füllgrabe, J., Jackson, A., Jimenez Sanchez, M., Karabiyik, C., Licitra, F., Lopez Ramirez, A., Pavel, M., Puri, C., Renna, M., Ricketts, T., Schlotawa, L., Vicinanza, M., Won, H., ... Rubinsztein, D. C. (2017). Autophagy and neurodegeneration: Pathogenic mechanisms and therapeutic opportunities. *Neuron*, *93*(5), 1015–1034. https://doi.org/10.1016/j.neuron.2017.01.022
- Meyer, M. J., Lapcevic, R., Romero, A. E., Yoon, M., Das, J., Beltrán, J. F., Mort, M., Stenson, P. D., Cooper, D. N., Paccanaro, A., & Yu, H. (2016). Mutation3d: Cancer gene prediction through atomic clustering of coding variants in the structural proteome. *Human Mutation*, 37(5), 447–456. https://doi.org/10.1002/humu.22963
- Oz-Levi, D., Ben-Zeev, B., Ruzzo, E. K., Hitomi, Y., Gelman, A., Pelak, K., Anikster, Y., Reznik-Wolf, H., Bar-Joseph, I., Olender, T., Alkelai, A., Weiss, M., Ben-Asher, E., Ge, D., Shianna, K. V., Elazar, Z., Goldstein, D. B., Pras, E., & Lancet, D. (2012). Mutation in TECPR2 reveals a role for autophagy in hereditary spastic paraparesis. *American Journal of Human Genetics*, 91(6), 1065–1072. https://doi. org/10.1016/j.ajhg.2012.09.015
- Oz-Levi, D., Gelman, A., Elazar, Z., & Lancet, D. (2013). TECPR2: A new autophagy link for neurodegeneration. *Autophagy*, *9*(5), 801–802. https://doi.org/10.4161/auto.23961

- Patwari, P. P., Wolfe, L. F., Sharma, G. D., & Berry-Kravis, E. (2020). TECPR2 mutation associated respiratory dysregulation: More than central apnea. *Journal of Clinical Sleep Medicine*, 16, 977–982. https:// doi.org/10.5664/jcsm.8434
- Popp, B., Ekici, A. B., Thiel, C. T., Hoyer, J., Wiesener, A., Kraus, C., Reis, A., & Zweier, C. (2017). Exome Pool-Seq in neurodevelopmental disorders. European Journal of Human Genetics: EJHG, 25(12), 1364–1376. https://doi.org/10.1038/ s41431-017-0022-1
- Popp, B., & Neuser, S. (2020). Tecpr2 protein models predicted by three algorithms. https://doi.org/10.5281/zenodo.4050472
- Rentzsch, P., Witten, D., Cooper, G. M., Shendure, J., & Kircher, M. (2019). Cadd: Predicting the deleteriousness of variants throughout the human genome. *Nucleic Acids Research*, 47(D1), D886–D894. https:// doi.org/10.1093/nar/gky1016
- Richards, S., Aziz, N., Bale, S., Bick, D., Das, S., Gastier-Foster, J., Grody, W. W., Hegde, M., Lyon, E., Spector, E., Voelkerding, K., & Rehm, H. L. (2015). Standards and guidelines for the interpretation of sequence variants: A joint consensus recommendation of the American College of Medical Genetics and Genomics and the Association for Molecular Pathology. *Genetics in Medicine*, 17(5), 405–424. https://doi.org/10.1038/ gim.2015.30
- Sobreira, N., Schiettecatte, F., Valle, D., & Hamosh, A. (2015). Genematcher: A matching tool for connecting investigators with an interest in the same gene. *Human Mutation*, 36(10), 928–930. https://doi.org/10.1002/humu.22844
- Stadel, D., Millarte, V., Tillmann, K. D., Huber, J., Tamin-Yecheskel, B. C., Akutsu, M., Demishtein, A., Anikster, Y., Perez, F., Dötsch, V., Elazar, Z., Rogov, V., Farhan, H., & Behrends, C. (2015). Tecpr2 cooperates with LC3C to regulate COPII-dependent ER export. *Molecular Cell*, 60(1), 89–104. https://doi.org/10.1016/j.molcel.2015. 09.010
- Teinert, J., Behne, R., Wimmer, M., & Ebrahimi-Fakhari, D. (2019). Novel insights into the clinical and molecular spectrum of congenital disorders of autophagy. *Journal of Inherited Metabolic Disease*, 43(1), 1–12. https://doi.org/10.1002/jimd.12084
- Wildeman, M., van Ophuizen, E., den Dunnen, J. T., & Taschner, P. E. M. (2008). Improving sequence variant descriptions in mutation databases and literature using the Mutalyzer sequence variation nomenclature checker. *Human Mutation*, 29(1), 6–13. https://doi. org/10.1002/humu.20654
- Xiong, H. Y., Alipanahi, B., Lee, L. J., Bretschneider, H., Merico, D., Yuen, R. K. C., Hua, Y., Gueroussov, S., Najafabadi, H. S., Hughes, T. R., Morris, Q., Barash, Y., Krainer, A. R., Jojic, N., Scherer, S. W., Blencowe, B. J., & Frey, B. J. (2015). Rna splicing. The human splicing code reveals new insights into the genetic determinants of disease. *Science*, 347(6218), 1254806. https://doi. org/10.1126/science.1254806
- Yang, J., Anishchenko, I., Park, H., Peng, Z., Ovchinnikov, S., & Baker, D. (2020). Improved protein structure prediction using predicted interresidue orientations. Proceedings of the National Academy of Sciences of the United States of America, 117(3), 1496–1503. https:// doi.org/10.1073/pnas.1914677117
- Zech, M., Jech, R., Boesch, S., Škorvánek, M., Weber, S., Wagner, M., Zhao, C., Jochim, A., Necpál, J., Dincer, Y., Vill, K., Distelmaier, F., Stoklosa, M., Krenn, M., Grunwald, S., Bock-Bierbaum, T., Fečíková, A., Havránková, P., Roth, J., ... Winkelmann, J. (2020). Monogenic variants in dystonia: An exome-wide sequencing study. *The Lancet, Neurology*, *19*(11), 908–918. https://doi.org/10.1016/ S1474-4422(20)30312-4
- Zhu, X., Petrovski, S., Xie, P., Ruzzo, E. K., Lu, Y. F., McSweeney, K. M., Ben-Zeev, B., Anikster, Y., Oz-Levi, D., Dhindsa, R. S., Hitomi, Y.,

WILEY-Human Mutation

Schoch, K., Spillmann, R. C., Heimer, G., Marek-Yagel, D., Tzadok, M., Han, Y., Worley, G., Goldstein, J., ... Oldstein, & D. B. (2015). Whole-exome sequencing in undiagnosed genetic diseases: Interpreting 119 trios. *Genetics in Medicine*, 17(10), 774-781. https://doi.org/10.1038/gim.2014.191

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

How to cite this article: Neuser, S., Brechmann, B., Heimer, G., Brösse, I., Schubert, S., O'Grady, L., Zech, M., Srivastava, S., Sweetser, D. A., Dincer, Y., Mall, V., Winkelmann, J., Behrends, C., Darras, B. T., Graham, R. J., Jayakar, P., Byrne, B., Bar-Aluma, B. E., Haberman, Y., ... Ebrahimi-Fakhari, D. (2021). Clinical, neuroimaging, and molecular spectrum of *TECPR2*associated hereditary sensory and autonomic neuropathy with intellectual disability. *Human Mutation*, 42, 762–776. https://doi.org/10.1002/humu.24206