

De Novo and Bi-allelic Pathogenic Variants in *NARS1* Cause Neurodevelopmental Delay Due to Toxic Gain-of-Function and Partial Loss-of-Function Effects

Andrea Manole,^{1,50} Stephanie Efthymiou,^{1,50} Emer O'Connor,^{1,50} Marisa I. Mendes,^{2,50} Matthew Jennings,^{3,50} Reza Maroofian,¹ Indran Davagnanam,⁴⁷ Kshitij Mankad,⁴ Maria Rodriguez Lopez,⁵ Vincenzo Salpietro,¹ Ricardo Harripaul,^{6,7} Lauren Badalato,⁸ Jagdeep Walia,⁸ Christopher S. Francklyn,⁹ Alkyoni Athanasiou-Fragkouli,¹ Roisin Sullivan,¹ Sonal Desai,¹⁰ Kristin Baranano,¹⁰ Faisal Zafar,¹¹ Nuzhat Rana,¹¹ Muhammed Ilyas,¹² Alejandro Horga,¹ Majdi Kara,¹³ Francesca Mattioli,¹⁶ Alice Goldenberg,¹⁵ Helen Griffin,³ Amelie Piton,¹⁶ Lindsay B. Henderson,¹⁷ Benyekhlef Kara,¹⁸ Ayca Dilruba Aslanger,¹⁸ Joost Raaphorst,^{19,20} Rolph Pfundt,¹⁹ Ruben Portier,²¹ Marwan Shinawi,²² Amelia Kirby,²³ Katherine M. Christensen,²³ Lu Wang,²⁴ Rasim O. Rosti,²⁴ Sohail A. Paracha,²⁵ Muhammad T. Sarwar,²⁵ Dagan Jenkins,⁴⁹ SYNAPS Study Group,²⁶ Jawad Ahmed,²⁵ Federico A. Santoni,^{27,28} Emmanuelle Ranza,^{27,29,30} Justyna Iwaszkiewicz,³¹ Cheryl Cytrynbaum,³² Rosanna Weksberg,³² Ingrid M. Wentzensen,¹⁷ Maria J. Guillen Sacoto,¹⁷ Yue Si,¹⁷ Aida Telegrafi,¹⁷

(Author list continued on next page)

Aminoacyl-tRNA synthetases (ARSs) are ubiquitous, ancient enzymes that charge amino acids to cognate tRNA molecules, the essential first step of protein translation. Here, we describe 32 individuals from 21 families, presenting with microcephaly, neurodevelopmental delay, seizures, peripheral neuropathy, and ataxia, with *de novo* heterozygous and bi-allelic mutations in asparaginyl-tRNA synthetase (*NARS1*). We demonstrate a reduction in *NARS1* mRNA expression as well as in *NARS1* enzyme levels and activity in both individual fibroblasts and induced neural progenitor cells (iNPCs). Molecular modeling of the recessive c.1633C>T (p.Arg545Cys) variant shows weaker spatial positioning and tRNA selectivity. We conclude that *de novo* and bi-allelic mutations in *NARS1* are a significant cause of neurodevelopmental disease, where the mechanism for *de novo* variants could be toxic gain-of-function and for recessive variants, partial loss-of-function.

Introduction

The attachment of tRNA to cognate amino acids is essential for protein translation. Aminoacyl-tRNA synthetases (ARSs) are a group of enzymes encoded by ancient

genes which are ubiquitously expressed and highly conserved.^{1–3} These enzymes play a fundamental role in the esterification of proteinogenic amino acids to cognate tRNA. In total, 37 genes encoding ARS enzymes have been described. Of these, 20 encode enzymes that function in

¹Department of Neuromuscular Disorders, University College London (UCL) Institute of Neurology, Queen Square, London, WC1N 3BG, UK; ²Metabolic Unit, Department of Clinical Chemistry, Amsterdam University Medical Centers, Vrije Universiteit Amsterdam, Amsterdam Neuroscience, Amsterdam Gastroenterology and Metabolism, Amsterdam, 1081 the Netherlands; ³Department of Clinical Neurosciences, University of Cambridge, Cambridge, CB2 0QQ UK; ⁴Department of Neuroradiology, Great Ormond Street Hospital for Children, London, WC1N 3JH, UK; ⁵Institute of Healthy Ageing, Department of Genetics, Evolution and Environment, University College London (UCL), London, WC1E 6BT, UK; ⁶Campbell Family Mental Health Research Institute, Centre for Addiction and Mental Health, ON, M5T 1R8, Canada; ⁷Institute of Medical Science and Department of Psychiatry, University of Toronto, Toronto, ON, M5T 1R8, Canada; ⁸Department of Pediatrics, Queen's University, Kingston, ON, K7L 2V7, Canada; ⁹Department of Biochemistry, University of Vermont College of Medicine, Burlington, VT 05405, USA; ¹⁰Department of Neurology and Pediatrics, Johns Hopkins School of Medicine, Baltimore, MD 21205, USA; ¹¹Department of Pediatrics, Multan Hospital, Multan, 60000, Pakistan; ¹²University of Islamabad, Islamabad, 45320, Pakistan; ¹³Department of Pediatrics, Tripoli Children's Hospital, Tripoli, Libya; ¹⁴University of Strasbourg, CNRS, GMGM UMR 7156, Strasbourg, 67083, France; ¹⁵Département de Génétique, centre de référence anomalies du développement et syndromes malformatifs, CHU de Rouen, Inserm U1245, UNIROUEN, Normandie Université, Centre Normand de Génomique et de Médecine Personnalisée, Rouen, 76031, France; ¹⁶Institute for Genetics and Molecular and Cellular Biology (IGBMC), University of Strasbourg, CNRS UMR7104, INSERM U1258, Illkirch, 67404, France; ¹⁷GeneDx, 207 Perry Parkway Gaithersburg, MD 20877, USA; ¹⁸Bezmialem Vakif Üniversitesi, Istanbul, 34093, Turkey; ¹⁹Department of Human Genetics, Donders Institute for Brain, Cognition and Behaviour, Radboud University Medical Center, 6500HB Nijmegen, the Netherlands; ²⁰Department of Neurology, Amsterdam Neuroscience Institute, Amsterdam University Medical Center, 1105AZ Amsterdam, the Netherlands; ²¹Department of Neurology, Medisch Spectrum Twente, 7512KZ Enschede, the Netherlands; ²²Department of Pediatrics, Divisions of Genetics and Genomic Medicine, Washington University School of Medicine, St. Louis, MO, 63110, USA; ²³Division of Medical Genetics, SSM Health Cardinal Glennon Children's Hospital, Saint Louis University School of Medicine, St. Louis, MO 63104, USA; ²⁴Howard Hughes Medical Institute, University of California San Diego and Rady Children's Hospital, La Jolla, CA 92130, USA; ²⁵Institute of Basic Medical Sciences, Khyber Medical University, 25100 Peshawar, Pakistan; ²⁶SYNAPS Study Group, see Supplemental Information for the study group members who contributed clinical cases and data; ²⁷Department of Genetic Medicine and Development, University of Geneva, 1206 Geneva, Switzerland; ²⁸Department of Endocrinology, Diabetes, and Metabolism, University Hospital of Lausanne, 1011 Lausanne, Switzerland; ²⁹Service of Genetic Medicine, University Hospitals of Geneva, 1205 Geneva, Switzerland; ³⁰Medigenome, The Swiss Institute of Genomic Medicine, Geneva, CH-1207, Switzerland; ³¹Swiss Institute of Bioinformatics, Molecular Modeling Group, Batiment Genopode, Unil Sorge, Lausanne, CH-1015, Switzerland; ³²Hospital for Sick

(Affiliations continued on next page)



Marisa V. Andrews,²² Dustin Baldrige,²² Heinz Gabriel,³³ Julia Mohr,³³ Barbara Oehl-Jaschkowitz,³⁴ Sylvain Debard,¹⁴ Bruno Senger,¹⁴ Frédéric Fischer,¹⁴ Conny van Ravenwaaij,³⁵ Annemarie J.M. Fock,³⁵ Servi J.C. Stevens,³⁶ Jürg Bähler,⁵ Amina Nasar,⁸ John F. Mantovani,⁴⁵ Adnan Manzur,⁴⁹ Anna Sarkozy,⁴⁹ Desirée E.C. Smith,² Gajja S. Salomons,² Zubair M. Ahmed,⁴⁶ Shaikh Riazuddin,³⁷ Saima Riazuddin,⁴⁶ Muhammad A. Usmani,⁴⁶ Annette Seibt,³⁸ Muhammad Ansar,^{27,48} Stylianos E. Antonarakis,^{27,29,39} John B. Vincent,^{6,7} Muhammad Ayub,⁸ Mona Grimmel,⁴⁰ Anne Marie Jelsing,⁴¹ Tina Duelund Hjortshøj,⁴¹ Helena Gásdal Karstensen,⁴¹ Marybeth Hummel,⁴² Tobias B. Haack,^{40,43} Yalda Jamshidi,⁴⁴ Felix Distelmaier,³⁸ Rita Horvath,³ Joseph G. Gleeson,²⁴ Hubert Becker,^{14,50} Jean-Louis Mandel,^{16,50} David A. Koolen,^{19,50} and Henry Houlden^{1,50,*}

the cytoplasm, and the remainder relate exclusively to mitochondrial enzymes. Despite the essential canonical function and ubiquitous expression of ARS enzymes, mutations in these genes have been implicated in a variety of human diseases with both recessive and dominant inheritance patterns.^{2,4-6} These mutations result in neurological disorders, ranging from mild late-onset peripheral neuropathy to severe multi-systemic neurodevelopmental disorders^{4,5,7-9} (Table S1).

Mutations in cytoplasmic ARS-encoding genes cause peripheral nervous system degeneration resulting in Charcot-Marie-Tooth neuropathies (*GARS1* and *AARS1* [MIM: 601065]) and brain stem and spinal cord hypomyelination (*DARS1* [MIM: 603084]). ARSs, and ARSs interacting genes, including *DARS1*, *RARS1* (MIM: 107820), *AIMP1* (MIM: 603605), and *AARS1*, have been implicated in neurodevelopmental disorders and epilepsies. Furthermore, mitochondrial *ARS2* mutations are often associated with leukoencephalopathy (*AARS2* [MIM: 615889] and *DARS2* [MIM: 611105]) or pontocerebellar hypoplasia (*RARS2* [MIM: 611524]). More recently, recessive mutations in *FARSA* (MIM: 602918), *VARSI* (MIM: 192150), *CARS1* (MIM: 123859), and *TARSI* (MIM: 187790), with subsequent partial loss of the ARS protein, have been linked to neurodevelopmental phenotypes.¹⁰⁻¹⁴ Modes of inheritance can be dominant or recessive; in cases such as *AARS1*, *YARSI* (MIM: 603623), *MARSI* (MIM: 156560), *HARSI* (MIM: 142810), and *GARS1*, both patterns can occur.⁶

The loss of function associated with mutations in ARSs is attributed to decreased aminoacylation efficiency or misfolding, causing protein instability with lower steady-state levels.¹³ However, in some cases (*GARS1*, *YARSI*, and

AARS1), it has not been possible to ascribe the phenotype to a loss of primary aminoacylation.¹⁵⁻¹⁷ Overall, the physiological functions of ARS genes and previously identified disease associations indicate an essential biological role for these proteins, implying that defects in all ARSs incur disease.⁶

Asparaginyl-tRNA^{Asn} is generated by asparaginyl-tRNA synthetase (*NARS1* [MIM: 108410; RefSeq accession number NM_004539.4]) in a reaction involving two steps. *NARS1* first catalyzes the ATP-dependent activation of asparagine (Asn) into Asn~AMP with the release of pyrophosphate, and then transfers the activated Asn onto tRNA^{Asn} with the release of AMP (Figure 1A). Here, we report the clinical phenotypes associated with *de novo* dominant and bi-allelic, autosomal recessive mutations in *NARS1* in 32 affected individuals from 21 families. We provide genetic proof for these mutations and analyze their impact through the use of individual cell lines, neural progenitor cells, and molecular modeling.

Subjects and Methods

Study Participants

Individuals were recruited via an international collaborative network of research and diagnostic sequencing laboratories. Samples and clinical information were obtained, with informed consent, from each institution using local institutional review board (IRB) ethics for functional analysis of human DNA and biomaterial. Clinical data collection involved a detailed review of medical records, photographs, videos, and phone interviews, as well as a clinical re-evaluation by a neurologist. Tables S2-S4 summarize the clinical and demographic details of the included cases.

Children, Division of Clinical and Metabolic Genetics, 555 University Ave., Toronto, M5G 1X8, Canada; ³³CeGaT GmbH and Praxis für Humangenetik Tuebingen, Tuebingen, 72076, Germany; ³⁴Biomedical Centre Cardinal-Wendel-Straße 14, 66424 Hamburg, Germany; ³⁵University of Groningen, University Medical Center Groningen, Department of Neurology, Groningen, 9713, the Netherlands; ³⁶Department of Clinical Genetics, Maastricht University Medical Centre, Maastricht, 6211, the Netherlands; ³⁷Jinnah Burn and Reconstructive Surgery Center, Allama Iqbal Medical College, University of Health Sciences, Lahore 54550, Pakistan; ³⁸Department of General Pediatrics, Heinrich-Heine-University, Moorenstr. 5, 40225 Düsseldorf, Germany; ³⁹iGE3 Institute of Genetics and Genomics of Geneva, 1211 Geneva, Switzerland; ⁴⁰Institute of Medical Genetics and Applied Genomics, University of Tuebingen, 72076 Tuebingen, Germany; ⁴¹Department of Clinical Genetics, University Hospital of Copenhagen, Rigshospitalet, 2100, Denmark; ⁴²Department of Pediatrics, Section of Medical Genetics, West Virginia University, Morgantown, WV 26506-9600, USA; ⁴³Centre for Rare Diseases, University of Tuebingen, 72076 Tuebingen, Germany; ⁴⁴Genetics Centre, Molecular and Clinical Sciences Institute, St George's University of London, London, SW17 0RE, UK; ⁴⁵Division of Child Neurology, Washington University School of Medicine, St. Louis, MO, 63110, USA; ⁴⁶Department of Biochemistry and Molecular Biology, Johns Hopkins School of Medicine, Baltimore, MD 21205, USA; ⁴⁷Department of Brain Repair and Rehabilitation, UCL Institute of Neurology, Queen Square, London, WC1N 3BG, UK; ⁴⁸Institute of Molecular and Clinical Ophthalmology Basel, Basel Switzerland; ⁴⁹Institute of Child Health, Guilford Street and Dubowitz Neuromuscular Centre, Great Ormond Street Hospital for Children, London, WC1N 3JH, UK

⁵⁰These authors contributed equally to this work

*Correspondence: h.houlden@ucl.ac.uk

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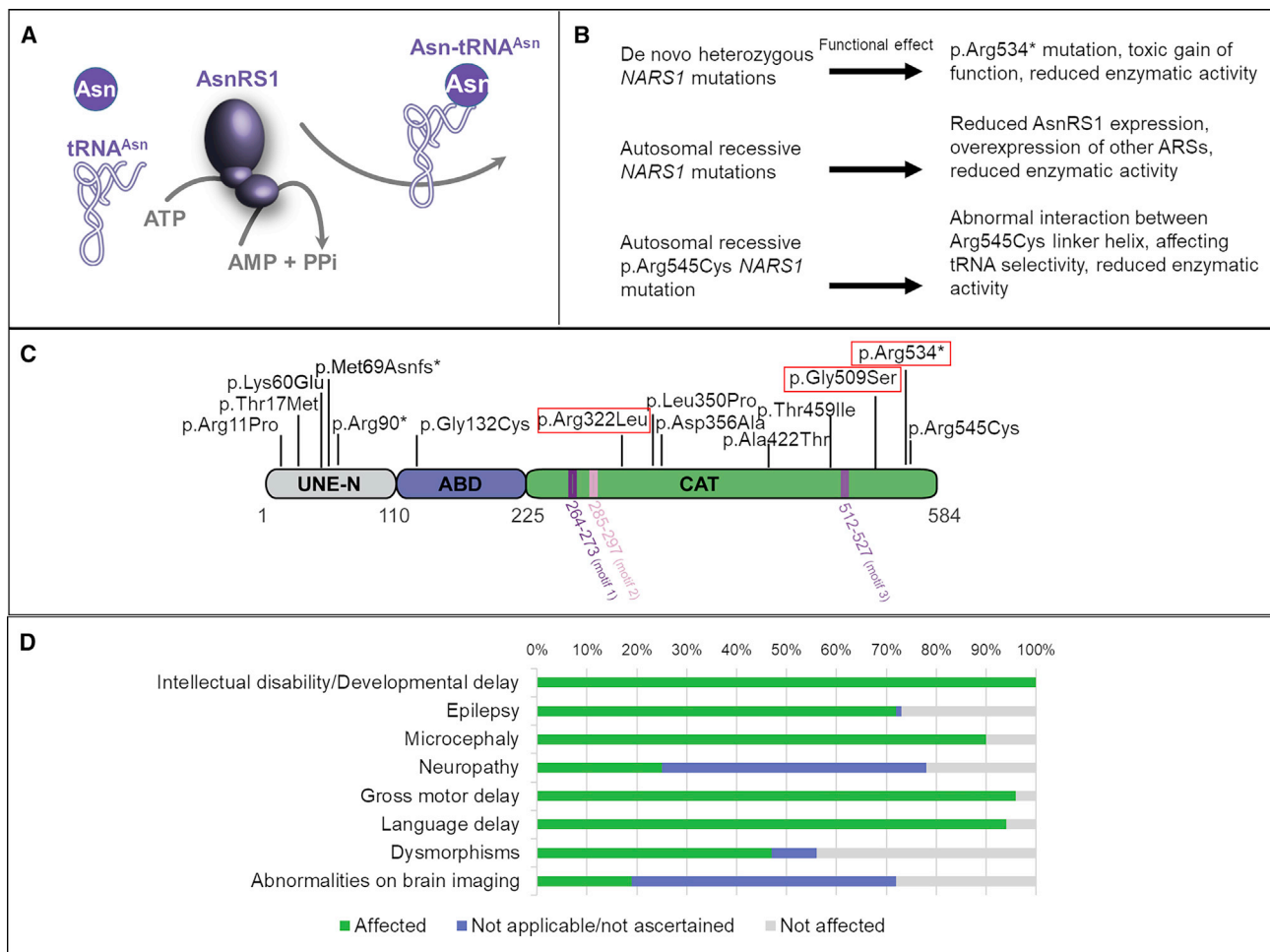


Figure 1. AsnRS1 Protein Structure and Function

(A) AA asparagine (Asn) is ligated to tRNA^{Asn} and catalyzed by AsnRS1 and ATP to produce Asn-tRNA (Asn), AMP, and pyrophosphate.

(B) NARS1 mutations and their predicted functional effect.

(C) Schematic representation of human ARS1 primary structure. Three main domains are depicted: the unique domain (UNE-N), the anticodon binding domain (ABD), and the catalytic domain (CAT). The nature and position of the mutants are shown above the primary structure, *de novo* boxed in red, and the positions of the domains are indicated below, including motif 1 (involved in AsnRS1 dimerization) and motifs 2 and 3 (which form the active site).

(D) Bar graph summarizing proportions of various clinical findings affecting individuals with NARS1 mutations.

Sequencing

Exome sequencing was carried out using a number of methods in different centers with different analysis platforms and pipelines used (see [Supplemental Methods](#), Section 2).

Bioinformatic Analysis

cDNA and protein sequence variants are described in accordance with the recommendations of the Human Genome Variation Society using Ensembl ENSG00000134440 and ENST00000256854.10 as the reference sequences. Evolutionary conservation of nucleotides was assessed using PhyloP (46 vertebrate species) and genomic evolutionary rate profiling (GERP) scores.¹⁸ These were accessed through the University of California—San Francisco (UCSC) Genome Browser¹⁹ using genomic coordinates from GRCh37/hg19. Grantham scores were used to assess the physicochemical nature of the amino acid (AA) substitutions. *In silico* analyses of sequence variants were performed using the pathoge-

nicity prediction tools SIFT, PolyPhen-2, and Mutation Taster version 2.

Our bioinformatics filtering strategy screened for exonic and donor/acceptor splicing variants. In accordance with the pedigree and phenotype, priority was given to rare variants (<0.01% in public databases, including 1000 Genomes Project; National Heart, Lung, and Blood Institute [NHLBI] Exome Variant Server; Complete Genomics 69; and Exome Aggregation Consortium [ExAC v0.2]) fitting a recessive (homozygous or compound heterozygous) or a *de novo* model and/or variants in genes previously linked to epilepsy, developmental delay, intellectual disability, and other neurological disorders. Upon whole-exome sequencing (WES) analysis of the index family (F9), the NARS1 variant c.1633C>T (p.Arg545Cys) was picked up according to its frequency and prediction tool scores (SIFT—damaging [score = 1], PolyPhen—damaging [score = 1], GERP—5.5, Mutation Taster—0.999992). All the candidate variants were further verified through the use of Sanger sequencing.

Generation of the *nrs1* Vector

The pJR1-41XU-*nrs1* expression vector was used to express *Schizosaccharomyces pombe nrs1* by amplifying the coding sequence of *nrs1* from *S. pombe* DNA with the Nrs1-PJR-F and Nrs1-PJR-R primers (all primers are provided in Table S5) through the use of Phusion HF polymerase from New England Biolabs (NEB). The PCR product was cloned into XhoI digested pJR1-41XU²⁰ through the use of CloneEZ from Genscript. Plasmids were sequenced to confirm the correct insertion of the fragment.

Deletion of *nrs1* Gene in *S. pombe* Cells

JB775 (*h- ade6-M216 ura4-D18 leu1-32*) cells were synchronized, made competent, and transformed as previously described.²¹ Cells were transformed using the plasmid containing the *nrs1* gene, pJR1-41XU-*nrs1*, and transformants were selected according to growth in Edinburgh minimal medium (EMM) + *ade* + *leu*, generating the strain MR397. The *nrs1* gene was deleted in MR397 cells through the use of the standard method via homologous recombination with the NatMx6 cassette^{22,23} using the primers Nrs1DelFw and Nrs1DelRv (Table S5). Transformants were selected in EMM + Nat with no thiamine to promote the expression of the *nrs1* gene from the plasmid. Deletions were checked via PCR using primers Nrs1ck-L and kanR and Nrs1ck-R and kanF (Table S5). The strain generated was named MR409. MR409 cells were synchronized, made competent, and transformed as previously described.²¹ The plasmids of the pJR-41XL series contained either the empty vector, wild-type *NARS1*, or the *NARS1* variants described. Transformants were selected in EMM + *ade* strains.

Cell Culture

Fibroblasts of affected individuals carrying the homozygous c.50C>T (p.Thr17Met), c.32G>C (p.Arg11Pro), and c.1633C>T (p.Arg545Cys) and compound heterozygous c.1067A>C (p.Asp356Ala) and c.203dupA (p.Met69Aspfs*4), as well as of corresponding controls, were grown in high-glucose Dulbecco's modified Eagle's medium (Sigma) supplemented with 10% fetal bovine serum and 1% penicillin and streptomycin.

Semiquantitative RT-PCR for Individual Lymphoblasts

Using TRIzol (Zymo research), as per manufacturer's instructions, total RNA was extracted from immortalized lymphoblasts available from P2 and parents. The concentration and purity of RNA was determined spectrophotometrically. 1 µg of RNA was reverse transcribed to first strand cDNA through the use of random primers and Moloney murine leukemia virus reverse transcriptase (Promega). GoTaq® Green Master Mix (Promega) was used and PCR reactions were performed with the following protocol: 95°C—2 min (95°C—30 s, 60°C—30 s, 73°C—1 min) for 35 cycles, 73°C—5 min, and 4°C hold. Two exponential curves representing the product formation were determined for both primer pairs. Cycles 28 and 29 were chosen for *NARS1* and *GAPDH*, respectively, so that amplification rates were in the linear range for semiquantitative comparisons. Reactions were repeated in triplicate.

Western Blotting

For western blotting analysis, protein lysates were obtained from cultured fibroblasts and total protein concentration was measured by means of a Bradford assay. Aliquots of total protein (15 µg) were loaded on 4%–12% sodium dodecyl sulfate (SDS)-polyacrylamide gels (NuPAGE 4%–12% Bis-Tris Protein Gels, ThermoFisher Scientific), transferred to polyvinylidene fluoride membranes, and

blocked and incubated overnight with a polyclonal antibody recognizing AsnRS1 (anti-rabbit 1:1000; Proteintech). Secondary antibody was added for 1 h, and signal was detected using enhanced chemiluminescence (ECL) reagents (Amersham Biosciences). Anti-beta-actin antibody (Sigma Aldrich, A3853; 1 in 5,000) was used as a loading control. Blots were repeated in triplicate and statistics were performed using Prism 6. Data are presented as mean ± standard error of the mean (SEM). The significance between the variables was shown based on the p value obtained (ns indicates $p > 0.05$, * $p < 0.05$, ** $p < 0.005$, *** $p < 0.0005$, **** $p < 0.00005$).

Blue Native Polyacrylamide Gel Electrophoresis (BN-PAGE)

Fibroblast pellets were lysed using 10mM Tris (pH 8), 150mM NaCl, 0.1% NP40 with physical agitation for 30 min, centrifuged at 8,000xg to remove debris. The supernatant was removed, and total protein was quantified with the Bradford assay. Protein concentrations were equalized and prepared to 20µl at 1µg/µl using NativePAGE sample buffer (Thermo) and 1ul of NuPAGE 5% G-250 Sample Additive (Thermo), and then loaded to a NativePAGE 3%–12% Bis-Tris Protein Gel (Thermo). Proteins were transferred to polyvinylidene fluoride (PVDF) membrane through the use of an iBlot2 PVDF Mini transfer stack (ThermoFisher Scientific) and probed with anti-NARS1 monoclonal antibody (Abcam ab129162, 1:5000) and glyceraldehyde 3-phosphate dehydrogenase (GAPDH; Santa Cruz). Blots were repeated in triplicate, and differences were analyzed using Welch-corrected t test.

Induced Neuronal Progenitor Cell (iNPC) Conversion

Based on the protocol published by Meyer et al.,²⁴ iNPCs were generated from primary fibroblasts by transduction with *Oct4*, *Klf4*, and *c-Myc*-Sendai virus, followed by culturing in neuronal progenitor cell (NPC) induction media (1:1 DMEM/F-12: Neurobasal, 2× N2, 2× B27, 1% GlutaMAX, 10ng/mL hLIF, 3µM CHIR99021, and 2 µM SB431542). Neuroepithelial colonies were formed after 3–4 weeks of culturing. These were then isolated and expanded before we extracted total RNA from individual fibroblasts, age and sex matched healthy control fibroblasts, and iNPCs cells using the mirVana miRNA Isolation Kit (Ambion) for gene expression analysis by qPCR to confirm iNPC lineage and RNAseq in control and individual iNPCs in order to identify differentially expressed genes.

qPCR

Cell pellets from individual fibroblasts and iNPCs were lysed using a Trizol reagent. Following the addition of chloroform, the aqueous phase was transferred to RNeasy spin column (QIAGEN) for RNA isolation and resuspension. cDNA was generated using the reverse transcriptase kit (Applied Biosystems) and qPCR (Applied Biosystems 7900HT) was performed in triplicates using SYBR Green PCR Master Mix (Invitrogen, 4309155). Samples were normalized to expression of GAPDH and β-actin and repeated in triplicate.

RNaseq

Libraries were prepared using Illumina TruSeq Stranded Total RNA with Ribo-Zero Human kit and were sequenced on an Illumina Hi-Seq 2500 using a paired-end protocol. Quality of sequencing reads were ensured using FastQC. Reads were aligned using STAR aligner, and variants were called using the two-pass protocol outlined in the GATK documentation (see Web Resources). The numbers of

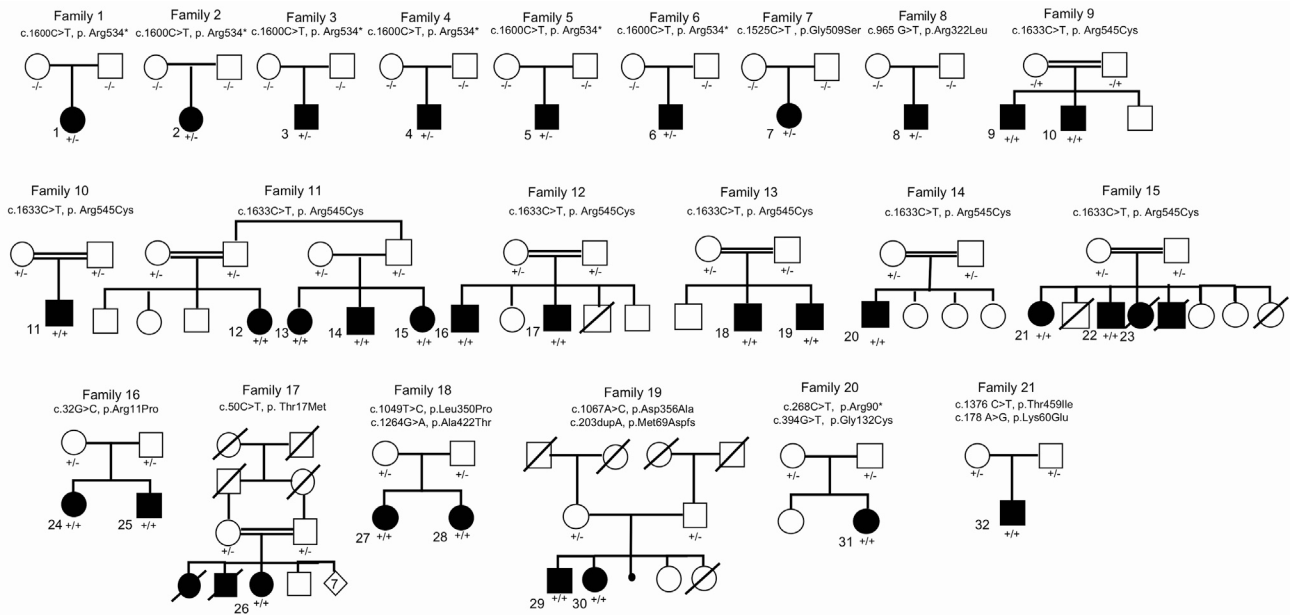


Figure 2. Pedigrees of the 21 Families and 32 Affected Individuals Identified in This Study with *de novo* and Bi-allelic Mutations in *NARS1*
 Filled symbols represent affected individuals and double bars represent consanguinity in the family. $-/-$, $+/-$, and $+/+$ represent wild-type, heterozygous, and homozygous variants, respectively.

reads were counted using HTSeq-count.²⁵ Differentially expressed genes were identified using the DESeq2 Bioconductor package.²⁶ Differentially expressed genes with a false discovery rate of ≤ 0.1 and a \log_2 (fold change) ≥ 1 were considered significant. Gene set enrichment analysis was performed using the CPDB web tool.

NARS1 Enzyme Assay

Aminoacylation was assessed by measuring *NARS1* activity in cultured fibroblasts and lymphoblasts. Cell lysates (cytosolic fraction) were incubated in triplicate at 37°C for 10 min in a reaction buffer containing 50mmol/L Tris buffer pH 7.5; 12mmol/L MgCl₂; 25mmol/L KCl; 1 mg/mL bovine serum albumin; 0.5mmol/L spermine; 1mmol/L ATP; 0.2mmol/L yeast total tRNA; 1 mmol/L dithiothreitol; and 0.3mmol/L [¹⁵N₂]-asparagine, [¹³C₄, ¹⁵N]-threonine, [D₂]-glycine, [¹⁵N₂]-arginine, and [D₄]-lysine. The reaction was terminated using trichloroacetic acid. Ammonia was then added to release the labeled AAs from the tRNAs. [¹³C₂, ¹⁵N]-glycine and [¹³C₆]-arginine were added as internal standards, and the labeled AAs were quantified via LC-MS/MS. Intra-assay variation was determined as $<15\%$ of TARS1, GARS1, RARS1 KARS1 activity which were simultaneously detected as control enzymes. AsnRS1 activities were measured blind, and testing was repeated in triplicate. Data are presented as mean \pm SEM. The statistical significance of the difference of AsnRS1 activity between controls and affected individuals and/or carriers was determined using a Student's *t* test with a 95% confidence interval through the use of SPSS 26.

Molecular Modeling Analysis

The crystal structure of *Brugia malayi* AsnRS1 with a 65% identity to human AsnRS1, stored under the 2XGT code in the Protein Data Bank, was used for the molecular modeling analysis. The homology model of the dimeric human AsnRS1 overlapped with the *S. cerevisiae* DARS-tRNAAsp ligase, co-crystallized with tRNA mole-

cule PDB 4WJ4, thus having a similar domain organization and sharing 27.5% of sequence identity with human AsnRS1. The protein was visualized with the University of California–San Francisco Chimera software.²⁷

Results

Genetic Analysis

We identified 21 families (F1–F21) and 32 affected individuals (P1–P32) with mutations in *NARS1* (Figure 1B shows *NARS1* variant schematic and Figure 2 illustrates pedigrees). Eight families had *de novo* heterozygous variants; six had c.1600C>T (p.Arg534*) (F1–F6, P1–P6); one had c.1525G>A (p.Gly509Ser) (F7, P7); and one had c.965G>T (p.Arg322Leu) (F8, P8). These variants were not present in our 652 normal brain series or in the gnomAD database.

Bi-allelic variants were found in thirteen families. Seven have homozygous c.1633C>T (p.Arg545Cys) variants (F9–F15, P9–P23); one has homozygous c.50C>T (p.Thr17Met) (F17, P26); and one has two siblings with homozygous c.32G>C (p.Arg11Pro) (F16, P24 and 25). For compound variants, one family has two siblings with compound heterozygous c.1067A>C (p.Asp356Ala) and c.203dupA (p.Met69Aspfs*4) (F19, P29 and P30). Two siblings had compound heterozygous c.1049T>C (p.Leu350Pro) and c.1264G>A (p.Ala422Thr) variants (F18, P27 and P28). There was one case with the compound heterozygous variants c.268C>T (p.Arg90*) and c.394G>T (p.Gly132Cys) (F20, P31) and a final individual with compound heterozygous c.1376C>T (p.Thr459Ile) and

c.178A>G (p.Lys60Glu) variants (F21, P32). In gnomAD, c.1264G>A (p.Ala422Thr) is present in six heterozygote individuals, whereas c.1633C>T (p.Arg545Cys) and c.50C>T (p.Thr17Met) were present in five and four heterozygotes, respectively. The c.100 A>T (p.Met34Leu), c.203dupA (p.Met69Aspfs*4), and c.1049T>C (p.Leu350Pro) variants were absent, while c.32G>C (p.Arg11Pro) was present in one individual. The c.1067A>C (p.Asp356Ala) variant in family 19 was present in 264 heterozygotes, suggesting that this variant may modify the phenotype and be pathogenic only when in *trans* with a severe variant such as c.203dupA (p.Met69Aspfs*4).

Clinical Characteristics

Table 1 summarizes the core clinical features of affected individuals with *NARS1* defects (see Tables S2–S4 for additional details). All individuals had global developmental delay (GDD) and intellectual disability, which varied in severity from moderate to profound. They had marked delays in language development. Motor development was also severely impaired, and one individual never acquired autonomous ambulation. Microcephaly was observed in the majority of cases (90%). These cases predominantly presented with primary microcephaly; however, secondary microcephaly was also noted. Epilepsy was highly associated with the phenotype, affecting 23 cases (74.2%), with six individuals experiencing seizures below the age of one. The semiology of these attacks varied, with a mixture of partial, myoclonic, and generalized tonic-clonic seizures described. An ataxic gait, poor balance, and dysarthria were frequently detected on examination; this suggests an additional neurodegenerative process; however, no structural abnormality of the cerebellum was observed on imaging. A demyelinating peripheral neuropathy occurred in eight individuals (25%) who had distal leg muscle atrophy. Dysmorphic features described included abnormal hands (e.g., clinodactyly, fetal finger pad, two-to-three-toe syndactyly, slender fingers) and/or feet (e.g., small feet, toe syndactyly, slender feet). Upslanting palpebral fissures was the most common facial dysmorphism reported. A broad forehead, wide mouth, wide-set teeth, and low-set ears with overfolded helices were also described. Skeletal abnormalities including scoliosis, pronounced thoracic kyphosis, and *pes-cavus* were also noted. Behavioral traits associated with the phenotype included impulsivity, stereotypies with repetitive speech and/or hand movements, and selective feeding rituals.

Genotype-Phenotype Correlations

Family 16, with the homozygous variant c.32G>C (p.Arg11Pro), had a particularly severe clinical picture comprised of severe developmental delay, progressive microcephaly, refractory seizures from infancy, and arrested myelination with pronounced cerebral atrophy on MRI (see Supplemental Note, Table S3, and Figure 3).

Otherwise, imaging was normal apart from microcephaly. There was no common structural change across

all cases. Individuals with the *de novo* c.1600C>T (p.Arg534*) variant showed severe microcephaly. In one family with this variant (F6), mild atrophy was observed (see Supplemental Information, Tables S2–S4, and Figure 3).

Individuals homozygous for c.1633C>T (p.Arg545Cys) demonstrated hypotonia and predominantly distal weakness. Spasticity was observed in individuals with the c.32G>C (p.Arg11Pro) or *de novo* variants.

A demyelinating polyneuropathy was documented in individuals homozygous for the c.1633C>T (p.Arg545Cys) variant (P9, P10, and P20), and in one case, this was confirmed with a sural nerve biopsy (F9, P9). It was also described in individuals with the *de novo* c.1600C>T (p.Arg534*) variant (P1, P2, and P5) and in the family with the compound heterozygous c.1049T>C (p.Leu350Pro) and c.1264G>A (p.Ala422Thr) variants (F18, P27 and P28).

Pathogenicity of *NARS1* Variants

NARS1 is intolerant to loss of function (missense variants constraint is $Z = 0.87$). We identified *de novo* *NARS1* mutations in eight families (F1–F8, P1–P8) with similar phenotypes. A variant at codon 534 recurred in six families (F1–F6, P1–P6). The two other *de novo* variants altered codons 322 and 509. The c.1600C>T (p.Arg534*) variant is located 15 AAs from the end of the 548-AA protein, representing a potential hotspot for pathogenic mutations. Arginine at codons 534 and 545 is universally conserved in AsnRS1 from all three major taxonomic groupings, implying a significant structural or functional role.

The homozygous c.1633C>T (p.Arg545Cys) variant was observed in seven families with recessive disease. This variant affects the same C-terminal catalytic stretch as does c.1600C>T (p.Arg534*), and therefore it might have a comparable mechanistic effect to c.1600C>T (p.Arg534*).

The c.1067A>C (p.Asp356Ala) variant was found in *trans* with the only recessive truncating allele observed thus far at c.203dupA (p.Met69Aspfs*4) (P29 and P30). Two missense variants (c.965G>T [p.Arg322Leu] and c.653T>C, p.Asn218Ser) were found in P8; however, because c.965G>T (p.Arg322Leu) occurred *de novo*, it could not be determined whether these variants were in *cis* or *in trans*. Moreover, the frequency of c.653T>C (p.Asn218Ser) in the gnomAD database (78 heterozygotes) suggests it is unlikely to be associated with a severe phenotype, leaving c.965G>T (p.Arg322Leu) as the most likely disease-causing variant. The Arg322 residue is essential for enzymatic activity and therefore is predicted to cause impaired enzyme activity. Both the c.50C>T (p.Thr17Met) and c.32G>C (p.Arg11Pro) variants are in the N-terminal UNE-N appended domain of AsnRS1, which is specific to eukaryotes, and has recently been shown to have chemokine activity.²⁸

Functional Characterization

Western Blotting and RT-PCR

Given the potential loss of function in homozygous *NARS1* individuals, we investigated gene expression levels

Table 1. Summary of NARS1 Variants and Clinical Features of Affected Individuals

Variant: Nucleotide, Protein	c.1600C>T, p.Arg534*	c.1525G>A, p.Gly509Ser	c.965G>T, p.Arg322Leu	c.1633 >T, p.Arg545Cys	c.32G>C, p.Arg11Pro	c.50C>T, p.Thr17Met	c.1049T>C c.1264G>A, p.Leu350Pro p.Ala422Thr	c.1067A>C c.203dupA, p.Asp356Ala p.Met69Aspfs*4	c.268 C>T c.394G>T, p.Arg90* p.Gly132Cys	c.1376 C>T, c.178 A>G, p.Thr459Ile, p.Lys60Glu
Variant type	de novo heterozygous	de novo heterozygous	de novo heterozygous	homozygous	homozygous	homozygous	compound heterozygous	compound heterozygous	compound heterozygous	compound heterozygous
Inheritance	AD <i>de novo</i>	AD <i>de novo</i>	AD <i>de novo</i>	AR	AR	AR	AR	AR	AR	AR
Family	1–6	7	8	9–15	16	17	18	19	20	21
Affected Individual(s)	1–6	7	8	9–23	24–25	26	27–28	29–30	31	32
Ethnicity/country of origin	European	UK	European	Pakistan/North India	Kosovo	Libya	German	Turkey	Canada	USA
Age at onset	birth	birth	birth	childhood	childhood	birth	birth	birth	birth	childhood
Consanguinity	no	no	no	yes	no	yes	No	no	no	no
Presentation	severe GDD	severe GDD	severe GDD	severe GDD	seizures	seizures	mod GDD	mod GDD	severe GDD	severe GDD
ID	yes	yes	yes	yes	yes	yes	yes	yes	yes	yes
Microcephaly	yes	no	NA	yes	yes	yes	yes	yes	yes	yes
Dysmorphic	yes	yes	yes	yes	no	NA	no	no	yes	no
Seizures Affected Individuals	yes 1, 2, 4, 5, 6	yes	yes	yes 9, 14, 15, 18, 19, 21, 22, 23	yes all individuals	yes	yes 27	yes all individuals	yes	yes
Spasticity Affected Individuals	yes 3, 4, 6	no	yes	no hypotonia in 9, 10, 16, 17	yes 24	na	no hypotonia	na	no hypotonia	yes
Neuropathy Affected Individuals	Yes 1, 2, 5	NA	NA	yes 9, 10, 20	NA	NA	yes	NA	NA	NA
Ataxia Affected Individuals	yes all individuals	NA	Yes	yes 9–12, 21	NA	NA	yes	NA	yes	yes

AD = autosomal dominant, AR = autosomal recessive, GDD = global developmental delay, ID = intellectual disability Mod = moderate, NA = not available.

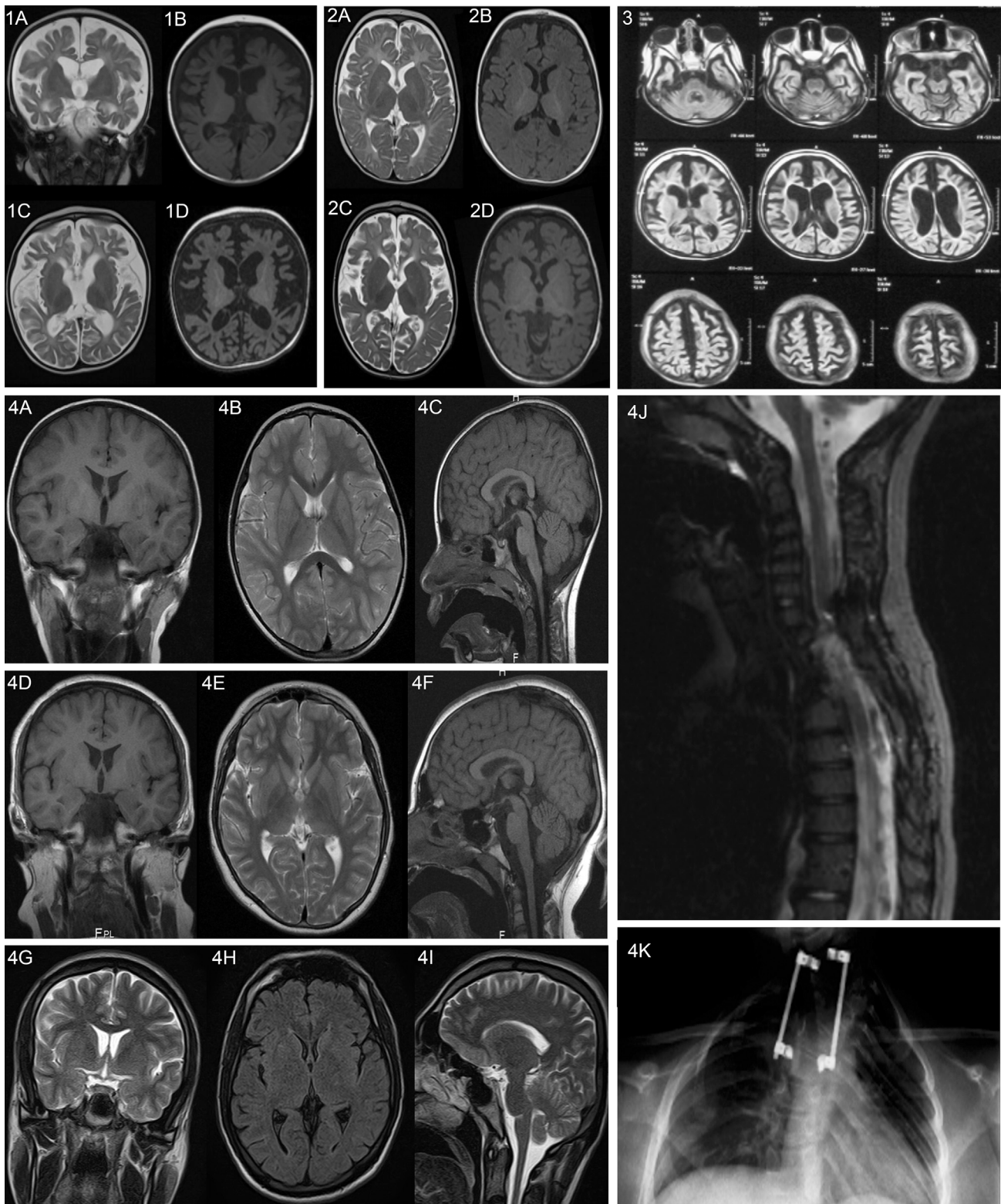


Figure 3. Radiological Findings of Individuals in Our Cohort

Set 1: Individual homozygous for c.32G>C (p.Arg11Pro). Upper row images (coronal T2-WI [1A] and axial T1-WI [1B]) at the age of 10 months show severely delayed myelination and fronto-temporal atrophy. Lower row images (axial T2-WI [1C] and axial T1-WI [1D]) repeated at the age of 18 months show progressive and global brain atrophy with an emerging pattern of severe hypomyelination. Set 2: An additional homozygous c.32G>C (p.Arg11Pro) individual. Upper row images (axial T2-WI [2A] and axial T1-WI [2B]) at the age of 8 months show mild fronto-temporal underdevelopment and severely delayed myelination. Lower row images (axial T2-WI [2C] and axial T1-WI [2D]) repeated at the age of 2 years shows progressive and global brain atrophy along with severe hypomyelination.

(legend continued on next page)

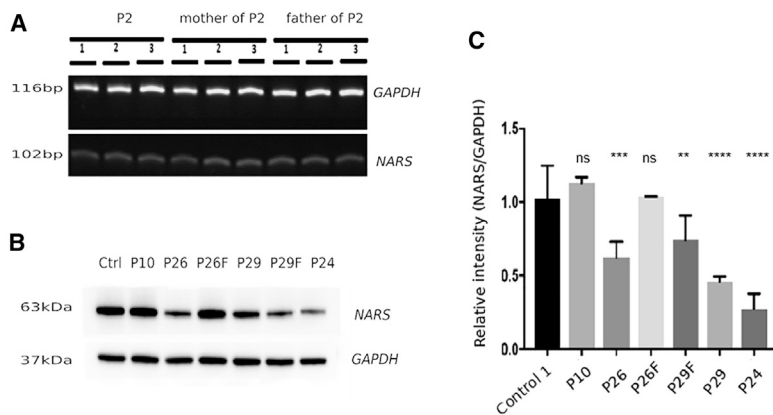


Figure 4. Protein Levels of AsnRS1 Are Reduced in Individual-Derived Cells

(A) RT-PCR of the *de novo* c.1600C>T (p.Arg534*) variant in P2 and parents (B) western blotting and (C) quantification graph of individuals with *NARS1* mutations compared with controls. Ctrl = control, P10 = homozygous c.1633C>T (p.Arg545Cys), P26 = homozygous c.50C>T (p.Thr17Met), P29 = compound heterozygous (c.1067A>C (p.Asp356Ala) and c.203dupA (p.Met69Aspfs*4) (F denotes father of individuals), P24 = homozygous c.32G>C (p.Arg11Pro).

through the use of semiquantitative PCR and protein levels through western blotting of AsnRS1 from lymphoblasts and fibroblasts from families harboring the p.Arg545Cys, p.Thr17Met, p.Asp356Ala, p.Met69Aspfs*4, and p.Arg11Pro variants. In all instances, both gene expression and protein levels are reduced (Figure 4A–4E).

iNPCs

iNPC colonies were produced and isolated from fibroblasts from P26 (c.50C>T), P29 (c.1067A>C), and P30 (c.203dupA). From the isolated colonies, gene expression was determined by using qPCR to select iNPC populations which presented decreased expression of fibroblast markers *COL1A1*, *COL3A1*, *TWIST2*, and *DKK3*, as well as increased numbers of NPC markers *NES*, *SOX1*, and *MSI1*, and the iNPC population was expanded to be subsequently used for RNA sequencing (RNaseq). RNaseq showed normal *NARS1* expression in iNPCs from affected individuals carrying the c.50C>T and c.1067A>C mutations, and decreased expression of the c.203dupA *NARS1* allele in the P29 cells. Interestingly, iNPCs from affected individuals show increased expression of several other ARSs (*DARS1*, *GARS1*, *RARS1*, *SARS1*, *TARS1*, *WARS1*, and *YARS1*) (Figure 5). This could be explained by the fact that the *NARS1* mutant(s) are inducing the integrated stress response (ISR), which activated a number of ARS genes as a result of the loss-of-function homozygous recessive variants. Impaired synthetase function may reduce the amount of charged tRNA available for translation elongation, with a possible increase in the levels of uncharged tRNA. Uncharged tRNAs produced as a result of AA deprivation have been reported to bind GCN2, leading to the activation of the ISR.²⁹ Analysis of the cellular pathways (Reactome, Gene Ontology) associated with genes with significantly altered mRNA levels showed that upregulated

genes were enriched (adjusted p value < 0.01) for pathways heavily associated with protein translation and processing such as endoplasmic reticulum (ER) and Golgi protein processing and ribosomal homeostasis. In addition, increased action of VEGFR1/2 (upregulated by ATF4, which is one of the key transcription factors in the ISR) was suggested.

Blue-Native Polyacrylamide Gel Electrophoresis (BN-PAGE)

Similar to most other disease-associated ARSs, AsnRS1 functions as a class II homodimer.³⁰ We showed severely reduced dimer formation in P26 (c.50C>T [p.Thr17Met]) and P29 (c.1067A>C [p.Asn356Ala] and c.203dupA [p.Met69Aspfs*4]) compared to healthy controls (Figure 5E). The unaffected parents carrying one heterozygous mutation each also appeared to show a decreased level of the AsnRS1 dimer. P10 (c.1633C>T [p.Arg545Cys]) showed an AsnRS1 dimer amount comparable to that healthy controls. The decreased AsnRS1 dimer formation observed in fibroblasts from P26 and P29 shown by BN-PAGE accounts for the apparent deficit in aminoacylation capacity, despite showing no consistent decrease in the levels of AsnRS1 monomers. This idea is further supported by the molecular model simulation (Figure 6) that predicts an unstable dimer for the p.Asn356Ala mutant because this substitution is located at the interface between the two AsnRS1 monomers.

ARS Enzymatic Assays

In comparison with controls, AsnRS1 enzymatic activity was reduced in proband-derived fibroblasts and lymphoblasts. The most dramatic decrease was observed for P2 (*de novo* c.1600C>T [p.Arg534*]), and the mildest decrease was observed for P24 (c.32G>C [p.Arg11Pro], 80% of the controls). AA residue Arg11 is located in the 5' end of the non-canonical UNE-N domain (Figure 7 and Figure S11), which has recently been shown to elicit cell migration of human immune cells via migration of CC chemokine

Set 3: Individual homozygous for c.50C>T (p.Thr17Met). Axial fluid-attenuated inversion recovery (FLAIR) images at the age of 9 months show global atrophy involving the cerebral and cerebellar hemispheres along with severe hypomyelination.

Set 4: MRI images of an individual with the homozygous c.1633C>T (p.Arg545Cys) variant. Coronal T1-WI (4A), axial T2-WI (4B), and sagittal T1-WI (4C) at the age of 4 years; coronal T1-WI (4D), axial T2-WI (4E), and sagittal T1-WI (4F) at the age of 11 years; and coronal T2-WI (4G), axial FLAIR (4H), and sagittal T2-WI (4I) at the age of 20 years. These demonstrate normal intracranial appearances across the three different ages. This individual had an upper thoracic scoliosis, which was operatively corrected at the age of 4, demonstrated on the sagittal T2-WI of the spine (4J) and frontal projection radiograph of the chest/thoracic spine (4K).

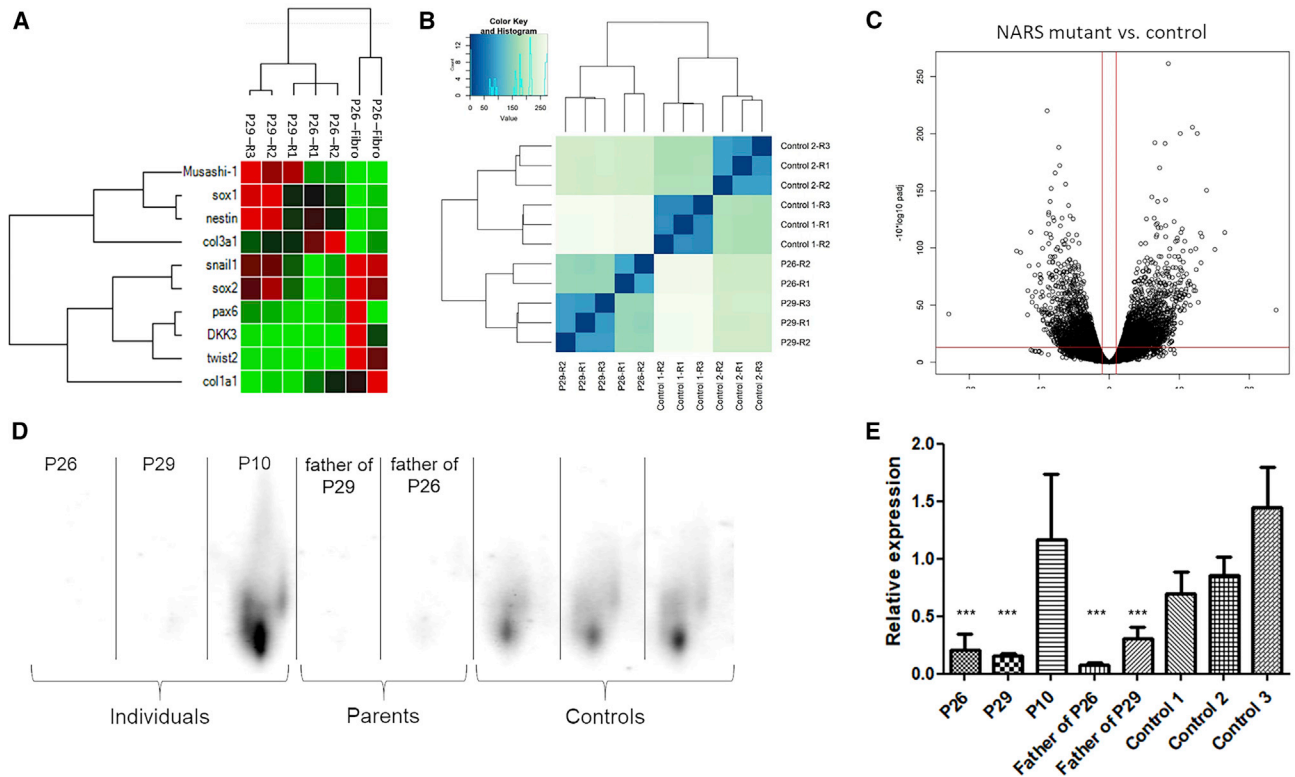


Figure 5. BN-PAGE and iNPC RNA-Sequencing

(A) iNPCs from P26 (c.50C>T [p.Thr17Met]) and P29 (c.203dupA [p.Met69Aspfs*4]) and c.1067A>C [p.Asp356Ala]) exhibit increased expression of most iNPC markers (sox1, sox2, nestin, snail1, pax6, DKK3, twist2, and Musashi-1) compared to fibroblast (fbb) as measured by qPCR, shown with hierarchical clustering.

(B) Heatmap with hierarchical clustering generated using all gene counts from RNaseq distinction of control (Ctrl1 a-c, Ctrl2 a-c) and individual-derived (P26 a-b, P29 a-c) iNPCs.

(C) Volcano plot showing log₂ of fold change in NARS mutant iNPCs compared to controls and -log₁₀ (adjusted p value).

(D) BN-PAGE western blot showing reduced levels of the AsnRS1 dimer in individuals P26 and P29 and fathers compared to control, but not for individual P10.

(E) Quantification of BN-PAGE western blot AsnRS1 dimer formation, showing significantly (***) reduced levels of the AsnRS1 in individuals P26 and P29 and fathers compared to control but not change for P10.

P26 = homozygous c.50C>T (p.Thr17Met), P29 = c.203dupA (p.Met69Aspfs*4), c.1067A>C (p.Asp356Ala), P10 = c.1633C>T (p.ArH545Cys), father of P26 = heterozygous c.50C>T (p.Thr17Met), father of P29 = c.1067A>C (p.Asp356Ala).

receptor 3 (CCR3) in an autoimmune disease associated with ARS genes.²⁸

Discussion

We identified *de novo* heterozygous and bi-allelic mutations in *NARS1* in 32 individuals with a neurodevelopmental phenotype. Mutations included recessive mutation hotspots affecting AA residues Arg534 and Arg545, respectively, both located in the last 40 AAs of the protein. Two homozygous variants identified at the 5' end, c.32G>C (p.Arg11Pro) and c.50C>T (p.Thr17Met), were associated with a severe clinical phenotype. Other mutations in *NARS1* were spread throughout and did not cluster in any particular region of the gene.

The clinical phenotypes associated with homozygous variants c.32G>C (p.Arg11Pro) and c.50C>T (p.Thr17Met) correlate with reduced protein levels and could reflect impaired protein stability as suggested by the structural

modeling of c.1633C>T (p.Arg545Cys) (Figure 6). Interestingly, MRI imaging of individuals harboring the c.32G>C (p.Arg11Pro) and c.50C>T (p.Thr17Met) variants showed atrophy and white matter abnormalities. In contrast, no such changes were identified in individuals with the p.Arg545Cys variant. The clustering of variants and associated phenotypes at the N and C termini suggests these regions are functionally important and disrupt the protein homodimer and ATP-binding and/or catalytic domain in *NARS1*. These two variants produced elevated AsnRS1 enzyme activity, which can be attributed to their location in the N-terminal extension domain. This domain has additional non-translational functions, enabling enzymatic activity of the modified protein. When we examine protein expression, protein synthesis, and the aminoacylation activity, it is clear that the non-translational functions of such ARS proteins, regulated by the newly evolved appended domains such as UNE-N, don't seem necessary for ARS activity (Figure 7 and Figure S11).

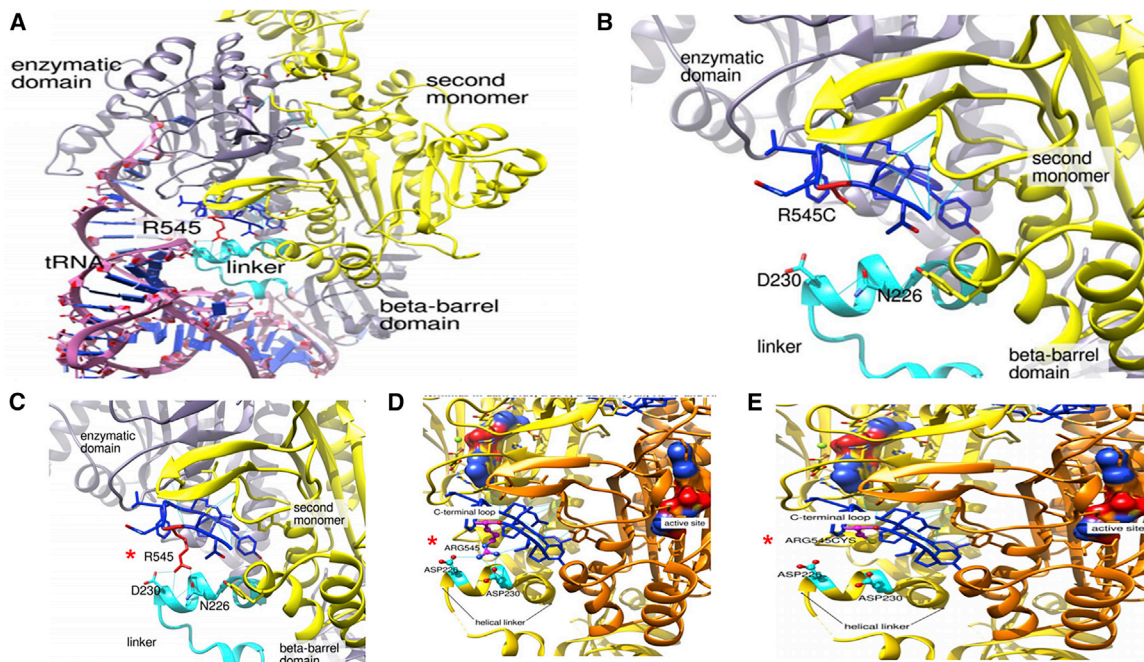


Figure 6. Molecular Modeling of the NARS1 p.Arg545Cys Homozygous Variant

The crystal structure is based on *B.malayi* AsnRS1. AsnRS1 is a homodimer; one AsnRS1 monomer is given in yellow and one in orange. Analog of the transition state presented in the surface representation, C terminus in dark blue, Asp230 and Asp226 in cyan, Arg545 and Arg545Cys in magenta.

(A) Interaction between *AsnRS1* and tRNA with residues on the helical linker.

(B–E) Zoom in on the C terminus helical linker region, (B) and (E) show loss of molecular interaction and folding of the p.Arg545Cys variant (*).

Our functional data, including fibroblasts and iNPCs transcriptomics, suggest that the majority of *NARS1* mutations cause a loss of the enzymatic protein by reduced expression and disruption of dimer formation. This results in abnormal protein synthesis and processing with a compensatory increase in expression of other ARSs (Figures 4, 5, and 7 and Figure S11). The increased activity of VEGFR1 and VEGFR 2 was of interest considering the reported actions of other ARSs, as in *GARS* and *EPRS* via GAIT complex, *SARS1*, *TARS1*, and mini *WARS1* on VEGF-related signaling.³¹ Pathways associated with downregulated genes were typically associated with cell cycle progression, DNA repair and replication such as G2/M checkpoint, homology directed repair, and telomere maintenance pathways; this suggests that this alteration of cellular proliferation could be a result of decreased protein synthesis (Figures S5–S10). In general, the mutations in *NARS1* resulted in loss of function in both studied iNPC cell lines (P26 and P29), leading to a transcriptomic signature of induced ISR, upregulation of protein translation and processing in the ER and Golgi, and altered ribosomal homeostasis. This is similar to the results of other studies in cells with reduced aminoacylation activity in disease-associated mutations in other cytosolic ARSs.^{5,31,32}

The recurrent homozygous c.1633C>T (p.Arg545Cys) variant in the western blot (Figure 4) and yeast model (Figure S12) showed near normal protein levels and an

increased yeast growth suggestive of a gain-of-function mechanism. However, protein modeling of this variant demonstrated loss of the helix linker, and this indicates reduced tRNA interaction and catalytic activity. This loss-of-function effect was evidenced by the reduced aminoacylation activity to 40% compared to controls (Figure 7 and Figure S11). This effect could potentially be more harmful for cells of the nervous system than for a unicellular organism. One of the possibilities is that the *NARS1* mutant mischarges a tRNA in the human cells that might be less conserved in fungi. Thus, the mischarging would be reduced in yeast, hence the better growth without side effects. Molecular modeling has shown that the p.Arg545Cys variant lies within a region that probably interacts with the sugar-phosphate backbone of the tRNA (at positions 68–69), close to the active site of the enzyme.³³ Replacing the bulky arginine with a cysteine does not seem to perturb the enzyme's overall structure (Figure 6). However, by disrupting the tRNA-enzyme contact, this variant may alter the enzyme selectivity toward tRNA, decreasing the overall affinity for tRNA. AsnRS1 enzyme activity for individuals homozygous for this variant (P9 and P20) showed decreased activity (Figure 7 and Figure S11). Similarly, the *de novo* c.1600C>T (p.Arg534*) variant, located adjacent to the end of the protein, has a gain-of-function effect that interferes with normal protein function. It is likely a protein that lacks the 15 AAs containing the ATP-binding domain is produced. This region is crucial for enzymatic function, and it escapes

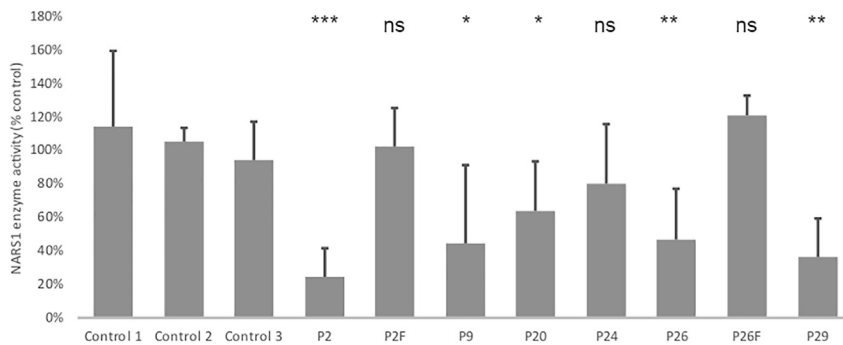


Figure 7. Reduced Asparaginyl-tRNA Synthetase Activity in Individuals with Homozygous *NARS1* Variants
 c.1600C>T (p.Arg534*) (P2), c.1633C>T (p.Arg545Cys) (P9 and P20), c.32G>C (p.Arg11Pro) (P24), c.50C>T (p.Thr17Met) (P26), and c.1067A>C (p.Asp356Ala)/c.203dupA (p.M69Aspfs*4) (P29) *NARS1* variants in comparison to the average of three unrelated fibroblast cell lines. (All cell lines are fibroblast except P2, which is a lymphoblast cell line. Control values for lymphoblast are similar to fibroblasts.) n = 9, p value FDR < 0.01.

mRNA decay, as shown by semiquantitative RT-PCR from family 2 (Figure 4A). We further developed a zebrafish model of this variant (Figure S13) which elicits a dominant negative effect on the wild-type allele, causing a dose-dependent phenotype, specifically cyclopia and gastrulation defects at 200–500pg. Similar cyclopia defects in zebrafish were reported for microcephaly gene *ORC1*.³⁴

Given the essential function and constraint metrics of *NARS1*, in conjunction with the clinical phenotypes of included individuals, we propose that genotypes with dominant heterozygous variants produce a toxic gain of function. This is compared with the homozygous recessive variants that probably experience a loss of function, though this can perhaps be least partially compensated for by other ARS genes. Taking into consideration the aminoacylation assay and yeast model for *de novo* mutations, and the western blot, aminoacylation assay, and modeling for homozygous recessive mutations, we have confirmed the pathogenicity of all *NARS1* mutations mentioned. Similar effects are seen in Aicardi-Goutières syndrome, which is caused by pathogenic variants in *ADAR*. There are relatively frequent alleles that are pathogenic when *in trans* to a null, but are never found in individuals with the homozygous state.³⁵ For distal C terminus mutations, such as the homozygous c.1633C>T (p.Arg545Cys) variant, the mechanism is likely due to abnormal protein structure and catalytic activity (Figures 1C and 6 and Figures S1 and S2).

Affected individuals had both central and peripheral nervous system involvement and a broad neurodevelopmental phenotype characterized by GDD, microcephaly, ataxia, neuropathy, and seizures. This is reflective of high *NARS1* expression in the cortex, cerebellum, and brainstem as demonstrated in mouse brains^{36,37} (Figure S3). Mutations have been reported for the majority of ARSs. AsnRS2, a mitochondrial ARS protein coded by *NARS2*, has recently been linked with an overlapping phenotype consisting of multisystem mitochondrial disorder (MID). Intellectual disability, epilepsy in childhood, hearing loss, and myopathy have also been seen in *NARS1* individuals.^{38–41} In addition, ARS interacting multifunctional proteins 1–3 (AIMP1–3) participate together with nine cytosolic ARSs to constitute the so-called multi-synthetase complex, and have also been associated with a variety of human diseases.⁴² In considering their critical cellular

functions, we expect that all ARSs will have a disease association.^{7–9} The *NARS1* data bring the number of characterized ARSs to 35 out of 37. On a modified Taylor's Venn diagram of AA properties, *NARS1* is placed in close proximity to other AAs with similar properties (*IARS1*, *LARS1*, *DARS1*, *EPRS1*, *NARS1*, *RARS1*, and *QARS1*) which also have more severe phenotypes⁴³ (Figure S4).

Our functional work supports the likelihood that there is a loss-of-function mechanism in homozygotes and has helped to further understand the role of *NARS1* mutations in disease. The development of CRISPR/Cas9 heterozygous knockin and homozygous knockout animal models is the next important step in understanding the molecular rationale of these *NARS1* variants. Considering the high number of individuals and variants identified here, the addition of *NARS1* to genetic testing panels for children and young adults presenting with NDD, epilepsy, and/or a demyelinating neuropathy may be of clinical benefit.

Data and Code Availability

The variants reported in this paper have been submitted to the Leiden Open Variation Database, and the accession numbers are: LOVD: 668185, LOVD: 668186, LOVD: 668187, LOVD: 668188, LOVD: 668189, LOVD: 668190, LOVD: 668191, LOVD: 668192, LOVD: 668193, LOVD: 668194, LOVD: 668195, LOVD: 668196, LOVD: 668197, and LOVD: 668198.

Supplemental Data

Supplemental Data can be found online at <https://doi.org/10.1016/j.ajhg.2020.06.016>.

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Declaration of Interests

Maria J. Guillen Sacoto, Lindsay B. Henderson, Yue Si, Aida Telegrafi, and Ingrid M. Wentzensen are employees of GeneDx. The other authors declare no competing interests.

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Web Resources

1000 Genomes Project, <https://www.genome.gov/27528684/1000-genomes-project>
Complete Genomics 69, <https://www.completegenomics.com/public-data/69-genomes/>
CPDB web tool, <http://cpdb.molgen.mpg.de/>
Ensembl, <http://www.ensembl.org/i>
FastQC, <http://www.bioinformatics.babraham.ac.uk/projects/fastqc/>
GATK documentation, <https://software.broadinstitute.org/gatk/>
Human Genome Variation Society, <http://www.hgvs.org>
Mutation Taster version 2, <http://www.mutationtaster.org/>
National Heart, Lung, and Blood Institute (NHLBI) Exome Variant Server, <https://evs.gs.washington.edu/EVS/>
Picard, <http://broadinstitute.github.io/picard/>
PolyPhen-2, <http://genetics.bwh.harvard.edu/pph2/>
SIFT, <http://sift.jcvi.org/>
University of California—San Francisco (UCSC) Genome Browser, <https://genome.ucsc.edu/>

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