Full title:

Natural history of Charcot-Marie-Tooth disease type 2A: a large international multicentre study

Short running title:

Phenotype and natural history of CMT2A

Menelaos Pipis1,#, Shawna ME Feely2,#, James M Polke1, Mariola Skorupinska1, Laura Perez2, Rosemary R Shy2, Matilde Laura1, Jasper M Morrow1, Isabella Moroni3, Chiara Pisciotta4, Franco Taroni5, Dragan Vujovic6, Thomas E Lloyd7, Gyula Acsadi8, Sabrina W Yum9, Richard Lewis10, Richard S Finkel11, David N Herrmann12, John W Day13, Jun Li14, Mario Saporta15, Reza Sadjadi16, David Walk17, Joshua Burns18, Francesco Muntoni19, Sindhu Ramchandren20, Rita Horvath21, Nicholas E Johnson22, Stephan Züchner23, Davide Pareyson4, Steven S Scherer6, Alexander M Rossor1, Michael E Shy2 and Mary M Reilly1 for the Inherited Neuropathies Consortium - Rare Disease Clinical Research Network (INC-RDCRN)*

1MRC Centre for Neuromuscular Diseases, UCL Queen Square Institute of Neurology, London, UK
2Department of Neurology, Carver College of Medicine, University of Iowa, Iowa City, Iowa, USA
3Department of Pediatric Neurosciences, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy
4Unit of Rare Neurodegenerative and Neurometabolic Diseases, Department of Clinical Neurosciences, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy
5Unit of Medical Genetics and Neurogenetics, Department of Diagnostics and Technology, Fondazione IRCCS Istituto Neurologico Carlo Besta, Milan, Italy
6Department of Neurology, The Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA
7Department of Neurology, The Johns Hopkins University School of Medicine, Baltimore, Maryland, USA
8Connecticut Children's Medical Center, Hartford, CT, USA
9The Children's Hospital of Philadelphia, and Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA
10Department of Neurology, Cedars-Sinai Medical Center, Los Angeles, CA, USA.
11Center for Experimental Neurotherapeutics, St. Jude Children’s Research Hospital, Memphis, TN, USA
12Department of Neurology, University of Rochester, Rochester, NY, USA
13Department of Neurology, Stanford Health Care, Stanford, CA, USA
14Department of Neurology, Wayne State University School of Medicine, Detroit, MI, USA
15Department of Neurology, University of Miami Miller School of Medicine, Miami, Florida, USA
16Massachusetts General Hospital, Boston, Massachusetts, USA
17Department of Neurology, University of Minnesota, Minneapolis, Minnesota, USA
18University of Sydney School of Health Sciences & Children's Hospital at Westmead, Sydney, Australia
19Dubowitz Neuromuscular Centre, NIHR Biomedical Research Centre at UCL Great Ormond Street Institute of Child Health and Great Ormond Street Hospital, London, UK
20PRA Health Services, Raleigh, NC, USA
21Department of Clinical Neurosciences, University of Cambridge, Cambridge Biomedical Campus, Cambridge, UK
22Virginia Commonwealth University, Richmond, VA, USA
23Dr. John T. Macdonald Foundation Department of Human Genetics and John P. Hussman Institute for Human Genomics, University of Miami Miller School of Medicine, Miami, FL, USA
# These authors contributed equally to this manuscript.

*Other members of this consortium that are not main authors but have contributed to this body of work are named in the appendix.

Corresponding Author: Mary M Reilly, MRC Centre for Neuromuscular Diseases, UCL
Queen Square Institute of Neurology, London, UK m.reilly@ucl.ac.uk
Abstract

Mitofusin-2 (MFN2) is one of two ubiquitously expressed homologous proteins in eukaryote cells, playing a critical role in mitochondrial fusion. Mutations in MFN2 (most commonly autosomal dominant; AD) cause Charcot-Marie-Tooth disease type 2A (CMT2A), the commonest axonal form of CMT, with significant allelic heterogeneity. Previous, moderately-sized, cross sectional genotype-phenotype studies of CMT2A have described the phenotypic spectrum of the disease, but longitudinal natural history studies are lacking.

In this large multicentre prospective cohort study of 196 patients with dominant and autosomal recessive (AR) CMT2A, we present an in depth genotype/phenotype study of the baseline characteristics of patients with CMT2A and longitudinal data (1-2 years) to describe the natural history. A childhood onset of AD-CMT2A is the most predictive marker of significant disease severity and is independent of the disease duration. When compared to adult onset AD-CMT2A, it is associated with significantly higher rates of use of ankle-foot orthoses, full-time use of wheelchair, dexterity difficulties and also has significantly higher CMTESv2 and CMTNSv2 scores at initial assessment.

Analysis of longitudinal data using the CMTESv2 and its Rasch-weighted counterpart, CMTESv2-R, show that over 1 year, the CMTESv2 increases significantly in AD-CMT2A (mean change $0.84 \pm 2.42$; two-tailed paired t-test $p = 0.039$). Furthermore, over 2 years both the CMTESv2 (mean change $0.97 \pm 1.77$; two-tailed paired t-test $p = 0.003$) and the CMTESv2-R (mean change $1.21 \pm 2.52$; two-tailed paired t-test $p = 0.009$) increase significantly with respective standardised response means (SRMs) of 0.55 and 0.48. In the paediatric CMT2A population (AD-CMT2A and AR-CMT2A grouped together), the CMTPedS increases significantly both over 1 year (mean change $2.24 \pm 3.09$; two-tailed paired t-test $p = 0.009$) and over 2 years (mean change $4.00 \pm 3.79$; two-tailed paired t-test $p = 0.031$) with respective SRMs of 0.72 and 1.06.
This cross-sectional and longitudinal study of the largest CMT2A cohort reported to date provides guidance for variant interpretation, informs prognosis and also provides natural history data that will guide clinical trial design.

Keywords

Mitofusin-2
Charcot-Marie-Tooth disease type 2A
Genotype-phenotype correlations
Charcot-Marie-Tooth Examination Score v2.0
Standardised response mean
## Abbreviations list

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AD-CMT2A</td>
<td>Autosomal dominant Charcot-Marie-Tooth disease type 2A</td>
</tr>
<tr>
<td>AR-CMT2A</td>
<td>Autosomal recessive Charcot-Marie-Tooth disease type 2A</td>
</tr>
<tr>
<td>CMTESv2</td>
<td>Charcot-Marie-Tooth Examination Score v2.0</td>
</tr>
<tr>
<td>CMTESv2-R</td>
<td>Rasch-modified Charcot-Marie-Tooth Examination Score v2.0</td>
</tr>
<tr>
<td>CMTNSv2</td>
<td>Charcot-Marie-Tooth Neuropathy Score v2.0</td>
</tr>
<tr>
<td>CMTNSv2-R</td>
<td>Rasch-modified Charcot-Marie-Tooth Neuropathy Score v2.0</td>
</tr>
<tr>
<td>CMTPedS</td>
<td>Charcot-Marie-Tooth disease Pediatric Scale</td>
</tr>
</tbody>
</table>
Introduction

Mitofusin-1 (MFN1) and mitofusin-2 (MFN2) are homologous mammalian proteins and members of the large mitochondrial transmembrane GTPase family, exhibiting ubiquitous expression in eukaryotic cells and playing a fundamental role in the dynamic mitochondrial remodelling process governed by mitochondrial fusion and fission (Chandhok et al., 2018). These two highly coordinated biological processes, amongst other functions, are considered critical in mitigating mitochondrial stress, contributing to mitochondrial quality control and facilitating cellular apoptosis in cases of extreme cellular stress (Youle and van der Bliek, 2012). MFN2 is a 757-amino acid long, nuclear encoded protein (Supplementary table 1), anchored to the outer mitochondrial membrane by two transmembrane domains (TM1 and TM2). Most of the protein, including its large dynamin-GTPase domain and two coiled-coil heptad repeat regions (HR1/cc1 and HR2/cc2), are cytosolic (Westermann, 2010; Filadi et al., 2018). MFN2 is essential for mitochondrial fusion even though the exact mechanism by which this occurs is not fully understood. A widely accepted hypothesis is that MFN2 proteins on opposing outer mitochondrial membranes mediate tethering through homologous interactions primarily of their HR2, but also HR1 regions (Filadi et al., 2018). Furthermore, evidence from cultured neurons obtained from Mfn2 knockout-mice and embryonic rat and mice neurons expressing known pathogenic Mfn2 variants, suggest a role of MFN2 in the bi-directional axonal transport of mitochondria (Baloh et al., 2007; Misko et al., 2010). MFN2 also mediates sites of endoplasmic reticulum-mitochondrial contact which are important for calcium homeostasis (de Brito and Scorrano, 2008; Merkwirth and Langer, 2008; Filadi et al., 2015, 2017; Leal et al., 2016).

Mutations in MFN2 are associated with Charcot-Marie-Tooth disease type 2A (CMT2A), accounting for 4-7% of all genetically diagnosed CMT and 30-40% of genetically diagnosed axonal CMT (CMT2) (Saporta et al., 2011; Murphy et al., 2012; Fridman et al.,
In comparison to CMT1A, the commonest form of CMT, CMT2A is associated with a more severe, motor-predominant phenotype which usually manifests earlier in life and carries a greater burden of disability (Feely et al., 2011; Bombelli et al., 2014). Additional features include optic nerve atrophy in up to 9% of patients (Chung et al., 2006; Verhoeven et al., 2006; Züchner et al., 2006; Bombelli et al., 2014), vocal cord involvement (Chung et al., 2010; Kanemaru et al., 2019), upper motor neuron dysfunction (Chung et al., 2006; Ajroud-Driss et al., 2009; Choi et al., 2015; Luigetti et al., 2016) and white matter lesions on MRI (Chung et al., 2010; Lee et al., 2017). Moreover, variants at specific amino acid positions, eg. p.Arg104, cause a more complex syndrome with learning difficulties (Brockmann et al., 2008; Del Bo et al., 2008; Genari et al., 2011; Tufano et al., 2015; Tomaselli et al., 2018).

Genotype-phenotype studies of 25-45 CMT2A cases show significant phenotypic and allelic heterogeneity (Chung et al., 2006; Verhoeven et al., 2006; Feely et al., 2011; Bombelli et al., 2014). Missense variants in the heterozygous state are frequently implicated in CMT2A and the majority of these reside within or adjacent to the dynamin-GTPase domain. GTPase domain dimerisation may occur during tethering and mitochondrial fusion (Filadi et al., 2018), and there is recent evidence suggesting that a dominant negative or gain of function pathomechanism may be responsible. In-vitro and in-vivo models of CMT2A have shown that certain pathogenic variants (p.Arg94Gln, p.Thr105Met) cause mitochondrial hypofusion (El Fissi et al., 2018; Rocha et al., 2018; Ueda and Ishihara, 2018) whilst others (p.Leu76Pro, p.Arg364Trp) cause mitochondrial hyperfusion (El Fissi et al., 2018; Ueda and Ishihara, 2018). Certain variants, such as the p.Arg94Trp/Gln and p.Arg364Trp are repeatedly observed to occur de-novo and occur in guanine and cytosine nucleotides that reside in CpG dinucleotide sequences (Chung et al., 2006; Verhoeven et al., 2006). Despite this, polymorphisms in MFN2 although uncommon do occur and hence the interpretation of novel MFN2 variants is challenging. Furthermore, autosomal recessive and semidominant cases of CMT2A have been
published further illustrating the allelic heterogeneity of the condition (Nicholson et al., 2008; Polke et al., 2011; Tomaselli et al., 2016). A recessive trait is inherited and causes disease in a recessive manner and heterozygous carriers of recessive traits are usually phenotypically normal. However, a recessive trait is considered to also be inherited in a semidominant manner when it can cause a late-onset, mild disease in the heterozygous state. Examples of variants showing semidominant inheritance in CMT2A include p.Thr362Arg (Tomaselli et al., 2016) and p.Thr362Met (Nicholson et al., 2008). Interestingly, the former, which is described in three further families in this study, is absent from the genome aggregation population database (gnomAD) (Karczewski et al., 2020) whereas the latter is present nine times in gnomAD. Nonetheless, these are very rare variants and using the minor allele frequency in population databases to distinguish between semidominant and likely benign heterozygous variants is challenging. Interestingly, multiple symmetrical lipomatosis, a rare and phenotypically distinct disease characterised predominantly by massive increase in upper body adipose tissue and the presence of lipomata, has been associated with biallelic MFN2 mutations (Sawyer et al., 2015; Rocha et al., 2017). In all cases, at least one of the MFN2 variants is p.Arg707Trp and phenotypically some patients also manifest a late-onset axonal neuropathy.

The mitofusin knockout mouse models illustrate the biological importance of mitofusins, since strains in which either Mfn1 or Mfn2 are completely knocked out die in utero (Chen et al., 2003). Furthermore, Mfn-depleted embryonic fibroblasts show fragmented mitochondria, most likely due to the severely impaired process of mitochondrial fusion (Chen et al., 2003). However, both Mfn1 and Mfn2 heterozygous knockout strains do not express a phenotype and demonstrate normal fertility (Chen et al., 2003). There are several transgenic mouse models of CMT2A (Mfn2R94Q; Mfn2R94W; Mfn2T105M) which, nonetheless, seem to provide conflicting evidence and none of which show progressive peripheral axonal
Aim of this study

This is a cross-sectional and longitudinal study of the largest CMT2A cohort reported to date which has been collected as part of the ongoing Inherited Neuropathy Consortium (INC-RDCRN) natural history study of CMT. The aim of the study is to provide genotype-phenotype correlations to aid variant interpretation, inform prognosis and to provide natural history data to guide clinical trial design.
Materials and Methods

Ethical approvals, study design and patient recruitment

Patients included in this study were enrolled in the INC-RDCRN 6601 natural history protocol (registered at ClinicalTrials.gov NCT01193075), and 6602 and 6603 research protocols which gained ethical approval from the institutional review boards and research ethics committees of the participating centres in the US, UK, Italy and Australia. All patients or their guardians signed the relevant consent/assent forms. Patients were evaluated at one of the 19 INC centres between 2009 and 2019 and at Wayne State University between 1996 and 2009. Antecedent clinical data was collected retrospectively from the patient history. Longitudinal follow-up data (clinical history and examination with or without neurophysiological studies) was collected prospectively during annual visits.

MFN2 variant curation and classification

For ease of reading, throughout this paper, the term autosomal dominant (AD) CMT2A (AD-CMT2A) refers to all the cases that carry a heterozygous variant in MFN2, irrespective if the variant was parentally inherited or occurred de-novo. In our study, we used pathogenicity criteria from both pathogenic and benign categories (Supplementary table 2) as published in the American College of Medical Genetics and Genomics and Association for Molecular Pathology (ACMG/AMP) guidelines (Richards et al., 2015), and classified all MFN2 variants into pathogenic (P), likely pathogenic (LP) and variants of uncertain significance (VUS) (Supplementary tables 3 and 4). The most recent Association for Clinical Genomic Science (ACGS) recommendations on variant classification (Ellard et al., 2020) were also taken into consideration to appropriately use downgraded or upgraded pathogenicity criteria in cases/pedigrees that could not be fully classified on the ACMG/AMP guidance alone. The ACMG/AMP guidelines suggest using the PP1 criterion (segregation data) as a stronger
evidence with increasing segregation data, however, they do not provide a quantification for this. To supplement this we used the Jarvik and Browning guidance (Jarvik and Browning, 2016) on how to quantify segregation data from multiple affected family members and to then appropriately assign pathogenicity criteria to that variant. Briefly, the product of all informative meioses across all affected members from all unrelated families is used to determine if the segregation criterion can be used as a supporting (PP1), moderate (PP1_moderate) or strong (PP1_strong) level of evidence based on probabilistic pre-set cut-offs between the three categories. We also used the REVEL meta-predictor tool which combines pathogenicity and conservation scores from individual in-silico prediction tools (Ioannidis et al., 2016). The REVEL-derived aggregate score is more accurate compared to the combination of individual tools which often assess overlapping subsets of evidence, thus inadvertently leading to ‘double-counting’ of evidence.

The presence of a variant in the gnomAD population database (Karczewski et al., 2020) was considered in the context of the disease prevalence, penetrance, genetic and allelic heterogeneity of CMT. A threshold of an allele count (AC) of 3 in gnomAD was used to distinguish between heterozygous variants that are plausible or not to be causal for CMT2A (Pipis et al., 2019). Furthermore, all the variants described in autosomal recessive CMT2A cases (AR-CMT2A) in this study (Supplementary table 5) had a gnomAD population allele frequency that would be compatible with the genetic architecture of autosomal recessive CMT as previously published (Pipis et al., 2019). We have also detailed the benign (B) and likely benign (LB) variants we have encountered in our study and the reasons for their classifications (Supplementary table 6).

Previously published case series and case reports of CMT2A were identified through an extensive PubMed literature review, the Inherited Neuropathy Variant Browser (Saghira et al., 2018) and ClinVar (Landrum et al., 2014).
CMT clinical outcome measures

The clinical outcome measures used in this study include the CMT Neuropathy Score version 2 (CMTNSv2) and the Rasch modified CMTNSv2 (CMTNSv2-R), both composite scores based on patients’ symptoms (3 items), examination findings (4 items), and electrophysiology (2 items) (Murphy et al., 2011; Sadjadi et al., 2014). The CMT Examination Score version 2 (CMTESv2) and Rasch modified version (CMTESv2-R) are subscores of the CMTNSv2 comprising 7 items from the patients’ symptoms and examination findings. The psychometrics of the Rasch weighting have only been performed in CMT1A patients (Sadjadi et al., 2014) and hence we used CMTESv2 as our primary clinical outcome measure for this study. The majority of assessment visits did not include neurophysiological studies and only the CMTESv2 and CMTESv2-R were obtained during these visits. Therefore, in order to maximise the sample size during the statistical analysis of the longitudinal data, the CMTESv2 and CMTESv2-R were primarily analysed; a similar analysis approach was also used in a CMT1A natural history study (Fridman et al., 2020). We also analysed data obtained from the CMT Pediatric Scale (CMTPedS), a well-validated composite score which assesses strength, hand dexterity, sensation, gait, balance, power and endurance in children with CMT from the age of 3 years (Burns et al., 2012). For each of these scales a higher score indicates a higher level of impairment. Clinical investigators at each site received training in the administration of the clinical outcome measures and were certified clinical investigators prior to use.

Statistical analysis

Data were analysed mostly on an available basis. The chi-squared goodness-of-fit test (skewness, kurtosis, median, mean) was used to ascertain the distribution of data. The baseline demographics and characteristics, clinical data from the history and examination and the physical disability were analysed using descriptive statistics. Throughout the manuscript values
describing continuous data represent the mean ± standard deviation (SD). Correlations between categorical data at baseline were assessed with chi-squared (χ²) or Fisher’s exact test as appropriate. Correlations between the CMTESv2 and the disease duration as calculated at the baseline visit were assessed with two-tailed, Spearman’s rank correlation coefficient. The longitudinal responsiveness of the CMTESv2, CMTESv2-R and CMTPedS was quantified as the standardised response mean (SRM = mean change / SD change). SRM values of 0.20-0.49, 0.50-0.79, and ≥0.80 correspond to small, moderate, and large responsiveness, respectively as suggested originally (Cohen, 1988). A two-tailed, paired Student’s t-test was used to compare longitudinal changes in the CMTESv2, CMTESv2-R and CMTPedS. A p-value (p) of 0.05 or less was considered statistically significant.

**Data availability**

The data that support the findings of this study are available from the corresponding author, upon reasonable request. The data are not publicly available since they contain information that could compromise the privacy of research participants.
Results

A total of 225 patients with MFN2 variants were recruited in 19 Inherited Neuropathy Consortium (INC) centres in the US, UK, Italy and Australia (100 males : 125 females). Eighty-seven of these were children under the age of 20 years (43 males : 44 females) and the average age at enrolment for the entire cohort was 31.30yrs ± 19.88yrs (standard deviation; SD). The age distribution was considered as a single mode with an arbitrary cut-off at the age of 20 years, rather than bimodal, as there are no distinct paediatric and adult presentations in CMT2A.

We identified 179 patients from 133 families with dominant pathogenic (ACMG class 5) or likely pathogenic (class 4) MFN2 variants (AD-CMT2A; Supplementary table 3), and 17 patients from 13 families with AR-CMT2A (Supplementary table 5); 13 out of 17 patients with AR-CMT2A harboured homozygous variants or compound heterozygous variants proven to be in trans phase. 29 patients from 23 families with VUSs are also reported (Supplementary table 4). For both the cross-sectional genotype-phenotype and longitudinal studies, only cases with pathogenic or likely pathogenic variants were taken into account.

Disease presentation and variant topology

The majority of patients with AD-CMT2A and AR-CMT2A first noticed symptoms in the first two decades of life, usually walking or balance difficulties. For both modes of inheritance, the correlation of the average age of onset with the genotype (amino acid position) is illustrated in Fig.1 with variants at the majority of amino acid positions being associated with symptom onset at or before the age of 15 years. The majority of these variants reside in the dynamin-GTPase domain even after standardising for the size of the domain. Although most of these are associated with early onset disease, there is considerable phenotypic heterogeneity within the domain with respect to the age of onset even for variants in adjacent positions. Furthermore, variants at specific amino acid positions that are usually associated with early
onset disease (p.Arg94, p.Arg104, p.Leu248, p.Arg364, p.Trp740) also seem to be associated with a tight time window during which the disease manifests. Patients with AR-CMT2A almost always had disease onset and first symptoms in childhood with an average age of onset of 8.06yrs ± 10.92yrs (SD); one exceptional case stands out who despite careful questioning could not time the onset of his symptoms before his late 40s. His brother who also carries the same variants in *trans* suffers from early onset AR-CMT2A and both suffer from a moderate burden of disease in their 6th decade of life (AR8 pedigree in Supplementary Table 5; Tomaselli et al., 2016).

In this cohort, and in line with previous publications, mutations in certain amino acid positions are always pathogenic, with no evidence of reduced penetrance or variable expressivity (range of phenotypic expression) despite different amino acid substitutions. Examples of different missense changes being observed at the same conserved amino acid position in our cohort include p.Arg94Trp/Gln, p.Arg104Glu/Trp, p.Ser249Thr/Cys and p.Trp740Ser/Arg. On the contrary, mutations in other amino acid positions are associated with phenotypic heterogeneity (early vs later onset of disease) dependent on the amino acid substitution at the same position. An example from our cohort is the group of 13 patients carrying the known p.Arg364Trp variant which is associated with early onset and severe disease and a single proband carrying the p.Arg364Gln variant who first presented with symptoms at the age of 32 years and had a recorded CMTESv2 of 8 at the age of 55 years, consistent with mild disease.

**Genotype-phenotype study**

The clinical characteristics of patients (Table 1a and 1b) were captured at their baseline visit and were analysed after classifying by inheritance pattern, age of onset, variant topology and the biological effect of variants. Inheritance pattern was defined as AD-CMT2A vs AR-
CMT2A and age of onset defined as childhood onset if disease presentation occurred between 1-20 years vs adult onset if the presentation occurred > 21 years (analysed in AD-CMT2A patients). Variant topology was defined as variants within the dynamin-GTPase domain vs those outside the domain and the biological effect of variants was defined as those variants shown to cause mitochondrial hypofusion in non-human disease models (p.Arg94Gln, p.Thr105Met) vs variants shown to cause mitochondrial hyperfusion (p.Leu76Pro, p.Arg364Trp).

Comparing AD-CMT2A to AR-CMT2A (Table 1a), both groups had a similar age of onset of symptoms (average 11.61yrs ± 15.07yrs (SD) vs average 8.06yrs ± 10.92yrs (SD), two-tailed Mann-Whitney U value 1291.5, p = 0.472) and at the baseline assessment had no significant difference in their disease duration (average 19.98yrs ± 14.84yrs (SD) vs average 25.35yrs ± 12.36yrs (SD), two-tailed Mann-Whitney U value 1057.5, p = 0.069). Disease duration was calculated from the age at the baseline visit minus the age of onset of symptoms. A minority of patients in both groups had delayed walking milestones (walked after the 15th month) but there was no significant difference between this percentage in the two groups (15% vs 30%, Fisher’s exact test p = 0.196). Despite a similar disease duration between the two groups, a significantly higher proportion of patients with AD-CMT2A had foot deformities (79% vs 53%, Fisher’s exact test p = 0.047), but a significantly higher proportion of patients with AR-CMT2A were using ankle-foot orthoses (63% vs 93%, χ² test p = 0.017) and had undergone foot surgery (25% vs 53%, Fisher’s exact test p = 0.032). Overall, patients with AR-CMT2A had a significantly higher mean CMTESv2 at baseline (10.75 ± 6.90 (SD) vs 14.57 ± 6.07 (SD), two-tailed Mann-Whitney U value 708.5, p = 0.028), as well as a higher mean CMTESv2-R indicating a higher burden of accrued disability over the same time-frame. Both groups had a moderate mean CMTPedS score (AD-CMT2A mean 26.45 ± 10.26 (SD) and AR-CMT2A mean 27.00 ± 11.31 (SD)) but with no significant difference between them. There
were no statistically significant differences between the two groups in the use of walking aids (29% vs 47%) or wheelchair (26% vs 20%), dexterity difficulties (64% vs 79%), optic nerve atrophy (7% vs 20%), hearing loss (both 7%) or scoliosis (12% vs 29%).

Of the 179 patients with pathogenic or likely pathogenic autosomal dominant variants (AD-CMT2A), 170 had available data on age of onset and the majority had onset in childhood (n=144; Table 1a). In both groups of patients, a minority had delayed walking milestones but there was no significant difference in this percentage between the two groups (14% vs 13%, Fisher’s exact test p = 1). Despite similar disease duration between childhood and adult-onset AD-CMT2A at the baseline visit (average 20.86yrs ± 15.56yrs (SD) vs average 15.15yrs ± 8.67yrs (SD); two-tailed Mann-Whitney U value 1592.5, p = 0.226), there was a significantly higher percentage of patients with childhood onset AD-CMT2A compared to adult onset AD-CMT2A, using ankle-foot orthoses (68% vs 40%, χ² test p = 0.007), wheelchair-dependent (31% vs 4%, χ² test p = 0.008) and with impaired dexterity (defined as either difficulties with cutlery or difficulty with buttons or both; 70% vs 46%, χ² test p = 0.021). Furthermore, patients with childhood-onset AD-CMT2A had a significantly higher mean of CMTESv2 (12.06 ± 6.82 (SD) vs 6.50 ± 3.42 (SD), two-tailed Mann-Whitney U value 787, p < 0.001), CMTESv2-R, CMTNSv2 (17.20 ± 9.79 (SD) vs 8.47 ± 4.93 (SD), two-tailed Mann-Whitney U value 249.5, p = 0.001) and CMTNSv2-R over a similar disease duration to adult-onset AD-CMT2A. Patients with childhood onset AD-CMT2A had a moderate mean CMTPedS score of 27.31 ± 9.57 (SD) at baseline. There was no significant difference in the prevalence of foot deformities between the two groups (80% vs 76%), the use of walking aids (29% vs 35%), the prevalence of previous foot surgery (29% vs 12%), hearing loss (6% vs 12%) or scoliosis (12% vs 13%).

In an analysis of AD-CMT2A cases with variants residing within the dynamin-GTPase domain (a.a. positions 93-342) versus AD-CMT2A cases with variants outside the domain (Table 1b), there were no significant differences in the age of onset of symptoms, baseline
mean CMTESv2, CMTESv2-R, CMTNSv2, CMTNSv2-R and CMTPedS scores and both groups had similar disease duration periods. Nonetheless, a slightly higher proportion of patients with variants in the dynamin-GTPase domain used ankle-foot orthoses (70% vs 53%, $\chi^2$ test $p = 0.026$), complained of dexterity difficulties (70% vs 55%, $\chi^2$ test $p = 0.047$) and developed scoliosis (16% vs 6%, $\chi^2$ test $p = 0.049$).

Analysis of patients harbouring variants that have been shown to cause mitochondrial hypofusion (p.Arg94Gln, p.Thr105Met) or hyperfusion (p.Leu76Pro, p.Arg364Trp), showed no significant difference in the age of onset, disease duration, mean CMTESv2, CMTESv2-R, CMTNSv2, CMTNSv2-R and CMTPedS scores (Table 1b).

In our cohort, optic nerve atrophy was reported in a total of 15 patients: 12 patients with AD-CMT2A (out of a total of 168 who had documented optic nerve assessment, 7.1%) and three patients with AR-CMT2A (out of a total of 15 who had documented optic nerve assessment, 20%). Despite the higher percentage in the AR-CMT2A group, this was not statistically significant (Fisher’s exact test $p = 0.111$). Furthermore, in patients with AD-CMT2A, optic atrophy was most frequently involved with variants at the amino acid position p.Arg104 (n=4), followed by p.Arg364Trp (n=2) and p.Leu248 (n=2). None of the category groups of age of onset of symptoms (child vs adult onset AD-CMT2A), variant topology (GTPase domain variants vs non-GTPase domain variants) or the biological effect of certain variants on mitochondrial fusion dynamics (hypofusion vs hyperfusion variants) had statistically significant enrichment with cases of CMT2A and concurrent optic atrophy.

**Disease progression**

CMT2A is a progressive disease with regards to length-dependent weakness and sensory loss and a cross-sectional analysis of the baseline data of patients with AD-CMT2A shows a statistically significant correlation between disease duration and the CMTESv2 (Fig.
this illustrates a worsening CMTESv2 as the disease progresses (two-tailed Spearman’s \( \rho = 0.44, p < 0.001 \)). Correlation of the CMTESv2-R with disease duration showed similar results with a parallel linear regression coefficient line (not shown). It is important to also note that this correlation includes patients with variants that are known to be associated with early and late onset disease, and with varying degrees of pace of progression.

According to the current study and previously published studies, the amino acid positions p.Arg94, p.Arg364 and p.Trp740 are the three commonest residues for the occurrence of missense variants in \( MFN2 \) causing CMT2A. Patients with variants in the amino acid position p.Arg94, which is the commonest of the three, show a significant correlation of baseline CMTESv2 with disease duration (two-tailed Spearman’s \( \rho = 0.65, p < 0.001 \)) (Fig. 3A). A detailed assessment of the disease impact over time in these 29 patients with regards to the use of ankle-foot orthoses, walking aids and wheelchair use is presented in Fig. 4. Almost all patients require ankle-foot orthoses in the first two decades of life, with the majority of prescriptions given in childhood and of the seven patients requiring regular use of a wheelchair, this was before the age of 40 yrs in six. Patients carrying the p.Arg364Trp variant (Fig. 3B) also show a significant correlation between baseline CMTESv2 and disease duration (two-tailed Spearman’s \( \rho = 0.72, p = 0.005 \)). However, these patients have more severe disease early on and throughout the entirety of the disease compared to p.Arg94 as illustrated by the higher CMTESv2 scores and have a slightly faster pace of progression. On the contrary, patients with variants at the p.Trp740 position (Fig. 3C) have milder disease early on and throughout the entirety of the disease compared to p.Arg94 and p.Arg364 as illustrated by the lower CMTESv2 scores. Patients with variants at the p.Trp740 position also show a significant correlation between baseline CMTESv2 and disease duration (two-tailed Spearman’s \( \rho = 0.58, p = 0.011 \)).

Out of the 179 patients with AD-CMT2A, 92 patients had longitudinal data, of which 38 had 1-year follow-up data and 34 had 2-year follow-up data. Eight patients with AR-
CMT2A had longitudinal data, of which six had 1-year follow-up data, four had 2-year follow-up data and five had 4-year follow-up data. The longitudinal data from the CMTESv2, the weighted CMTESv2-R and the CMTPedS of all AD-CMT2A and AR-CMT2A patients was used to calculate the mean change over 1 and 2 years for each group respectively. Subsequently the aggregated data were used to calculate the standardised response mean (SRM) of the CMTESv2, CMTESv2-R and CMTPedS over 1 and 2 years (Table 2). The SRM is a metric that describes how sensitive a particular outcome is to change. In patients with AD-CMT2A, the CMTESv2 increased significantly over 1 year (mean change 0.84 ± 2.42 (SD), two-tailed paired t-test p = 0.039); the equivalent 1-year SRM at 0.35 showed small responsiveness. Surprisingly, the CMTESv2-R did not show a significant change over 1 year (mean change 0.63 ± 3.19 (SD), two-tailed paired t-test p = 0.230). Over 2 years, the CMTESv2 increased significantly (mean change 0.97 ± 1.77 (SD), two-tailed paired t-test p = 0.003), as did the CMTESv2-R (mean change 1.21 ± 2.52 (SD), two-tailed paired t-test p = 0.009), and their 2-year SRM values of 0.55 and 0.48 reflect moderate and small responsiveness, respectively. Analysis of the CMTPedS in all the paediatric AD-CMT2A and AR-CMT2A cases grouped together, showed that it increased significantly over 1 year (mean change 2.24 ± 3.09; two-tailed paired t-test p = 0.009) and over 2 years (mean change 4.00 ± 3.79; two-tailed paired t-test p = 0.031) with respective SRMs of 0.72 and 1.06. There was no significant change in the CMTESv2 or CMTESv2-R in the AR-CMT2A group and this is likely to be due to the small sample size of patients with available follow-up data from baseline.
Discussion

In this large international prospective study of 196 patients with CMT2A, the majority of pathogenic and likely pathogenic variants occur in the dynamin-GTPase domain of MFN2 which plays a central role in mitochondrial fusion (Westermann, 2010; Chandhok et al., 2018). In the gnomAD population database (Karczewski et al., 2020), of the 30 most frequent missense variants observed in MFN2, each with an allele count (AC) of greater than nine (range AC 9-1942), only three are situated in the dynamin-GTPase domain (p.Gly298Arg AC=606, p.Cys281Ser AC=92, p.Arg250Gln AC=62).

There are earlier reports associating nonsense MFN2 variants (E65X, R418X) in the heterozygous state with CMT2A and since such truncated transcripts are expected to undergo nonsense mediated decay, this would suggest that MFN2 may be intolerant of haploinsufficiency. However, evidence from the Mfn2 heterozygous knockout mice that do not express a phenotype (Chen et al., 2003) and more recent evidence from human models of the disease would argue against this. The mother of a proband with AR-CMT2A, who is a heterozygous carrier of the p.Glu308X variant and is unaffected by history and neurophysiology (EMG) at the age of 39, had RT-PCR transcript analysis from blood which indicated that the truncated transcript undergoes nonsense mediated decay (NMD) (Polke et al., 2011). The lack of a phenotype in her (at least until mid-adulthood) would suggest that MFN2 tolerates haploinsufficiency. An exception to this are nonsense MFN2 variants in the last exon, which are expected and have been shown to escape NMD (Kawarai et al., 2016) and in line with this, heterozygous carriers of the variants p.751X and p.752X develop an early onset severe CMT2A phenotype (Verhoeven et al., 2006; Feely et al., 2011; Kawarai et al., 2016). Ascertaining the pathogenicity of splice donor or acceptor site variants, which may cause exon skipping, is challenging. For these reasons and in the absence of transcriptomic and variant-specific functional data, nonsense, frameshift and splice donor and acceptor site...
variants residing in NMD-insensitive regions of the transcript have been classed as VUSs in this cohort.

Care should be taken when interpreting novel variants within specific domains and using cross-sectional correlations. For example, the difference both in the average age and range of disease onset between patients carrying a variant at the p.Arg94 amino acid position (average age of onset 4.7yrs ± 2.5yrs (SD)) and those carrying a variant at the adjacent p.Arg95 amino acid position (average age of onset 31.8yrs ± 23.6yrs (SD)) is significant. Similarly, patients carrying the p.Arg364Trp variant, often present with an early onset and progressive disease, in contrast to patients carrying the nearby p.Thr362Arg variant which usually presents with symptoms in adulthood and has a more indolent course. Specific MFN2 variants have been previously described to exhibit interfamilial variability with regards to age of onset, such as the p.Leu741Trp described in two unrelated families, one with average age of onset in the 3rd decade of life and the other with onset of disease in the 5th decade in most members (Dankwa et al., 2018; Lin et al., 2019). Other variants have been reported to show intrafamilial variability such as the p.Arg95Gly with variable clinical severity and significantly different age of onset of symptoms in different affected family members (Dankwa et al., 2019) and p.Leu146Phe with age of onset of disease ranging from childhood to late adulthood for different family members (Klein et al., 2011). A further family in our cohort carrying the dominant p.Ala100Ser variant exhibited intrafamilial heterogeneity, since both siblings had onset of disease in early adulthood and moderate CMTESv2 scores in their 40s-50s, whereas their mother had a late onset neuropathy with a CMTESv2 of 6 at the age of 81.

Our baseline genotype-phenotype correlations, illustrate how CMT2A is an early and severe form of CMT2 with the majority of patients having foot deformities, requiring ankle-foot orthoses and complaining of impaired dexterity at their first visit. At the first visit, the mean CMTESv2 score was 11.06 ± 6.90 (SD) (sample mean age 33.70 ± 19.42 (SD)), the mean
CMTNSv2 score was 15.68 ± 9.56 (SD) (sample mean age 36.00 ± 19.78 (SD)) and the mean CMTPedS score was 26.47 ± 10.17 (SD) (sample mean age 10.3 ± 4.36 (SD)), all scores indicative of a moderate burden of disability with CMT2A early in life. A considerable proportion require use of walking aids and are wheelchair-dependent at their first visit and this degree of physical impairment is not encountered in CMT1A (Reilly et al., 2011), CMT1B (Sanmaneechai et al., 2015) or CMTX1 (Panosyan et al., 2017). Interestingly, the majority of patients with CMT2A seem to achieve their gross motor developmental milestones normally as the majority walk at or before the 15th month, yet go on to develop early, and in most cases, severe CMT which progresses faster than other subtypes. This is in contrast to CMT1B, a demyelinating form of CMT, in which the vast majority of patients with an infantile onset show a delay in walking independently, beyond the 15th month (Sanmaneechai et al., 2015). Taking into account our comparison of baseline characteristics across a range of groups, a childhood onset of disease in AD-CMT2A seems to be the most reliable predictor of significant physical disability accrued and is independent of the disease duration.

The SRM is a popular effect size index used to estimate the responsiveness of outcome measures to clinical change. Based on data from two clinical trials of ascorbic acid in CMT1A (Pareyson et al., 2011; Lewis et al., 2013; Picosquito et al., 2015), the 1yr SRM of the CMTESv1 was 0.17 indicating minimal responsiveness. Natural history data from a large multicentre CMT1A cohort, also showed a minimally responsive CMTESv2 and CMTESv2-R over 2 years with SRMs of 0.11 and 0.17, respectively (Fridman et al., 2020). By comparison, CMT2A is a more rapidly progressive disease and this is reflected in a 1yr SRM of 0.35 for the CMTESv2 when used in AD-CMT2A. This means that a hypothetical CMT2A double blinded, randomised placebo-controlled trial, powered to detect a complete cessation in disease progression as measured by the CMTESv2 over a 12-month period with 80% power at p<0.05 significance, would require 131 individuals in each arm. For a treatment trial with a duration
of 24 months and using a 2-year SRM of 0.55, the number of individuals needed in each arm would be 53. Complementing clinical assessment tools, biomarkers such as MRI-quantified intramuscular fat accumulation at calf-level are showing promise as a sensitive outcome measure with two studies showing a highly responsive 1-year SRM of 0.83 and 1.04 in CMT1A (Morrow et al., 2016, 2018). Given that CMT2A is a more progressive disease, it is probable that MRI-quantified intramuscular fat accumulation in CMT2A will prove to be an even more sensitive outcome measure in CMT2A and this is currently being investigated. Surprisingly, the Rasch-weighted CMTESv2-R was not sensitive to change at 1 year and had a lower SRM compared to CMTESv2 at a 2yr follow-up. This perceived insensitivity of the CMTESv2-R may have arisen because the psychometrics of the Rasch weighting were performed using CMT1A data which is a more slowly progressive disease compared to CMT2A (Sadjadi et al., 2014). Despite a small sample size of our paediatric CMT2A cohort, analysis of the longitudinal CMTPedS data showed a significant mean change over 1 and 2 years, corresponding to a respective moderate (1yr SRM 0.72) and large responsiveness (2yr SRM 1.06) of this clinical outcome measure. Furthermore, some severe paediatric CMT2A cases reach the ceiling of the outcome score by their early teens and the subsequent plateauing of the clinical scores would give a false impression of disease stabilisation. Ultimately, this may make the overall rate of progression seem smaller than it actually is and therefore, a larger paediatric CMT2A cohort is needed to delineate more accurately the progression of CMT2A in childhood.

With the use of next-generation sequencing panels now commonplace, more patients with CMT receive a genetic result than ever before. This has also led to the identification of large numbers of novel variants in MFN2, the significance of which are unknown. Large CMT2A cohort studies such as ours are valuable to help investigators curate variants. Moreover, with genetic therapies in development and clinical trials on the horizon, we need to have responsive clinical outcome measures in order to be trial-ready. This study provides
evidence that CMTESv2 is a responsive outcome measure for a 2 year clinical trial which together with the concurrent development of responsive biomarkers means we are in a good position to perform clinical trials as candidate therapies become available for CMT2A.
Acknowledgements

This study makes use of data generated by the DECIPHER community. A full list of centres who contributed to the generation of the data is available from http://decipher.sanger.ac.uk and via email from decipher@sanger.ac.uk. Funding for the project was provided by the Wellcome Trust.

Funding

MP is funded by the NIH and the Inherited Neuropathy Consortium (INC; grant U54NS065712) as well as the BMA (Vera Down grant). MMR is grateful to the National Institutes of Neurological Diseases and Stroke and office of Rare Diseases (U54NS065712) and to Muscular Dystrophy Association (MDA510281) for their support. This research was also supported by the National Institute for Health Research University College London Hospitals Biomedical Research Centre. The U54 award also receives funding from the MDA and CMTA. IM, CP and DP are members of the European Reference Network for Rare Neuromuscular Diseases (ERN EURO-NMD). RH was supported by the European Research Council (309548), the Wellcome Investigator Award (109915/Z/15/Z) the Medical Research Council (UK) (MR/N025431/1), the Wellcome Trust Pathfinder Scheme (201064/Z/16/Z) and the Newton Fund (UK/Turkey, MR/N027302/1).

Competing interests

The authors report no competing interests.

Supplementary material

Supplementary tables 1-6 included.
Appendix – Other Inherited Neuropathy Consortium (INC) working group members not in author list

Silvia Baratta, Paula Bray, Daniela Calabrese, Kayla Cornett, Gabrielle Donlevy, Katy Eichinger, Maria Foscan, Silvia Genitrini, Natalie Rose Grant, Tara Jones, Diana Lee, Brett A. McCray, Stefania Magri, Manoj Menezes, Krista Mullen, Tina Nanji, Sara Nuzzo, Emanuela Pagliano, Roy Poh, Eun Park, Saba Sadaf, Paola Saveri, Carly Siskind, Janet E. Sowden, Charlotte J. Sumner, Simone Thomas
References


Burns J, Ouvrier R, Estilow T, Shy R, Laurá M, Pallant JF, et al. Validation of the Charcot-


Murphy SM, Herrmann DN, McDermott MP, Scherer SS, Shy ME, Reilly MM, et al. 


Figure 1. Graph displaying the average age of onset of symptoms at each amino acid position for all the pathogenic and likely pathogenic variants with dominant inheritance and AR-CMT2A cases (far right of graph) identified in this study. For each amino acid position, the cumulative variant count irrespective of the amino acid substitution, is also listed (n; for example ‘p.Arg94Trp, n=31’ contains all 31 cases of p.Arg94Trp, p.Arg94Gln, p.Arg94Gly and p.Arg94Leu). On the X-axis, the variants are equidistant from one another for graphical purposes; the distant between the variants displayed is not to scale. However, the primary structure of the MFN2 protein has been drawn below the graph in a skewed fashion to account for this and highlights which functional domains the variants reside in. The average age of onset of symptoms for each amino acid position is displayed with a horizontal red bar and the standard deviation bars, indicative of the spread in age of onset for each position, are also shown for all amino acid positions with variant counts of 2 or more.
Figure 2. Correlation of CMTESv2 and disease duration as surrogate evidence of disease progression. The disease duration was calculated by subtracting the age of onset from the age at assessment at the baseline visit. The dashed line represents the linear regression coefficient ($R^2$). 148 patients with a dominant pathogenic or likely pathogenic $MFN2$ variant had CMTESv2 data at their baseline visit; the correlation between CMTESv2 and disease duration is statistically significant (two-tailed Spearman’s $\rho 0.44$, $p < 0.001$).

Figure 3. Correlation of CMTESv2 and disease duration in the three commonest missense variants causing CMT2A. The disease duration was calculated by subtracting the age of onset from the age at assessment at the baseline visit. The dashed line represents the linear regression coefficient ($R^2$). (A) 24 patients with a heterozygous mutation at the p.Arg94-- amino acid position had CMTESv2 data at their baseline visit. This subgroup includes patients with the variants p.Arg94Trp, p.Arg94Gln, p.Arg94Gly and p.Arg94Leu; the correlation between CMTESv2 and disease duration in this group is statistically significant (two-tailed Spearman’s $\rho 0.65$, $p < 0.001$). B) 13 patients carrying the heterozygous variant p.Arg364Trp had CMTESv2 data at their baseline visit and the correlation between CMTESv2 and disease duration is statistically significant (two-tailed Spearman’s $\rho 0.72$, $p = 0.0054$). C) 18 patients with a heterozygous variant at the p.Trp740-- amino acid position had CMTESv2 data at their baseline visit and the correlation between CMTESv2 and disease duration is statistically significant (two-tailed Spearman’s $\rho 0.58$, $p = 0.011$).
Figure 4. Genotype-physical impact correlation in 29 patients with CMT2A due to pathogenic and likely pathogenic variants at the p.Arg94-- amino acid position. Each horizontal set of data-points plotted over time (age in years) corresponds to one of the 29 patients (1-29 on the Y-axis). Patients 1-16 carry the p.Arg94Trp variant, patient 17 carries the p.Arg94Leu variant, patients 18 and 19 carry the p.Arg94Gly variant and patients 20-29 carry the p.Arg94Gln variant. In each set of data-points, the red square represents the age at which the patient was first seen and enrolled in the study (baseline visit): the dashed line to its left represents retrospective information gathered by history and the solid line to its right represents prospective information gathered during the entirety of the study. The green diamond represents the last age with available clinical data. The majority of the patients have only been seen once and hence the red square and green diamond coincide. The blue circle represents the age of onset of symptoms in the patient, the yellow triangle represents the age at which ankle-foot orthoses were needed and prescribed, the purple star represents the age at which walking aids (stick, walker) were required and the blue cross represents the age at which the patient reverted to the regular use of a wheelchair.