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# The role of combined SNV and CNV burden in patients with distal symmetric polyneuropathy

Davut Pehlivan, MD<sup>1,2</sup>, Christine R. Beck, PhD<sup>1</sup>, Yuji Okamoto, MD, PhD<sup>1</sup>, Tamar Harel, MD, PhD<sup>1</sup>, Zeynep H.C. Akdemir, PhD<sup>1</sup>, Shalini N. Jhangiani, MS<sup>3</sup>, Marjorie A. Withers, BS<sup>1</sup>, Meryem Tuba Goksungur, MD<sup>4</sup>, Claudia M.B. Carvalho, PhD<sup>1</sup>, Dirk Czesnik, MD<sup>5</sup>, Claudia Gonzaga-Jauregui, PhD<sup>1</sup>, Wojciech Wiszniewski, MD, PhD<sup>1</sup>, Donna M. Muzny, MS<sup>3</sup>, Richard A. Gibbs, PhD<sup>3</sup>, Bernd Rautenstrauss, PhD<sup>6,10</sup>, Michael W. Sereda, MD<sup>5,7</sup>, and James R. Lupski, MD, PhD, DSc(hon)<sup>1,3,8,9</sup>

<sup>1</sup>Department of Molecular and Human Genetics, Baylor College of Medicine, Houston, Texas, USA

<sup>2</sup>Section of Pediatric Neurology, Texas Children's Hospital, Houston, Texas, USA

<sup>3</sup>Human Genome Sequencing Center, Baylor College of Medicine, Houston, Texas, USA

<sup>4</sup>Department of Neurology, Istanbul University, Istanbul Medical Faculty, Istanbul, Turkey

<sup>5</sup>Department of Clinical Neurophysiology, University Medical Center Göttingen, Göttingen, Germany

<sup>6</sup>Medizinisch Genetisches Zentrum, Munich, Germany

<sup>7</sup>Max-Planck Institute of Experimental Medicine, Göttingen, Germany

<sup>8</sup>Department of Pediatrics, Baylor College of Medicine, Houston, Texas, USA

<sup>9</sup>Texas Children's Hospital, Houston, Texas, USA

# Abstract

**Purpose**—Charcot-Marie-Tooth (CMT) disease is a heterogeneous group of genetic disorders of the peripheral nervous system. Copy-number variants (CNVs) contribute significantly to CMT, as duplication of *PMP22* underlies the majority of CMT1 cases. We hypothesized that CNVs and/or single-nucleotide variants (SNVs) might exist in patients with CMT with an unknown molecular genetic etiology.

**Methods**—Two hundred patients with CMT, negative for both SNV mutations in several CMT genes and for CNVs involving *PMP22*, were screened for CNVs by high-resolution

SUPPLEMENTARY MATERIAL

Supplementary material is linked to the online version of the paper at http://www.nature.com/gim DISCLOSURE

Correspondence: James R. Lupski (jlupski@bcm.edu).

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oligonucleotide array comparative genomic hybridization. Whole-exome sequencing was conducted on individuals with rare, potentially pathogenic CNVs.

**Results**—Putatively causative CNVs were identified in five subjects (~2.5%); four of the five map to known neuropathy genes. Breakpoint sequencing revealed *Alu-Alu*-mediated junctions as a predominant contributor. Exome sequencing identified *MFN2* SNVs in two of the individuals.

**Conclusion**—Neuropathy-associated CNV outside of the *PMP22* locus is rare in CMT. Nevertheless, there is potential clinical utility in testing for CNVs and exome sequencing in CMT cases negative for the CMT1A duplication. These findings suggest that complex phenotypes including neuropathy can potentially be caused by a combination of SNVs and CNVs affecting more than one disease-associated locus and contributing to a mutational burden.

#### Keywords

array comparative genomic hybridization; Charcot-Marie-Tooth disease; exome sequencing; MPZ; PRICKLE1

## INTRODUCTION

Charcot-Marie-Tooth (CMT) disease is a clinically and genetically heterogeneous group of disorders of the peripheral nervous system, with a prevalence of ~1 in 2,500, making it among the most common inherited neurologic disorders. CMT is characterized by distal weakness, muscular atrophy, foot deformities, gait disturbance, and loss of tendon reflexes accompanied by a relatively sharply demarcated loss of sensation presenting as a distal symmetric polyneuropathy.<sup>1</sup> The classification was initially developed according to neuropathological and electrophysiological observations, with CMT type 1 and CMT type 2 representing demyelinating and axonal neuropathies, respectively. Gene discoveries within the past two decades have challenged this simplistic classification of CMT. Approximately 70 genes and 10 additional genomic loci have been linked to variants that can cause a CMT-like peripheral neuropathy, making it one of the most genetically diverse neurological phenotypes. All forms of Mendelian inheritance patterns (autosomal dominant, X-linked, and autosomal recessive) have been observed in CMT.<sup>2</sup>

A recurrent 1.4-Mb duplication of a chromosomal segment involving peripheral myelin protein 22 (*PMP22*) is mediated by nonallelic homologous recombination and is the major molecular genetic subgroup of CMT, accounting for ~70–80% of CMT1 and ~50% of all CMT cases (OMIM 118220).<sup>3,4</sup> The reciprocal 1.4-Mb deletion underlies the more clinically mild hereditary neuropathy with liability to pressure palsies (OMIM 162500).<sup>5</sup> Diagnostic testing identified a *PMP22* duplication or deletion in >78% of ~17,000 patients with clinical neuropathy for whom a molecular diagnosis was concluded. Either copy-number variants (CNVs) or single-nucleotide variants (SNVs) involving four genes—*PMP22, GJB1 (Cx32), MPZ*, or *MFN2*—accounted for ~95% of disease-causative mutations.<sup>6</sup>

We previously reported nonrecurrent, rare, unique CNVs involving the *PMP22* region in patients with CMT1A or hereditary neuropathy with liability to pressure palsies.<sup>7</sup> Underlying molecular mechanisms for rare, nonrecurrent *PMP22* CNVs differ from the recurrent events mediated by nonallelic low copy repeats acting as ectopic homologous

recombination substrates.<sup>8</sup> These nonrecurrent rearrangements are potentially mediated by replicative mechanisms such as fork-stalling, template switching, microhomology-mediated, break-induced replication,<sup>9,10</sup> *Alu-Alu*-mediated recombination events,<sup>11,12</sup> and nonhomologous end joining. CNVs involving CMT genes other than *PMP22* have rarely been identified. We previously reported three different deletion CNVs in *Cx32* in patients with X-linked CMT; breakpoint junction studies showed either nonhomologous end joining or *Alu-Alu*-mediated events underlying CNV formation.<sup>13</sup> We also independently described a homozygous intragenic duplication CNV in *NDRG1* leading to autosomal recessive CMT4D in our current CMT cohort.<sup>14</sup> In addition, two different CNVs involving amplification of *MPZ* have been identified.<sup>15,16</sup>

With recent advances in technologies such as array comparative genomic hybridization (aCGH) and genomic sequencing, it is now appreciated that CNVs may encompass more nucleotide variation than SNVs when comparing a given genome with the haploid reference.<sup>17</sup> We investigated the frequency of CNVs involving non-PMP22 CMT genes through the study of a large cohort of subjects with molecularly undefined CMT. We also sought to identify possible novel candidate genes by examining linkage regions. Finally, we investigated the hypothesis that a combination of CNV and a rare SNV mutation can contribute to patient phenotypes by performing whole-exome sequencing (WES) in the subjects in whom a candidate CMT-associated CNV was identified. Five of 200 individuals with phenotypically characterized CMT that was molecularly undefined contained CNVs in genes potentially responsible for their neuropathy, generating an overall rate of ~2.5%. One of these genes was previously unlinked (KIF24) to peripheral neuropathy, and another had been associated with seizures and a sensory neuropathy, but not a motor neuropathy (PRICKLE1). WES elucidated MFN2 variants in two individuals in whom a CNV in a CMT gene was found. These findings suggest the possibility that either SNV or CNV mutations in a CMT gene, and potentially a combination of both SNVs and CNVs in multiple CMT genes through a mutational burden mechanism,<sup>18</sup> may contribute to complex neurological phenotypes.

## MATERIALS AND METHODS

#### Subjects

This study was approved by the institutional review board at Baylor College of Medicine. Two hundred DNA samples from unrelated subjects with CMT were studied. Informed consent was obtained and peripheral blood then was drawn from patients and family members. DNA was isolated either directly from blood or transformed lymphoblastoid cell lines using a Gentra Puregene purification kit (Qiagen, Valencia, CA). Patients were screened previously for *PMP22* CNVs. Most patients were screened for point mutations or SNVs in many of the known CMT genes, including *PMP22, MPZ*, and *GJB1*.

#### Array comparative genomic hybridization

We designed a high-density oligonucleotide aCGH microarray ( $8 \times 60$ K format) that covered 67 neuropathy genes (±10-kb flanking regions) and 10 known linkage regions (Supplementary Table S1 online) using the Agilent eArray system (http://

earray.chem.agilent.com/earray). This targeted aCGH design for investigating CNVs in CMT has an average genomic resolution of ~1 probe/200 bp for the 67 gene regions and ~1 probe/5 kb for 10 linkage regions. The design did not allow evaluation far upstream of *MPZ* (in BAB7807 we detected a CNV larger than our array coverage); therefore we designed a specific tiling-path  $4 \times 44$ K aCGH spanning 1 Mb to further define the *MPZ* CNV. Arrays were conducted according to the Agilent oligonucleotide aCGH protocol (version 6.0) and previously described methods.<sup>19</sup> Gender-matched male (NA10851) and female (NA15510) control DNAs were used for the hybridization procedure (obtained from Coriell Cell Repositories; http://ccr.coriell.org).

#### **Breakpoint junction studies**

Based on aCGH data, we designed primers flanking the boundaries of each deleted/ duplicated segment. For duplicated segments, primers were embedded within the duplicated segment in an outward-facing orientation (tandem duplication), and for deleted segments primers were placed outside the deleted segment in an inward-facing direction. Both longrange and conventional polymerase chain reaction (PCR) amplification methods were used to detect the breakpoint junctions.

Primers for patient-specific breakpoint junction amplification are presented in Supplementary Table S2 online. Amplicons from patients BAB392, BAB1219, and BAB7807 were further analyzed by primer walking. We aligned breakpoint sequences manually or using MultAlin (http://multalin.toulouse.inra.fr/multalin/).<sup>20</sup>

#### **Reverse-transcription PCR and sequencing**

Lymphoblastoid cells from BAB1415 and BAB392 were cultured using standard practices. RNA was extracted using a Qiagen RNeasy Plus miniprep kit. RNA was quantified via NanoDrop, and integrity was determined by agarose electrophoresis.

Reverse transcription (RT) was carried out using qScript cDNA super mix (Quanta Biosciences, Gaithersburg, MD) with 0.5  $\mu$ g of RNA. PCR amplification of RT products was performed by TaKaRa LA taq using the primers listed in Supplementary Table S2 online. Amplification was performed for 32 cycles, and resultant amplicons were sequenced to ensure they were on target. Multiple RNA and complementary DNA reactions were used for each individual; results were consistent across three separate replicates.

#### Exome sequencing

We applied WES to samples BAB7807, BAB1219, BAB392, BAB1415, and BAB3724 at the Baylor College of Medicine Human Genome Sequencing Center through the Baylor-Hopkins Center for Mendelian Genomics research initiative. We used publicly available databases such as the 1000 Genomes Project (http://www.1000genomes.org) and other large-scale exome sequencing projects, including the Exome variant server for the National Heart, Lung, and Blood Institute's GO Exome Sequencing Project, Seattle, WA (http://evs.gs.washington.edu/EVS/), and the Atherosclerosis Risk in Communities Study Database (http://drupal.cscc.unc.edu/aric/) as well as our in-house-generated exomes (from ~4,000 individuals) at Baylor College of Medicine for analyses of candidate variants/mutations. All

experiments and analyses were performed according to previously described methods.<sup>21</sup> SNVs and insertion/deletion mutations in known neuropathy genes were included in Table 1 if they had both an Exome Sequencing Project minor allele frequency 1% and an internal database minor allele frequency 1%. Sanger sequencing verified all the candidate mutations presented in Table 1.

We then determined whether we could detect potential CNVs from WES using the program CoNIFER.<sup>22</sup> This program predicts duplications and deletions via differences in WES read depth that deviate from a large number of genomes ascertained using the same exome capture and sequencing methodology.

### RESULTS

#### **Clinical information**

Patient BAB7807 presented at about 23 years of age with progressive weakness, myoclonus, fatigue, gait disturbance, and paraesthesias. Physical examination revealed distal accentuated polyneuropathy with mild atrophy of the hand and foot muscles, absent ankle deep tendon reflex, hypoesthesia of the hands and feet, and reduced heat and cold sensation in the distal extremities. Cranial nerves were grossly intact. Myography revealed no positive sharp waves and showed mild chronic denervations of the right tibialis muscle anterior. Nerve conduction velocities were consistent with demyelinating neuropathy (Table 1). Nerve biopsy was suggestive of a mild chronic degenerative demyelinating neuropathy, with no signs of vasculitis or neuritis. Muscle biopsy showed mild neurogenic changes without signs of myopathy or myositis.

BAB1219 is a 35-year-old woman. She presented in elementary school with ankle weakness and distal lower limb paresthesias, in addition to leg pain/cramping after walking long distances. She also reported mild hand weakness. The patient was one of five siblings, and family history was negative for neuromuscular complaints. Physical examination showed bilateral distal muscle weakness in her lower extremities and symmetrically diminished sensation in both distal upper and lower extremities. Electromyography study showed the presence of severe, generalized distally prominent, mixed sensory and motor neuropathy with prominent demyelinating features.

BAB392 was clinically evaluated at 4 years of age because of weakness in his legs. The patient's right foot was pointing inward at birth, and his left foot soon followed. Motor development was consistent with his age; he walked at ~9 months but tended to fall frequently. Generalized tonic-clonic seizures developed between the ages of 7 and 13 years and were treated with phenobarbital. He also had mild intellectual disability. Follow-up evaluation at 17 years of age revealed that the patient was wheelchair-bound with prominent muscle atrophy in the hands and distal lower extremities. Deep tendon reflexes were absent throughout. Electromyography studies showed neuropathic changes of the muscles, most marked distally.

BAB1415 had weakness in her legs starting at around 2 years 9 months of age. Hand weakness was progressive and became apparent 2 years later. Physical examination at age 9

years showed bilateral claw hands. Muscle strength was 4/5 in the upper and lower extremity muscle groups. Deep tendon reflexes were normal and symmetric in the upper extremities and absent in the lower extremities. Sensation to light touch was mildly decreased in the distal lower extremities. Gait was abnormal and characterized by bilateral foot drop. Muscle biopsy at 3.5 years was consistent with a chronic demyelinating neuropathy.

Nerve conduction velocity study results for all five patients in this study were conducted (Table 1). Study results for patient BAB3724 have been published.<sup>14</sup>

#### High-resolution aCGH analyses

To examine for potential disease-associated CNV in non-CMT1A patients, we performed whole-genome-targeted aCGH (see Materials and Methods) in 200 families with CMT who were known to be negative for the most common CMT-causative mutations. Our approach revealed five novel CNVs (in four known CMT genes and one candidate CMT gene) ranging in size from 4.2 to 30 kb (Figure 1). An intragenic homozygous exon 6 to 8 duplication in *NDRG1* was previously published in a family with recessive CMT.<sup>14</sup> We now report the characterization of CNVs in the remaining four subjects with neuropathy.

In two subjects we detected CNVs of *MPZ* consisting of a 20.2-kb deletion and a 4.2-kb duplication. Subject BAB7807 had a 20.2-kb deletion involving the first coding exon of *MPZ* and the first three exons of the *SDHC* gene (Figure 2a,c). Our specific aCGH design allowed us to evaluate the upstream deletion region and to detect the breakpoint junction interval (Figure 2b). Subject BAB1219 had a 4.2-kb duplication involving all coding exons, the 5' untranslated region and a portion of the 3' untranslated region of *MPZ* (Figure 2d,f). This *MPZ* duplication was independently confirmed by multiplex ligation-dependent probe amplification (data not shown). Patient BAB392 had an intragenic 29.7-kb deletion involving the first exon of *PRICKLE1* (RefSeq gene ID NM\_153026.2) (Figure 3a,e). Individual BAB1415 has a 16.5-kb deletion involving exons 5 and 6 of *KIF24* (RefSeq gene ID NM\_194313.2) (Figure 4a,d).

#### Breakpoint junction sequencing

Breakpoint junction sequencing was performed in all cases to fine-map CNVs and to gain potential insights into the underlying mechanisms for rearrangement formation. PCR amplification of junctions was achieved for all five CNVs. Patient BAB7807 harbors a deletion with *Alu* elements at both proximal and distal breakpoints (*FLAM\_C* within intron one of *MPZ* and *AluSz6-D* at distal junction). Although the lengths of the repeats differ (137 bp for *FLAM\_C* and 312 bp for the *AluSz6-D*), the two short interspersed elements share 23 bp of perfect sequence identity at the breakpoint junction (Figure 2b). The two short interspersed elements are highly divergent, yet the junction creates a chimeric *FLAM/AluS* element.

BAB1219 contained a 4.2-kb tandem *MPZ* duplication involving all coding regions of the gene and the 5' untranslated region; the distal breakpoint lies 10 bp upstream of the transcription start site. Breakpoint analysis revealed a 2-bp microhomology (CC) at the junction, consistent with a microhomology-mediated mechanism such as fork stalling

template switching underlying this duplication.<sup>10</sup> No repetitive sequence was observed at either the proximal or the distal breakpoint junctions (Figure 2e).

Breakpoint amplification of BAB392 also revealed short interspersed elements at both proximal and distal regions of the deletion (*AluSp* is proximal and 304 bp in length, and *AluSx* is distal and 310 bp in length). The *Alu* elements shared 16 bp of microhomology, whereas overall they share only 78% sequence identity (Figure 3b). This junction creates a chimeric *Alu* element between the two *AluS* elements. As parental samples were available for BAB392, we also were able to determine that this deletion is de novo in the proband (Figure 3c).

The deletion of subject BAB1415 also was mediated by two *Alu* elements, an *AluYa5* (310 bp) at the proximal region and an *AluY* (306 bp) at the distal region. At the breakpoint junction, the two *Alu* elements shared 18 bp of perfect sequence identity, yet overall, they share 85% homology (Figure 4b).

In summary, breakpoint analysis showed that three of the four subjects presented here contained junctions that generate chimeric short interspersed elements. The three *Alu-Alu* CNVs contained between 16 and 22 bp of microhomology at the junction; the size of the deletions ranged from 16.5 to 29.7 kb. The last patient (BAB1219) contained a duplication that was not mediated by repetitive elements; there were 2 bp of microhomology at the junction.

#### Aberrant splicing and transcript levels

For both BAB392 and BAB1415, cultured lymphoblastoid cell lines enabled RNA studies to be performed. In BAB392, the first exon of *PRICKLE1* was deleted along with the region directly upstream of the gene. Therefore, we predicted that transcription of *PRICKLE1* could be negatively affected by this mutation. RT-PCR amplification of BAB392 RNA suggests decreased levels of *PRICKLE1* messenger RNA in relation to both BAB1415 and a control subject without CMT. This difference was seen for two different primer pairs (exon 2-1 forward to 5 reverse and a second exon 2-2 forward primer to 4 reverse) (Figure 3d), was consistent across three different RNA preparations and complementary DNA reactions. *GAPDH* and *TBP* amplification showed no change in transcript levels in RNAs from multiple patients and controls (data not shown).

For subject BAB1415, the CNV is predicted to alter RNA structure by skipping two *KIF24* exons (5 and 6). Independent RT-PCR reactions from this patient showed the predicted pattern, resulting in peaks on peaks at the end of exon 4 and the beginning of exon 7 (Figure 4c). Two control RNAs showed the standard pattern of splicing from exons 4 to 5 and 6 to 7 (data not shown). The skipping of exons 5 and 6 in BAB1415 predicts a messenger RNA that upon translation will terminate one amino acid downstream of exon 4 (Figure 4c).

#### Exome sequencing analyses

The axonal neuropathy present in BAB392 could potentially represent a phenotypic expansion of the sensory neuropathy associated with *PRICKLE1* heterozygous mutations. Additionally, the *KIF24* deletion in BAB1415 may possibly be unrelated to the neuropathy

present in the patient. Because of the uncertainty of genotype-phenotype correlations in these two patients, we conducted WES on all five patients with rare CNVs in CMT genes or linkage regions. With these data, we sought to determine SNV or insertion/deletion mutations that might contribute to the phenotypes of the patients. Sequencing resulted in the discovery of Mitofusin 2, or MFN2, mutations in two patients. The mutations result in an NM\_001127660 p.V244L (Chr1:12059066 G>T) and p.S249F (Chr1:12059082 C>T) change in BAB392 and BAB1415, respectively (Table 1). Alterations of these two amino acid residues are reported to represent causative pathogenic mutations in the Human Gene Mutation Database (http://www.hgmd.cf.ac.uk); however, the mutation previously reported at p.V244 was a G>A transition resulting in a methionine residue. This V244M mutation is present twice in our in-house database ( $\sim 1/2,400$ ), both in patients with CMT.<sup>23</sup> The V244L change in BAB392 is also likely to be pathogenic, because the amino acid residue is highly conserved and the mutation is de novo and predicted to be deleterious by various bioinformatic algorithms (Supplementary Figure S1 online). Two of the three other patients also contained a nonsynonymous neuropathy-associated SNV with minor allele frequency <1%; however, the phenotypes present in BAB1219, BAB3724, and BAB7807 seem to be consistent with gene dosage changes caused by the CNVs present in their genomes. Nevertheless, a contribution of these SNV to a potential mutational burden cannot be excluded.<sup>18</sup> Parental samples were available only for BAB392 and BAB3724.

Utilizing the WES data, we then determined whether sequencing could predict the CNVs in the five patients. The software CoNIFER<sup>22</sup> was applied to the WES data from the patients; however, none of the experimentally discovered and verified CNVs was bioinformatically predicted from WES data.

### DISCUSSION

Our aCGH approach detected nonrecurrent CNVs in *MPZ* (two), *PRICKLE1, KIF24*, and *NDRG1* (ref. 14) in 200 molecularly unresolved CMT cases. Deletion and duplication sizes ranged from 4 to 30 kb (Figure 1), encompassing anywhere from one exon to the entire gene. Breakpoint junction studies revealed that *Alu-Alu*-mediated rearrangements (present in three of five junctions) are the predominant molecular mechanism in CNVs present in CMT cases negative for CMT1A duplication.

Zhang et al.<sup>24</sup> analyzed nonrecurrent CNVs in the 17p12 region and found that mechanisms other than nonallelic homologous recombination, for example, fork-stalling, templateswitching, microhomology-mediated, break-induced replication, nonhomologous end joining, and *Alu*-mediated recombination events, contributed to nonrecurrent CNV formation in CMT1A. Recently, 57 deletion and duplication breakpoints at the *SPAST* locus were examined and ~70% of these events involved directly-oriented *Alu* elements, illustrating the predominance of *Alu*-mediated mechanisms in nonrecurrent rearrangements.<sup>11</sup> Early studies showed two of four nonrecurrent CNVs associated with Smith-Magenis syndrome were due to *Alu*-mediated mechanisms.<sup>25</sup> Additionally, a recent study showed that ~80% of simple deletions and ~50% of tandem duplications at 17p13.3 are mediated by *Alu-Alu* events.<sup>12</sup> These data show that the *Alu-Alu*-mediated mutational

mechanism is prevalent in nonrecurrent rearrangements and is responsible for over 50% of the CNVs presented here.

Our approach reveals that 4 of 200 individuals (2%) have a CNV in a known neuropathy gene (*MPZ* (two individuals), *PRICKLE1*, and *NDRG1*). However, the sizes of the CNVs detected herein may be too small to identify by some currently used commercial molecular diagnostic techniques. Methods including locus-specific tests such as fluorescence in situ hybridization (FISH) or genome-wide assays such as SNP arrays may be insufficient, and high-resolution aCGH may be required in cases negative for CMT1A duplication or SNV mutations. Huang et al.<sup>26</sup> performed aCGH targeting 34 hereditary peripheral neuropathy regions in 97 patients who were screened for the most common CMT genes. They found an exonic and an intronic CNV in *ARHGEF10*, an intronic loss in *SPTLC1*, and an exonic loss in *MTMR2*; these were interpreted as polymorphic variants because of their frequency or intronic location. A smaller, nonrecurrent duplication of *PMP22* was also identified. Among our 200 samples we found 31 CNV polymorphisms in addition to the five variants reported here (Supplementary Table S3 online). The five disease-associated and candidate CNVs are not present in the database of genomic variants.

Deletion and duplication of the *MPZ* gene were found in BAB7807 and BAB1219, respectively. Interestingly, although SNV mutations of this gene are common, CNVs involving *MPZ* are not prevalent in CMT. Mutations in *MPZ* are associated with CMT1B, an autosomal dominant form of the disease.<sup>27</sup> The deletion of the first exon of *MPZ* in BAB7807 continues to contribute to the range of mutations that lead to perturbation of *MPZ* expression.

Duplication of all coding exons of MPZ was recently reported.<sup>15</sup> The duplication cosegregated with CMT in 12 tested individuals; the coordinates presented indicate that the BAB1219 mutation is probably different and therefore a novel MPZ duplication. Additionally, a three-generation family with a quadruplication of MPZ and neighboring SDHC and Clorf192 genes (six members affected and four unaffected) was recently reported.<sup>16</sup> Those authors further demonstrated the increased expression of MPZ in lymphoblastoid cell lines ( $5 \times$  wild-type) and sural nerves ( $3 \times$ ) in the affected individuals. The phenotype of BAB1219 is consistent with previous clinical findings of MPZ duplication cases, which have a demyelinating sensorimotor neuropathy phenotype that starts during childhood. As the number of copies of MPZ increases, myelination of nerve fibers decreases gradually.<sup>28</sup> These data support the contention that MPZ, like PMP22, is dosage sensitive, and that rearrangement leading to either increased or decreased copy number can cause peripheral neuropathy. Mutations (point mutations and copy-number gains) in MPZ are associated with a broad phenotypic spectrum, ranging from severe onset during the neonatal period to late onset of milder symptoms resulting from hypomyelination.<sup>15</sup> The MPZ duplication in BAB1219 seems to be more deleterious than the deletion present in BAB7807 in terms of both age at onset and nerve conduction velocity (Table 1). This seems consistent with the previously reported observation that haploinsufficiency of MPZ, generated by nonsense-mediated decay of a truncated transcript, is associated with a less severe phenotype than aggregation of MPZ protein within the endoplasmic reticulum.<sup>29,30</sup>

Patient BAB392 had a deletion of the 5' untranslated region of *PRICKLE1*. Homozygous mutations in *PRICKLE1* (OMIM 608500) were first described in three consanguineous families with progressive myoclonus epilepsy-ataxia syndrome; one family presented with a mild neuropathy (OMIM 612437).<sup>31</sup> Heterozygous missense mutations of both *PRICKLE1* and *PRICKLE2* and deletions of *PRICKLE2* were found in patients with myoclonic/ generalized tonic-clonic seizure responsive to valproic acid and/or developmental delay in humans, mice, and flies.<sup>32</sup> Our patient had similar findings, including generalized tonic-clonic seizures that responded to treatment with phenobarbital (valproic acid and phenobarbital have antiepileptic effects through  $\gamma$ -aminobutyric acid receptors). Bosoi et al.<sup>33</sup> recently reported heterozygous variants in *PRICKLE1* in human subjects with neural tube defects.

We also found a two-exon deletion CNV in *KIF24* in BAB1415. Furthermore, RNA sequencing showed heterozygous exon 5 and 6 skipping in the affected individual (Figure 4). *KIF24* is a member of the kinesin gene family and is a ubiquitously expressed gene associated with microtubules. No mutation has been reported as causative of diseases in *KIF24*. Mutations in other members of the kinesin family have been associated with peripheral neuropathies, including *KIF1B* and CMT type 2A1 (OMIM 118210) and *KIF5A* and spastic paraplegia with peripheral neuropathy.<sup>34,35</sup> A loss-of-function mutation in *KIF1B* was identified in a family with autosomal dominant CMT2 family; however, mutations in *MFN2* are the primary pathologic segregating mutation in CMT2A.<sup>34,36</sup>

WES was conducted to determine whether *KIF24* was a likely CMT candidate gene in BAB1415 as well as the severity of the neuropathy in BAB392. This led to the discovery of *MFN2* mutations in the two individuals. The likely molecular diagnosis in BAB1415 is a previously reported *MFN2* mutation, and the contribution of the *KIF24* CNV remains to be determined. By contrast, BAB392 presented with a composite phenotype consisting of seizures and neuropathy. It is likely that the *PRICKLE1* CNV combined with the *MFN2* SNV led to the patient's manifestations. Interestingly, when parental samples were examined, both the *PRICKLE1* deletion CNV and the *MFN2* SNV were de novo in the patient. This depicts the necessity of considering all possible rationales for a patient phenotype, even with a likely causative new mutation. Blended phenotypes have been observed previously with two SNV point mutations in an individual, comprising ~5% of diagnostic dilemmas referred for potential molecular diagnosis by exome sequencing.<sup>37,38</sup>

In summary, CNVs are a relevant cause of CMT in patients with unknown molecular genetic etiology (responsible for ~2% of patients without CMT1A), but such variants are rare in genes other than *PMP22*. Our approach allowed detection of, to our knowledge, the first disease-associated autosomal recessive duplication CNV in a rare neuropathy gene, *NDRG1* (ref 14), as well as an *MPZ* deletion associated with neuropathy. Furthermore, WES combined with CNV analysis was critical in unraveling the etiology of complex phenotypic presentations and composite neurological phenotypes potentially representing a mutational burden involving more than one gene, and it helped to define candidate mutations.

# Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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J.R.L. has stock ownership in 23andMe, is a paid consultant for Regeneron Pharmaceuticals, has stock options in Lasergen, Inc., is a member of the Scientific Advisory Board of Baylor Miraca Genetics Laboratories, and is a coinventor on US and European patents related to molecular diagnostics for inherited neuropathies, eye diseases, and bacterial genomic fingerprinting. The Department of Molecular and Human Genetics at Baylor College of Medicine derives revenue from the chromosomal microarray analysis and clinical exome sequencing offered in the Baylor Miraca Genetics Laboratory (http://www.bmgl.com/BMGL/Default.aspx).

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# Figure 1. Summary of copy-number variants (CNVs) found in Charcot-Marie-Tooth genes and linkage regions

Identification numbers are shown in the left column. Genes are depicted by black horizontal arrows with the arrowhead showing their direction of transcription; cen, centromere; tel, telomere (all genes shown with respect to the + strand). Green boxes depict the position of deletion CNVs, and red boxes depict the position of duplication CNVs. Depiction of the gene structures, including deleted or duplicated exons (vertical bars), is shown below the schematic of the regions. Size of the CNVs is indicated below the colored boxes.

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# Figure 2. Array comparative genomic hybridization (CGH) and breakpoint junction studies of the *MPZ* gene deletion (in BAB7807) and duplication (in BAB1219)

(a) Chromosome 1 ideogram with G bands indicated (top). The location of MPZ is shown with a red vertical line at 1q23.3. Array CGH results of the 20-kb deletion involving upstream parts of both MPZ and SDHC. (b) Breakpoint sequence analysis for MPZ deletion using the MultAlin alignment tool. Proximal and distal reference sequences are mapped within FLAM C and AluSz6-D sequences, respectively. The patient's breakpoint sequence that matches the reference sequence is shown in blue, mismatched sequence is shown in black, and perfect matching between all three sequences are shown in red. The breakpoint junction where the transition occurred from the proximal to the distal reference sequence is specified with a black circle. (c) Graphic representation of MPZ (vertical bars; deleted exon 1 is shown in green); size and orientation of the gene are shown above the exons. (d) Chromosome 1 ideogram with G bands as described in Figure 2a. Array CGH results reveal genomic duplication. The red line indicates MPZ on chromosome 1q23.3. (e) Breakpoint sequence analysis for MPZ duplication. The proximal and distal sequences are from hg19/ GRCh37. Proximal reference sequence and patient breakpoint sequences that match the proximal reference sequence are shown in red, the distal reference sequence and patient breakpoint sequences that match the distal reference sequence are shown in green, and 2-bp microhomology (CC) at the junction is shown in purple. (f) Graphical representation of six

exons of *MPZ* (vertical bars; duplicated exons are shown in red); the size and orientation of the gene are shown above the exons.

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# Figure 3. Array comparative genomic hybridization (CGH) and breakpoint junction studies of the *PRICKLE1* deletion in BAB392

(a) Chromosome 12 ideogram with G bands indicated (top). The location of *PRICKLE1* is shown with a red vertical line at 12q12. Array CGH results of the ~30-kb deletion involving the 5' untranslated region and start of the *PRICKLE1* coding sequence. (b) Breakpoint sequence analysis for the PRICKLE1 deletion using the MultAlin alignment tool. Proximal and distal reference sequences are mapped within *AluSp* and *AluSx* sequences, respectively. Sequences matching the consensus Alu sequence are shown in blue, the mismatched sequence is shown in black, and concordance between the three Alu sequences is depicted in red. The breakpoint junction is specified with a black circle where the transition occurred from the proximal to the distal reference sequence. (c) Polymerase chain reaction (PCR) confirmation of the junction present in the BAB392 sample. Amplification of PSRC1 gene serves as a control. Breakpoint amplification is successful in the patient (~1.5-kb band) and not parental (BAB393 and BAB394). (d) Reverse-transcriptase PCR depicting relatively decreased transcript levels of PRICKLE1 in BAB392 is shown. The amount of amplification from BAB392 is lower than two controls who lack CNV in PRICKLE1. This is consistent with the 5' untranslated region deletion possibly leading to decreased transcription of the gene or nonsense-mediated decay of the transcript. (e) Graphic view of eight exons of PRICKLE1 (vertical bars, deleted first exon is shown in green); the size and orientation of the gene are noted above the exons.

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# Figure 4. Array comparative genomic hybridization (CGH) and breakpoint junction studies of the KIF24 gene deletion in BAB1415

(a) Chromosome 9 ideogram with G bands indicated (top). The location of *KIF24* is shown with a red vertical line at 9p13.3. Array CGH results of the ~16.5-kb deletion involving exons 5 and 6 of *KIF24*. (b) Breakpoint sequence for *KIF24* deletion using the MultAlin alignment tool. Proximal and distal reference sequences are mapped within *AluYa5* and *AluY* sequences, respectively. The patient's breakpoint sequence that matches the reference sequence is shown in blue, the mismatched sequence is shown in black, and perfect matching between all three sequences is depicted in red. The breakpoint junction is specified with a black circle where the transition occurred from the proximal to the distal reference sequence. (c) Skipping of exons 5 and 6 because of the deletion is confirmed in the complementary DNA. Peaks on peaks at the end of exon 4 and the beginning of exon 7 show the two species of RNA in BAB1415, both wild-type and deletion-containing. The lack of exons 5 and 6 in the deletion-containing RNA lead to a premature termination codon at the beginning of exon 7. (d) Graphic view of 13 exons of *KIF24* (vertical bars, deleted exons 5 and 6 are shown in green); the size and orientation of the gene are shown above the exons.

Table 1

Motor nerve conduction studies of four patients in the study.

								Patie	nt									
		BAB392		ł	3AB1219		E	8AB1415		$\mathbf{B}_{\ell}$	AB3724 <sup>4</sup>	ı	В	AB7807				
Age at testing (years)		4			35			3.5			30			24				
Age of onset (years)		3-4		С	hildhoot	Į		2–3			3			23				
Family history		Negative			Negative			Vegative		4	Vegative		4	Vegative		Nor	mal valı	ies
Motor NCS	DL (ms)	V (m/s)	Amp (mV)	DL (ms)	V (m/s)	Amp (mV)	DL (ms)	V (m/s)	Amp (mV)	DL (ms)	V (m/s)	Amp (mV)	DL (ms)	V (m/s)	Amp (mV)	DL (ms)	V (m/s)	Amp (mV)
R. median	3.1	44	2.5	8.9	23.3	8.51	2.5	39	N/A	NR	NR	NR	N/A	41.4	N/A	<4.0	>50	5.0
R. ulnar	3.0	42	2.5	5.3	29.9	7.57	1.8	67	*	NR	NR	NR	N/A	N/A	N/A	<3.5	>50	5.0
R. common peroneal	4.9	28	0.75	NR	NR	NR	NR	NR	NR	NR	NR	NR	N/A	22.3	N/A	<4.5	>40	2.0
R. tibialis	4.8	22	1.5	NR	NR	NR	NR	NR	NR	NR	NR	NR	N/A	29.2	N/A	<5.0	>40	4.0
Clinical diagnosis	CMT , develo	; Tonic-c seizures; pmental	lonic delay		CMT1			CMT1		CMT; hei g	sensorin aring loss laucoma	eural ;;		CMT				
CNV	PRICI (	KLEI de de novo)	letion	MP2	c duplica	tion	KII	724 deleti	on	l <b>np</b>	NDRG1 plication	a	, MP	Z first ex leletion	uoy			
SNV	Chr G> p.V2	1:12059 •T; <i>MF</i> 44L; de 1	<b>066</b> V2 novo	Chr1 G>A; J	4:10518( <i>INF2</i> p.R	)633 1045Q	Chr1:1 MF.	2059082 N2 p.S24	C>T; 9F	Chr1 <sup>2</sup> A>C; P.	4:102481 ; <i>DYNC</i> 1 K2401N	630 1H1		N/A				
	:																	

Boldface type indicates likely pathogenic mutations.

Amp, amplitude; CMT, Charcot-Marie-Tooth; CNV, copy-number variant; DL, distal latency; NCS, nerve conduction studies; N/A, not available; NR, not recordable; R., right; SNV, single-nucleotide variant; V, conduction velocity.

<sup>a</sup>Previously published *NDRGI* CNV is homozygous and is a known recessive locus. All others are heterozygous and have known or postulated autosomal dominant mechanisms.