Myelin is dependent on the Charcot-Marie-Tooth Type 4H disease culprit protein FRABIN/FGD4 in Schwann cells

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Abstract: Studying the function and malfunction of genes and proteins associated with inherited forms of peripheral neuropathies has provided multiple clues to our understanding of myelinated nerves in health and disease. Here, we have generated a mouse model for the peripheral neuropathy Charcot-Marie-Tooth disease type 4H by constitutively disrupting the mouse orthologue of the suspected culprit gene FGD4 that encodes the small RhoGTPase Cdc42 guanine nucleotide exchange factor Frabin. Lack of Frabin/Fgd4 causes dysmyelination in mice in early peripheral nerve development, followed by profound myelin abnormalities and demyelination at later stages. At the age of 60 weeks, this was accompanied by electrophysiological deficits. By crossing mice carrying alleles of Frabin/Fgd4 flanked by loxP sequences with animals expressing Cre recombinase in a cell type-specific manner, we show that Schwann cell-autonomous Frabin/Fgd4 function is essential for proper myelination without detectable primary contributions from neurons. Deletion of Frabin/Fgd4 in Schwann cells of fully myelinated nerve fibres revealed that this protein is not only required for correct nerve development but also for accurate myelin maintenance. Moreover, we established that correct activation of Cdc42 is dependent on Frabin/Fgd4 function in healthy peripheral nerves. Genetic disruption of Cdc42 in Schwann cells of adult myelinated nerves resulted in myelin alterations similar to those observed in Frabin/Fgd4-deficient mice, indicating that Cdc42 and the Frabin/Fgd4-Cdc42 axis are critical for myelin homeostasis. In line with known regulatory roles of Cdc42, we found that Frabin/Fgd4 regulates Schwann cell endocytosis, a process that is increasingly recognized as a relevant mechanism in peripheral nerve pathophysiology. Taken together, our results indicate that regulation of Cdc42 by Frabin/Fgd4 in Schwann cells is critical for the structure and function of the peripheral nervous system. In particular, this regulatory link is continuously required in adult fully myelinated nerve fibres. Thus, mechanisms regulated by Frabin/Fgd4-Cdc42 are promising targets that can help to identify additional regulators of myelin development and homeostasis, which may crucially contribute also to malfunctions in different types of peripheral neuropathies.

DOI: https://doi.org/10.1093/brain/aws275
Jan; Suter, Ueli (2012). Myelin is dependent on the Charcot-Marie-Tooth Type 4H disease culprit protein FRABIN/FGD4 in Schwann cells. Brain. 135(Pt 12):3567-3683.
DOI: https://doi.org/10.1093/brain/aws275
Myelin is dependent on the Charcot–Marie–Tooth Type 4H disease culprit protein FRABIN/FGD4 in Schwann cells

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Studying the function and malfunction of genes and proteins associated with inherited forms of peripheral neuropathies has provided multiple clues to our understanding of myelinated nerves in health and disease. Here, we have generated a mouse model for the peripheral neuropathy Charcot–Marie–Tooth disease type 4H by constitutively disrupting the mouse orthologue of the suspected culprit gene FGD4 that encodes the small RhoGTPase Cdc42-guanine nucleotide exchange factor Frabin. Lack of Frabin/Fgd4 causes dysmyelination in mice in early peripheral nerve development, followed by profound myelin abnormalities and demyelination at later stages. At the age of 60 weeks, this was accompanied by electrophysiological deficits. By crossing mice carrying alleles of Frabin/ Fgd4 flanked by loxP sequences with animals expressing Cre recombinase in a cell type-specific manner, we show that Schwann cell-autonomous Frabin/Fgd4 function is essential for proper myelination without detectable primary contributions from neurons. Deletion of Frabin/Fgd4 in Schwann cells of fully myelinated nerve fibres revealed that this protein is not only required for correct nerve development but also for accurate myelin maintenance. Moreover, we established that correct activation of Cdc42 is dependent on Frabin/Fgd4 function in healthy peripheral nerves. Genetic disruption of Cdc42 in Schwann cells of adult myelinated nerves resulted in myelin alterations similar to those observed in Frabin/Fgd4-deficient mice, indicating that Cdc42 and the Frabin/Fgd4–Cdc42 axis are critical for myelin homeostasis. In line with known regulatory
roles of Cdc42, we found that Frabin/Fgd4 regulates Schwann cell endocytosis, a process that is increasingly recognized as a relevant mechanism in peripheral nerve pathophysiology. Taken together, our results indicate that regulation of Cdc42 by Frabin/Fgd4 in Schwann cells is critical for the structure and function of the peripheral nervous system. In particular, this regulatory link is continuously required in adult fully myelinated nerve fibres. Thus, mechanisms regulated by Frabin/Fgd4–Cdc42 are promising targets that can help to identify additional regulators of myelin development and homeostasis, which may crucially contribute also to malfunctions in different types of peripheral neuropathies.

Introduction

Charcot–Marie–Tooth disease, also known as hereditary motor and sensory neuropathy, is one of the most common inherited neurological disorders (Ske, 1974). Charcot–Marie–Tooth disease is a clinically and genetically heterogenous disease, usually characterized by distal muscle weakness and atrophy, distal sensory loss and limb deformities (Shy et al., 2005). Clinically, Charcot–Marie–Tooth disease is divided based on electrophysiological and histopathological characteristics into demyelinating or dysmyelinating (CMT1, CMT3 and CMT4) and axonal forms (CMT2). The demyelinating or dysmyelinating neuropathies are commonly assumed to start with damage to Schwann cells and are mainly associated with reduced nerve conduction velocity. However, as demyelinating Charcot–Marie–Tooth disease progresses and axonal degeneration becomes an additional pathological feature, reduced amplitudes and dispersion of compound muscle action potentials become the main correlate to clinical disability (Krajewski et al., 2000; Suter and Scherer, 2003). Axonal forms of Charcot–Marie–Tooth disease are thought to originate on the neuronal side and are linked to decreased compound muscle action potential amplitudes. The distinction with regard to the initially affected cell type is blurred in intermediate forms of Charcot–Marie–Tooth disease (Nicholson and Myers, 2006). As was recently shown, neuron-specific ablation of the prion protein in mice may cause a demyelinating neuropathy (Bremer et al., 2010). This observation reiterated that experimental proof is required to unambiguously determine the originally affected cell type(s) in peripheral neuropathies, as peripheral nerves are controlled by continuous reciprocal Schwann cell–neuron interactions (Nave, 2010; Pereira et al., 2012). Such knowledge is critical to understand the underlying disease mechanisms in the different forms of Charcot–Marie–Tooth disease and provides a rational basis for potential treatment strategies.

Mutations in >30 genes are associated with Charcot–Marie–Tooth disease variants, with autosomal dominant, autosomal recessive or X-linked inheritance patterns (Reilly et al., 2011). The subtype CMT4H belongs to the autosomal recessive demyelinating forms of Charcot–Marie–Tooth disease (De Sandre-Giovannoli et al., 2005). Several different mutations in FRABIN/FGD4, a member of a family of Cdc42-specific guanine nucleotide exchange factors (GEFs) (Nakanishi and Takai, 2008), have been associated with CMT4H (Delague et al., 2007; Stendel et al., 2007; Fabrizi et al., 2009; Houlden et al., 2009). These include missense, nonsense, frame-shifting and splice mutations. The relatively small number of patients does not allow a well-founded genotype–phenotype correlation. Generally, patients with the CMT4H subtype are clinically affected in early childhood, with disease onset occasionally as early as the first year of life (Baets et al., 2011). In most cases, CMT4H is characterized by a slowly progressive neuropathy with increasing distal sensory loss and distal weakness. Electrophysiological examinations revealed markedly reduced nerve conduction velocity. Nerve biopsies show redundant myelin and myelin infoldings and outfoldings as striking pathological hallmarks. Furthermore, severely hypomyelinated fibres indicate demyelination and remyelination consistent with the classification of CMT4H as demyelinating Charcot–Marie–Tooth disease (De Sandre-Giovannoli et al., 2005; Stendel et al., 2007; Fabrizi et al., 2009). Reduced density of myelinated large calibre fibres, likely caused by axonal degeneration, appears as an additional feature.

The available knowledge from human genetics suggests a strict dependence of myelinated peripheral nerves on Frabin/Fgd4. However, the current information concerning Frabin/Fgd4 function is limited and has been obtained from cell culture studies in mainly non-neural cells (Nakanishi and Takai, 2008). We wished to examine Frabin/Fgd4 function in vivo. For this purpose, we used mouse genetics focusing on nerve development and nerve homeostasis. We reasoned that these experiments would also be instrumental for our understanding of the disease mechanism involved in CMT4H and related neuropathies. Our analysis led to the establishment of an animal model for CMT4H and revealed that Schwann cells are the main cell type initially affected by the loss of Frabin/Fgd4 function. Furthermore, we found that the critical role of Frabin/Fgd4 in peripheral nerves is linked to its molecular function as GEF for Cdc42 and most likely involves regulation of endocytosis.

Materials and methods

Mice

Frabin mutant mice were produced by standard technology (Mouse Clinical Institute, Strasbourg, France). Cdc42 mutants (Thurnherr et al., 2006; Wu et al., 2006; Benninger et al., 2007), Hb9-Cre (Arber et al., 1999) and Dhh-Cre (Jaegle et al., 2003; Pereira et al., 2009) mice...
have been described. In experiments involving the Plp-CreERT2 transgene (Leone et al., 2003), Cre recombinase was activated by repeated injection of tamoxifen (100 mg/kg, intraperitoneally) in 10-week-old mice on 5 consecutive days. Genotypes of mice were determined by PCR on genomic DNA derived from ear biopsies using appropriate primer pairs (Supplementary material). Mice were kept under standard housing conditions on Lignocel bedding (Provimi). Experiments followed approved protocols (Veterinary Office, Canton Zurich, Switzerland).

**Behavioural analysis**

SHIRPA (SmithKline, Harwell, Imperial College, Royal London Hospital, phenotype assessment) was carried out as described (Rogers et al., 1997). Sensory and motor tests followed established protocols (Bremer et al., 2010).

**Electrophysiology**

Motor nerve conduction in 30- and 60-week-old anaesthetized mice was measured as described (Zielasek et al., 1996; Bremer et al., 2010). In brief, upon supramaximal stimulation (i.e. at least 30% above the current needed to obtain a maximal compound muscle action potential) of the tibial nerve at the ankle (‘distal’) and stimulation of the sciatic nerve at the sciatic notch (‘proximal’), compound muscle action potentials were recorded with a pair of steel needle electrodes in the foot muscles. Nerve conduction velocities were calculated in metres per second from distal and proximal latencies. Ten successive F-waves were recorded, and the shortest latencies were taken upon stimulation at the ankle (Zielasek et al., 1996).

**Electron microscopy and histological analysis**

Tissue embedding and electron microscopy were performed as described (Pereira et al., 2010; Somandin et al., 2012). Ultrathin sections were collected on carbon-coated Formvar grids (Electron Microscopy Sciences) and analysed in a Morgagni 268 transmission electron microscope. Complete transverse sections of plantaris, quadriceps and saphenous nerves were captured at ×1000 magnification and were reconstructed using Adobe® Photoshop®. All affected fibres carrying aberrant myelin features were quantified per nerve section. Myelin thickness was measured in Adobe® Photoshop® on electron microscopy pictures.

**RhoGTPase activity assay**

Cdc42 activity was measured as described (Sander et al., 1998; Benninger et al., 2007). The glutathione-S-transferase–p21-activated kinase–cricb (Cdc42–Rac1 interactive binding) domain construct was provided by Dr J. Collard (The Netherlands Cancer Institute, Amsterdam). In brief, the material to be analysed (sciatic nerves or Schwann cells) was homogenized in 10% glycerol, 50 mM Tris–HCl, pH 7.4, 100 mM NaCl, 1% Nonident P-40, 2 mM MgCl₂ and protease inhibitor cocktail (Sigma-Aldrich) and was centrifuged for 5 min at 21 000g at 4°C. Aliquots of 10% of the volume were taken from the supernatant to determine the total protein amount. The remaining supernatant was incubated with the bait protein bound to glutathione-coupled Sepharose™ beads (GE Healthcare) at 4°C for 60 min. The beads and protein bound to the fusion protein were washed three times in an excess of homogenization buffer, eluted in Laemmli sample buffer and analysed for bound Cdc42 by western blotting.

**Cell culture and expression silencing**

RT4 rat schwannoma cells were grown in Dulbecco’s modified Eagle’s medium containing 10% foetal calf serum (Gibco) and were transfected using Lipofectamine® 2000 (Invitrogen) according to the manufacturer’s protocol. Small interfering RNA transfections were carried out repeatedly at 72, 48 and 24 h before assays were performed. Frabin/Fgd4 expression silencing: Fgd4 small interfering RNA (SIO1512574; Qiagen) with the targeting sequence 5'-CTG AAT GGA GTA AGA AAC GAA-3'. Control small interfering RNA: AllStars Negative Control siRNA (1027292; Qiagen). For fluorescence-activated flow cytometry and microscopy analysis, small interfering RNAs were labelled with Alexa Fluor® 488. Short hairpin RNA transfections were done 48 h before performing the assays, using the pSicoR vector, carrying a green fluorescent protein expression cassette and containing the short hairpin RNA sequence of choice. Targeting sequences for Fgd4: 5'-GCA GCA AGC CAT TCT AAT-3' (shRNA1) and 5'-GAA GAA GAG GAT ATT GTA-3' (shRNA2). Control short hairpin RNA (targeting dsred2): 5'-AGT TTC AGT ACC GCT CCA A-3' (Ozcelik et al., 2010).

**Transferrin assays**

Transferrin assay for fluorescence-activated flow cytometry was performed as described (Sidirooulos et al., 2012) using short hairpin or small interfering RNA-transfected RT4 cells. Fluorescence intensity of internalized transferrin was measured for 2000 green fluorescent protein- (short hairpin RNA transfections) or Alexa Fluor® 488-positive (small interfering RNA transfections) cells. High expressing cells were subjected to comparative analysis. For fluorescence microscopy, the identical assay was performed with the exceptions that Alexa Fluor®-labelled transferrin (Invitrogen; 20 μg/mL in serum-free Dulbecco’s modified Eagle’s medium) was used and cells were not detached before fixation. Images were captured with an epifluorescence Zeiss Axiosvert microscope equipped with a Zeiss High Resolution Monochromatic camera.

**Western blotting and antibodies**

Sciatic nerves from adult mice were isolated and separated from the epineurium. Nerves were homogenized with a chilled mortar and pestle in lysis buffer (0.1% SDS, 10 mM Tris–HCl, 150 mM NaCl, 50 mM NaF, 1 mM NaVO₄, 1 mM EDTA, 0.5% sodium-deoxycholate, protease inhibitor mixture; Sigma). Extracts were processed using standard SDS-PAGE and western blotting procedures. The following antibodies were used: AKT-P Serine 473 (Cell Signaling Technology, 1:1000), AKT (Cell Signaling Technology, 1:1000), ErbB2-P Tyrosine 1248 (Abcam, 1:1000), ErbB2 (Abcam, 1:1000), disc large homologue 1 (Dlg1) (BD Transduction Laboratories, 1:1000), myelin basic protein (AbD Serotec, 1:1000), JNK (Cell Signaling Technology, 1:1000), AKT-P Serine 473 (Cell Signaling Technology, 1:1000), myelin basic protein (AbD Serotec, 1:1000), JNK-P (Cell Signaling Technology, 1:1000), NKG2D (BD Transduction Laboratories, 1:1000), myelin basic protein (AbD Serotec, 1:1000), JNK-P, ErbB2-P Tyrosine 1248 (Abcam, 1:1000), ErbB2 (Abcam, 1:1000), disc large homologue 1 (Dlg1) (BD Transduction Laboratories, 1:1000), myelin basic protein (AbD Serotec, 1:1000). Secondary antibodies were obtained from Promega and Southern Biotech. Bands were quantified using Quantity One® software (Bio-Rad).
Cryo-embedding and immunostaining of cryosections

Sciatic nerves were removed from mice, fixed in 4% paraformaldehyde for 1 h, incubated overnight in 30% sucrose and frozen in OCT (Tissue Tek). Sections (10 μm) were cut on a HM560 Cryostat (Microm), post-fixed in 4% paraformaldehyde for 5 min, blocked in blocking solution (10% goat serum in PBS with 0.3% Triton® X-100) for 1 h at room temperature and incubated with primary antibodies against S100β (Sigma, mouse monoclonal, 1:200) and Frabin/Fgd4 (Pineda, rabbit polyclonal, 1:200) overnight at 4°C in blocking solution. Sections were washed with PBS and incubated with secondary antibodies against mouse coupled to Alexa Fluor® 488 (Invitrogen, 1:500) and rabbit coupled to Cy3 (Jackson ImmunoResearch, 1:500) for 1 h at room temperature. Sections were washed with PBS, incubated for 5 min with 4',6-diamidino-2-phenylindole and mounted with Immu-Mount™ (Thermo Scientific).

Statistical analysis

Data show the mean ± standard error of the mean. Two-tailed Student’s t-test was used with significance set to *P < 0.05, **P < 0.01 or ***P < 0.001; n = number of independent experiments.

Results

Ubiquitous loss of Frabin/Fgd4 induces features of a demyelinating peripheral neuropathy in mice

We have generated a conditional mouse null allele for Frabin/Fgd4 using standard embryonic stem cell technology. To achieve this goal, exon 4 of Fgd4 was flanked with LoxP sites by homologous recombination, and mice with this allele were obtained (Fig. 1A).

Figure 1  Loss of Frabin/Fgd4 leads to electrophysiological characteristics of demyelinating peripheral neuropathies. (A) Ablation of exon 4 in the Fgd4 locus generates a premature stop codon in exon 5 because of a frame shift (filled triangles = introduced loxP sites; START = translational start codon; STOP = conventional translational stop codon; STOP (bold) = premature translational stop codon generated after Fgd4 exon 4 ablation), resulting in B, loss of Frabin/Fgd4 protein (western blot; asterisk: unspecific signal). (C and D) At 60 weeks, Fgd4−/− mice show longer distal latency, disperse compound muscle action potentials with mild reduction in amplitude, longer F-wave latency and reduced nerve conduction velocity in sciatic nerve compared with wild-type mice; in C, a representative original recording is shown. Arrowhead indicates stimulus artefact; open arrow indicates onset of compound muscle action potential (distal latency); filled arrow indicates onset of F-wave. Data represent the mean ± standard error of the mean. Wild-type mice, n = 7; Fgd4−/− mice, n = 10. P-values in D: *P < 0.05; ***P < 0.001; Student’s t-test (two-tailed).
In our initial experiments, we bred and analysed a mouse line that lacks Frabin/Fgd4 ubiquitously. This approach mimics the genetic situation in patients with CMT4H and was aimed to assess whether FRABIN/FGD4 mutations are definitively responsible for CMT4H, and mice lacking Frabin/Fgd4 constitute an appropriate animal model for this disease. In addition, we anticipated that Frabin/Fgd4-deficient mice would be suitable to study Frabin/Fgd4 function in vivo. For this purpose, we bred mice carrying conditional Fgd4 alleles (Fgd4 flox) with mice expressing Cre recombinase in the germline. This procedure resulted in offspring with a constitutively inactivated Fgd4 allele. As expected, mice homozygous for this deletion allele (Fgd4<sup>−/−</sup>) are lacking Frabin/Fgd4 protein expression as verified by western blot analysis of sciatic nerve lysates (Fig. 1B). Fgd4<sup>−/−</sup> mice are viable, fertile and were born at expected Mendelian ratios. We found no major abnormal behaviour by SHIRPA (SmithKline Beecham, Harwell, Imperial College, Royal London Hospital, phenotype assessment) analysis up to the age of 60 weeks. Frabin/Fgd4 mutants displayed normal body size and weight and were inconspicuous in sensory examinations (Supplementary Table S1). In a grip strength test, 60-week-old Fgd4<sup>−/−</sup> mice exhibited mildly impaired performance compared with wild-type mice. However, we found no significant differences in rotaod performance and footprint analysis (step length, step width and foot angle), collectively indicating a mild phenotype at this age.

As CMT4H belongs to the demyelinating forms of peripheral neuropathies, we expected characteristic electrophysiological alterations in myelinated nerves of Fgd4<sup>−/−</sup> mice. Indeed, when analysing the function of motor fibres, we detected moderately increased distal latencies and mildly increased F-wave latencies, reduced nerve conduction velocity and dispersed compound muscle action potentials with mildly reduced amplitudes in 60-week-old Fgd4<sup>−/−</sup> mice compared with wild-type mice (Fig. 1C and D), whereas at 30 weeks of age no abnormalities were observed (data not shown).

These electrophysiological alterations are consistent with a demyelinating phenotype caused by loss of Frabin/Fgd4. To corroborate our findings, we performed histological examinations of distal peripheral nerves with variable motor and sensory contributions (plantar, sciatic, quadriceps and saphenous) and of proximal parts of motor and sensory nerves [lumbar spine nerve roots 4 (L4) ventral and dorsal roots]. An overview revealed aberrant myelin structures in all analysed nerves of Fgd4<sup>−/−</sup> mice, albeit with different frequencies (Fig. 2 and Supplementary Fig. 1). The predominant irregular features were myelin outfoldings and infoldings of various complexities, in combination with redundant myelin loops, both around nodes of Ranvier and along internodes. In addition, we found axons in older animals without myelin or with abnormally thin myelin, indicating demyelination and incomplete remyelination. Myelin debris and polyaxonal myelination were occasionally observed.

Based on this survey, we performed quantitative temporal examinations in developing and adult mice. Our initial qualitative investigations in the sciatic nerve provided evidence that aberrant myelin features might already be increased in early Fgd4<sup>−/−</sup> nerve development compared with wild-type mice (Fig. 2A–D). Thus, we analysed and quantified, at the ultrastructural level by electron microscopy, the number of fibres that displayed aberrant myelin structures on complete reconstructions of post-natal Day 5 sciatic nerves. A subtle, but significant surplus of affected fibres was found in Fgd4<sup>−/−</sup> mice compared with wild-type mice (Fig. 3), confirming an early onset of this characteristic morphological phenotype. To examine phenotype progression over time, we tracked alterations in the plantaris nerve, as this distal nerve is a sensitive indicator of nerve changes in mouse models of hereditary motor and sensory neuropathies (Frei et al., 1999; our unpublished observations). Fgd4<sup>−/−</sup> mice displayed significantly more fibres with aberrant myelin structures compared with wild-type mice at all time points analysed from post-natal Day 14 to 80 weeks of age (Fig. 4A and B). These alterations became progressively more severe with ~6% of affected fibres at 2 weeks of age and reaching 40% in 80-week-old Fgd4<sup>−/−</sup> mice. In parallel, the complexity and extent of aberrant myelin structures associated with individual axons increased with age as judged by qualitative examinations (Fig. 4A). Myelinated axons without overt myelin aberrations had normal myelin thickness as assessed by growth ratio (axon diameter/fibre diameter) measurements in 10-week-old animals (Fig. 4C). In 60-week-old animals, however, we observed a tendency towards an increased growth ratio, indicating thinner myelin (Fig. 4D). In line with these findings, the number of fibres with signs of demyelination and partial remyelination revealed a tendency towards an increase in 60-week-old Fgd4<sup>−/−</sup> mice compared with wild-type mice, reaching statistical significance in 80-week-old animals (Fig. 4E). Remarkably, we found a rather steep rise in demyelination/remyelination features between 60- and 80-week-old Fgd4<sup>−/−</sup> animals, suggesting that demyelination proceeds rapidly when a threshold of damage is reached. On the axonal side, quantification of myelinated axons of entire plantaris nerves revealed no significant difference between Fgd4<sup>−/−</sup> and wild-type mice at all ages examined up to 80 weeks, indicating no major primary or secondary axonal loss (Fig. 4F). In summary, our findings show that loss of Frabin/Fgd4 in the mouse leads to electrophysiological and morphological abnormalities resembling the hallmarks of CMT4H, thus establishing Fgd4<sup>−/−</sup> mice as an animal model for this disorder.

### Schwann cells critically depend on Frabin/Fgd4 function

Intimate and reciprocal Schwann cell–axon interactions are a major regulatory hallmark of peripheral nerves in health and disease (Suter and Scherer, 2003; Jessen and Mirsky, 2005; Nave and Trapp, 2008; Salzer, 2008; Taveggia et al., 2010; Pereira et al., 2012). Thus, our findings in Fgd4<sup>−/−</sup> mice may be because of a primary requirement of Frabin/Fgd4 function in either Schwann cells or neurons, or both the cell types. The answer to this question is important to understand why and how peripheral nerves rely on Frabin/Fgd4, and this knowledge will provide critical insights into the CMT4H disease mechanism. Thus, we bred mice carrying the conditional Frabin/Fgd4 null allele (Fig. 1A) with established mouse lines that express Cre recombinase specifically in the Schwann cell lineage Dhh-Cre (Jaegle et al., 2003; Pereira et al., 2009) or in the motor neuron lineage (Hb9-Cre; Arber
et al., 1999) (Fig. 5A). Dhh-Cre Fgd4 flox/flox animals (Dhh-Cre; Fig. 5B) and Hb9-Cre Fgd4 flox/flox animals (Hb9-Cre; Fig. 5C) displayed loss of Frabin/Fgd4 in the targeted cell type as expected. Qualitative histological analysis of various adult peripheral nerves revealed prominent aberrant myelin features in conditional Fgd4 mutants with Schwann cell-specific Frabin/Fgd4 deletion, comparable with those observed in Fgd4−/− mice (Fig. 5D). In contrast, nerves of motor neuron-specific Frabin/Fgd4 deletion mutants were not different from the wild-type mice. For quantitative analysis, we examined fully reconstructed sections of

![Figure 2](image-url)

**Figure 2** Frabin/Fgd4-deficient mice form aberrant PNS myelin. Fgd4−/− mice display aberrant myelin features during early steps of myelination (A–D: post-natal Day 5; sciatic nerve) and in myelin maintenance (E–G, I and K: 80 weeks old mice; H, J and M–P: 60 weeks old mice; L: 10 weeks old mice; plantaris nerve), including simple myelin outfoldings (A), redundant myelin (B), complex myelin outfoldings (E) and highly complex myelin outfoldings (C and D), redundant myelin loops outside (F) and protruding into the axon (G), degradation of myelin (I), signs of demyelination (K) and remyelination (H and J) and rarely polyaxonal myelination (L). Aberrant myelin features tend to be located in the vicinity of nodes of Ranvier and Schmidt–Lanterman incisures (M, N and P). (A–L) Cross-sections. (M–P) Longitudinal sections. Scale bars = 1 μm (A–L); 5 μm (M–P).
alleles with an established Plp-CreERT2 mouse line (Leone et al., 2003). In the offspring, Cre-mediated recombination in Schwann cells was induced by tamoxifen injections in young adult Pnp-CreERT2 Fgd4 flox/flox animals (10 weeks old) (Fig. 6A). Specific loss of Frabin/Fgd4 in Schwann cells was verified by immunostaining (Fig. 6B). Qualitative analysis of plantaris nerves derived from such adult-induced Frabin/Fgd4 mutants at the age of 30 weeks revealed aberrant myelin features similar to age-matched Fgd4−/− animals (Fig. 6C). Quantifications of whole-nerve reconstructions at the electron microscopy level showed that adult-induced Frabin/Fgd4 mutants had slightly fewer affected plantaris nerve fibres compared with Fgd4−/− mice (~10 versus 15%). However, there was still a strong increase in myelin alterations in adult-induced Frabin/Fgd4 mutants compared with wild-type or tamoxifen-injected control mice (~10 versus 2%; Fig. 6D). These data demonstrate a critical function of Frabin/Fgd4 in myelin maintenance and imply that the observed phenotype in Fgd4−/− mice reflects both a developmental and a maintenance component.

Frabin/Fgd4 function in Schwann cells is required for myelin maintenance

The availability of a conditional Frabin/Fgd4 null allele allowed us to also address the physiologically and pathophysiologically important issue whether Frabin/Fgd4 is required for the maintenance of properly developed myelin, independent of the role of Frabin/Fgd4 in myelinization during development. To answer this question, we bred mice carrying the conditional Frabin/Fgd4 null allele with an established Plp-CreERT2 mouse line (Leone et al., 2003). In the offspring, Cre-mediated recombination in Schwann cells was induced by tamoxifen injections in young adult Pnp-CreERT2 Fgd4 flox/flox animals (10 weeks old) (Fig. 6A). Specific loss of Frabin/Fgd4 in Schwann cells was verified by

Frabin/Fgd4 controls Cdc42 activity in peripheral nerves

Next, we addressed whether loss of Frabin/Fgd4 alters signalling pathways known to be critical in peripheral nerve myelination (Pereira et al., 2012). In particular, alterations in neuregulin signalling and the AKT pathway have been associated with hypermyelination, myelin outfoldings and demyelination (Cotter et al., 2010; Goebels et al., 2010, 2012). However, we found no significant differences of total and active levels of AKT and ErbB2 and of the amounts of the regulatory phosphatase and tensin homologue (PTEN)-interactor Dlg1 in sciatic nerve lysates derived from 10-week-old Fgd4−/− mice compared with wild-type mice by western blot analysis (Fig. 7A and B). JNK levels and JNK
**Figure 4** *Fgd4*−/− mice develop a progressive demyelinating neuropathy with aberrant myelin formation. (A) Transverse sections of plantaris nerves show a temporally progressive accumulation of aberrant myelin features in *Fgd4*−/− mice compared with wild-type (wt/wt) mice aged 2–80 weeks. (B) Quantification of aberrant myelin features on entire transversal nerve reconstructions at electron microscopy level reveals a significant and progressive increase in the amount of affected fibres in *Fgd4*−/− mice aged 2–80 weeks. (C and D) G-ratio is not significantly changed between wild-type and *Fgd4*−/− mice aged 10 and 60 weeks (at least 100 fibres quantified per animal). (E) Signs of demyelination and remyelination accumulate significantly between the age of 60 and 80 weeks in *Fgd4*−/− mice, the difference to wild-type (wt/wt) mice reaching significance by 80 weeks. (F) No loss in the overall number of myelinated fibres was detectable up to the age of 80 weeks. Arrows indicate fibres displaying myelin alterations. Arrowhead indicates demyelinated fibre. Scale bars = 5 µm; *P < 0.05, **P < 0.01, ***P < 0.001; Student’s t-test (two-tailed). Three animals were analysed for each genotype per time point.
phosphorylation were also not altered. In addition, the amount of myelin basic protein that we examined as a representative myelin protein was unchanged.

On the molecular level, Frabin/Fgd4 has been reported to act as a Cdc42-specific GEF (Umikawa et al., 1999), and loss of Cdc42 in Schwann cells is incompatible with proper developmental myelination (Benninger et al., 2007). Thus, we analysed Cdc42 and found that its active form Cdc42-GTP was substantially and significantly reduced in sciatic nerves of adult Fgd4−/− mice compared with age-matched wild-type mice (Fig. 7C and D). As total levels of Cdc42 remained unchanged, the ratios of active-Cdc42/total-Cdc42 and active-Cdc42/glyceraldehyde-3-phosphate dehydrogenase (as an indirect measure of active Cdc42 per cellular unit) were also reduced. These findings support the hypothesis that Frabin/Fgd4 acts as a GEF for Cdc42 in peripheral nerves in vivo, as RhoGEFs shift the equilibrium between active and total amounts of RhoGTPases towards the active forms (Etienne-Manneville and Hall, 2002).

Cdc42 is required for peripheral nerve myelin maintenance

Our data indicated that Frabin/Fgd4 in Schwann cells is essential for maintenance of proper myelin, and the activation of Cdc42 in Fgd4−/− nerves is strongly decreased. Based on these findings, we hypothesized that if the critical function of Frabin/Fgd4 in myelin maintenance involves correct Cdc42 activation, Schwann cell-specific elimination of Cdc42 at adult stages should also cause myelin deficiencies. To address this question, we combined mice carrying floxed Cdc42 alleles, which were previously used to analyse the role of Cdc42 in Schwann cell development (Benninger et al., 2007), with mice carrying the Plp-CreERT2 transgenic allele. The experimental setting used was analogous to that described earlier for Frabin/Fgd4 deletion in adult myelinating Schwann cells (Fig. 8A). Cdc42 deletion was induced in 10-week-old animals, and western blot analysis revealed subsequent loss of Cdc42 as expected (Fig. 8B). Thereafter, we analysed sciatic nerves of 10-month-old animals by standard electron microscopy and FIB-SEM (serial electron microscopy coupled with in situ focused ion beam milling; Pereira et al., 2010). Prominent myelin foldings were present in nerves with adult-onset Cdc42 deletion (Fig. 8C).
Frabin/Fgd4 regulates endocytosis in Schwann cells

We then asked which cellular functions of Schwann cells might be critically dependent on Frabin/Fgd4. Thus, we analysed migration and process extensions of cultured rat Schwann cells after efficient short hairpin RNA-mediated Frabin/Fgd4 expression knock-down. However, both processes were not altered compared with control cells (data not shown). As myelination depends strongly on endocytosis (Stendel et al., 2010; Sidiropoulos et al., 2012), we also tested whether Frabin/Fgd4 regulates this crucial cellular process. First, we examined whether knock-down of Frabin/Fgd4 affects Cdc42 in cultured Schwann cells. Indeed, reduced Frabin/Fgd4 levels led to lower amounts of active Cdc42 in the rat schwannoma cell line RT4 (Fig. 9A and B), consistent with our in vivo data (Fig. 7C). Next, we examined endocytosis efficiency with an established transferrin uptake assay in Frabin/Fgd4-silenced RT4 cells (Sidiropoulos et al., 2012). Reduced transferrin uptake was found in such cells with ∼20% decrease compared with control cells (Fig. 9C and D). We conclude that regulation of Schwann cell endocytosis by Frabin/Fgd4 is likely to contribute to the disease mechanism in CMT4H.

Discussion

Frabin-deficient mice as an animal model for CMT4H

During the past 15 years, transgenic animal models for inherited peripheral neuropathies have become essential tools for dissecting...
pathological mechanisms involved in various subtypes of Charcot–Marie–Tooth disease (Suter and Nave, 1999; Nave et al., 2007). Such models also provided the basis for evaluating treatment strategies (Fledrich et al., 2012a, b) and contributed significantly to our current understanding of the molecular mechanisms that govern myelination (Pereira et al., 2012). With the Fgd4 knockout mouse line, we present the generation and analysis of an animal model for CMT4H because of loss of Frabin/Fgd4. As in CMT4H patients, Fgd4−/− mice are affected by a recessive, dysmyelinating and demyelinating peripheral neuropathy with early onset and progressive course upon histological examination. Mouse nerves revealed myelin infoldings and outfoldings, redundant myelin loops and signs of demyelination and remyelination, analogous to neuropathological features seen in CMT4H biopsies. Taking advantage of the fact that animal models allow detailed spatial and temporal analyses, we found that both sensory and motor fibres are similarly affected if Frabin/Fgd4 is missing. Moreover, nerve fibres were distally strikingly more affected than proximally, in line with the distally accentuated neuropathy in CMT4H. The reasons for this rather generally observed phenomenon in Charcot–Marie–Tooth disease pose a persisting question, still lacking completely satisfactory answers. Potential explanations include higher susceptibility to mechanical stress of distal fibre segments or transport impairments along the axons that are reflected in this manner. Of major conceptual importance, temporal quantitative analysis of pathological features revealed significant myelin alterations already at early stages of myelination (post-natal Day 5), continuously increasing with age. Thus, loss of Frabin/Fgd4 harms nerves early in development suggesting that early-onset dysmyelination contributes to CMT4H. Furthermore, the observed increase in pathological features with age is consistent with the slowly progressive course of CMT4H. Interestingly, we found the classical hallmarks of demyelinating Charcot–Marie–Tooth disease, demyelination and remyelination associated with reduced nerve conduction velocity, only in older animals. In addition, these features were generally milder than in patients with CMT4H, in agreement with other mouse models for demyelinating Charcot–Marie–Tooth disease showing milder neurological pathology than the corresponding patients (Martini, 2000). This discrepancy is particularly striking when Myotubularin-related protein-2 or Myotubularin-related protein-13/Set-binding factor-2-deficient mice and their human counterparts, the autosomal recessive Charcot–Marie–Tooth disease subtypes CMT4B1 or CMT4B2, are compared (Bolino et al., 2000; Azzedine et al., 2003; Senderek et al., 2003; Bolino et al., 2004; Bonneick et al., 2005; Tersar et al., 2007; Robinson et al., 2008). Incidentally, the neuropathological

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Figure 7 Ablation of Frabin/Fgd4 reduces activation of the RhoGTPase Cdc42 in peripheral nerves in vivo. (A and B) Western blot analyses demonstrating that total and active levels of AKT, ErbB2 receptor and JNK, and total levels of MBP and Dig1, are not changed in sciatic nerve of Fgd4−/− mice compared with wild-type (wt/wt) mice at the age of 10 weeks. (C and D) Active, but not total levels of Cdc42 are significantly reduced in sciatic nerves of adult Fgd4−/− mice compared with age-matched wild-type mice. Tissues from four wild-type and Fgd4−/− mice were analysed. ***P < 0.001; Student’s t-test (two-tailed). GADPH = glyceraldehyde-3-phosphate dehydrogenase.
features in these mutants with regards to the observed myelin abnormalities are virtually identical compared with Frabin/Fgd4 mutants.

Loss of Frabin/Fgd4 function in Schwann cells is sufficient to induce a CMT4H-like phenotype

Demyelinating Charcot–Marie–Tooth disease neuropathies are usually attributed to an initial Schwann cell-specific damage, as myelin constitutes a Schwann cell compartment. However, the possibility of axonal contributions cannot be excluded, as establishment and maintenance of the myelin sheath depend on axonal signals and continuous bidirectional Schwann cell-axon communication (Pereira et al., 2012). We found that Frabin/Fgd4 is expressed in mouse Schwann cells and PNS neurons, compatible with a functional role of Frabin/Fgd4 in both cell types. Thus, we dissected the cell type-specific implications of Frabin/Fgd4 loss by genetically ablating Frabin/Fgd4 exclusively either in Schwann cells or in motor neurons. This experimental strategy also allowed us to determine which cell type is primarily dependent on Frabin/Fgd4 function in myelination. Schwann cell-specific ablation of Frabin/Fgd4 alone was sufficient to fully replicate the myelin aberrations observed in Fgd4−/− mice. In contrast, we found no differences in motor neuron-specific Frabin/Fgd4-deficient mice compared with wild-type mice. Although we cannot fully exclude a subtle role of Frabin/Fgd4 in motor neurons that escaped our attention or species-specific differences in this context, our results suggest that the disease-initiating event in CMT4H is loss of a Schwann cell-specific cellular function of Frabin/Fgd4.

Frabin/Fgd4 function in Schwann cells is distinctively required in myelin maintenance

Establishing the PNS myelin sheath is a complex and expanded process, which roughly starts at birth and is completed in young
adulthood (Jessen and Mirsky, 2005). CMT4H has an early onset and a slowly progressive course, attaining a severe clinical phenotype during the second half of life when myelination has long been completed. Thus, we asked whether Frabin/Fgd4 function is only necessary in development when the myelin sheath is being established. This pre-existing myelin defect would then passively aggravate during adulthood, leading to the observed progressive pathology, without requiring Frabin/Fgd4 function during myelin maintenance. Alternatively, Frabin/Fgd4 function may be actively required for both myelin development and maintenance. If true, lack of Frabin/Fgd4 function in the adult may also contribute to the progressive course of CMT4H. We addressed these mechanistically important questions using a mouse line that allows inducible Schwann cell-specific Frabin/Fgd4 deletion in adult animals. We found that loss of Frabin/Fgd4, induced in young adults when myelination was virtually completed, still resulted in histopathological myelin alterations as seen in Fgd4−/− animals and in CMT4H. Quantitative comparisons revealed only a slightly reduced number of affected fibres compared with age-matched constitutive Fgd4−/− mutants. We conclude that Frabin/Fgd4
function is not only required for correct developmental myelination but also specifically and prominently for myelin maintenance. Thus, one might speculate that a potential therapeutic approach based on reconstitution of Frabin/Fgd4 function in CMT4H might possibly work in adult individuals, as an active process requiring Frabin/Fgd4 in adult myelinating Schwann cells seems to be majorly impaired.

The Frabin–Cdc42 axis in Schwann cells and myelination

Frabin/Fgd4 is a member of the faciogenital dysplasia family of proteins and has been described as a GEF for RhoGTPases, in particular for Cdc42 (Nakanishi and Takai, 2008). The available evidence to support this claim is based mainly on the ability of Frabin/Fgd4 to activate downstream effectors of Cdc42, to influence cell morphology and to recruit Cdc42 to the plasma membrane, when Frabin/Fgd4 was overexpressed in cultured cells (Nakanishi and Takai, 2008). Furthermore, biochemical evidence for a GEF function of Frabin/Fgd4 on Cdc42 has been obtained in a cell-free system (Umikawa et al., 1999). To examine the Frabin/Fgd4–Cdc42 axis in peripheral nerves more specifically, we measured active and total Cdc42 levels in sciatic nerve lysates from wild-type and Fgd4−/− mice. We found a substantial and significant reduction in active Cdc42 when Frabin/Fgd4 was absent, while total Cdc42 levels were not changed. These findings provide evidence for a functional link between Frabin/Fgd4 and active Cdc42 in vivo and further support GEF activity of Frabin/Fgd4 on Cdc42. Although we consider it unlikely, we cannot exclude contributions by indirect activation mechanisms. A definitive analysis would require direct in vivo measurements of specific GTP exchange rates in peripheral nerves. Currently, this procedure is technically not feasible. To further support our findings in other ways, we switched to cell culture. Both in Frabin/Fgd4-silenced RT4 cells and Frabin/Fgd4-silenced primary rat Schwann cells (data not shown), we found prominently reduced active levels of Cdc42 compared with control cells. These data are in agreement with our in vivo results and support the hypothesis that the key molecular function of Frabin/Fgd4 involves activation of Cdc42 in the context of crucial Schwann cell functions. Interestingly, a second GEF for RhoGTPases, ARHGEF10, has also been implicated in the control of myelination. A missense mutation associated with slowed nerve conduction velocity and thin PNS myelin sheaths was found (Verhoeven et al., 2003). In contrast to Frabin/Fgd4 mutations, this mutant seems to act through hyperactivated GEF activity on RhoGTPases, in line with the dominant segregation of reduced nerve conduction velocity within the ARHGEF10 mutation-carrying family (Chaya et al., 2011).

Our data support the hypothesis that the initial trigger in CMT4H is loss of Frabin/Fgd4 in Schwann cells. Frabin/Fgd4 function is still actively required to maintain a correctly structured myelin sheath in the adult and is paralleled by reduced active levels of Cdc42. Indeed, we show that eliminating Cdc42 from adult myelinating Schwann cells phenocopies the consequences of loss of Frabin/Fgd4. These experiments further strengthen the argument for a critical connection between Frabin/Fgd4 and Cdc42 in the PNS throughout life and demonstrate that Cdc42 is required for myelin maintenance, in addition to the critical role of this RhoGTPase and associated signalling pathways in developmental myelination (Benninger et al., 2007; Pereira et al., 2012). Taken together, these results provide strong evidence that the reduction in active Cdc42 levels is at least contributing to the pathology of CMT4H.

In our quest to understand the cellular consequences of the functional link between Frabin/Fgd4 and Cdc42 in Schwann cell biology, we faced the extraordinary pleiotropy of Cdc42 function (Etienne-Manneville and Hall, 2002; Jaffe and Hall, 2005; Melendez et al., 2011). Specificity in cellular function is reached by strict regulation of Cdc42 activity, and GEFs play a decisive role in this regulatory process. They recruit and activate Cdc42 at specific subcellular locations, therefore exerting distinct mechanistic influences on Cdc42 effectors, which finally result in particular cellular functions (Garcia-Mata and Burridge, 2007). For example, the faciogenital dysplasia family member Fgd1, which is mutated in Aarskog–Scott syndrome, a rare X-linked disorder characterized by typical facial dysmorphism and skeletal and genital anomalies (Pasteris et al., 1994), is involved in Cdc42-dependent processes, such as cell migration (Oshima et al., 2011), vesicular transport (Egorov et al., 2009) and podosome formation (Daubon et al., 2011). Similarly, Fgd2 has been linked to Cdc42-dependent vesicle trafficking (Huber et al., 2008), and Fgd3 influences cell motility and cellular morphology in a Cdc42-dependent manner (Hayakawa et al., 2008). Moreover, a unique clinical form of Charcot–Marie–Tooth disease with glomerulopathy results from specific allelic variants of inverted formin 2 (Boyer et al., 2011). This form protein interacts with Cdc42 and myelin and lymphocyte protein), implicated in proper myelination (Schaeren-Wiemers et al., 2004; Buser et al., 2009), suggesting that myelinopathy and glomerulopathy in this special form of Charcot–Marie–Tooth disease may represent particular dysfunction of cell types (Schwann cells and podocytes), both with specialized membrane biology. We reasoned that Frabin/Fgd4 might be involved in similar cellular processes and found that this GEF is required for efficient endocytosis. There is increasing evidence that endocytic transport is critical in myelination. Disturbed Rab11-dependent endocytic recycling is a potential disease mechanism for CMT4C (Lupo et al., 2009; Roberts et al., 2010; Stendel et al., 2010) and dynamin 2 mutations, associated with dominant-intermediate Charcot–Marie–Tooth disease type B, cause disturbed clathrin-mediated endocytosis (Sidirooulos et al., 2012). Cdc42 is involved in endocytosis (Ridley, 2006; Doherty and McMahon, 2009) and most likely enables clathrin-mediated endocytosis (Yang et al., 2001; Bu et al., 2010; Shen et al., 2011) by directed actin polymerization and therefore rearrangement of the actin cytoskeleton in the vicinity of clathrin-coated pits (Kaksonen et al., 2006). We have observed co-localization of Frabin/Fgd4 and Cdc42 after overexpression in RT4 cells (data not shown), consistent with the hypothesis that Frabin/Fgd4 may be involved in recruiting and activating Cdc42 at the plasma membrane of Schwann cells, thereby contributing to the cellular process of endocytosis. Endocytosis is critical for the regulation of protein and lipid homeostasis in the plasma membrane and a defect in endocytosis may result in irregular accumulation of proteins and lipids at the plasma membrane. Thus, we suggest that altered myelin dynamics may contribute
to the myelin pathology observed in Frabin/Fgd4-mutant mice and in CMT4H. Alternatively, receptor molecules may accumulate on the cellular surface resulting in altered cellular signalling, which could be, at least partially, responsible for the observed CMT4H pathology. Nevertheless, given the broad spectrum of functions of Cdc42, other Cdc42-dependent cellular processes are likely to be also involved in CMT4H.

Convergent and divergent signalling pathways and the formation of redundant myelin folds

Some autosomal recessive demyelinating forms of Charcot–Marie–Tooth disease, most prominently CMT4B, CMT4H and to a minor degree CMT4F (culprit gene periaxin; Boerkoel et al., 2001; Takashima et al., 2002; Marchesi et al., 2010), are characterized by comparable histopathological features, including myelin thickening, redundant myelin loops (focally folded myelin with tomacula formation), a characteristic feature also found in hereditary neuropathy with liability to pressure palsy; Suter and Scherer, 2003) and myelin infoldings and outfoldings (Nave et al., 2007). Interestingly, when analysing myelin thickness in animal models for CMT4B and CMT4H, no altered growth ratio of unaffected fibres (fibres without histopathological features) was detected (Bonneick et al., 2005; Tersar et al., 2007; this study). These findings indicate that the neuropathological features in these animal models do not reflect a major overshooting in radial growth of myelin, but rather a lateral surplus of myelin. The observed excess of myelin seemed to arise mainly in the neighbourhood of nodes of Ranvier and Schmidt–Lanterman incisures where the main addition of myelin is thought to occur. This speculation based on histological results raises the question of how a possible lateral surplus of myelin occurs. One possible answer to this question consists in the perspective of persisting myelin growth over the normal level. Myelination in the PNS is stimulated by axonal neuregulin-1 type III through mainly PI3Kinase/AKT and extracellular-signal regulated kinase/mitogen-activated protein kinase signalling (Pereira et al., 2012). Accordingly, mice with myelinating glia-specific loss of PTEN, a lipid phosphatase that inhibits AKT signalling, exhibit mammalian target for rapamycin-dependent overmyelination (Goebbels et al., 2010, 2012). Furthermore, interactions between mammalian Dlg1 and PTEN are required for the stabilization of PTEN acting as myelination brake in Schwann cells to prevent AKT-dependent overmyelination (Cotter et al., 2010). In the PNS of Myotubularin-related protein-2−/− mice, the absence of this myelination brake attained by reduction in Dlg1 levels has been suggested to be causative for the observed pathological over-myelination phenotype (Cotter et al., 2010). In our analysis of Fgd4−/− mice, we did not detect hyperactivation of known myelination-driving pathways. Provided that these results are not because of transient- or low-signal phenomena that escaped detection, we favour the hypothesis that there is molecular heterogeneity in the different pathways leading to hypermyelination, in particular in those cases where aberrant myelin folds are observed. Given the virtual identical pathology in CMT4H and CMT4B, phosphoinositide signalling pathways that are not immediately linked to AKT regulation need to be considered (Suter, 2007). Frabin/Fgd4 contains a FYVE (Fab1, YOTB, Vac1 and EEA1) and two pleckstrin homology domains that most likely bind to phosphoinositides, suggesting potential cooperation with the CMT4B culprits Myotubularin-related protein-2 and Myotubularin-related protein-13/Set-binding factor-2, which regulate synthesis of specific phosphoinositides (Hnia et al., 2012) as does the CMT4J culprit protein FIG4 (Chow et al., 2007).

Conclusions

With the generation and characterization of an animal model for CMT4H, we show that loss of Frabin/Fgd4 causes a recessive, distally pronounced, demyelinating peripheral neuropathy with early onset and progressive course. In addition, we have provided novel insights into cellular and molecular mechanisms altered in CMT4H. This knowledge sets a firm basis to dissect the underlying disease mechanisms further, to elucidate common and divergent pathways leading to hereditary neuropathies, and eventually to provide hints as to novel molecularly targeted treatment strategies.

Acknowledgements

The authors thank the members of the Suter lab for many fruitful discussions, S. Arber for Hb9-Cre mice and D. Meijer for Dhh-Cre mice, J. Collard for reagents and R. Martini for advice. The authors are also grateful for the support from the Electron Microscopy Center of the ETH Zurich and acknowledge the generation of Frabin/Fgd4 mutant mice by the Mouse Clinical Institute (Strasbourg, France).

Funding

The Swiss National Science Foundation and the National Centre of Competence in Research (NCCR), Neural Plasticity and Repair (to U.S.); Research funds of the University of Würzburg Medical Center (to C.W. and K.V.T.); The Deutsche Forschungsgemeinschaft (SE 1839/1-1 to J.S.).

Supplementary material

Supplementary material is available at Brain online.

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