The CMT4B disease-causing proteins MTMR2 and MTMR13/SBF2 regulate AKT signalling

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Abstract
Charcot-Marie-Tooth disease type 4B is caused by mutations in the genes encoding either the lipid phosphatase myotubularin-related protein-2 (MTMR2) or its regulatory binding partner MTMR13/SBF2. Mtmr2 dephosphorylates PI-3-P and PI-3,5-P2 to form phosphatidylinositol and PI-5-P, respectively, while Mtmr13/Sbf2 is an enzymatically inactive member of the myotubularin protein family. We have found altered levels of the critical signalling protein AKT in mouse mutants for Mtmr2 and Mtmr13/Sbf2. Thus, we analysed the influence of Mtmr2 and Mtmr13/Sbf2 on signalling processes. We found that overexpression of Mtmr2 prevents the degradation of the epidermal growth factor receptor (EGFR) and leads to sustained Akt activation whereas Erk activation is not affected. Mtmr13/Sbf2 counteracts the blockage of EGFR degradation without affecting prolonged Akt activation. Our data indicate that Mtmr2 and Mtmr13/Sbf2 play critical roles in the sorting and modulation of cellular signalling which are likely to be disturbed in CMT4B.

Keywords: Charcot-Marie-Tooth disease • neuropathy • myotubularin • EGFR • Akt

Introduction

The family of myotubularin-related proteins (Mtmrs) consists of 14 members in human beings and orthologues have been found in all eukaryotes but not in bacteria. Mutations in three family members are associated with human diseases. Mutations in myotubularin, the founding member of the Mtmr family, lead to X-linked myotubular myopathy [1]. X-linked myotubular myopathy is a severe muscle disease characterized by hypotonia and generalized muscle weakness in affected newborn males. Histopathological studies of the skeletal muscle in these patients reveal the presence of small rounded muscle fibres that contain centrally located nuclei. Since these fibres resemble foetal myotubes, it was proposed that the terminal differentiation of the muscle fibre is blocked [2]. Mutations in MTMR2 or MTMR13/SBF2 lead to Charcot-Marie-Tooth (CMT) diseases type 4B1 or CMT4B2, respectively. These recessive peripheral neuropathies are clinically indistinguishable and are characterized by focally folded myelin sheets and demyelination in peripheral nerves [3–5].

The Mtmr proteins are defined by a phosphoinositide-binding GRAM-pleckstrin homology domain (GRAM-PH), a phosphatase domain and a coiled coil. Active members dephosphorylate PI-3-P to phosphatidylinositol and PI-3,5-P2 to PI-5-P. Although initially described as a protein tyrosine/dual specificity phosphatase based on the primary sequence, no phosphoprotein has been identified as substrate so far [6]. Mtmr2, an active member of the family, is a 73 kD protein that exists as a dimer in eukaryotic cells. It contains a C-terminal Post-synaptic density protein-95, Drosophila disc large tumour suppressor, and Zonula occludens-1 protein (PDZ) binding motif in addition to the standard Mtmr domains. The GRAM-PH domain binds to several phosphoinositides and thereby mediates membrane binding [7]. Interestingly, six of the 14 members have substitutions in the catalytic site of the phosphatase domain rendering them inactive. Several inactive myotubularins interact with active Mtmrs [8]. Alternatively, inactive Mtmrs can also exist independently within cells [9, 10]. Mtmr13/Sbf2, an inactive member, is a 210 kD protein that interacts with Mtmr2 and enhances its catalytic activity. The formation of this complex is crucial for the integrity of the peripheral nervous system since loss of either of the proteins leads to CMT4B. Mtmr2 contributes the disease-relevant catalytic activity to this complex while the important functions or domains of Mtmr13/Sbf2 have not yet been identified [11]. It is likely that Mtmr13/Sbf2 guides the catalytic activity to the appropriate place within the cell. Putative domains for this purpose are the N-terminal differentially expressed in neoplastic versus normal cells (DENN) domain which influences the proliferation of cells or the C-terminal PI-3,4,5-P3 binding pleckstrin homology domain [12]. Mtmr12/3-PAP, another
inactive member, also interacts with Mtmr2 [13]. In contrast to Mtmr13/Sbf2, Mtmr12/3-PAP contains only the standard Mtmr domains indicating that the functional mechanisms are likely to be different from Mtmr13/Sbf2.

The correct metabolism of phosphoinositides is a critical issue for the integrity of peripheral nerves [14]. Several disease-associated proteins have phosphoinositide binding domains (MTMR2, MTMR13/SBF2, DYNAMIN2, N-myc downstream regulated gene 1 [NDRG1] and FRABIN) or dephosphorylate phosphoinositides as MTMR2 and the Sac type phosphatase FIG4 (pheromone-regulated or induced gene 4) [15–19]. Phosphoinositides are fixed to membranes by acyl chains and expose their phosphorylated head to the cytoplasm. Seven different phosphoinositides are generated by the reversible phosphorylation and dephosphorylation of the inositol ring of phosphatidylinositol at positions D3, D4 and D5. Particular phosphoinositides are enriched in certain subcellular compartments. The exposed head groups are anchor sites for phosphoinositide-binding domains which allow the assembly of multiprotein effector complexes with compartment-specific cellular functions [20, 21]. PI-3-P and PI-3,5-P2, the substrates of Mtmr2, localize mainly to the endosomal compartment pointing to a function in protein trafficking, sorting and degradation. PI-3,5-P2 is produced from PI-3-P by PI3Ktye, an enzyme that is found on early and late endosomes [22, 23]. It is transiently formed after the stimulation of cells with different treatments [24–26]. Several effector proteins including Mtmrs, sorting nexins (SNX), Synp1p and vps24 can bind to PI-3,5-P2 with low affinity and specificity [27–29]. Depending on the bound effector, the cargo of the corresponding vesicle is either directed to the lysosome or to the trans-Golgi network [30, 31].

This report is based on the observation that the overexpression of myotubularin blocks the degradation of the epidermal growth factor receptor (EGFR) [26]. We show that Mtmr2 has a comparable effect and that the adaptor unit Mtmr13/Sbf2 can counteract this outcome. In addition, we demonstrate that the blocked degradation leads to sustained Akt, but not Erk activation. Mtmr13/Sbf2 can counteract this effect if its C-terminal PI-3,4,5-P3 binding pleckstrin homology domain is removed. Active and inactive Mtmrs therefore integrate the PI-3-kinase-derived PI-3,4,5-P3 and the PIKfyve-derived PI-3,5-P2 signal to generate sustained Akt, but not Erk activation. Mtmr13/Sbf2, Mtmr12/3-PAP contains only the standard Mtmr domains indicating that the functional mechanisms are likely to be different from Mtmr13/Sbf2.

EGFR degradation

Three 10 cm dishes were transfected with 2 μg pcDNA-hEGFR plasmid (kindly provided by Gordon Gill, San Diego) and 6 μl Fugene6. One day after transfection, cells were distributed in twelve 3.5 cm dishes for triplicates at four time-points. The transfection rate was between 10% and 20% and did not vary between cell lines. After an additional 24 hrs in culture, the cells were starved overnight in 1% bovine serum albumin (BSA)/Dulbecco’s modified Eagle medium (DMEM)/Penicillin/Streptomycin. Cells were stimulated with 50 ng/ml EGF for 0/20/40/180 min. and then lysed in 0.5% Triton-X100/100 mM NaCl/50 mM Tris-HCl pH 7.5. The supernatant was used for Western blot analysis after sonication and centrifugation. The following antibodies were used: 1:1000 rabbit anti-EGFR (Neomarkers, Fremont, CA, USA; RB-1417); 1:2000 mouse anti-actin (Sigma, Buchs, Switzerland; A5316), rabbit anti-Akt (Cell Signaling, Danvers, MA, USA; 9272), mouse anti-P-Akt (Cell Signaling; 4051), rabbit anti-Erk (Cell Signaling, 9102), mouse anti-P-Erk (Cell Signaling; 9106). As secondary antibodies, alkaline phosphatase-coupled goat anti mouse and horse radish peroxidase (HRP)-coupled goat anti-rabbit were used, followed by chemiluminescence detection. Quantification was performed with ImageJ. The experiment was repeated three times for each cell line.

HHR uptake

FlpIn293 cells stably expressing Mtmrs were plated in triplicates in 3.5 cm dishes 2 days prior to the experiment. Cells were starved for 30 min. in starvation medium (1% BSA/DMEM) and then pulsed with 2 mg/ml HRP in starvation medium for the indicated time. Cells were immediately put on ice, washed two times with ice cold PBS/1% BSA followed by three washes with ice cold PBS and then scraped from the plate with PBS. Cells were lysed by sonication in 0.15% Triton-X100/PBS/protease inhibitors and centrifuged. HRP activity was measured with o-Dianisidine and H2O2 and normalized to the total protein content.

Immunofluorescence microscopy

COS cells were transfected in 3.5 cm dishes with Fugene6 or FugeneHD according manufacturers recommendations. Equal amounts of Mtmr plasmids (His-Mtmr2 or HA-Sbf2) and myc-tagged SNX1/2/5 (kindly provided by Jo-Ann Trejo, Chapel Hill and Rohan Teasdale, St. Lucia) were taken for the transfections. The cells were fixed 24 or 36 hrs after transfection with 2% paraformaldehyde in PBS. Cells were blocked in blocking buffer (10% foetal calf serum/0.05% Saponin/PBS) for 30 min. prior to incubation with antibodies. Antibodies were diluted as follows in blocking buffer: rabbit anti-Mtmr2 (1:1000; [11]), rabbit anti-Mtmr13/Sbf2 (1:1000; [10]), mouse anti-myc 9E10 (1:10) and rabbit anti-EGFR (Neomarkers, RB-1417, 1:500).

Materials and methods

Plasmids and cell lines

The previously described His-tagged Mtmr2 and HA-tagged Sbf2 cDNAs were transferred from pcDNA3.1 zeo(+) into pcDNA5/FRT [10]. These constructs were then used to generate stable cell lines with the Flp-in system according to manufacturer’s recommendations (Invitrogen, Paisley, UK). Two independent lines were used for the experiments (except for Mtmr2 CA17S). Transient transfection was not used since the cotransfection efficiency with these long plasmids was too low. Cell lines expressing two Mtmrs were obtained by random integration of a second, pcDNA3.1-based construct and selection with zeocin. Cell lines were checked by immunofluorescence for homogenous expression. More than 95% of the cells express the indicated Mtmrs (see Fig. S1).
Appropriate Cy3 and ALEXA-488 secondary antibodies were used for visualization. Single optical sections with a thickness of 0.3 μm were taken with a confocal laser scanning microscope (Leica SP2 or SP5, Leica Microsystems, Wetzlar, Germany) for the analysis. Pictures of Figs 1 and S1 were taken with an Olympus IX81 microscope (Olympus Corporation, Tokyo, Japan).

Transgenic mice

Animal experiments were approved by the veterinary office of the Canton of Zurich. The generation and genotyping of Mtmr2 and Mtmr13/Sbf2 knock-out mice was described previously [32, 33]. Sciatic nerves from three adult animals (P60) were pooled, powdered in liquid nitrogen, and then lysed in 0.5% Triton-X100, 100 mM NaCl, 50 mM Tris-HCl pH 7.5. Twenty micrograms protein was used for Western blot analysis as described above.

Results

Influence of Mtmr overexpression on receptor trafficking and fluid phase endocytosis

The degradation of the EGFR is a well-established paradigm for the switching off of activated receptors. A previous study revealed that overexpression of myotubularin blocks the degradation of the EGFR after ligand stimulation [26]. We were interested if Mtmr2 and Mtmr13/Sbf2 have a similar effect. We therefore generated FlpIn293 cell lines stably overexpressing Mtmr2 and/or Mtmr13/Sbf2 or mutants thereof to manipulate the intracellular levels and ratios of these enzymes. We did not use an RNAi approach because compensatory effects of the highly homologues and ubiquitously expressed proteins Mtmr1 and Mtmr5/Sbf1 may interfere with the effect and we used stable cell lines to overcome the problems of transient transfections (see ‘Materials and methods’ for details). The expression levels of Mtmr2 and its mutated forms were always higher than the expression level of Mtmr13/Sbf2 (based on Coomassie stained immunoprecipitation experiments and Western blotting, data not shown).

Overexpression of Mtmr2 blocked the degradation of the EGFR in a similar way as previously described for myotubularin (Fig. 1A, B). Immunostaining of these cells revealed an enhanced surface staining of the EGFR indicating that the EGFR is either not internalized or more efficiently recycled (Fig. 1C). This effect depends on the phosphatase activity since an Mtmr2 mutant lacking the phosphatase activity (C417S) has no effect. Similarly, overexpression of Mtmr13/Sbf2, a phosphatase-dead member of the myotubularin family, has also no effect on degradation. Interestingly, EGFR degradation is not blocked in cells overexpressing both Mtmr2 and Mtmr13/Sbf2 indicating that Mtmr13/Sbf2 acts as an antagonist of Mtmr2. This antagonistic function seems to be mediated by the N-terminal Mtmr2-binding part of Mtmr13/Sbf2, since the deletion of the C-terminal pleckstrin homology domain of Mtmr13/Sbf2 does not reverse this dominating effect. Mtmr2 has therefore the same effect on EGFR degradation as myotubularin and this effect is regulated by the adaptor unit Mtmr13/Sbf2.

Next, we were interested whether Mtmrs influence only the sorting of membrane proteins or if fluid phase endocytosis is also affected. Fluid phase endocytosis was monitored by measuring the uptake of HRP, a marker that reaches the lysosomes by endocytosis. We did not observe significant differences in HRP uptake by overexpressing Mtmrs indicating that general vesicle trafficking is not affected (Fig. 2). This is in line with the finding that overexpression of Mtmrs does not induce vacuoles under resting condition, a
Mtmrs do not change fluid phase uptake. Stably transfected FlpIn293 cells were pulsed with HRP and its uptake was determined after 0/1/10/60 min. (mean ± S.D., n = 3).

**Colocalization with sorting nexins**

SNX are a family of proteins which bind to receptor tyrosine kinases and phosphoinositides [35]. Several members of this family regulate EGFR degradation in a positive or negative manner. They are therefore a putative link between the phosphoinositide-metabolizing Mtmrs and the observed block in EGFR degradation. Consequently, we tested if Mtmr2 and Mtmr13/Sbf2 colocalize with SNX. COS cells were cotransfected either with Mtmr2 or Mtmr13/Sbf2 and with SNX1, SNX2 or SNX5. The phosphoinositide binding properties and the influence on EGFR degradation of these SNX are well established. Whereas overexpression of SNX1 and SNX2 promote EGFR degradation, overexpression of SNX5 plays an inhibitory role. The phosphoinositide binding specificities of these SNX are broad, but they bind to PI-3,5-P2, a substrate of Mtmr2. SNX5 also binds to PI-5-P, which can be produced by Mtmr2 [30, 36, 37]. Mtmrs and SNX show a diffuse cytoplasmic, vesicular or plasma membrane staining under resting conditions (Fig. S2). Cells were stimulated with EGF and analysed after 30 min. when the EGFR passes the endosomal compartment and when PI-3,5-P2 peaks [26]. The cells respond at this time-point with an enlargement of endosomes allowing an accurate colocalization analysis. Mtmrs are able to colocalize with SNX suggesting that they influence sorting of receptors by modulating the phosphoinositide composition of membranes (Fig. 3).

**Influence on downstream signalling**

Stimulation of the EGFR leads to a transient activation of the Akt and Erk pathways. We wanted to test if the blocked degradation has consequences on signal output by the EGFR. We observed a transient phosphorylation of Akt, Erk1 and Erk2 in wild-type FlpIn293 cells after 20 and 40 min. (Fig. 4). The phosphorylation decreases to basal levels after 180 min. In contrast, sustained activation of Akt was observed in FlpIn293 cells overexpressing Mtmr2. Sustained phosphorylation of Akt depends on phosphatase activity since the phosphatase-dead mutant of Mtmr2 (C417S) and Mtmr13/Sbf2 have no effect on signal output. The effect is specific for Akt since Phospho-Erk levels decrease to basal level as in wild-type cells. Interestingly, the sustained Akt activation was also observed in cells overexpressing Mtmr2 and Mtmr13/Sbf2 together which degrade the EGFR with normal efficiency (Fig. 1). These findings suggest that sustained Akt activity is not simply due to the lack of EGFR degradation but due to specific sorting. If Mtmr2 is overexpressed with a form of Mtmr13/Sbf2 which lacks the C-terminal PI-3,4,5-P3 binding pleckstrin homology domain, phospho-Akt returns to basal level after 180 min. similar to wild-type cells. We conclude that sustained Akt activation is only achieved when Mtmr2 phosphatase activity and the PI-3,4,5-P3-binding pleckstrin homology domain of Mtmr13/Sbf2 are available.

**Signalling in knockout animals**

We were also interested if the observed in vitro influence of Mtmr2 overexpression on Akt signalling is relevant in vivo. For this purpose, we used our previously described Mtmr2 and Mtmr13/Sbf2 knockout animals that, in contrast to the acute stimulation experiments described so far, reflect a steady state condition. These animals show a similar but milder pathology than human beings affected with CMT 4B1 or CMT 4B2 [32, 33]. The sciatic nerves of these animals were analysed because the peripheral nervous system is the only affected tissue identified so far. We expected a down-regulation of phospho-Akt in mice lacking both Mtmr2 and Mtmr13/Sbf2 since Mtmr2 overexpression leads to sustained Akt activation. Indeed a reduction of the steady state ratio of phospho-Akt to total Akt was observed in the mutant. The total level of phospho-Akt in mutant animals, however, is comparable to the levels in wild-type animals while the level of total Akt is significantly increased (Fig. 5). Mutant mice therefore appear to compensate for the inefficient phosphorylation of Akt by up-regulating the total level of this kinase. Neither total-Erk nor phospho-Erk differ between wild-type and mutant mice. Akt is a complex regulator of peripheral nerve development and function [38]. How our data integrate with the critical role of Akt in this tissue remains to be elucidated.

**Discussion**

Mtmrs play a crucial role in human health and disease. Active family members dephosphorylate PI-3-P and PI-3,5-P2 in vitro and in vivo, but the consequences for the phosphoinositide-regulated cellular processes are poorly understood. We have overexpressed Mtmr2 to negatively regulate the levels of these two phosphoinositides in vivo. A more specific regulation is not possible since there is no mutant Mtmr2 available which specifically
dephosphorylates only one of the two substrates. Thus, whether PI-3-P or PI-3,5-P2 or both are the relevant substrates \textit{in vivo} cannot be distinguished. Whereas the level of PI-3-P is high within cells, the level of PI-3,5-P2 is low under resting conditions and increases transiently after stimulation of cells [24–26, 39]. The observation that Mtmr2 relocalizes within cells under conditions with elevated PI-3,5-P2 levels, \textit{e.g.} under hypoosmotic conditions or after EGFR stimulation, point to PI-3,5-P2 as an important \textit{in vivo} substrate [10, 26]. In addition, one might speculate that the binding pocket for the D5 phosphate would have been closed during evolution if the activity towards PI-3,5-P2 is not necessary \textit{in vivo}. This has been partially achieved \textit{in vitro} with the artificial mutation H357N in Mtmr2 which reduces the activity towards PI-3,5-P2 but not towards PI-3-P [40].

\bf{Fig. 3} Colocalization of Mtmr2 or Mtmr13/Sbf2 with SNX1, SNX2 or SNX5. COS cells were cotransfected with the indicated plasmids, starved overnight 24 hrs later and then pulsed for 20 min. with EGF. Colocalization on vesicles was seen for all combinations of Mtmrs with SNX. Note that Mtmr13/Sbf2 localizes only to subdomains of vesicles (arrows). Scale bars: 20 \mu m (upper panel), 4 \mu m (lower panel).
Mtmr2 and Mtmr13/Sbf2 can acquire three different structural states in vitro and in vivo which affect the signalling of cell in different ways: (i) as a Mtmr2 dimer, (ii) as a Mtmr13/Sbf2 dimer and (iii) as a Mtmr2//Mtmr13/Sbf2 heterotetramer. Our results indicate, that the concentration, ratios and localization of these two Mtmrs have an important impact in which compartments the receptors are guided. Overexpression of Mtmr2 blocks the degradation of EGFR. Since this effect depends on phosphatase activity, it can either result from (i) the dephosphorylation of PI-3-P to phosphatidylinositol (ii) the dephosphorylation of PI-3,5-P2 or (iii) the production of PI-5-P. PI-3,5-P2 is produced in the endosomal compartment and assembles protein complexes for the trafficking into other subcellular locations of the cell [41, 42]. Vps24 is part of the ESCRT-III complex and one of these adaptors [29]. Previous studies revealed that the knockdown of vps24 with RNAi or the overexpression of dominant-negative forms of vps24 block the degradation of the EGFR [43, 44]. PI-3,5-P2 is therefore an important metabolite for the degradation of the EGFR and PI-3,5-P2 dephosphorylation by Mtmr2 is one of the ways to interfere with EGFR degradation. PI-5-P is the least characterized phosphoinositide. SNX5 has a PX domain that binds PI-5-P. Interestingly, overexpression of SNX5 prevents the degradation of the EGFR indicating that it guides the EGFR away from lysosomes [37]. Consequently, reduction of PI-3,5-P2 levels and the production of PI-5-P are both possible explanations for the observed block in EGFR degradation.

It is therefore reasonable to propose that the phosphatase activity of Mtmrs block downstream pathways depending on PI-3,5-P2 and redirects receptors to pathways depending on PI-5-P (Fig. 6). Overexpression of Mtmr13/Sbf2 can counteract this function of Mtmr2. We have previously described that Mtmr13/Sbf2 exists in double transfected cells in a complex with Mtmr2 or separately [10]. It is therefore not clear if Mtmr13/Sbf2 fulfils this counteracting function in a complex with Mtmr2 or independently.

Overexpression of Mtmr2 leads to sustained Akt activation. Surprisingly, this effect on downstream signalling of the EGFR is limited to Akt whereas Erk shows the normal transient activation observed in growth factor stimulated cells. The same specific activation pattern was observed after infection of HeLa cells with *Shigella flexneri*. Phosphorylation of Akt depends in this case on IpgD, a phosphatidylinositol-4,5-bisphosphate-4-phosphatases which is secreted into the cytoplasm of host cells leading to increased PI-5-P levels [45]. The generation of PI-5-P from PI-4,5-P2 seems to be a widespread mechanism of pathogenic bacteria to activate the Akt pathway to increase survival of host cells [46]. Bacteria probably use PI-4,5-P2 as a substrate because its level is constitutively high within cells, whereas PI-3,5-P2 levels are low.
and only available transiently. Enhanced PI-5-P levels were also observed when 
myotubularin was overexpressed in Jurkat cells [47]. Mtmm13/Sbf2 appears to play a different role in downstream signalling than in receptor degradation. Although Mtmm13/Sbf2 counteracts the protective function of Mtmm2 in EGFR degradation, it does not interfere with sustained Akt activation. Sustained Akt activation is therefore due to different trafficking of the receptor and not due to increased levels of the EGFR. The permissive function on sustained MTMR2-induced Akt activation of Mtmm13/Sbf2 is lost, when its C-terminal pleckstrin homology domain is deleted. This suggests that the sustained activation of Akt by the Mtmm2 // Mtmm13/Sbf2 complex is only possible when PI-3,4,5-P3-binding pleckstrin homology domain of Mtmm13/Sbf2 is present. PI-3,4,5-P3 as well as PI-3,5-P2 are available after stimulation of the EGFR [26, 48]. Mtmm2 and Mtmm13/Sbf2 recognize both lipids with GRAM-PH domain and pleckstrin homology domains and are able to modulate the PI-3,5-P2 level in the endosomal compartment (Fig. 6). Receptors may recognize the phosphoinositide composition of membranes indirectly via their bound SNX and are then transported to distinct compartments where they fulfill their function. Mtmm13/Sbf2 is a catalytically inactive Mtmr. Dead Mtmrs emerged only in higher eukaryotes and do not exist in yeast [6]. PI-3,4,5-P3 and Akt signalling are also absent in yeast suggesting that the function of dead Mtmrs might have coevolved with PI3-kinase signalling in higher eukaryotes.

What are the potential implications for human diseases? CMT4B is a severe demyelinating peripheral neuropathy with myelin outfoldings as the pathological hallmark [49]. The overall structure and thickness of the myelin sheath appear normal in patients and in animal models pointing to an increased synthesis or impaired membrane degradation in regions of active myelin turnover. Here we show that Mtmrs have an influence on protein turnover in vitro. In the peripheral nerve, Mtmrs may either be involved in the turnover of structural myelin proteins or indirectly influence the signal output of an unidentified receptor. Pharmacological targeting of Mtmm2 or Mtmm13/Sbf2 might be promising since curcumin, an inhibitor of Pikfyve, shows a positive effect in some animal models of peripheral neuropathies. Curcumin prevents the accumulation of mutant
peripheral myelin protein 22 (PMP22) or protein zero (P0) in the endoplasmic reticulum in vitro suggesting that the levels of PI-3-P and/or PI-3,5-P2 play a general role in the development of peripheral neuropathies [50–52]. Future studies may elucidate which disease-associated proteins play a role in this pathway leading to peripheral neuropathies (reviewed in [53, 54]).

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

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