Mitofusin 2 is necessary for transport of axonal mitochondria and interacts with the Miro/Milton complex

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Mitofusins (Mfn1 and Mfn2) are outer mitochondrial membrane proteins involved in regulating mitochondrial dynamics. Mutations in Mfn2 cause Charcot-Marie-Tooth disease (CMT) type 2A, an inherited disease characterized by degeneration of long peripheral axons, but the nature of this tissue selectivity remains unknown. Here, we present evidence that Mfn2 is directly involved in and required for axonal mitochondrial transport, distinct from its role in mitochondrial fusion. Live imaging of neurons cultured from Mfn2 knock-out mice or neurons expressing Mfn2 disease mutants shows that axonal mitochondria spend more time paused and undergo slower anterograde and retrograde movements, indicating an alteration in attachment to microtubule-based transport systems. Furthermore, Mfn2 disruption altered mitochondrial movement selectively, leaving transport of other organelles intact. Importantly, both Mfn1 and Mfn2 interact with mammalian Miro (Miro1/Miro2) and Milton (OIP106/GRIF1) proteins, members of the molecular complex that links mitochondria to kinesin motors. Knockdown of Miro2 in cultured neurons produced transport deficits identical to loss of Mfn2, indicating that both proteins must be present at the outer membrane to mediate axonal mitochondrial transport. In contrast, disruption of mitochondrial fusion via knockdown of the inner mitochondrial membrane protein Opa1 had no effect on mitochondrial motility, indicating that loss of fusion does not inherently alter mitochondrial transport. These experiments identify a role for mitofusins in directly regulating mitochondrial transport and offer important insight into the cell type specificity and molecular mechanisms of axonal degeneration in CMT2A and dominant optic atrophy.

Introduction

As principal mediators of ATP production and calcium buffering, mitochondria actively distribute to areas of high energy demand and calcium flux (Berthold et al., 1993; Li et al., 2004; Misgeld et al., 2007). Based on this principle, disruption of axonal transport has long been proposed as a potential mechanism in human neurodegenerative diseases (Duncan and Goldstein, 2006; De Vos et al., 2008).

Mitochondrial transport is regulated by a series of molecular adaptors that mediate the attachment of mitochondria to molecular motors (Hollenbeck and Saxton, 2005; Kang et al., 2008; Li et al., 2009). In Drosophila, connection of mitochondria to kinesin motors involves the outer mitochondrial membrane (OMM) protein dMiro, which indirectly attaches to kinesin heavy chain via the adaptor protein Milton (Guo et al., 2005; Glater et al., 2006). In mammals, two isoforms of Miro (Miro1 and Miro2) and Milton (OIP106 and GRIF1) have been identified and are proposed to act in a similar manner (Fransson et al., 2006).

In addition to constant movement, mitochondria also undergo frequent fission and fusion events (Chan, 2006). The mitochondrial fusion apparatus consists primarily of mitofusin 1 and mitofusin 2 (Mfn1 and Mfn2) located on the OMM, and Opa1 located on the inner mitochondrial membrane (Rojo et al., 2002; Chen et al., 2003; Eura et al., 2003; Wong et al., 2003; Cipolat et al., 2004). Both the mitofusins and Opa1 are dynamin family GTPases and are believed to tether opposing membranes and facilitate their fusion. Although the precise reason for mitochondrial fusion remains unclear, interruption of mitochondrial fusion (via loss of Opa1 or Mfn1/2) leads to loss of mitochondrial potential and diminished oxidative phosphorylation, indicating it is critical for maintaining mitochondrial function (Olichon et al., 2003; Chen et al., 2005).

Dominantly inherited point mutations in MFN2 are the most commonly identified cause of axonal Charcot-Marie-Tooth (CMT) disease (designated CMT2A); however, it is still not clear how mutations in this ubiquitously expressed protein lead to relatively selective degeneration of long peripheral axons (Zuchner et al., 2004; Lawson et al., 2005; Verhoeven et al., 2006). Mutations in OPA1 disrupt mitochondrial fusion and lead to degeneration of optic nerve axons, referred to as dominant optic atrophy (Votruba et al., 2003; Newman, 2005). In some cases, CMT2A mutant forms of MFN2 are unable to mediate mitochondrial fusion in fibroblasts, whereas others are completely competent to promote fusion (Detmer and Chan, 2007). Interestingly, exogenous expression of disease mutant forms of MFN2 in cultured sensory neurons leads to a profound abnormality mitochondrial transport (Baloh et al., 2007), and mice overexpressing disease
mutant MFN2 in motor neurons show markedly abnormal axonal mitochondrial distribution (Detmer et al., 2008). These data support that MFN2 influences mitochondrial transport; however, the molecular basis of this effect has not been defined, and it has been observed only in the context of MFN2 overexpression.

Here, we show that loss of Mfn2 profoundly and selectively disrupts axonal mitochondrial transport and that Mfn2 directly interacts with OMM proteins and adaptors involved in mitochondrial transport. Furthermore, we show that disrupting mitochondrial fusion via Opal1 knockdown does not itself disrupt axonal mitochondrial transport. These data indicate that Mfn2 plays an integral role in the regulation of mitochondrial transport and has important implications for understanding the pathophysiology of CMT2A and dominant optic atrophy.

Materials and Methods

Plasmids and silencing RNAs. Human wild-type MFN2 (wtMFN2) and MFN2 disease mutant constructs were previously described (Baloh et al., 2007). For small interfering RNA (siRNA) constructs, targeting sequences were generated using the Dharmacon siDesign Center and cloned into the H14 vector (Dharmacon) according to the manufacturer’s specifications to introduce a mitochondria-targeting sequence. All transfections were performed using the Dharmafekt transfection reagent (Dharmacon), and plated directly onto 24-well plates coated with laminin and poly-D-lysine. Cultures were maintained in Neurobasal medium with 2% supplement (Invitrogen), 50 ng/ml NGF, 5 mM glutamine, and antibiotics. Addition of 5-fluorouracil for the first 4 d in culture was used as an antimiticotic treatment.

Lentivirus production and infection. Lentiviruses were produced as described previously (Baloh et al., 2007). Briefly, human embryonic kidney (HEK) 293T cells were plated onto six-well plates and transduced using Mirus reagent with a packaging vector (Δ8.91), envelope vector (vesicular stomatitis virus-glycoprotein), and transfer vector encoding the gene to be expressed. Media was changed once at 12 h and collected at 24 and 48 h, pooled, and applied directly to DRG cultures resulting in >95% infection efficiency.

Transfection of DRG cultures. To label mitochondria in a small subset of neurons within a culture, Lipofectamine 2000 (Invitrogen) was used according to the manufacturer’s specifications to introduce a mitochondria-targeted DsRed. Briefly, 0.4 μg of plasmid DNA diluted in 25 μl of transfection reagent diluted in 25 μl. The mixture was incubated for 20 min and added to cells. Medium was changed 6 h later to remove the transfection reagent.

Measurement of mitochondrial transport. For imaging of rat DRG neurons expressing Mfn2 wild-type or disease constructs or silencing RNA constructs, cultures were infected at 4 d in vitro (DIV4) and subsequently transfected with mitochondrial marker on DIV7. Imaging of mitochondrial movements was possible on DIV10–13. All imaging was performed in a climate-controlled chamber (In Vivo Scientific) at 37°C and 5% CO2 and images were acquired with a Cool Snap HQ2 CCD camera (Photometrics) mounted on a Nikon Eclipse Ti-U microscope. Images were acquired at 40× magnification every 1 s for 5 min. Kymographs were then generated and analyzed using MetaMorph software (Molecular Devices). Mitochondria were classified as either moving or stationary based on whether they achieved a displacement of >2 μm. Constant velocity movement segments identified on kymographs were measured for mitochondria that were classified as moving. A velocity cutoff of 0.1 μm/s was established, and all movements below this velocity were treated as time spent paused. Measurements were pooled, and the time that mitochondria spent moving at given velocities was binned and represented on a histogram as a percentage of the total time that mitochondria spent moving above the 0.1 μm/s
either endosomes or peroxisomes in single axons of DRG neurons, we performed sequential time-lapse imaging of mitochondria and other organelles accompanying kymographs were processed using the “emboss” filter in Adobe Photoshop for clarity.

Coimmunoprecipitation. HEK 293T cells were transfected using Mirus with constructs containing His-Mfn1, EGFP-Mfn2, myc-Miro1, myc-Miro2, Flag-GRIF1, Flag-OIP106, or Bluescript plasmid as a vector control. Cells were lysed in 1% Triton X-100, 5 mM EDTA, 300 mM NaCl, and 50 mM Tris-HCl, pH 7.5 (Glater et al., 2006), with complete protease inhibitor mixture (Roche). Cells were lysed at 4°C for 30 min and clarified by spinning at 10,000g for 30 min, and protein concentration was determined using the BCA protein assay (Thermo Scientific). Equal amounts of protein from each lysate were resolved on SDS-PAGE gel, transferred to nitrocellulose membranes, and probed with antibodies to Mfn2 (Sigma) or anti-Mfn1 (Novus Biologicals), at a 1:1000 dilution in 5% nonfat milk/TBS-Tween 20.

Results
CMT2A-associated MFN2 mutants disrupt both anterograde and retrograde mitochondrial transport
Previous studies indicate that CMT2A-associated MFN2 mutants produce a marked decrease in overall mitochondrial mobility in axons of cultured sensory neurons (Baloh et al., 2007), and alter mitochondrial distribution in motor axons of transgenic mice (Dettmer et al., 2008). To clearly define the abnormality in mitochondrial transport observed in CMT2A disease mutants, we introduced wtMFN2 or mutant MFN2 constructs using lentivirus (>99% infection) that were expressed at equal levels (supplemental Fig. S1, available at www.jneurosci.org as supplemental material), followed by transfection with a mitochondrially targeted RFP that labels only a small number of neurons, allowing precise analysis of anterograde and retrograde movements in single axons. In wtMFN2-expressing cells, kymograph analysis of mitochondrial movements consistently depicted fast persistent movements in both anterograde and retrograde directions, accompanied by slower moving and stationary mitochondria. In contrast, CMT2A disease mutant MFN2 (R94Q)-expressing neurons showed a striking absence of the fast persistent movements, with the amount of time spent paused between anterograde and retrograde movements significantly greater in mutant-expressing neurons than in wtMFN2-expressing controls (Fig. 1A, B). Additionally, mitochondria from mutant-expressing neurons moved at slower velocities in the anterograde and retrograde directions (Fig. 1C). These findings indicate that mitochondria in MFN2 mutant-expressing cells were unable to either initiate or sustain fast processive movements, suggesting a disruption of microtubule-based mitochondrial transport. We also observed that expression of the R94Q mutant produced smaller fragmented axonal mitochondria, consistent with the previously reported inability of this mutant to mediate fusion in fibroblasts (Dettmer and Chan, 2007). Together, these findings indicate that mutant MFN2 expression influences both transport and fusion of axonal mitochondria.

CMT2A-associated MFN2 mutants do not alter transport of other organelles
To determine whether this was a direct effect on mitochondrial mobility or a global disruption of all organelar transport, we performed sequential time-lapse imaging of mitochondria and either endosomes or peroxisomes in single axons of DRG neurons expressing wtMFN2 or MFN2 disease mutants (R94Q and H361Y) (Fig. 2). Axons expressing wild-type or mutant MFN2 were coinfected with a mito-RFP (A) and RhoB-GFP (B) mitochondrial and endosomal markers, respectively. Mutant-expressing neurons revealed diminished mitochondrial mobility (A, kymograph) in the same axons that showed normal endosomal transport (B, kymograph). C, D, The percentage of mobile mitochondria is significantly decreased in R94Q- and H361Y- compared with wtMFN2-expressing neurons (*p < 0.001), whereas the percentage of mobile endosomes in these axons was normal. E, F, Overlay images of phase contrast and EGFP-SKL (which labels peroxisomes) in wtMFN2 (E) and H361Y (F)-expressing DRG axons. G, Similar to endosomes, the percentage of mobile peroxisomes was unaltered by the expression of MFN2 mutants.

Mfn2 is required for the proper transport of axonal mitochondria
The specific disruption of mitochondrial transport by MFN2 disease mutants suggests that MFN2 may directly regulate microtubule-based mitochondrial transport. To determine whether MFN2 is
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Mfn2 is required for normal axonal transport of mitochondria. To assess the effects of loss of Mfn2 on mitochondrial mobility, we analyzed mitochondrial transport in DRG neurons cultured from Mfn2 knock-out animals. A–C, DRG cultures from wild-type (A) or Mfn2−/− (B) mice infected with mito-RFP and imaged 4 d after infection at 10× magnification coupled with corresponding kymographs from single axons. Corresponding axons are presented above each kymograph as an embossed image for clarity. B, Diminished numbers of mitochondria are observed in Mfn2−/− axons due to a delay in the overall outward migration of mitochondria that is corrected by reintroduction of wild-type human MFN2 (C, Mfn2 rescue). D, Size–frequency histogram of mitochondria shows a decrease in the lengths of mitochondria observed in Mfn2−/− axons that is restored with reintroduction of MFN2 (Mfn2 rescue). E–G, Kymograph analysis of mitochondrial movements in individual MFN2+/− axons showed a profound abnormality in transport, resembling the deficit observed with expression of MFN2 disease mutants. The moving mitochondria from Mfn2−/− axons spent more time paused between movements (E, F) (*p < 0.001, t test; n is number of axons from which kymographs were generated), moved at slower velocities in the anterograde and retrograde directions (G), and were completely rescued by expression of either Mfn2 or Mfn1.

necessary for mitochondrial transport, we cultured DRG neurons from Mfn2-null (Mfn2−/−) embryos (supplemental Fig. S2, available at www.jneurosci.org as supplemental material). Lentivirus infection of DRG neurons with mito-RFP showed a marked delay in the migration of mitochondria out into axons of Mfn2−/− neurons compared with DRG neurons from wild-type littermates (Fig. 3A, B). As expected, mitochondria from Mfn2−/− axons were smaller than controls, consistent with a mitochondrial fusion defect (Fig. 3D). Live imaging and kymograph analysis revealed abnormal mitochondrial movement patterns in Mfn2−/− neurons that mirrored the movement abnormalities seen in MFN2 disease mutant-expressing neurons (Fig. 3A, B, kymographs). This effect was again specific to mitochondrial transport, as peroxisomal transport was normal in these neurons (supplemental Fig. S3, available at www.jneurosci.org as supplemental material). Reintroduction of wild-type MFN2 by lentiviral infection into Mfn2−/− cultures fully rescued both the delay in migration and the abnormal mitochondrial movement patterns (Fig. 3C, kymograph). Similar to the Mfn2 disease mutant-expressing neurons, mitochondria in Mfn2−/− neurons spent a greater amount of time paused between anterograde and retrograde movements (Fig. 3E, F) and more time at slower velocities in both anterograde and retrograde directions than in controls (Fig. 3G).

Interestingly, overexpression of Mfn1 also restored mitochondrial pause time and velocity distributions to normal levels (Fig. 3E, F). Mfn2 and Mfn1 are highly homologous and are known to have overlapping functions in mediating mitochondrial fusion (Chen et al., 2003). Analysis of Mfn1 and Mfn2 expression in embryonic fibroblasts and DRG neurons showed that both Mfn1 and Mfn2 are expressed in DRG neurons, although Mfn2 is higher in neurons than in fibroblasts (supplemental Fig. S1, available at www.jneurosci.org as supplemental material). Therefore, although both are able to mediate mitochondrial transport, the severe transport abnormalities in Mfn2−/− DRG neurons suggest that Mfn1 levels in these cells are not sufficient to compensate for the loss of Mfn2.

These findings indicate that Mfn2 is necessary for proper mitochondrial transport and that CMT2A-associated Mfn2 mutants disrupt this function in a dominant-negative manner. The increased pause time of mitochondria observed in Mfn2−/− axons suggests that Mfn2 may either directly promote mitochondrial attachment to microtubules or maintain the processivity of the microtubule motors kinesin and dynein.
microtubule-based transport motors. Miro1 and Miro2 are OMM Rho-like GTPases that link mitochondria to kinesin, and Drosophila Miro is essential for mitochondrial transport in flies (Fransson et al., 2006; Glater et al., 2006; Macskill et al., 2009b; Russo et al., 2009). Coimmunoprecipitation experiments demonstrated that Mfn2 and Mfn1 are capable of interacting with either Miro1 or Miro2 when exogenously expressed in HEK 293T cells, with an apparently stronger interaction observed between Mfn2:Miro2 than Mfn2:Miro1 (Fig. 4A,B). We next examined OIP106 and GRIF, homologues of the Drosophila Milton protein that interacts with both Miro proteins and kinesin (Fransson et al., 2006). We found that Mfn2 and Mfn1 also coimmunoprecipitated with OIP106 and GRIF1 (Fig. 4C–F). Finally, we investigated whether the mitofusins directly interact with kinesin. Because Kif5C, a conventional kinesin-1 superfamily member, is highly expressed in neurons, we used exogenously expressed Kif5C for our coimmunoprecipitation experiments but found that neither Mfn2 nor Mfn1 directly interacted with Kif5C (Fig. 4G,H). Similarly, no interaction between Mfn2 and kinesin light chain was observed (data not shown). To investigate whether Mfn2 could play a role in “anchoring” mitochondria, we also examined whether Mfn2 could interact with syntaphilin (Synph). Synph is an OMM protein that prevents mitochondria from moving by anchoring them to microtubules (Kang et al., 2008). However, neither wt-MFN2 nor R94Q coimmunoprecipitated with Synph (Fig. 4I).

These experiments demonstrate that Mfn2 and Mfn1 interact with components of the kinesin-based mitochondrial molecular transport apparatus, providing an explanation for how loss of Mfn2 disrupts mitochondrial transport. Of note, the CMT2A-associated MFN2 mutants (R94Q and H361Y) maintained the ability to interact with Miro1/2 and OIP106/GRIF1 (supplemental Fig. S4, available at www.jneurosci.org as supplemental material). Furthermore, mutant MFN2 proteins did not disrupt either the interaction between OIP106 and kinesin or the interaction between Miro2 and OIP106 (supplemental Fig. S4, available at www.jneurosci.org as supplemental material). Therefore, the disruption of mitochondrial transport observed with disease mutant MFN2 expression is likely through altered regulation of the kinesin transport apparatus (processivity, attachment to microtubules) rather than simple disruption of Mfn:Miro:OIP106 complex formation.

Knockdown of Miro2 produces a mitochondrial transport abnormality similar to loss of Mfn2

Drosophila Miro is required for both anterograde and retrograde transport of axonal mitochondria (Guo et al., 2005; Russo et al., 2009). Although the mammalian Miro proteins interact with Milton homologues (Fransson et al., 2006), the effect of loss of Miro proteins on axonal mitochondrial transport in mammalian
that loss of mitochondrial fusion leads to dissipation of mitochondrial motility (Fig. 5). The pattern of mitochondrial movements on kymograph analysis was similar to that seen in Mfn2−/− neurons, with loss of fast, sustained movements (Fig. 5A, B). Unlike loss of Mfn2, which produced concurrent mitochondrial fusion and transport defects with smaller fragmented mitochondria, loss of Miro2 did not alter the length of axonal mitochondria (Fig. 5C). Mitochondria in Miro2-depleted neurons spent more time paused between anterograde and retrograde movements, and the velocity distributions were skewed toward slower velocities (Fig. 5D, E). Of note, overexpression of Miro2 in Mfn2−/− cultures was not able to rescue transport deficits (data not shown). Together, the fact that Mfn2 interacts directly with the Miro proteins, and that loss of Mfn2 or Miro2 disrupts axonal mitochondrial transport in a nearly identical manner, suggests that both Mfn2 and Miro2 must be present for proper axonal mitochondrial transport.

Disruption of mitochondrial fusion by knockdown of Opa1 does not affect the axonal transport of mitochondria

Some of the CMT2A disease mutant MFN2 alleles are unable to mediate mitochondrial fusion (Detmer and Chan, 2007). Given that loss of mitochondrial fusion leads to dissipation of ΔΨ and diminished oxidative phosphorylation (Lodi et al., 2004; Chen et al., 2005), it is possible that the observed alterations in mitochondrial transport could be secondary to loss of mitochondrial fusion. To investigate whether diminishing mitochondrial fusion directly or indirectly disrupts axonal mitochondrial transport, we used lentivirally delivered siRNA to knockdown Opa1 (supplemental Fig. S1, available at www.jneurosci.org as supplemental material) (∼90% transcript knockdown) in cultured DRG neurons. Knockdown of Opa1 severely disrupts mitochondrial fusion in non-neuronal cells, leading to a decrease in mitochondrial size similar to that of loss of Mfn1/ Mfn2 (Cipolat et al., 2004; Chen et al., 2005). Measurement of mitochondrial lengths confirmed that the mitochondria were significantly shorter in Opa1 knockdown DRG axons (Fig. 6C), consistent with the expected decrease in mitochondrial fusion (Cipolat et al., 2004; Chen et al., 2005). However, time-lapse imaging and kymograph analysis revealed that patterns of mitochondrial movement in Opal knockdown neurons were indistinguishable from those of controls (Fig. 6A, B). Mitochondria in neurons depleted of Opa1 spent equal amounts of time paused between movements in anterograde and retrograde directions and moved at similar velocities compared with controls (Fig. 6D, E).

CMT2A-associated MFN2 mutants cannot compensate for the mitochondrial transport abnormality in Mfn2−/− neurons

As described above, studies in fibroblasts have shown that some CMT2A-associated MFN2 mutants are unable to mediate mitochondrial fusion (R94Q), whereas others are fusion competent (L76P, W740S) (Detmer and Chan, 2007). However, exogenous expression of all of these mutants produced a mitochondrial transport defect in DRG neurons (Baloh et al., 2007), suggesting that the ability of CMT2A mutants to mediate fusion and transport may be separable. To determine whether the role of Mfn2 in mitochondrial fusion could be dissociated from its role in transport, we expressed the L76P and W740S mutants (fusion competent) and the R94Q mutant (fusion incompetent) in cultured Mfn2−/− neurons and assessed their ability to rescue the transport defect. In contrast to wtMFN2, none of the CMT2A-associated mutants were able to completely normalize the time mitochondria spent paused between movements or to fully restore the faster movement segments seen on velocity distributions (Fig. 7A–C), with the fusion-incompetent R94Q allele being least capable of rescuing the defect in transport in Mfn2−/− neurons. Therefore, CMT2A MFN2 mutants were not able to normalize axonal mitochondrial transport in DRG axons whether they are fusion competent (L76P, W740S) or incompetent (R94Q). This finding indicates that disruption of axonal mito-
mitochondrial transport correlates better with pathogenicity of CMT2A-associated MFN2 mutants than with the loss of the ability to mediate mitochondrial fusion.

Discussion
These experiments provide evidence for a direct role for Mfn2 in mediating mitochondrial transport, distinct from its role in mitochondrial fusion. Both loss of Mfn2 and expression of CMT2A-associated MFN2 disease mutants specifically alter mitochondrial movement patterns, causing mitochondria (but not other organelles) to move at slower velocities and pause for greater lengths of time, consistent with an inability to attach/move via the microtubule-based kinesin and dynein transport systems. Together with the finding that mitofusins interact with Miro proteins and OIP106/GRIF1, these data indicate that Mfn2 is a key component of the linker/adaptor complex between mitochondria and kinesin/microtubules.

Mfn2 as a multifaceted regulator of mitochondrial dynamics and function
Outer mitochondrial membrane proteins are well positioned to play a key role in coordinating and regulating diverse aspects of mitochondrial function. Perhaps it should not be a surprise then that mitofusins have been reported to perform numerous functions, including the regulation of mitochondrial fusion (Chan, 2006), endoplasmic reticulum (ER)–mitochondrial tethering (de Brito and Scorrano, 2008), apoptotic cell death and outer membrane permeability (Suen et al., 2008), oxidative phosphorylation and mitochondrial function. Perhaps it should be not a surprise then that mitofusins have been reported to perform numerous functions, including the regulation of mitochondrial fusion (Chan, 2006), endoplasmic reticulum (ER)–mitochondrial tethering (de Brito and Scorrano, 2008), apoptotic cell death and outer membrane permeability (Suen et al., 2008), oxidative phosphorylation and gradient coupling (Pich et al., 2005), and microtubule-based mitochondrial transport.

Our data indicate that regulation of mitochondrial transport by Mfn2 is achieved by interacting with the Miro: Milton complex. The increased pause time and slower movement velocities in both anterograde and retrograde directions due to loss of Mfn2 closely resemble the mitochondrial movement abnormalities seen in Miro2-depleted DRG neurons and in dMiro knock-out flies (Russo et al., 2009). Our data indicate that the regulation of mitochondrial transport by Mfn2 is achieved by interacting with the Miro:Milton complex. The increased pause time and slower movement velocities in both anterograde and retrograde directions due to loss of Mfn2 closely resemble the mitochondrial movement abnormalities seen in Miro2-depleted DRG neurons and in dMiro knock-out flies (Russo et al., 2009). Other reports found that overexpression of Miro proteins can either increase (Guo et al., 2005) or decrease (Russo et al., 2009) axonal mitochondrial transport. Importantly, we observed that overexpression of Miro2 could not rescue the transport deficit in Mfn2−/− neurons, indicating that both Mfn2 and Miro2 are required to cooperatively mediate mitochondrial transport.

While Miro proteins have only been shown to interact with anterogradely moving kinesin motors, loss of either Mfn2 or Miro2 led to disruption of both anterograde and retrograde mitochondrial transport, similar to loss of dMiro in flies (Russo et al., 2009). The Mfn2:Miro complex may also bind to dynein and direct alternating engagements of either motor with microtubule tracks. However, this does not have to be the case, as anterograde and retrograde organelar transports are coordinately regulated, and selective disruption of either kinesins or dynein/dynactin typically diminishes bidirectional transport, though the exact reason remains unclear (Brady et al., 1990; Waterman-Storer et al., 1997; Martin et al., 1999; Deacon et al., 2003). The increased time mitochondria spend pausing in axons from neurons lacking Mfn2 or Miro2 may also indicate a disruption of motor processivity. Together, this suggests that the loss of Mfn2 in Mfn2−/− neurons did not alter mitochondrial movement patterns. C, Mitochondrial size–frequency histogram showed that loss of Opa1 significantly decreased mitochondrial length similar to that seen in Mfn2−/− cultures, reflective of decreased mitochondrial fusion. D, E, Pause times between anterograde or retrograde movements (D) and velocity of movements (E) were not altered by Opa1 knockdown (n is the number of axons from which kymographs were generated), unlike loss of Mfn2, indicating that loss of mitochondrial fusion alone does not alter mitochondrial transport.

Alterations in mitochondrial dynamics in human disease
Mitochondrial dysfunction is involved in numerous neurodegenerative and metabolic diseases; however, whether mitochondrial dysfunction plays a primary role in disease pathogenesis or is simply a consequence of other aspects of cellular dysfunction (disruption of protein homeostasis, environmental toxins, etc) is unclear. In this context, studying the molecular mechanisms by which mutations in mitochondrial proteins (such as MFN2 or
OPA1) cause human disease is helpful, as altered mitochondrial function likely plays a primary role in disease. One of the fundamental mysteries of inherited neurodegenerative diseases is the selective vulnerability of particular populations of neurons to mutations in universally expressed proteins. CMT2A and dominant optic atrophy are associated with mutations in MFN2 and OPA1, respectively; both are ubiquitously expressed proteins essential for mitochondrial fusion (Alexander et al., 2000; Delettre et al., 2000; Zuchner et al., 2004). However, instead of leading to identical human diseases, mutations in MFN2 cause a severe peripheral neuropathy affecting the longest axons in the body, whereas mutations in OPA1 (typically via haploinsufficiency) predominately lead to degeneration of optic nerve axons. Our findings suggest that the fundamental difference in the pathogenesis of these diseases lies in the fact that loss of OPA1 disrupts mitochondrial fusion without altering mitochondrial transport. Interestingly, early changes in mitochondrial dynamics are seen in neurons expressing mutant forms of neurofilament that cause CMT2E, suggesting that alterations in mitochondrial transport and distribution may be a final common pathway leading to axonal degeneration in multiple types of CMT (Baloh, 2008; Tradewell et al., 2009). A viable working hypothesis is that the long peripheral nerves affected in CMT are more vulnerable to disruption of mitochondrial transport and distribution than other cell types. Defining evidence for or against this hypothesis from patient samples or animal models remains to be demonstrated and is a focus of ongoing research.

Although our studies indicate that Mfn2 is clearly required for mitochondrial transport, the precise molecular mechanism by which Mfn2 disease mutants disrupt this process remains unclear, as they maintain the ability to interact with Miro1/2 and with OIP106/GRIF1. These data indicate the CMT2A disease mutants do not simply disrupt formation of the molecular adaptor complex linking mitochondria to kinesins, but instead disrupt the proper function of this complex and, hence, further studies will be needed to understand exactly how Mfn2 disease mutants alter the properties of the adaptor complex on a molecular level.

The role of Mfn1 and Mfn2 in controlling mitochondrial dynamics

Although Mfn2 and Mfn1 are highly homologous proteins with overlapping functions, several lines of evidence have shed light on properties that are divergent between them. Mfn1 more effectively tethers mitochondrial membranes and alone is sufficient to mediate fusion with Opa1 (Cipolat et al., 2004; Ishihara et al., 2004). Though both Mfn1- and Mfn2-null mice show early embryonic lethality due to a placental defect (Chen et al., 2003), loss of Mfn1 outside of the placenta (using a Cre-lox system) led to no detectable abnormalities, whereas similar nonplacental loss of Mfn2 caused severe multisystem defects and early postnatal death (Chen et al., 2007). Additionally, recent evidence suggested that Mfn2 but not Mfn1 is involved in ER–mitochondrial tethering (de Brito and Scorrano, 2008).

In our study, loss of Mfn2 alone is sufficient to produce a severe mitochondrial transport deficit in DRG neurons that could be restored by expression of either Mfn2 or Mfn1, consistent with the fact that both can interact with the Miro:Milton complex. Indeed, we found that exogenous expression of Mfn2 but not Mfn1 is involved in ER–mitochondrial tethering (de Brito and Scorrano, 2008).

| Figure 7. Fusion-competent, CMT2A-associated MFN2 mutants cannot compensate for the mitochondrial transport abnormality in Mfn2−/− neurons. To compare the ability of fusion-competent (L76P, W740S) and fusion-incompetent (R94Q) MFN2 mutants to mediate mitochondrial transport, we reintroduced these mutants into Mfn2−/− neurons and assessed their ability to rescue the mitochondrial transport abnormality. As compared with wtMFN2, neither the fusion-competent (L76P, W740S) nor fusion-incompetent (R94Q) mutants were able to fully restore mitochondrial pause time (A, B) (n is number of axons from which kymographs were generated) or mitochondrial velocities (C). |
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