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Reducing dynamin 2 expression rescues X-linked centronuclear myopathy

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Centronuclear myopathies (CNM) are congenital disorders associated with muscle weakness and abnormally located nuclei in skeletal muscle. An autosomal dominant form of CNM results from mutations in the gene encoding dynamin 2 (DNM2), and loss-of-function mutations in the gene encoding myotubularin (MTM1) result in X-linked CNM (XLCNM, also called myotubular myopathy), which promotes severe neonatal hypotonia and early death. Currently, no effective treatments exist for XLCNM. Here, we found increased DNM2 levels in XLCNM patients and a mouse model of XLCNM (Mtm1−/−). Generation of Mtm1−/− mice that were heterozygous for Dnm2 revealed that reduction of DNM2 in XLCNM mice restored life span, whole-body strength, and diaphragm function and increased muscle strength. Additionally, classic CNM-associated histological features, including fiber atrophy and nuclei mispositioning, were absent or reduced. Ultrastructural analysis revealed improvement of sarcomere organization and triad structures. Skeletal muscle–specific decrease of Dnm2 during embryogenesis or in young mice after disease onset revealed that the rescue associated with downregulation of Dnm2 is cell autonomous and is able to stop and potentially revert XLCNM progression. These data indicate that MTM1 and DNM2 regulate muscle organization and force through a common pathway. Furthermore, despite DN2M being a key mecha-noenzyme, its reduction is beneficial for XLCNM and represents a potential therapeutic approach for patients.

Introduction

Centronuclear myopathies (CNM) are a group of congenital myopathies characterized by muscle weakness, fiber atrophy, predominance of type I fibers, and increased centralization of nuclei not secondary to muscle regeneration (1, 2). Three main forms of CNM have been characterized: X-linked CNM (XLCNM; OMIM 310400), due to mutations in the phosphoinositides phosphatase myotubularin (MTM1) (3); autosomal recessive CNM (ARNCM, OMIM 255200), caused by mutations in the membrane remodeling protein amphiphysin 2 (BIN1) (4), and autosomal dominant CNM (ADCNM, OMIM 160150), due to mutations in dynamin 2 (DNM2) (5). The relationship between the implicated genes in muscle is not known, and potent therapeutic approaches are lacking.

XLCNM, also called myotubular myopathy, is the most common and severe form of CNM, with neonatal onset and a poor prognosis (1). To date, more than 200 different mutations in MTM1 have been reported in about 450 families, most of which lead to a strong reduction of MTM1 protein (6–9). Mtm1−/− knockout mice or mice with knockin of a patient mutation both recapitulate the CNM phenotype with classical histological features including abnormal organelle positioning, mislocalization of nuclei, and muscle atrophy, associated with a corresponding reduction in muscle strength (10–12). A defect in triad structures associated with abnormal excitation-contraction coupling has been reported in several animal models and patients with various forms of CNM (13–15), identifying a common defect in all CNM forms. This is consistent with a proposed role of MTM1 in the regulation of phosphoinositides level on the sarcoplasmic reticulum component of the triads (14, 16).

Dynamins are large GTPase proteins that play important roles in membrane trafficking and endocytosis (17–20) and in actin cytoskeleton assembly (19–21). Dynamins can form polymerized rings around membrane tubules (22) and are involved in membrane fusion events (20, 23, 24), providing a paradigm of a mechanoenzyme. Three dynamins have been identified in humans: dynamin 1, expressed mainly in neurons, dynamin 3, expressed predominantly in brain and testis, and DNM2, which is ubiquitously expressed. Different heterozygous DNM2 mutations have been identified in tissue-specific diseases; ADCNM affecting skeletal muscle (5) and autosomal dominant Charcot-Marie-Tooth (CMTDIB, OMIM 606482) peripheral neuropathy (25).

Recent biochemical studies have indicated that some CNM-causing DNM2 mutations increase dynamin oligomer stability and GTPase activity (26, 27). This was supported in vivo by either knock-in or overexpression of the most common CNM-DNM2 patient mutation in mice, which induced CNM-like features at adulthood (28, 29), indicating that the disease is not due to haploinsufficiency. Overexpression of WT DNM2 also caused similar CNM-like perturbation to the muscle, albeit to a lesser extent (28, 30).

The main goal of this study was to validate a rescue approach for XLCNM. Due to their common implication in CNM, we suspected that MTM1 and DNM2 might function in a common pathway,
where either MTM1 loss-of-function or DNM2 gain-of-function lead to the CNM phenotype. We hypothesized that MTM1 might be a negative regulator of DNM2 function in muscle and that reduced DNM2 expression in an XLCNM model may rescue the XLCNM phenotype. To test this hypothesis, we produced Mtm1–/y mice that were heterozygous for DNM2 (Mtm1–/yDnm2+/–). We found that reducing DNM2 expression in Mtm1–/y mice is sufficient to rescue the early XLCNM lethality and most features of the disease. We also showed that reducing DNM2 expression only in muscle from embryogenesis, or even after the onset of the disease, was sufficient to increase the life span of Mtm1–/y mice. Finally we provide what we believe is the first evidence that MTM1 is a regulator of DNM2 in skeletal muscle and propose a rationale disease for treatment.

**Results**

**Characterization of Dnm2 heterozygous (Dnm2+/–) mice.** Constitutive knockout of Dnm2 was previously shown to be lethal early during embryogenesis (31). We created Dnm2+/– mice by target exon 8 (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI71206DS1). From 100 pups, we did not identify any Dnm2+/– mice, confirming that Dnm2+/– is embryonically lethal. Dnm2+/– pups were identified at expected Mendelian ratios and were further analyzed under the EUMODIC phenotyping program (32). Basic blood chemistry tests indicated no difference between WT and Dnm2+/– mice for urea (indicating normal kidney function), calcium (implying osmotic homeostasis), and total cholesterol (suggesting absence of cardiovascular disease) levels (Supplemental Figure 2A). Normal ECG measurements suggested unaltered electrical activity in the heart (Supplemental Figure 2B). Overall, there was no difference in body weight (Supplemental Figure 2C) or in lean tissue or fat content between WT and Dnm2+/– mice (Supplemental Figure 2D). An electromyography test for basic muscle function revealed no difference in single nerve conduction velocity (Supplemental Figure 2E). Tibialis anterior (TA) muscle mass was similar between WT and Dnm2+/– mice (Supplemental Figure 2F), and no difference in absolute or specific maximal force or fatigability of the TA was detected (Supplemental Figure 2, G–I), indicating overall that Dnm2+/– mice are clinically and physiologically similar to WT mice, with no detectable difference in muscle function.

**DNM2 levels in XLCNM.** Before investigating the therapeutic potential of downregulation of DNM2 in XLCNM, we measured DNM2 protein levels by Western blot on XLCNM patient muscle lysates (Figure 1A). We identified a 1.5-fold increase in DNM2 protein expression from 5 neonatal XLCNM muscle biopsies when compared with control age-matched biopsies (Figure 1B).
We then determined whether the increase in DNM2 expression was also observed in an animal model of XLCNM. In this study, we used Mtm1−/y mice that faithfully reproduce XLCNM (10, 15). TA muscle lysates from 5-week-old Mtm1−/y mice exhibited a significant increase in DNM2 levels compared with WT littermates (Figure 1, C and D), suggesting that increased DNM2 is linked to the XLCNM phenotype. An increase in DNM2 expression was also observed in the diaphragm (Figure 1, E and F), which appeared affected histologically, with more atrophic fibers containing mislocalized nuclei (Figure 1G). This finding suggests respiratory insufficiency as a cause of death in Mtm1−/y mice, similar to what occurs in patients.

Reducing DNM2 expression greatly prolongs the life span of Mtm1−/y mice. (A) Life span of all mice, represented as percentage survival. All mice in groups WT, Dnm2+/−, and Mtm1−/yDnm2+/− survived to at least 12 months. (B) Mouse whole-body weight. (C–F) Relative level of DNM2 protein was determined by densitometry of DNM2 standardized to GAPDH (blots on Supplemental Figure 5). DNM2 level is represented as fold difference from WT lysate. Expression was determined in 8-week-old, 16-week-old, and 6-month-old mice from diaphragm (DIA) (C), gastrocnemius (GAS) (D), TA (E), and soleus (SOL) (F) muscles. n = 2–8 mice. (G) mRNA levels were quantified by qRT-PCR analysis, with DNM2 levels expressed relative to GAPDH. Graph represents 3 independent experiments. Gastrocnemius (H), TA (I), and soleus (J) muscle weights (n = 5–13 mice). All graphs depict mean ± SEM. *P < 0.05; **P < 0.01; ***P < 0.001. w, weeks of age; m, months of age.

Figure 2
Reduced DNM2 expression greatly rescues the life span of Mtm1−/y mice. (A) Life span of all mice, represented as percentage survival. All mice in groups WT, Dnm2+/−, and Mtm1−/yDnm2+/− survived to at least 12 months. (B) Mouse whole-body weight. (C–F) Relative level of DNM2 protein was determined by densitometry of DNM2 standardized to GAPDH (blots on Supplemental Figure 5). DNM2 level is represented as fold difference from WT lysate. Expression was determined in 8-week-old, 16-week-old, and 6-month-old mice from diaphragm (DIA) (C), gastrocnemius (GAS) (D), TA (E), and soleus (SOL) (F) muscles. n = 2–8 mice. (G) mRNA levels were quantified by qRT-PCR analysis, with DNM2 levels expressed relative to GAPDH. Graph represents 3 independent experiments. Gastrocnemius (H), TA (I), and soleus (J) muscle weights (n = 5–13 mice). All graphs depict mean ± SEM. *P < 0.05; **P < 0.01; ***P < 0.001. w, weeks of age; m, months of age.

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Reducing DNM2 expression greatly prolongs the life span of Mtm1−/y mice. To determine whether reducing DNM2 expression may rescue XLCNM due to MTM1 mutations, we crossed Mtm1−/y mice with Dnm2+/− mice to produce male offspring that were Mtm1−/yDnm2+/−. Most Mtm1−/y mice died between 1 and 3 months of age as previously reported (10), whereas 100% of Mtm1−/yDnm2+/− mice survived for at least 1 year (Figure 2A), with no significant difference in body weight compared with WT mice (Figure 2B), indicating that reduced expression of DNM2 can rescue the early lethality observed in Mtm1−/y mice. Mtm1−/yDnm2+/− mice that were not sacrificed reached 2 years of age. At 8 weeks, when approximately 60% of Mtm1−/y mice were still alive, Mtm1−/yDnm2+/− mice were not distinguishable from WT mice upon general inspection, whereas Mtm1−/y mice displayed a significant decrease in movement and activity (Supplemental Video 1).

We determined the level of DNM2 protein expression on lysates from several muscles at different ages. In the diaphragm, DNM2 levels were reduced approximately 50% in Dnm2+/− and
Mtm1–/yDnm2+/– mice compared with WT mice, as expected, at both 8 weeks and 6 months of age (Figure 2C). Mtm1–/y mice exhibited an increase in DNM2 levels in diaphragm compared with WT mice at 8 weeks, consistent with results from 5-week-old mice (Figure 1F). In the gastrocnemius (Figure 2D), TA (Figure 2E), and soleus (Figure 2F) muscles, the same trend was seen at 8 weeks, 16 weeks, and 6 months (Supplemental Figure 5). Therefore, DNM2 levels are consistently increased in Mtm1–/y mice and consistently reduced in Dnm2+/– and Mtm1–/yDnm2+/– mice compared with WT mice in different muscles at different ages.

To determine whether the varied DNM2 protein level is due to altered protein synthesis, we performed quantitative RT-PCR (qRT-PCR) analysis on 8-week-old TA muscle lysates. The mRNA Dnm2 levels in both Dnm2+/– and Mtm1–/yDnm2+/– mice were significantly reduced compared with WT and Mtm1–/y mice (Figure 2G), correlating with DNM2 protein expression (Figure 2E). No significant increase was seen in Dnm2 mRNA expression in Mtm1–/y muscles, indicating that increased DNM2 protein expression in Mtm1–/y muscles may be due to increased stabilization of DNM2 or reduced degradation rather than increased transcription.
As the TA was one of the most affected muscles in Mtm1–/y mice (10, 15), we investigated the localization of MTM1 and DNM2 by immunofluorescence analysis in TA from 8-week-old mice. The Z-line, identified by α-actinin staining, appeared relatively undisturbed (Supplemental Figure 3A). DNM2 colocalized with α-actinin at the Z-line and appeared relatively unperturbed in Mtm1–/y and Mtm1–/yDnm2+/– mice (Supplemental Figure 3A). MTM1 was barely detectable in Mtm1–/y and Mtm1–/yDnm2+/– mice, as expected (Supplemental Figure 3B). These data show that reducing expression of DNM2 rescues the life span and body weight of Mtm1–/y mice to WT level.

The 3′ UTR region of DNM2 contains a site for miR-133a binding, resulting in repression of protein expression, and lack of miR-133a was shown to cause CNM-like features in mice, corresponding with an increase in DNM2 expression (30). qRT-PCR analysis performed on 8-week-old TA lysates from Mtm1–/y mice identified no significant difference in miR-133a expression (Supplemental Figure 4A), despite the increase in DNM2 level (Figure 2E). miR-133a was, however, significantly increased in Dnm2+/– and Mtm1–/yDnm2+/– mice (Supplemental Figure 4, A and B). The 1.5-fold increase in miR-133a, however, did not lead to a stronger decrease in DNM2 level than expected in Dnm2+/– mice (50%, Figure 2E). The molecular mechanisms causing increased miR-133a expression are unclear at present.

**Muscle atrophy in Mtm1–/y mice is rescued by reducing DNM2 expression.** To analyze further the effect of reducing DNM2 expression in Mtm1–/y mice, we measured muscle mass. The fast-twitch gastrocnemius muscle was atrophied in Mtm1–/y mice; however, no atrophy was observed in Mtm1–/yDnm2+/– mice analyzed at up to 1 year of age (Figure 3H). Likewise, other fast-twitch muscles, the extensor digitorum longus (EDL) and plantaris muscle, did not exhibit atrophy in Mtm1–/yDnm2+/– mice at up to 1 year of age compared with WT mice (Supplemental Figure 4, A and B). The TA showed a strong atrophy in Mtm1–/y mice compared with Mtm1–/yDnm2+/–, WT, and Dnm2+/– mice at 8 weeks (Figure 2I). Mtm1–/yDnm2+/– TA weights were indistinguishable from those of WT mice at this age. At 16 weeks, unlike Mtm1–/y mice, Mtm1–/yDnm2+/– mice were still alive (Figure 2A) and presented with some TA atrophy compared with WT and Dnm2+/– mice, indicating that TA atrophy in Mtm1–/yDnm2+/– is delayed. Interestingly the slow-twitch soleus muscle exhibited atrophy in Mtm1–/y mice at 8 weeks, whereas there was no atrophy in Mtm1–/yDnm2+/– mice at up to 1 year of age compared with WT mice (Figure 2J). Similar results were seen when whole muscle mass was measured relative to body weight (Supplemental Figure 4, D–F). No difference was observed in liver or heart weights between Mtm1–/yDnm2+/– and WT mice (Supplemental Figure 4, G and H). Therefore, muscle atrophy was fully rescued in gastrocnemius and soleus muscles and strongly delayed in TA muscle following the reduction of DNM2 expression in Mtm1–/y mice.

**CNM histology is greatly rescued in Mtm1–/y mice by reducing DNM2 expression.** CNM presents histologically with mislocalized internal nuclei and muscle fiber hypertrophy. We analyzed 2 main time points: early (8 weeks), when the majority of the Mtm1–/y mice were still alive, and late (16 weeks), when 95% of Mtm1–/y mice have died. At 8 weeks, Mtm1–/y TA muscle exhibited characteristic mislocalization of nuclei (Figure 3, A and F), reduced fiber size (Figure 3, A and D), and abnormal succinate dehydrogenase (SDH) staining with subsarcolemmal and central accumulations (Figure 3A). Mtm1–/yDnm2+/– TA muscles were histologically similar to those of WT and Dnm2+/– mice, as observed for the gastrocnemius and soleus muscles (Supplemental Figure 4, I and J), with only a few abnormal fibers on SDH staining (Figure 3A). Fiber hypertrophy was rescued and internal and central nuclei significantly reduced compared with Mtm1–/y mice (Figure 3, A, D, and F). In addition, membrane accumulations around the nucleus were reduced (Figure 3B). By 16 weeks, the TA phenotype from Mtm1–/yDnm2+/– was mixed, as some areas appeared healthy and other areas resembled 8-week-old Mtm1–/y muscle, with muscle fiber hypertrophy, mislocalization of nuclei, and abnormal SDH staining (Figure 3, C, E, and F). The localization of desmin, an MTM1-binding partner shown to be disrupted in Mtm1–/y mice (33), was barely altered in Mtm1–/yDnm2+/– mice (Supplemental Figure 6A), indicating that normal desmin localization is restored in Mtm1–/yDnm2+/– mice. Overall, these results indicate the CNM phenotype in different muscles in Mtm1–/y mice is rescued or strongly delayed by reducing DNM2 expression.
Reducing DNM2 expression improves muscle strength and performance of Mtm1–/– mice. We next determined whether reducing DNM2 expression rescues the functional phenotype of Mtm1–/– mice. The string test requires mice that are suspended by their front paws to lift and hold their hind limbs on the wire. While Mtm1–/– mice fell off on several trials and were unable to perform the test by 8 weeks, Mtm1–/–Dnm2+/– mice performed the test similarly to WT and Dnm2+/– mice (Figure 4A and Supplemental Videos 2–4), indicating a rescue of whole-body strength. At 8 weeks, TA muscles from Mtm1–/–Dnm2+/– mice exhibited extremely weak absolute and specific maximal muscle force, whereas Mtm1–/–Dnm2+/– mice performed the test similarly to WT and Dnm2+/– mice (Figure 4B and C). At 16 weeks, the maximal force of the TA muscle of Mtm1–/–Dnm2+/– mice was reduced, consistent with histological data. Furthermore, no change in fatigability was observed at 8 weeks; however, at 16 weeks, Mtm1–/– Dnm2+/– TA muscles fatigued faster than controls (Figure 4D).

Notably histologically and physiologically, 16-week-old Mtm1–/– Dnm2+/– mice performed better than Mtm1–/– mice at 8 weeks, indicating either a slower progression of the disease or rescue in some but not all muscle fibers, as indicated by the mixed phenotype observed histologically in the TA at 16 weeks (Figure 3C).

Figure 5
 Improved muscle ultrastructure of Mtm1–/– mice with reduced DNM2 expression. TA muscles from 8-week-old (A) and 16-week-old (B) mice were imaged by TEM. Scale bars: 0.5 μm (A); 1 μm (B).

Overall, atrophy and decreased muscle force of the TA muscle in Mtm1–/–Dnm2+/– mice is reduced and strongly delayed compared with that of Mtm1–/– mice.

Improved muscle ultrastructure in Mtm1–/–Dnm2+/– mice. We next analyzed the ultrastructure of Mtm1–/–Dnm2+/– muscle by transmission electron microscopy (TEM). Eight-week-old Mtm1–/–Dnm2+/– TA morphology resembled that of WT and Dnm2+/– muscles, with aligned Z-lines and sarcomeres and no obvious mitochondrial structural abnormalities, whereas Mtm1–/– muscle displayed abnormal mitochondria shape, membrane accumulations, Z-line misalignment, and altered myofibrillar width (Figure 5A), as reported previously (10, 15, 34). Notably, muscle from Mtm1–/–Dnm2+/– mice at 16 weeks was heterogeneous; some regions appeared healthy, while other areas appeared disturbed (Figure 5B). Furthermore, mitochondrial abnormalities that were not evident at 8 weeks were detectable in some regions of 16-week-old Mtm1–/–Dnm2+/– mice. This supports our previous histological and physiological results, indicating that the CNM phenotype in Mtm1–/–Dnm2+/– TA muscle is partially albeit substantially rescued at different time points.

Triad structures are normalized in Mtm1–/–Dnm2+/– mice. A common feature shared between CNM patients and animal models of CNM is a disruption of the structure and position of triads within skeletal muscle (13–15, 35). DHPRα, a voltage-dependent calcium channel found on T-tubules of mature muscles, was localized in punctuate structures within TA myofibers of WT, Dnm2+/–, and Mtm1–/–Dnm2+/– mice, consistent with normal T-tubule localization (Supplemental Figure 6B). However, in Mtm1–/– muscle, this specific staining was lost, indicating severe disruption of the T-tubules. This was confirmed by staining for ryanodine receptor (RyR1), a calcium channel localized specifically at the sarcoplasmic reticulum of triads. Transverse and longitudinal images showed a rescue of RyR1 localization in most fibers from Mtm1–/–Dnm2+/– mice, with only a few fibers exhibiting RyR1 accumulations or perturbed localization, as seen extensively in Mtm1–/– mice (Figure 6A). High-magnification TEM images confirmed strong disruption of T-tubule/triad structures in Mtm1–/– mice compared with WT and Dnm2+/– mice, whereas well-positioned triads were clearly visible in Mtm1–/–Dnm2+/– mice (Figure 6B). Analysis of the triads confirmed no difference in the number of triads per sarcomere in WT, Dnm2+/–, or Mtm1–/–Dnm2+/– mice, whereas Mtm1–/– mice exhibited a reduced number of triads per sarcomere (Figure 6C), as shown previously (34). Caveolin 3 is found at T-tubules during muscle development and regeneration and at the sarcolemma in mature muscle (36). While WT and Dnm2+/– muscle showed caveolin 3 localizing to the sarcolemma as expected, many fibers from Mtm1–/– mice exhibited a strong internal staining pattern of caveolin 3 (Figure 6D). This phenotype was largely rescued in Mtm1–/–Dnm2+/– muscle, with only occasional fibers showing internal localization of caveolin 3. Therefore, the localization and structure of triads was rescued in 8-week-old Mtm1–/–Dnm2+/– mice.

Long-term physiological phenotype of Mtm1–/–Dnm2+/– mice. XLCNM presents with very severe muscle weakness in patients and lethality in Mtm1–/– mice within 1 to 3 months; however, reducing DNM2 expression in these mice rescued the life span and greatly improved muscle strength at 8 and 16 weeks. We determined the phenotype of Mtm1–/–Dnm2+/– mice at later time points to assess the extent of muscle function compatible with a normal lifespan. Mtm1–/–Dnm2+/– mice aged 6 months and 12 months were able to move and perform basic tasks (Supplemental Video 5).

Aged Mtm1–/–Dnm2+/– mice looked similar to WT mice (Figure...
7A), but walked with hind feet pointing outwards (Figure 7B and Supplemental Figure 7A). This feature was not progressive from 6 to 12 months. To determine overall maximal leg strength, the grip-strength test was performed using a dynamometer. No difference in strength of the 2 front paws was observed between WT and Mtm1−/yDnm2+/− mice during the first 12 months (Supplemental Figure 7B). A small reduction in strength was observed in Mtm1−/yDnm2+/− mice compared with WT mice when all 4
paws were measured, at both 6 months and 12 months (Figure 7C), indicating the hind limbs of Mtm1−/−Dnm2+/− mice exhibit reduced maximal muscle force compared with WT mice, which was not progressive. The rotarod test for general motor coordination, strength, and endurance was performed, with no difference observed at 6 or 12 months (Figure 7D), confirming the general coordination and overall strength of Mtm1−/−Dnm2+/− mice was not severely perturbed. The hanging test is a
strenuous test that requires mice to be suspended from a cage lid for 60 seconds. The severely affected $Mtm1^{-/-}$ mice were unable to perform this test after 1 month (Figure 7E). In comparison, $Mtm1^{-/-}Dnm2^{+/–}$ mice could perform this test up to the last age tested (12 months), albeit to a lesser extent than WT mice (Figure 7E and Supplemental Videos 6–8). As $Mtm1^{-/-}Dnm2^{+/–}$ mice were able to successfully perform basic motor strength tests at up to 12 months of age, we conclude that the disease progression was stopped over time and life span and basic motor function were rescued.

Normal long-term diaphragm function in $Mtm1^{-/-}Dnm2^{+/–}$ mice. We have shown that individual muscles appear to be differently affected in $Mtm1^{-/-}$ mice, and the corresponding rescue by reduction of DNM2 is varied between muscles (Figure 2, C–J, and Supplemental Figure 4). The main concern for the longevity of patients with XLCNM is the diaphragm's function, as they have life-threatening respiratory failure (1). In addition, we found the histology of the diaphragm in 5-week-old $Mtm1^{-/-}$ mice was strongly altered (Figure 1G). We thus tested the function of the diaphragm in 6-month-old.
mice using the plethysmograph test to measure the spontaneous breathing pattern under resting conditions. Mtm1\(^{-/-}\)Dnm2\(^{-/-}\) mice performed similarly to WT and Dnm2\(^{-/-}\) mice, with no significant difference detected (Figure 7F and Supplemental Figure 8). No significant difference in the force-frequency relationship was detected in isolated strips of diaphragm; however, a reduction in specific maximal force was observed (Figure 7G). Histologically, Mtm1\(^{-/-}\)Dnm2\(^{-/-}\) diaphragm resembled that of WT and Dnm2\(^{-/-}\) muscle, with no major alterations to nuclei positioning or fibrosis observed (Figure 7H). The diaphragm of Mtm1\(^{-/-}\)Dnm2\(^{-/-}\) mice sustained reduced DNM2 protein levels at 6 months (Figure 2C), compared with Mtm1\(^{-/-}\) mice in which DNM2 expression was elevated at 5 weeks (Figure 1, E and F) and 8 weeks (Figure 2C). Overall, the diaphragm muscle of Mtm1\(^{-/-}\)Dnm2\(^{-/-}\) mice was indistinguishable from that of control mice, supporting the extensive phenotypic amelioration of the XLCNM phenotypes upon DNM2 reduction.

Muscle-specific reduction of DNM2 is sufficient to rescue the phenotype and improve the life span in Mtm1\(^{-/-}\) mice. In this study, we were able to fully rescue the life span of Mtm1\(^{-/-}\) mice and most of the clinical and histological features of the disease by reducing DNM2 expression in utero in all
tissues. To determine whether the rescue of the muscle phenotype was cell-autonomous, we crossed HSA-Cre (where HSA indicates human skeletal muscle c-tactin) and HSA-Cre-ER\(^{TK}\) mice (37) with floxed Dnm2 mice to produce Dnm2\(^{2\text{lox/}}\) (Cre positive) and Dnm2\(^{2\text{lox/}}\) (Cre-ER\(^{TK}\)) mice. These mice were then crossed with Mtm1\(^{-/}\) mice to produce tissue-specific excision of Dnm2. When Dnm2 expression was reduced in muscle (Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\)), HSA promoter active from 9 dpc ref. (38), we were able to increase the life span of Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice, with 75% surviving until at least 16 weeks, while no Mtm1\(^{-/}\) mice survived to this age (Figure 8A), consistent with our results from Mtm1\(^{-/}\)Dnm2\(^{+/}\) mice (Figure 2A). Four WT and 4 Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice have been kept alive for future long-term analysis, and all mice were 12 months old at the time of publication. A corresponding increase in body weight was also observed in Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice compared with Mtm1\(^{-/}\) mice (Figure 8B). No difference in mass of the gastrocnemius or soleus muscles was observed between Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) and WT mice (Figure 8C). At 16 weeks, when all Mtm1\(^{-/}\) littermates had died, the TA of Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice exhibited atrophy, with a reduction in muscle mass and fiber size compared with WT littermates, associated with increased central and internal nuclei and some abnormal SDH staining (Figure 8, D–F), similar to that seen in the TA muscle from 16-week-old Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice (Figure 2). These alterations were less pronounced than those seen in Mtm1\(^{-/}\) mice at 8 weeks. Importantly, the gastrocnemius and soleus of Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice did not exhibit a significant difference in fiber size or nuclei position compared with WT mice (Supplemental Figure 9, A–D), indicating the phenotype is rescued differently in different muscles. DNM2 protein levels were measured in different muscles at 16 weeks, and there was a significant reduction in DNM2 expression in the diaphragm, but not in other muscles measured, compared with WT (Figure 8G and Supplemental Figure 9, E–H). Importantly, Mtm1\(^{-/}\) mice were not alive at this age to compare the levels of DNM2, which were already increased in these muscles at younger ages (Figure 2). As the diaphragm is a vital muscle required for breathing, we propose that reduced DNM2 expression in the diaphragm muscle is critical for increased survival of Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice and to rescue XLCNM.

Muscle-specific reduction of DNM2 after birth is sufficient to rescue Mtm1\(^{-/}\) mice. We crossed Mtm1\(^{-/}\) mice with Dnm2\(^{+/}\) mice under the HSA-Cre ER\(^{TK}\) system to allow excision of DNM2 after birth in muscle, induced by tamoxifen injection (37). Importantly, tamoxifen injections were performed when mice were 3 weeks old, after the onset of muscle atrophy (Figure 2) and centralized nuclei (15). We found that 70% of the injected Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice survived to 16 weeks (Figure 9A). A higher body weight was observed compared with that of Mtm1\(^{-/}\) mice; however at 16 weeks, body weight was still significantly reduced compared with that of WT mice (Figure 9B). No difference in normalized mass of the gastrocnemius, soleus, or TA muscles was observed compared with WT mice at 16 weeks (Figure 9C), unlike Mtm1\(^{-/}\) mice at earlier time points (Figure 2). Further analysis of TA muscles from Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice showed some fiber hypotrophy (Figure 9, D and E), associated with increased central and internal nuclei, and abnormal SDH staining (Figure 9, D and F), similar to the TA muscle from 16-week-old Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice (Figure 3). We noted a decrease in DNM2 protein expression in gastrocnemius from Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice at 16 weeks compared with WT mice; however, an increase in DNM2 protein expression was observed in TA and diaphragm muscles (Figure 9G and Supplemental Figure 10). These differences may be due to differential efficiency in DNM2 excision upon tamoxifen-mediated activation of the Cre recombinase. The increased DNM2 expression in the diaphragm at 16 weeks may be correlated with the reduced survival rate in Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice from 8 to 16 weeks. Therefore, we conclude that reduction of DNM2 levels in muscle after birth, after the onset of symptoms, is sufficient to improve the life span and the CNM phenotype observed in Mtm1\(^{-/}\) mice.

Discussion

In this study, we showed that reducing DNM2 in Mtm1\(^{-/}\) mice rescues most features of XLCNM and greatly prolongs life span and long-term muscle and motor performance. Furthermore, reduction of DNM2 in muscle alone, and after the onset of symptoms, is sufficient to rescue the phenotype.

DNM2 and MTM1 function in a common pathway. Loss-of-function mutations in MTM1 and dominant mutations in DNM2 lead to CNM in human and CNM phenotypes in mice models (1, 9). Moreover, biochemical analysis suggests several DNM2 CNM mutants increase DNM2 GTPase activity and oligomerization (26, 27). Furthermore, overexpression of WT DNM2 by AAV or by miR-133a inhibition resulting in increased DNM2 expression promoted muscle defects and centralization of nuclei in mouse muscle (28, 30). We thus hypothesized DNM2-CNMM mutations are gain-of-function. However, no link between MTM1 and DNM2 functions was previously established. As lowering the genetic dosage of DNM2, either systemically or in skeletal muscle, strongly ameliorated the XLCNM phenotype of Mtm1\(^{-/}\) mice, we have identified MTM1 and DNM2 as being linked specifically in muscle, in a common pathway controlling muscle mass and maximal force. Moreover, we suggest MTM1 is a negative regulator of the DNM2 pathway in muscle, as decreased DNM2 can compensate for absence of MTM1.

These data represent a first link between MTM1 and DNM2 in the pathogenesis of CNM. Moreover, the rescue data presented here support that DNM2-CNMM mutations are gain-of-function in skeletal muscle. We thus hypothesize that several forms of CNM are due to increased expression and/or function of DNM2. In particular, MTM1 loss in mice, and potentially in human, may lead to CNM phenotypes through overexpression and/or over-activation of DNM2.

Interestingly, we noted increased DNM2 protein expression in XLCNM patients and in Mtm1\(^{-/}\) muscle (Figure 1), supporting the hypothesis that increased DNM2 levels lead in part to the CNM phenotypes. No significant increase was observed in Dnm2 mRNA in Mtm1\(^{-/}\) muscle, indicating that the increase in DNM2 protein expression is due to either increased stability of the protein or reduced degradation. While genetic deletion of a Dnm2 allele led to a 50% decrease at the protein level in Dnm2\(^{+/}\) mice, we noted different DNM2 levels depending on the muscles investigated, following the same genetic deletion in the Mtm1\(^{-/}\) mice. DNM2 protein levels were monitored at 50% of WT levels in the diaphragm (Figure 2C) and 70% in TA muscles of Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) mice at 16 weeks (Figure 2E), and between 130% and 170% in Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) and Mtm1\(^{-/}\)Dnm2\(^{2\text{lox/}}\) TA muscles (Figures 8 and 9). Importantly, the DNM2 levels correlated with the alteration of these muscles, as assessed by histology. As Mtm1\(^{-/}\) mice died before this age, data for this genotype were not available, but we noted a consistent DNM2 increase in all muscles tested in 8-week-old Mtm1\(^{-/}\) mice. Thus these data sustain the hypothesis that increased DNM2 levels led in part to the CNM phenotype and that decrease or normalization of DNM2 levels rescues the XLCNM phenotype.
It was recently shown that increased flexibility of dynamin together with the tilting of the membrane-inserted pleckstrin homology (PH) domain is a prerequisite for normal membrane fission (39). In ADCNM, DNM2 mutations are mostly clustered at the interface between the middle and PH domains (24, 40), suggesting they affect the stability of a closed conformation and lead to increased flexibility of this region, tilting of the PH domain, and membrane fission (24). Overactivation of dynamin-mediated membrane fission either by mutations or increased DNM2 levels may thus alter the proper balance for membrane remodeling. While T-tubule defects are a hallmark of CNM and thus regulation of membrane tubulation at this site a tempting hypothesis for explaining the MTM1/DNM2 pathway, dynamin was never observed on T-tubules in normal adult muscle (28, 29). At the physiological level, decreased DNM2 strongly reduced or delayed signs of CNM in Mtm1−/− mice, including muscle mass, maximal force, and triad and nuclei positioning. This indicates a role for endogenous DNM2 in these different processes. Future studies will need to address the site of action of DNM2 in muscle.

**DNM2 reduction as a potential therapy for XLCNM.** We identify here a cross-therapy strategy, whereby we downregulate a CNM gene (DNM2) to rescue the severe phenotype due to a downregulation of another CNM gene (MTM1). This technique differs from therapeutic strategies that aim to compensate for the lack of a protein by reexpression of the corresponding gene or overexpression of a homolog or by modification of the mutated gene to create a functional protein (such as exon skipping).

A main goal of this study was to validate a rescue approach for XLCNM. We first reduced DNM2 levels during embryogenesis in all tissues of the Mtm1−/− mice; this experimental setup led to an extensive delay in the appearance of CNM phenotypes, a strong amelioration of motor behavior, and a normal life span, while Mtm1−/− mice died mostly in the first 2 months from progressive muscle weakness.

Both total and muscle-specific Mtm1−/− induced CNM phenotypes in mice, indicating XLCNM arises from a primary defect in skeletal muscle (10). We obtained a phenotypic rescue of XLCNM in Mtm1−/− mice by reducing DNM2 levels specifically in skeletal muscle, supporting the concept that XLCNM derives from a primary skeletal muscle defect and that specific treatment of muscle will be enough to ameliorate this disease in humans. In addition, our rescue data strongly support that rescuing myofiber intracellular organization, including myofibril organization, triad structure, and to some extent, nuclei positioning has a direct beneficial outcome on muscle function and life span and sustain the idea that these defects are primary causes of the XLCNM pathology.

However, muscle-restricted downregulation of DNM2 from embryonic stages or from 3-week-old mice rescued only 70%–75% of mice, compared with 100% rescue of the Mtm1−/− Dnm2−/− mice. Muscle-specific knockdown of DNM2 may not be sufficient to rescue all Mtm1−/− mice if other organs are also affected. The HSA promoter is expressed differently in different fiber types, with highest expression in type IIB fibers (41), whereas a key feature of CNM is increased type 1 fibers, which may account for the reduced survival rate.

Unlike the Mtm1−/− mice, in which motor behavior of neonates appears normal and deteriorates from 2 to 3 weeks of age, followed by a rapid progressive muscle weakness, patients are born with a severe hypotonia and muscle weakness that do not appear progressively. In addition, several reports indicate decreased fetal movements. Using time-inducible heterozygous deletion of DNM2, we found decreasing DNM2 expression after birth and after the appearance of some CNM symptoms was sufficient to improve life span and muscle weight in Mtm1−/− mice. Thus, decreasing DNM2 levels after birth appears sufficient to revert the deterioration of muscle histology and progression of the disease. This is indeed an important advantage for potential application of similar strategies in patients. To translate this proof-of-concept toward clinical trials, targeted downregulation of DNM2 or drugs inhibiting DNM2 function will have to be validated. Compounds inhibiting the function of DNM2 may be valuable for this aim, assuming their delivery and biodistribution in skeletal muscle is effective. In conclusion, we have identified DNM2 as a potential therapeutic target for XLCNM.

**Methods**

An expanded Methods section can be found in the Supplemental Methods.

**Generation of Dnm2 heterozygous mice**

The targeting vector was created with LoxP sites flanking exon 8 of Dnm2 (Supplemental Figure 1), then linearized, and electroporated into ES cells. Recombinant ES cells were injected into C57BL/6 blastocysts that were implanted in pseudopregnant females and germline transmission determined. Mice bred and analyzed were 129pases strain (CMV promoter). HSA-Cre C57BL/6 and HSA Cre-ERT2 mice were a kind gift from Daniel Metzger (IGBMC) (37, 38) and were used to produce mice with muscle-specific reduction of DNM2.

**Generation of Mtm1−/−Dnm2 heterozygous mice**

Female Mtm1−/− mice (129pase strain) (10, 15) were bred with male Dnm2 heterozygous mice to produce 4 possible genotypes in male offspring: Mtm1−/−Dnm2−/− (WT); Mtm1−/−Dnm2+/- (Dnm2+/-); Mtm1−/−Dnm2−/- (Mtm1−/−); and Mtm1−/−Dnm2−/−. All mice analyzed in this study were male.

**Animal experiments**

Animals were housed in a temperature-controlled room (19–22°C) with a 12-hour light/12-hour dark cycle. Mice were humanely sacrificed by CO2 inhalation followed by cervical dislocation, according to national and European legislation on animal experimentation. Muscles and other tissues were dissected and frozen in nitrogen-cooled isopentane and liquid nitrogen for histological and immunoblot assays, respectively.

**Phenotyping of Dnm2−/− mice**

Dnm2−/− male and female mice aged 10 to 15 weeks were phenotyped under the EUMODIC phenotyping program (see Supplemental Methods).

**String, grip, hanging, rotarod, and footprint tests**

**String test.** Mice were suspended on a wire by their forelimbs and allowed 20 seconds to climb, bringing their hind limbs onto the wire.

**Grip test.** Maximal strength measured from 2 or 4 paws using a dynamometer (Biosed).

**Hanging test.** Mice were suspended from a cage lid for a maximum of 60 seconds, and the time the mouse fell off the cage was recorded.

**Rotarod test.** Coordination and whole-body muscle strength and fatigability were tested using an accelerated rotating rod test (Panlab).

**Footprint test.** The angle of the hind limbs was measured by analyzing the footprint pattern using the ImageJ analysis program (n = 5–8 mice per group).

**Plethysmograph measurements**

The spontaneous breathing pattern in nonrestrained unstimulated mice was measured using a whole-body barometric plethysmograph (EMKA Technologies) at the Institut Clinique de la Souris (ICS) (Illkirch, France) (n = 3–5 mice per group).
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TA and diaphragm muscle contractile properties

TA muscle force measurements were evaluated by measuring in situ muscle isometric contraction in response to nerve and muscle stimulation (n = 5–11 mice per group). Isometric contraction was assessed on muscle strips from the ventral part of the costal diaphragm (n = 3–5 mice per group).

Western blotting

Western blotting was performed as described previously (28). Primary antibodies were as follows: home-made DNM2-2R680 (1:500), DNM2-2R865 (1:500), MTM1-R2827 (1:500), and GAPDH (1:10,000, MAB374; Chemicon); secondary antibodies were as follows: anti-rabbit HRP or anti-mouse HRP (1:10,000, Jackson ImmunoResearch Laboratories).

qRT-PCR

Total RNA was extracted from 8-week-old TA muscle lysates using Tri reagent (Molecular Research Center), reverse transcribed, and amplified using Oligo dT primers. Real-time qRT-PCR was performed using a Lightcycler 480 (Roche Diagnostics) with DNM2 primers (forward, CCAAGGCCATCCTCCCT; reverse, TGGTGAGTAGACCCGGAAGT) and GAPDH primers as a standard, with the SYBR Green 1 Master kit (Roche Diagnostics). Results were standardized to GAPDH (n = 2–3 mice per group, performed in triplicate).

Histology and immunofluorescence

Longitudinal and transverse cryosections (8 μm) of mouse skeletal muscles were prepared, fixed, and stained with the indicated antibodies and viewed using a laser-scanning confocal microscope (TCS SP5; Leica Microsystems). H&E and SDH sections were imaged with a slide scanner NanoZoomer 2HT equipped with the fluorescence module L1600-21 (Hamamatsu Photonics). Cross-sectional area (CSA) and nuclei position were analyzed in H&E sections from TA mouse skeletal muscle, using FIJI image analysis software (n = 4–7 mice per group).

TEM

TEM was performed on TA muscle biopsies as described previously (10, 28). The ratio of triads to sarcomere was calculated as described previously (34).

Microscopy and statistical analysis

All microscopy was performed at the IGBMC Imaging Centre. All samples for microscopy were mounted in Fluorosave reagent (Merck). Light microscopy was performed using a fluorescence microscope (DM4000; Leica Microsystems) fitted with a color CCD camera (CoolSNAP cf color; Photometrics). Confocal microscopy was performed using a confocal laser-scanning microscope (TCS SP2 or SP5; Leica Microsystems). ImageJ and FIJI analysis software were used for image analysis. Statistical analysis was performed using an unpaired Student’s t test (2-tailed) unless stated otherwise. P < 0.05 was considered significant.

Study approval

Animal experimentation was approved by the institutional ethical committee Com’Eth IGBMC-ICS (2012-128). All human biopsies were taken after informed consent was obtained. The use of patient biopsies for this study was authorized by the ethical committee of Pitie-Salpetriere Hospital (CCPPRB).

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