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Reprogramming the Dynamin 2 mRNA by Spliceosome-mediated RNA Trans-splicing

Delphine Trochet¹, Bernard Prudhon¹, Arnaud Jollet¹, Stéphanie Lorain¹ and Marc Bitoun¹

Dynamin 2 (DNM2) is a large GTPase, ubiquitously expressed, involved in membrane trafficking and regulation of actin and microtubule cytoskeletons. DNM2 mutations cause autosomal dominant centronuclear myopathy which is a rare congenital myopathy characterized by skeletal muscle weakness and histopathological features including nuclear centralization in absence of regeneration. No curative treatment is currently available for the DNM2-related autosomal dominant centronuclear myopathy. In order to develop therapeutic strategy, we evaluated here the potential of Spliceosome-Mediated RNA Trans-splicing technology to reprogram the Dnm2-mRNA in vitro and in vivo in mice. We show that classical 3′-trans-splicing strategy cannot be considered as accurate therapeutic strategy regarding toxicity of the pre-trans-splicing molecules leading to low rate of trans-splicing in vivo. Thus, we tested alternative strategies devoted to prevent this toxicity and enhance frequency of trans-splicing events. We succeeded to overcome the toxicity through a 5′-trans-splicing strategy which also allows detection of trans-splicing events at mRNA and protein levels in vitro and in vivo. These results suggest that the Spliceosome-Mediated RNA Trans-splicing strategy may be used to reprogram mutated Dnm2-mRNA but highlight the potential toxicity linked to the molecular tools which have to be carefully investigated during preclinical development.

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Subject Category: siRNAs, shRNAs, and miRNAs

Introduction

Dynamin 2 (DNM2) is a large GTPase involved in endocytosis, intracellular membrane trafficking and cytoskeleton organization.¹ DNM2 is ubiquitously expressed with higher abundance in heart and skeletal muscle.² DNM2 oligomerizes around the neck of nascent vesicles and regulates membrane fission events from plasma membrane³⁴ and intracellular compartments.⁵⁶ The DNM2 transcript encodes a protein composed of 5 domains: a N-terminal catalytic GTPase domain responsible for GTP binding and hydrolysis, a middle domain (MID) involved in DNM2 self-assembly, a pleckstrin homology (PH) domain that interacts with phosphoinositides and targets dynamin to membranes, a GTPase effector domain required for GTPase activation and DNM2 self-assembly, and a C-terminal Proline rich-domain (PRD) that mediates protein–protein interactions.

Heterozygous mutations in DNM2 gene are responsible for the autosomal dominant centronuclear myopathy (MIM# 160150).⁷ Autosomal dominant centronuclear myopathy is a rare congenital myopathy exhibiting a large clinical spectrum, and characterized by the presence of centrally located nuclei in a large number of muscle fibres.⁷⁸ No curative treatment is available and the pathophysiological mechanisms of the disease are still poorly understood. The human DNM2 (OMIM #602378) gene is composed of 22 exons encoding four major splice isoforms by using a combination of two alternative splice sites.⁹ To date, more than 20 mutations, affecting six exons spread over the MID, PH, and GTPase effector domain domains, have been identified in centronuclear myopathy patients.⁸ The most frequent mutation located in exon 11 (p.Arg465Trp in the MID) accounts for 30% of patients.⁸ The same mutation, also harbored by exon 11 in mouse, was used to develop a Knock-In-Dnm2²⁴⁶⁵⁶⁵ mouse model of the disease.¹⁰ The knock-out of Dnm2 in mouse is lethal during embryonic stage while heterozygous Dnm2 knock-out mice do not develop phenotype.¹¹,¹² Besides, overexpression of wild-type Dnm2 is also deleterious in mouse tissues.¹³ These data argue for a tight regulation of the DNM2 expression level compatible with cell homeostasis. Consequently, future therapeutic strategies for DNM2-linked autosomal dominant centronuclear myopathy will require maintenance of the DNM2 gene product in physiological range. Spliceosome-mediated RNA trans-splicing (SMarT) appears suitable to reach such a requirement.

Trans-splicing is a natural process which consists in splicing reaction between two independently transcribed pre-mRNAs using the cellular spliceosome machinery.¹⁴ SMarT has been developed to correct mutated mRNAs by trans-splicing with an exogenous RNA provided by an engineered pre-trans-splicing molecule (PTM).¹⁵ SMarT converts the mutant mRNA into wild-type mRNA and maintains natural regulation of the target mRNA providing a spatially and quantitatively appropriate level of expression.¹⁶,¹⁷ Additionally, SMarT strategy can cover distinct mutations with a unique molecular tool as it allows the replacement of large part of the targeted RNA. SMarT technology was used to reprogram...
5',19 3'19 or internal20 mRNA sequences. In this study, we assessed the feasibility of 3'- and 5'-trans-splicing strategies to reprogram Dnm2 transcripts in vitro and in vivo. We show that 5'-SmarT is a suitable approach for reprogramming the Dnm2 transcript. However, assessment of optimized strategies to increase efficiency and avoid the PTM self-translation will be crucial for further preclinical development.

**Results**

**Design of Dnm2 3'-PTM**

Given that the majority of the published reports has been concentrated on the 3'-trans-splicing approach, we first designed 3'-PTM targeting intron 10 in order to reprogram Dnm2 transcript from the exon 11 to the last exon. PTM contain: (i) a cytomegalovirus (CMV) promoter, (ii) antisense sequences (AS) complementary to intron 10 except for the negative control lacking the AS sequence (noAS-PTM), (iii) a conserved yeast branch point sequence (BP), (iv) a poly(A) tract, (v) a strong canonical 3' acceptor splice site (3'-SS),15 and (vi) the wild-type cDNA sequence of the mouse Dnm2 (exons 11 to 22) fused with a Flag epitope motif. The sequence from exon 12 to exon 22 was modified by codon optimization. These nucleotide modifications, preserve the translated protein sequence, increase the transgene expression, and allow detection of chimeric trans-spliced sequence by reverse transcriptase-polymerase chain reaction (RT-PCR). Sequence of exon 11 was unchanged in order to retain putative endogenous Exonic Splice Enhancer sites (Figure 1a). We designed five distinct AS (AS2 to AS6) of ~150 nucleotides in the 3' part of intron 10 (from -450 to +2 relative to exon 11). The AS3 masks the endogenous BP, poly(A) tract, 3'-SS, and the 3’ intron/exon boundary of exon 11 (Figure 1b). The targeted intron is involved in alternative splicing. Therefore, the expected trans-spliced product would contain the Dnm2 endogenous sequence from exon 1 to 11 including either exon 10 or 10 bis followed by Dnm2 optimized sequence (exons 12 to 22) and the Flag sequence.

3'-trans-spliced transcripts were detectable in cells despite unexpected multiple translation from the PTM

3'-PTM plasmids were transiently transfected into NIH3T3 fibroblasts and trans-splicing events were analyzed by RT-PCR 48 hours after transfection. Taking advantage of the Dnm2 modified sequence in PTM, we used oligonucleotides which amplify specifically trans-spliced transcripts (E8-F/E14Opt-R) or both cis- and trans-spliced transcripts (E9-F/E11-R) (Figure 1b). Trans-spliced mRNA was amplified after two rounds of PCR in cells transfected with AS2- and AS3-3'-PTM but not in untransfected cells (Figure 2a). Sequencing confirmed trans-splicing in both cases occurring either after endogenous exon 10 (isoform 1) or 10 bis (isoform 2) (not shown).

Trans-splicing events were also assessed by western blot using an anti-Flag antibody. We did not detect any protein at the expected molecular weight (~100 kDa). Dnm2 immunoprecipitation performed to increase the amount of Dnm2 in protein extracts, also failed to detect trans-spliced protein whatever the 3'-PTM (data not shown). However, western blot detected several additional bands migrating at molecular weight ranging from ~25 to 50 kDa (Figure 2b). Their presence for all the 3'-PTM, with or without AS, suggested that these bands came from the PTM itself rather than illegitimate trans-splicing events. We identified 11 putative open reading frames (ORFs) in-frame with the Flag epitope in the 3'-PTM (Figure 2c), four of them with a predicted molecular weight compatible with our observations (i.e., 39, 35, 33, and 26 kDa) (Figure 2a,b). Finally, Flag immunofluorescence detects these peptides everywhere in the cytoplasm preventing the discrimination of trans-spliced protein by this technique (not shown).

The 3'-PTM self-translation is associated with muscle toxicity in vivo

Recombinant adeno-associated viruses (AAV) expressing the 5 AS-3'-PTM and the noAS-PTM were produced and injected in Tibialis anterior (TA) skeletal muscles of three month-old wild-type mice (~ 4.1011 viral genomes (vg)/muscle). Muscle integrity was evaluated by Hematoxylin-Eosin staining on TA transverse sections 1.5 months later. Saline-injected control TAs showed normal myofibres except in small areas showing nuclear centralization compatible with muscle regeneration induced by needle injury. The number of fibres with nuclear centralization was drastically more important in TAs injected with AAV-PTM regardless of the AS sequence (reaching the totality of the muscle for the AS5 and AS6) (Figure 3a and Supplementary Figure S1a,b). To ensure that the nuclear centralization resulted from fibres regeneration, we looked at muscle integrity earlier after the injection. Indeed, a massive necrosis occurred 15 days after injection (Figure 3a). An intermediate level of regeneration was observed with AAV-AS2-3’-PTM and AAV-noAS-3’-PTM (see Supplementary Figure S1a,b).

The monitoring of the amount of AAV-AS5-3’-PTM particles over time by qPCR showed a relatively high level after 1 week (812 vg/ng of DNA), which then dropped at 2 weeks (333 vg/ng) to reach values close to the range of saline-control after 1.5 months (Figure 3b). These observations attested the virus loss in accordance with the regeneration level observed in Hematoxylin and Eosin (Figure 3b and Supplementary Figure S1a,b). However, 1.5 months after the injection, low level of viral particles was still detectable for AAV-AS2-3’-PTM and AAV-AS4-3’-PTM (58 and 21 vg/ng, respectively) (See Supplementary Figure S1c). Consequently, further RT-PCR and western blot analyses were performed to determine if these low remaining amounts of viral particles sustain trans-splicing events. The trans-splicing specific primers detect trans-spliced mRNA at the expected size for TA injected with AAV-AS2-3’-PTM and AAV-AS4-3’-PTM (Figure 3c). Even after a second round of PCR, no trans-spliced mRNA was detected for the other AS, the noAS and saline-injected TA (not shown). Sequencing of the trans-spliced products obtained for AS2 and AS4 confirmed the trans-splicing reaction in both cases. Trans-splicing occurred either after the Dnm2 endogenous exon 10 (isoform 1) or 10 bis (isoform 2) for both AAV-AS2-3’-PTM and AAV-AS4-3’-PTM (Figure 3d and Supplementary Figure S2a). The unexpected amplicon obtained for the AAV-AS5-3’-PTM (15 days after the injection) corresponds to an illegitimate trans-splicing event joining exon 8 to exon 11 (see Supplementary Figure S2b).
A sequence alignment between intron 8 and AS5 has shown that the largest region of similarity encompasses only 28 bp with 67% of identity (see Supplementary Figure S2c). This suggests that illegitimate trans-splicing reaction does not necessarily require high degree of similarity between untargeted regions and the binding sequence.

No trans-spliced protein was detected in muscle extracts even after Dnm2 immunoprecipitation. PTM expression was detectable by western blot only for AAV-AS4-3'-PTM (barely detectable for AS2 before immunoprecipitation) (see Supplementary Figure S2d and not shown) in accordance with the vg loss secondary to muscle necrosis. Consequently, alternative strategies to avoid translation from PTM and bypass PTM-related toxicity were developed.

Design of modified 3'-PTM succeeded to avoid PTM translation but not the toxicity

With the objective to force the translation of nontoxic molecule, the Green Fluorescent Protein (GFP) ORF was inserted between the antisense sequence and the splice site of our previous 3' constructs. The GFP ORF starts with a Kozak initiation context and ends with a series of three successive STOP codons to optimize the translation ending. It was therefore expected that only the GFP (and not the toxic peptides) is translated from the mRNA exported to the cytoplasm, with no consequence on translation but not the toxicity -PTM succeeded to avoid PTM expression is assessed using anti-Flag western blot (see Supplementary Figure S3a).

AAV viruses expressing AS2- and NoAS-GFP-inserted-3'-PTM were injected into TA muscle of three-month-old wild-type mice (1.1011 vg/muscle). After 1 month, Hematoxylin-Eosin staining showed a toxicity similar to the one observed for the 3'-classical strategy. Indeed, a large majority of fibres presented a nuclear centralization reflecting muscle regeneration (Figure 4c,d). In addition, no trans-splicing event was detected neither by RT-PCR nor western blot in muscles (data not shown).

Design of Dnm2 5'-PTM

Therefore, we developed 5'-trans-splicing strategy allowing the deletion of the polyadenylation site of the PTM (supplied by the endogenous RNA when trans-splicing occurs). Indeed, the polyadenylation contributes to mRNA stability, nuclear export and translation. We produced 5'-PTM expression plasmids containing: (i) a CMV promoter, (ii) an optional 150 base-pair-length sequence mimicking presence of intronic sequence known to enhance expression of the constructs and called “intron” thereafter, (iii) the 5' part of optimized wild-type Dnm2 cDNA sequence (exons 1 to 13 including exon 10 bis) fused in 5' to Flag tag, (iv) a strong 5' donor splice site (5'-SS) followed by a downstream intronic splicing enhancer

Figure 1 The 3' trans-splicing strategy. (a) Schematic illustration of the 3'-PTM constructs containing CMV promoter, the AS sequence, the 3' splice site, the wild-type Dnm2 cDNA (optimized sequence of exons 11 to 22 with exon 13 bis) in frame with Flag epitope, and the polyadenylation site. (b) Schematic illustration of 3' trans-splicing reaction between the endogenous Dnm2 mRNA and the 3'-PTM correcting the mutation in exon 11. Locations of AS sequences are shown in the dotted line square. Primers used for trans-spliced mRNA amplification by RT-PCR are depicted by arrows on the trans-spliced mRNA. CMV, cytomegalovirus promoter; AS, antisense sequence; SS, splice site; pA, polyadenylation site; Dnm2, Dynamin 2; PTM, pre-trans-splicing molecules; RT-PCR, reverse transcription-polymerase chain reaction.
sequence previously reported to enhance the 5’ trans-splicing efficiency, (v) a 150 nucleotides AS complementary to intron 13, and (vi) an optional 3'-UTR with polyadenylation site (Figure 5a). We designed three consecutive AS starting from the exon intron junction (AS1, AS2, and AS3). The AS1 masks the endogenous 5'-SS and the 3' exon/intron boundary of exon 13 (Figure 5b). For each AS, four constructs were made including or not the intron and polyadenylation site (i.e., ±pA and ±int). These constructs were named AS-5'-PTM, AS+int-5'-PTM, AS-ΔpA-5'-PTM, AS+int-ΔpA-5'-PTM (Figure 5a). Since Dnm2 alternative splicing occurs in the targeted region, the expected trans-spliced products would contain the Flag sequence and Dnm2 optimized sequence (exons 1 to 13) followed by endogenous Dnm2 sequence starting from exon 13 bis or 14 to exon 22.

5'-PTM are not translated and trans-spliced transcripts are detectable in vitro

The four constructs for each AS were transfected in NIH3T3 cells. No PTM translation was detected by western blot using anti-Flag antibody, regardless of the presence of polyadenylation site or intron (Figure 6a and not shown). In addition, trans-spliced protein was detected at the expected molecular weight for the AS3+int-5'-PTM construct after Dnm2 immunoprecipitation (Figure 6a and Supplementary Figure S4a).

Expression of Dnm2 trans-spliced transcripts was assessed by nested RT-PCR 48 hours after transfection. We amplified bands at the expected size for most of the 5'-PTM constructs using trans-spliced Dnm2 specific primers (Figure 6b). No product was amplified in untransfected cells and with the two noAS control PTM (with or without pA). Semi-quantitative RT-PCRs amplifying cis-spliced endogenous Dnm2-mRNA, PTM-RNA, and Gapdh mRNA showed similar expression level in the different conditions (Figure 6b). Amplification sequencing confirmed trans-splicing events between optimized exon 13 and endogenous Dnm2 sequence (Figure 6c and data not shown). Moreover, sequencing revealed that the natural alternative splicing of exon 13 bis is preserved since we observed exclusively heterozygous sequences corresponding to isoform 2 (including exon 13 bis) mixed with isoform 4 (without exon 13 bis) (Figure 6c and not shown), except for cells transfected with AS2-ΔpA-5'-PTM showing only isoform 2.

We detected flag expression by immunofluorescence in 3T3 cells transfected with AS1-, AS2-, and AS3-5'-PTM containing or not the polyadenylation site (+pA-PTM: ~1–4% of cells; ΔpA-PTM:<1% of cells, Supplementary Figure S5).
Rare cells are flag-positive with the noAS-PTM containing pA site suggesting a weak PTM translation but no flag-positive cell was detected using the noAS-∆pA-PTM. We thus assume that flag staining observed in cells transfected with AS-∆pA-PTM represents trans-spliced Dnm2 protein. The flag staining pattern is compatible with normal Dnm2 subcellular localization and colocalization between trans-spliced and endogenous Dnm2 was confirmed by confocal imaging (Figure 6d).

**5′-PTM is not toxic and trans-splicing events occurred in vivo**

The AS1-, AS2-, AS3-, and noAS-5′-PTMs and their counterpart devoid of polyadenylation site (∆pA) were produced in recombinant AAV. We injected 1.1011 vg or phosphate-buffered saline (PBS) into TA muscles of 3-month-old wild-type mice that were analyzed 1 month later. Hematoxylin-Eosin staining showed that areas containing fibres with internalized nuclei were similar for PBS-injected TAs (i.e., ~5% of the muscle section) and TAs injected with AAV-5′-PTM with or without pA site (i.e., 0.75-13% of the muscle section) (Figure 7a).

Nested RT-PCR was performed on RNA extracted from muscles using cis- and trans-splicing specific primers described above (Figure 5b). We detected Dnm2 trans-spliced transcripts at the expected size only in muscles injected with AS1, AS1-∆pA, and AS2-∆pA (Figure 7b). Semi-quantitative RT-PCR amplifying endogenous Dnm2 mRNA, PTM-RNA, and Gapdh mRNA showed comparable level of expression in all conditions (Figure 7b). Sequencing of trans-spliced transcripts showed expected chimeric sequence with splice between optimized exon 13 and endogenous sequence. RT-PCR amplified specifically either isoform 4 (without exon 13 bis) or isoform 2 (with exon 13 bis) for muscle transduced with AAV-AS1- and AAV-AS2-5′-PTMs, respectively (Figure 7c).

Flagged and endogenous Dnm2 colocalized in restricted area of transversal TA-sections injected with AAV-AS1-5′-PTM and AAV-AS1-∆pA-5′-PTM (1–3 groups of around 20
fibres per muscle section) (Figure 7d). There was no flag-positive area in muscle injected with the noAS constructs. The presence of Dnm2 trans-spliced protein in muscle injected with AAV-AS1-5′-PTM was further confirmed by western blot after Dnm2 immunoprecipitation (see Supplementary Figure S4b).

**Discussion**

Among the existing platform technologies developed for gene therapy, SMarT combines major advantages including maintenance of endogenous regulation of the corrected mRNA, absence of overexpression, and correction of multiple mutations using a single therapeutic molecule. SMarT was successfully used in vitro and in vivo to repair mRNA in several context of genetic diseases. In particular, functional improvement was reached in cystic fibrosis,24 hemophilia A,28 X-linked immunodeficiency and Spinal muscular atrophy.30,35 With the objective to correct DNM2 mutations causing centronuclear myopathy, our aim was here to evaluate the feasibility of SMarT-induced reprograming of the mouse Dnm2-mRNA in vitro and in vivo.

The most striking result obtained in the 3′-replacement strategy is the expression of several peptides fused with the flag epitope. The presence of 11 putative AUG-ORF in frame with the flag in the PTM coding sequence argues for PTM self-translation as already indicated in 3′-replacement studies.15,33,34 Monjaret and collaborators recently demonstrated undesirable ORFs translation as well as PTM internal cis-splicing from Titin- and Dysferlin-PTMs.33 Given that ORF prediction does not account for all the bands observed with the Dnm2 3′-PTM, similar cis-splicing events cannot be ruled out in this study despite of absence of putative donor splice site. It may be hypothesized that some observed peptides come from the use of cryptic CUG start codons (see Figure 1b) as already reported in mammals37 or from post-translational modifications of canonical AUG-ORF.

Regardless of their mechanism of production, the major concern relative to the 3′-PTM expression is its huge toxicity in vivo. Similar observations using the noAS-PTM, suggest toxicity linked to PTM translation rather than illegitimate trans-splicing. Parts or totality of Dnm2 PH, GTPase effector domain, and proline-rich domain domains can be translated from the 3′-PTM coding sequence depending of the start codon used (Figure 1b). Therefore, we can hypothesize that truncated peptides can trap Dnm2 partners or act as dominant negative mutants to impair Dnm2 functions. In addition, muscle toxicity could be linked to other mechanisms such
as proteasome dysregulation or expression of undetectable peptides from other phases. Despite of massive viral loss secondary to necrosis-regeneration, low level of 3′ trans-spliced mRNAs was detected indicating that intron 10 is an accurate targetable region for Dnm2 3′-trans-splicing. This result led us to assess alternative ways to avoid PTM translation. An “GFP-inserted 3′-PTM” strategy was developed based on the monocistronic expression of two alternative splice sites (exons 10 and 10 bis) containing the Flag, the Dnm2 GTPase and Middle domain (525 aa) plus few amino acids encoded by the linker and a part of the AS sequence (59 to 64 kDa) is shown below the PTM. Primers used for RT-PCR mRNA amplification are depicted by arrows below the TS mRNA. CMV, cytomegalo virus promoter; AS, antisense sequence; SS, splices site; DISE, Downstream intronic splice enhancer; pA, polyadenylation site. PTM, pre-trans-splicing molecules; RT-PCR, reverse transcription-polymerase chain reaction; TA, Tibialis anterior.

Figure 5 The 5′ trans-splicing strategy. (a) Schematic illustration the 5′-PTM constructs containing a CMV promoter, the wild-type Dnm2 cDNA (optimized sequence of exons 1 to 13 including exon 10 bis), a 5′ splice donor site followed by the AS and with or without polyadenylation site and intron. (b) Schematic illustration of the 5′ trans-splicing reaction between the 5′-PTM and the endogenous Dnm2 mRNA correcting the mutation in exon 11. Location of AS sequences is shown above the dotted line square. The predicted translation of the PTM (59 to 64 Kda PTM peptide) is shown below the PTM. CMV, cytomegalo virus promoter; AS, antisense sequence; SS, splices site; DISE, Downstream intronic splice enhancer; pA, polyadenylation site.

In conclusion, we provide here the first evidence that trans-splicing inducing molecular tools. Four major DNM2 isoforms are expressed using a combination of two alternative splice sites (i.e., exons 10 and 10 bis are alternatively spliced and an exon 13 bis can be included or not). These four isoforms are ubiquitously expressed but exhibit different intracellular distribution and support distinct cellular functions. Consequently, their preservation in a therapeutic approach is an important issue. The two intronic regions targeted in our study are involved in alternative splicing (i.e., intron 10 and 13 in the 3′- and 5′-replacement strategies, respectively). In the 3′-replacement study, an exclusive alternative splicing after either exon 10 or 10 bis is observed in vitro and in vivo suggesting that the location of the AS guides the spliceosome in favor of one event. Conversely, the chosen AS for 5′-replacement do not bias the natural splicing, since the large majority of trans-spliced product sequences shows both splice isoforms in vitro. However, contrasting again with our in vitro results, the splicing is altered in vivo, since either trans-spliced isoform 4 or isoform 2 are produced. The reasons of this discrepancy between in vitro and in vivo contexts are unclear and could rely on differences between splicing regulations between 3T3 cells and muscle. Therefore, maintenance of natural alternative splicing in region targeted by SMarT needs to be carefully considered.
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to increase efficiency of DNM2-mRNA reprogramming to correct pathogenic DNM2 mutations in patient-derived cells and animal models of the autosomal dominant centronuclear myopathy, two important steps of preclinical development.

Materials and methods

**PTM constructs.** The 5′- and 3′-PTM coding sequences were optimized by GeneArt technology (GeneArt, Regensburg, Germany). For the 3′-PTM constructs, the Dnm2 coding sequence (i.e., endogenous exon 11 followed by exons 12 to 22 optimized sequence) was generated by a PCR-Fusion from mouse cDNA and Dnm2 optimized cDNA (GeneArt) and cloned into pGEMT-Easy vector (Promega, Charbonnieres, France). The linker including the branching point, the polypurimidine tract and the splice acceptor site (5′-ccgcsccgacttataagctgctgaTCT AACTGATATCTTTTTTTTTTTTTTTTTTTTTAAATACAG-3′) preceded by a KpnI restriction site was added by PCR in 5′ of the PTM coding sequence.

**AS2–AS6 in intron 10.** AS2 (IVS10 −300 to −150); AS3 (IVS10 −150 to +2); AS4 (IVS10 −450 to −298), AS5 (IVS10 −252 to −96), AS6 (IVS10 −320 to −96) (where nucleotide +1 is the first nucleotide of exon 11).

**For the GFP-inserted 3′-PTM constructs,** the full length GFP sequence was amplified from a GFP plasmid using a forward primer introducing a Kosak sequence and a reverse primer introducing three stop codons (both primers also

Figure 6 Detection of 5′ trans-splicing in vitro. (a) Left: Anti-Flag Western Blot on total proteins from 3T3 cells transfected with 5′-PTM or a plasmid expressing full length Dnm2-Flag fusion protein as control (Dynaflag). NT: nontransfected. Right: Dnm2 immunoprecipitation followed by Flag western blot on 3T3 cells transfected with 5′-PTM. The image is overexposed and over contrasted to enhance the detection of the band indicated by an asterisk. (b) RT-PCR analysis on mRNA extracted from 3T3 cells transfected with 5′-PTM. Dyna-endo: endogenous Dnm2 transcripts amplified using E10EndoF/E12endoR primers, TS: trans-spliced RNA amplified using E8OptiF/Ex17endoR and Ex11OptiF/Ex14endoR primers; PTM are amplified using E8OptF/Ex11OptiR primers. C−: negative PCR control. The isoform of Dnm2 identified after sequencing for each band is indicated below (i.e., isoform 2 or 4 or both). 3T3 cells were transfected three times. (c) Top: Schematic representation of Dnm2 alternative splicing leading to isoforms 2 and 4, locations of primers used for PCR amplification are depicted. Bottom: Example of heterozygous trans-spliced sequence showing Ex13 optimized sequence followed by ex13 bis and ex14 endogenous sequence. The orange squares show the optimized nucleotides. (d) Anti-Flag and anti-DNM2 immunofluorescence on 3T3 cells 48 hours after transfection with AS2-, AS3-, noAS-ΔpA-5′-PTM and full length Dnm2-Flag fusion constructs (scale bar = 10 µm). PTM, pre-trans-splicing molecules; RT-PCR, reverse transcription-polymerase chain reaction.
contain SacII restriction site). The PCR product was then cloned into the pSMD2-3′-PTM between the AS and the linker sequence after SacII digestion.

For the 5′-PTM constructs, the Dnm2 coding sequence (i.e., Dnm2 exons 1 to 13) was generated by PCR from Dnm2 optimized cDNA. The forward primer contains a KpnI restriction site followed by an ATG in a Kosak context, the Flag sequence and the first 27 nucleotides of the optimized Dnm2 exon 1. The reverse primer contains a SalI restriction site, the downstream intronic splicing enhancer element from the rat FGFR2 gene20 and the strong canonical 5′ donor splice site sequence (GTAAGAACAG) followed by the last 25 nucleotides of Dnm2 exon 13 (optimized). The PCR product was introduced into the pSMD2 vector between the CMV promoter and the polyA signal by ligation after KpnI-SalI digestion. The AS for 5′-PTM were amplified by PCR from mouse genomic DNA using primers both containing a SalI restriction site and introduced in pSMD2-5′-PTM in SalI site downstream to the linker sequence. The locations of AS in intron 13 are as follow: AS1 (IVS13 from −3 to +147); AS2 (IVS13 from +126 to +285); AS3 (IVS13 from +264 to +416) (where nucleotide −1 is the last nucleotide of exon 13). To remove the SV40 polyA signal, a supplementary

**Figure 7 Detection of 5′ trans-spliced Dynamin 2 mRNA and protein in muscles.** (a) Muscle integrity after AAV-mediated AS1-5′-PTM transduction; Histogram showing the area of muscle sections with internal or central nuclei (%). The area of fibres with internal nuclei was normalized by the total area of the muscle section (Mean ± SD, n = 3 sections counted for each condition). (b) RT-PCR analysis on mRNA extracted from transduced muscles using specific primers for trans-spliced and cis-spliced mRNA TS: Trans-spliced PCR amplicon, Endo: endogenous. (c) Electropherogram of trans-spliced amplicons showing Ex13 optimized sequence followed by ex13 bis (isoform 2, top) or ex14 endogenous sequence (isoform 4, bottom). The orange squares show the optimized nucleotides (d). Immunofluorescence analysis on TA transverse sections from mice injected of AAV-AS1-5′-PTM. Fibres were double-stained with anti-FLAG (red) and anti-DNM2 (green) antibodies. Scale bar = 50 µm. AAV, adeno-associated viruses; DNM2, dynamin 2; PTM, pre-trans-splicing molecules; RT-PCR, reverse transcription-polymerase chain reaction; TA, Tibialis anterior.
AAV-PTM (1.10^{11}–4.10^{11} vg/muscle) were performed in two intramuscular injections (30 µl/TA, within 24 hours interval) of a mix of 100 three-month-old C57/BL6 mice. Anesthesia was achieved by transfection in 293 cells as described previously using the CMV promoter to the end of the AS sequence. The latter fragment was then reinserted in a pSMD2 vector previously Xbal digested. For the others 5′-PTM sequences (i.e., AS1, AS2, and noAS) the coding sequence followed by the splicing sites and AS sequence were directly replaced in the latter ΔpA construct using HindIII/Xhol digestion. All constructs were validated by DNA sequencing. Sequences of primers used for cloning are available on request.

Cell culture and transfection. Mouse embryonic fibroblast NIH3T3 cells (from ATCC) were grown in Dulbecco’s modified Eagle’s media (Life Technologies, Villebon sur Yvette, France) supplemented with fetal bovine serum 10%, penicillin 100 U/ml, and streptomycin 100 µg/ml. The cells were maintained in 37°C humidified incubators containing 5% CO2 and ambient oxygen. For transfection, cells were grown to 70% confluency in six-well plates and transfected with 1.5 µg of pSMD2-Dnm2-PTM plasmids using Exgen 500 transfection reagent according to the manufacturer’s protocol (Euromedex, Strasbourg, France). Cells were transfected three times per condition and harvested 48 hours later for RNA or Protein analysis.

In vivo AAV injection. Experiments were performed on adult three-month-old C57BL6 mice. Anesthesia was achieved by transfection in 293 cells as described previously41 using the use of animals for research and were approved by an external Ethical committee (approval No. 00351.02 delivered by the French Ministry of Higher Education and Scientific Research).

RNA isolation and cDNA analysis. Total RNA was isolated using RNA easy or RNA easy Fibrous tissue mini kits (Qia gen, Courtaboeuf, France) for cells and muscles, respectively, according to the manufacturer’s protocol. Cells or tissue sections were passed through a 22G syringe several times for DNA isolation in the lysis buffer. Total RNA (1 µg) was submitted to reverse transcription using the Superscript III reverse transcriptase kit (Life Technologies) using random primers. cDNA were amplified by PCR under the following conditions: 94°C for 5 minutes, cycles of 30 seconds at 94°C, 30 seconds at the appropriate temperature (58 to 61°C), 30 seconds to 1 minute at 72°C, and a final step of 7 minutes at 72°C. Thirty five cycles were performed to amplify trans-spliced Dnm2 transcripts, 28 for endogenous Dnm2, 23 for Gapdh, and 27 for 18S strategies, 28 for endogenous Dnm2, 23 for Gapdh, and 27 for 18S for the 5′ and 3′ strategies are: E8endoF: 5′-GGGTTCTGTTTGTAGTGA CGACTGC-3′, and Ex17Endo-R: 5′-CCCAGGCCGCGAGGA ACGAG-3′. The 5′-PTM is amplified using E8OptiF and ex11OptiR (5′-TGTCAGATCGCTGTTTTTCTTC-3′). Gapdh primers are F: 5′-ACCACAGTCCATGCCCAC-3′ and R: 5′-TCCACACCCCTGTTGCTGTA-3′.

For nested PCR or when two rounds of PCR were done, the first PCR was diluted to 1/50 and 1µl was used for the second round for 35 cycles. DNA sequencing was performed by Eurofins (Paris, France).

Western blot and immunoprecipitation. For in vitro experiments, cell pellets were homogenized in lysis buffer containing 50 mmol/l of Tris–HCl pH 7.5, 150 mmol/l NaCl, 1mmol/l ethylenediaminetetraacetate (EDTA, NP40 1%) supplemented with protease inhibitor cocktail 1% (Sigma-Aldrich, Saint Quentin Fallavier, France) by sonication (10 seconds). For in vivo experiments, 50 muscle tissue sections (15 µm thickness) or one TA for immunoprecipitation were mechanically homogenized in the lysis buffer using Fastprep Lysing Matrix D and Fastprep apparatus (MP Biomedical, Illkirch, France). After centrifugation (14,000×g, 4°C, 15 minutes), protein concentration in the supernatant was determined with the BCA Protein Assay Kit (Thermo Scientific Pierce, Villebon sur Yvette, France). 40 µg of protein were mixed with loading buffer (50 mmol/l Tris–HCl, sodium dodecyl sulfate (SDS) 2%, glycerol 10%, beta-mercaptoethanol 1% and bromophenol blue) and denatured at 90°C for 5 minutes. Protein samples were separated on sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) 10% and transferred onto polyvinylidene difluoride (PVDF) membranes (0.45 µm pore size, Life Technologies) overnight at 100 mA at 4°C. Membranes were blocked for 1 hour at room temperature in PBS containing nonfat dry milk (5%) and Tween20 (0.1%) and then exposed to the following primary antibodies: mouse monoclonal anti-Flag (Sigma F1804), rabbit polyclonal anti-DNM2 (Abcam ab3457, Cambridge, UK) in PBS-Tween20 (0.1%), milk (1%) overnight at 4°C. Membranes were rinsed in PBS-Tween20 (0.1%) and incubated 1 hour with secondary horseradish peroxidase-conjugated antibodies (antimouse or antirabbit from Jackson ImmunoResearch, Suffolk, UK) in PBS-Tween20 (0.1%). Chemiluminescence was detected using ECL detection Kit (Merck-Millipore, Darmstadt, Germany). Acquisition was performed on G-Box (Ozyme, Montigny-le-Bretonneux, France).

For immunoprecipitation, cell or muscle protein extracts (1,200 µg) were diluted in 400 µl of lysis buffer (containing 50 mmol/l of Tris–HCl pH 7.5, 150 mmol/l NaCl, 1mmol/l EDTA, NP40 1%, protease inhibitor 1%) and precleared with 30 µl washed protein G–Sepharose (PGS 4 fast flow; GE Healthcare, Orsay, France). Precleared lysates were then gently rolled overnight at 4°C with 3 µg anti-DNM2 antibody (Abcam ab3457). Then, 30 µl washed PGS was added and gently rolled for 1 hour at 4°C. Pelleted PGS was taken up in loading sample buffer, boiled at 95°C for 5 minutes and submitted to western blot.

AAV production. AAV2/1 pseudotyped vectors were prepared by transfection in 293 cells as described previously using the

CCGAGAAGCTGTCGCCAGC-3′, Ex10endoF: 5′-GGTTGGTCA A GCTGAAGAGAC -3′, E12endoR: 5′-GGTTTGTGGTTGATGA CGACTGC-3′, and Ex17Endo-R: 5′-CCCAGGCCGCGAGGA ACGAG-3′. The 5′-PTM is amplified using E8OptiF and ex11OptiR (5′-TGTCAGATCGCTGTTTTTCTTC-3′). Gapdh primers are F: 5′-ACCACAGTCCATGCCCAC-3′ and R: 5′-TCCACACCCCTGTTGCTGTA-3′.
pSMD2-Dnm2-PTM plasmid, the pXX6 plasmid coding for the adenoviral sequences essential for AAV production, and the pRepCap plasmid coding for AAV1 capsid. Vector particles were purified on iodixanol gradient and concentrated on Amicon Ultra-15 100K columns (Merck-Millipore). The particle titer (number of vg/ml) was determined by quantitative PCR.

**Immunocytochemistry.** For immunostaining, transfected 3T3 cells were fixed 10 minutes at −20°C in methanol 48 hours after transfection. Cells were blocked 1 hour at room temperature in Tween 0.1%, Goat Serum 5%, Bovine serum albumin (BSA) 5% in PBS. Cells were then exposed to primary antibodies: rabbit anti-C-terminal DNM2 (Abcam ab34577) and mouse anti Flag (Sigma F1804) in PBS containing BSA 1% and Tween 0.1% overnight at 4°C. Cells were rinsed in PBS and incubated 1 hour at room temperature with the secondary antibodies (Goat antiamouse Alexa-Fluor 568 and Goat anti-rabbit Alexa-Fluor 488, Life Technologies) and mounted using VECTASHIELD mounting medium with 4’,6-diamindino-2-phenylindole (DAPI) (Vector Laboratories, Peterborough, UK). For immunofluorescence on TA sections, cryosections (8 µm thick) were fixed in paraformaldehyde 4% (15 minutes at room temperature). After washing in PBS, cryosections were permeabilized in Triton X-100 (0.5%) in PBS for 10 minutes at room temperature and blocked in PBS-Triton X-100 (0.1%), BSA (5%) and Donkey serum (5%) for 30 minutes. Sections were incubated with primary antibodies: rabbit anti-C-terminal DNM2 (Abcam) and Goat anti Flag (Abcam) overnight at 4°C, in PBS with Triton X-100 (0.1%) and BSA (1%). After PBS-Triton X100 (0.1%) washes, sections were incubated with secondary antibodies (Donkey anti-Goat Alexa 568 and Donkey anti-rabbit Alexa 488, Life Technologies) added simultaneously for 60 minutes at room temperature. The slides were mounted with VECTASHIELD mounting medium (Vector Laboratories). Images were acquired using either axiophot microscope (Zeiss) or confocal microscope (Olympus FV-1000).

**Histological analysis.** Muscle cryosections (8 µm thickness) were stained with Hematoxylin-Eosin by standard method. Light microscopy was performed using an upright microscope (DMR, Leica, Nanterre, France). Images were captured using a monochrome camera (DS-Ri1; Nikon) and NIS-Elements BR 3.2 software (Nikon). The area of fibres with internal nuclei was measured using the NIS-Elements BR 3.2 software (Nikon). The area of fibres with internal nuclei was measured using the NIS-Elements BR software and divided by the total area of the muscle section (n = 3 cryosections for each condition).

**Vg quantification.** Genomic DNA was extracted from mouse muscles sections using DNA purification kit (Promega) according to the manufacturer’s protocol. Copy numbers of AAV genome were measured on 100 ng of genomic DNA by quantitative real-time PCR on a LightCycler480 (Roche diagnostic, Meylan, France) by using TaqMan probe. A plasmid containing a part of the titin cDNA were tenfold serially diluted (from 10^2 to 10^6 copies) and used as a control to establish the standard curve for absolute quantification. The sequences of primers and Taqman probes are available on request. AAV genome copy number was expressed as an absolute value per ng of genomic DNA. All genomic DNA samples were analyzed in triplicates.

**Supplementary material**

**Figure S1.** Toxicity of 3'-PTM constructs in muscles.

**Figure S2.** Events identified after 3'-PTM transduction in muscles.

**Figure S3.** Events identified after expression of GFP-inserted-3’-PTM in cells.

**Figure S4.** Detection of 5’-trans-spliced DNM2 protein in cells and muscles.

**Figure S5.** Anti-Flag and anti-dynamin 2 immunofluorescence on 3T3 transfected cells.

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