CRISPR gene editing in pluripotent stem cells reveals the function of MBNL proteins during human in vitro myogenesis

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ABSTRACT

Alternative splicing has emerged as a fundamental mechanism for the spatiotemporal control of development. A better understanding of how this mechanism is regulated has the potential not only to elucidate fundamental biological principles, but also to decipher pathological mechanisms implicated in diseases where normal splicing networks are mis-regulated. Here, we took advantage of human pluripotent stem cells to decipher during human myogenesis the role of MBNL proteins, a family of tissue-specific splicing regulators whose loss of function is associated with Myotonic Dystrophy type 1 (DM1), an inherited neuromuscular disease. Thanks to the CRISPR/Cas9 technology, we generated human-induced pluripotent stem cells (hiPSCs) depleted in MBNL proteins and evaluated the consequences of their losses on the generation of skeletal muscle cells. Our results suggested that MBNL proteins are required for the late myogenic maturation. In addition, loss of MBNL1 and MBNL2 recapitulated the main features of DM1 observed in hiPSC-derived skeletal muscle cells. Comparative transcriptomic
analyses also revealed the muscle-related processes regulated by these proteins that are commonly mis-regulated in DM1. Together, our study reveals the temporal requirement of MBNL proteins in human myogenesis and should facilitate the identification of new therapeutic strategies capable to cope with the loss of function of these MBNL proteins.

**INTRODUCTION**

Alternative splicing has emerged in the post-genomic era as the main driver of proteome diversity. The identification that at least 95% of human genes produce multiple spliced RNA via alternative exon usage has revealed the prevalence of this additional layer of gene expression regulation (1, 2). The regulation of alternative splicing varies with cell type, during development or upon cellular differentiation, thereby participating in the fine-tuning of a gene signature temporally and spatially (3). To ensure this developmental and tissue-specific regulation, alternative splicing relies on complex mechanisms involving an intricate protein-RNA network. This leads to the precise and sequential recognition of cis-acting sequences on the pre-mRNA by trans-acting RNA-binding factors. It has become apparent that these coordinated splicing networks have important physiological functions in developmental processes and that disruption of these networks negatively impacts health and contributes to human diseases.

According to some estimates, up to 50% of all pathogenic mutations may affect splicing in some way (4). Illustrating this, Myotonic Dystrophy Type 1 (DM1) represents one paradigmatic example of genetic disease in which loss of MBNL splicing factors is considered as the central pathological event. DM1 is an autosomal dominant neuromuscular disease caused by abnormal expansion of CTG repeats in the 3’untranslated region (UTR) of the *Dystrophia*
myotonia protein kinase (DMPK) gene (5, 6). The pathogenesis of DM1 involves an RNA gain-of-function mechanism caused by the expression of mutant RNAs containing hundreds to thousands of CUG repeats that sequester Muscleblind-like RNA binding proteins (MBNL) involved in the regulation of alternative splicing (7, 8). This functional loss of MBNL proteins results in several alternative splicing mis-regulation events that have been associated for some of them to clinical symptoms such as myotonia, muscle weakness and insulin resistance (7, 9). Three different genes encode human MBNL homologs: MBNL1, MBNL2 and MBNL3 which all share structural similarities (8, 10, 11). MBNL1 and MBNL2 are ubiquitously expressed but while MBNL1 is found in similar levels in all tissues, MBNL2 is more abundant in the brain (10, 11). In a majority of tissues MBNL1 and MBNL2 expression rises during differentiation. In contrast, MBNL3 is expressed at much lower levels, with peak expression in placenta, and its functions are poorly defined compared to its two other paralogs (11, 12). According to this differential expression over time, MBNL1 and MBNL2 proteins have been shown to modulate the splicing profile of a large number of genes during fetal-to-adult transitions (13). At the functional level, knockout Mbnl1 mice display skeletal muscle defects such as myotonia or central nuclei only at an adult stage (10) whereas loss of Mbnl2 mice leads predominantly to brain phenotypes with no major defects in skeletal muscles (14). However, the exploitation of these loss-of-function models was hampered by compensatory mechanisms which result in an increased expression of Mbnl2 as observed in knock-out Mbnl1 mice (15). In contrast, Mbnl1(-/-); Mbnl2(-/-) double knockouts are embryonic lethal while Mbnl1(-/-); Mbnl2(+/-) mice are viable but develop severe phenotypes such as reduced lifespan, heart conduction block, severe myotonia and progressive skeletal muscle weakness (15, 16). Concerning Mbnl3(-/-) mice, results showed decreased skeletal muscle regeneration and reduced grip strength associated with age despite low expression of Mbnl3 in adult muscles. Moreover, all the phenotypes described in Mbnl3 (-/-) mice were not associated with significant changes in alternative
splicing (17, 18). Altogether, these studies have highlighted the developmental function of Mbnl proteins in mice, but they also raise the question of whether MBNL proteins are similarly involved in human development as well as the precise timing at which these proteins intervene.

In this study, we focused on the role of MBNL proteins during human skeletal myogenesis. Due to their capacity to differentiate into any cell types of an organism, we based our approach on the combined use of human induced pluripotent stem cells (hiPSCs) and CRISPR/Cas9 technology to generate hiPSCs depleted in MBNL homologs and to therefore evaluate the impact of MBNL proteins loss at the cellular and molecular level during the different developmental steps of myogenesis.
RESULTS

CRISPR/Cas9 strategy to generate MBNL KO(s) hiPSCs

To generate MBNL knockout hiPSC lines, sgRNAs were specifically designed to target one of the first coding exons as well as the presence of an enzymatic restriction site at the PAM sequence (Figure 1A and 1B) (19). Each sgRNA was complexed with SpCas9 protein to perform ribonucleoprotein transfection of dissociated hiPSCs (20). The selection of gene-edited clones was based on a two-step selection process. Thanks to the use of sgRNA targeting sequences within a restriction endonuclease site (Figure 1B; framed in purple) at the PAM level (Figure 1B; NGG in red), restriction fragment length polymorphism was first analyzed to rapidly detect CRISPR gene editing according to total or partial loss of restriction sites (Table S1). The second selection was based on Sanger DNA sequencing to select clones with INDELS that have generated STOP codons (Figure 1C). For MBNL1, we obtained an editing efficiency of 25% with 24 pre-selected clones out of 96 clones tested (10 with a homozygous profile and 14 heterozygous). After the sequencing of these 24 clones, 3 homozygous and 2 heterozygous clones were validated (Figure 1C and 1D and Supplementary Figure 1A). Concerning MBNL2, 9 clones were pre-selected out of 96 (editing efficiency of 9%) and 3 were finally validated by sequencing. We generated two heterozygous clones and obtained only one MBNL2(-/-) homozygous knockout clone (Figure 1C and Supplementary Figure 1A). Because of the compensatory phenomena described between Mbnl2 and Mbnl1 in knockout mice (15), double knockout MBNL1(-/-); MBNL2(-/-) (DKO) were generated (Figure 1C and 1D and Supplementary Figure 1A). The condition MBNL1(-/-); MBNL2(+/-) was also generated as Mbnl1(-/-); Mbnl2(-/-) double mice knockouts are embryonic lethal whereas Mbnl1(-/-); Mbnl2(+/-) mice are viable but develop severe phenotypes (15, 16). Finally, to decipher the contribution of MBNL3, we generated triple knockout of MBNL1(-/-); MBNL2(-/-); MBNL3(-/-). We obtained an editing efficiency of 22% and validated 5 different triple knockout (TKO)
clones, two of which were considered for further analyses (Figure 1C and 1D and Supplementary Figure 1A). For each generated clone, pluripotency was validated by flow cytometry of the pluripotent markers SSEA3 and TRA1-81. Genomic integrity was also verified by a karyotype analysis and SNP genotyping (Supplementary Figure 1B and 1C). To determine off-target activity of our gRNAs, we analyzed the most likely off-target sites, and no mutation induced by genome editing was observed (Table S2).

**No major impairment of the early myogenic commitment in the absence of MBNL proteins**

The capacity of MBNL-depleted human pluripotent stem cells to differentiate into skeletal muscle cells was evaluated by using a transgene-free protocol which can be schematically divided into three steps (21) (Figure 2A). We first monitored the expression of the different isoforms of MBNL at the protein and mRNA level along this myogenic differentiation process. As previously reported (22), MBNL1 and MBNL2 were not found to be expressed in undifferentiated human pluripotent stem cells. Their level of expression gradually increased as differentiation progresses to reach a maximum expression after ~20 days of differentiation, corresponding to the beginning of the final myogenic maturation step (Figure 2B and Supplementary Figure 2A and 2D).

In contrast, MBNL3 expression slightly increased during the first days of differentiation followed by its decreased expression at later time points (Figure 2B and Supplementary Figure 2A-D). As previously described in human primary myoblasts, our results also confirmed that MBNL1 is the most abundant MBNL proteins in hiPSC-derived skeletal muscle cells although MBNL1 and MBNL2 proteins follow a similar kinetic of expression (Figure 2B).

The analysis at the protein level demonstrated a specific lack of expression of MBNL paralogs in the different gene-edited hiPSC lines differentiated into skeletal muscle cells (Figure...
As reported in mouse model of Mbnl loss of function (15), we also observed a compensatory increase in MBNL1 and MBNL2 expression in MBNL2(-/-) and MBNL1(-/-), respectively (Figure 2C and 2D and Supplementary Figure 2E). MBNL3 level was found not to be increased as a compensatory response to MBNL1 and MBNL2 loss (Supplementary Figure 2C and 2E).

To evaluate the functional impact of MBNL proteins loss during myogenic differentiation, the expression of stage-specific muscle markers was evaluated at different time points. As MBNL proteins are known to be important players in DM1 pathogenesis (23, 24), we used DM1 hiPSC as a reference (Figure 3A and 3B). At 5 days of differentiation, approximately 80% of cells expressed the myogenic progenitor marker PAX3, with no significant difference between the different cell lines indicating that muscle lineage commitment is not affected by MBNL depletion (Figure 4A and Supplementary Figure 3A). Similar observation was made at 16 days of differentiation by the quantification of cells positive for DESMIN, a muscle-specific intermediate filament. Approximately 75% cells were DESMIN+ with no significant difference between conditions showing that the myogenic precursors developed into skeletal myoblasts in the absence of MBNL proteins (Figure 4B and Supplementary Figure 3B).

Despite the absence of phenotypes at the cellular level, we sought to evaluate the molecular consequences of MBNL depletion. We chose to focus on several gene transcripts important for skeletal muscle function and previously shown to be mis-spliced in DM1 patients’ biopsies (9, 25). While these mis-splicing events were identified in adult tissues, RT-PCR analysis showed that the splicing profiles of DMD exon 78, INSR exon 11 and MBNL1 exon 5 were significantly deregulated at the myoblast stage in DM1, MBNL1(-/-), DKO and TKO compared to non-DM1 control (WT) (Figure 4C and Supplementary Figure 3C). DKO showed more pronounced splicing changes than MBNL1(-/-) and DM1 conditions. A similar
observation was made with the depletion of the three paralogs (Figure 4C). In contrast, the loss of MBNL2 alone did not significantly lead to mis-splicing events when compared to non-DM1 control (WT) myoblasts illustrating the compensation of MBNL2 by MBNL1 as previously reported (23) (Figure 4C). These observations suggested that MBNL proteins are dispensable for myogenic commitment even if MBNL loss of function led to early mis-splicing events that could interfere with later stages of myogenesis.

MBNL1 and MBNL2 depletion impacts late myogenesis and reproduces the main features associated with DM1

We next evaluated the consequences of MBNL loss of function on the capacity of hiPSC-derived myoblasts to further differentiate and fuse into myotubes (21). As a point of reference, we first characterized this late differentiation by using DM1 hiPSC-derived myoblasts. Primary or immortalized muscle cell cultures derived from muscle biopsies of DM1 patients have been shown to present altered myogenic fusion as well as DM1-associated molecular features including mutated DMPK mRNAs aggregates that sequester MBNL1 and subsequently lead to alternative splicing defects (26). Accordingly, we also observed that DM1 hiPSC-derived myoblasts under differentiation condition form smaller myotubes and present a significantly reduced fusion capacity when compared to WT hiPSC-derived myoblasts (Figure 5A and Supplementary Figure 4A). Moreover, the main DM1-associated molecular features including mutated DMPK mRNAs aggregates, nuclear sequestration of MBNL1 and MBNL2 and presence of several mis-spliced events were detected in these cells (Figure 3C-E, Figure 5B-C and Supplementary Figure 4B). These data demonstrate that DM1 hiPSC-derived skeletal myotubes recapitulate phenotypic and molecular traits similar to those found in skeletal muscle cell cultures isolated from DM1 patients.
In comparison, loss of either MBNL1 or MBNL2 resulted in a moderate effect on myogenic fusion which is likely related to the compensatory phenomenon existing between the two paralogs (Figure 5A and Supplementary Figure 4A). Immunofluorescence analysis revealed that this upregulation was mainly localized in the nucleus (Figure 5B), in agreement with previous observations made in murine models of Mbnl loss of function (15). In contrast to single MBNL knock-out lines, MBNL1(-/-); MBNL2(+/-) and DKO hiPSC-derived myotubes showed an aggravated fusion defect (Figure 5A). Concordant with the absence of MBNL3 protein at this stage, depletion of MBNL3 in addition to the two paralogs did not exacerbate this phenotype in TKO lines when compared to DKO (Figure 5A).

At the molecular level, and concordant with the observations made at the myoblast stage, several DM1-associated mis-splicing events, known to contribute to muscle function and maintenance such as BIN1 exon 7 and DMD exon78, were found to be mis-spliced in DM1 hiPSC-derived myotubes (Figure 5C). For all targets examined, dramatic mis-splicing events were also observed in DKO myotubes (Figure 5C). These data strongly suggest that MBNL1 and MBNL2 proteins regulate RNA processing essential for normal late myogenesis.

**Comparative analysis of the molecular pathways misregulated between DM1, MBNL1 KO and DKO hiPSC-derived myotubes**

To better evaluate the consequences of MBNL1 and MBNL2 depletions during myogenesis, we sought to conduct a comparative transcriptomic analysis between DKO and WT hiPSC-derived myotubes. We also included MBNL1(-/-) hiPSC-derived myotubes to evaluate the effect of MBNL2 compensation. For this purpose, we used AmpliSeq targeted sequencing technology allowing analysis of over 20,000 distinct human RNA targets simultaneously using a highly multiplexed amplification method. The principal component analysis (PCA) showed a distinct transcriptomic profile for each condition (Figure 6A). The
comparison between MBNL1(-/-) and WT hiPSC-derived myotubes led to the identification of 126 genes differentially expressed (false discovery rate (FDR) < 0.05; Log2FoldChange > 0.4), with 31 being up-regulated and 95 down-regulated in MBNL1(-/-) myotubes compared to WT (Figure 6B). In the same order of magnitude, 122 differentially expressed genes (DEGs) were found when comparing DKO to WT hiPSC-derived myotubes, 40 up-regulated and 82 down-regulated in DKO myotubes. A more pronounced difference was observed between DKO and MBNL1(-/-) hiPSC-derived myotubes with 258 differentially expressed genes (Figure 6B). All the differentially expressed genes (DEGs) are listed in table S3. Strikingly, only 12 genes were commonly deregulated in MBNL1(-/-) versus WT myotubes and in DKO versus WT myotubes (Figure 6C). Hierarchical clustering on the 122 DEGs detected in DKO compared to WT hiPSC-derived myotubes showed that MBNL1(-/-) hiPSC-derived myotubes were comparable to WT hiPSC-derived myotubes (Figure 6D). Accordingly, gene ontology (GO) analysis of DEGs in MBNL1(-/-) versus WT myotubes revealed enrichment in multiple biological processes, but few of these are related to skeletal muscle (Figure 6E and Table S4). These observations could be explained, at least, by the functional compensation by MBNL2 in MBNL1(-/-) myotubes.

Concerning DEGs in DKO compared to WT hiPSC-derived myotubes, gene ontology analysis revealed enrichment in several biological processes related to skeletal muscle such as “striated muscle contraction” (adjusted pvalue: 0.00001), “actomyosin structure organization” (adjusted pvalue: 0.02341) or “muscle filament sliding” (adjusted pvalue: 0.02348) (Figure 6E). Strikingly, most of these muscle related-biological processes were found to be down-regulated in DKO hiPSC-derived myotubes (Figure 6F). In the same way, gene ontology analyses of DEGs in DKO compared to MBNL1(-/-) hiPSC-derived myotubes showed similar results with multiple down-regulated biological processes related to skeletal muscle (Supplementary Figure 5A and 5B). Among the list of down-regulated genes identified in DKO hiPSC-derived
myotubes, we highlighted two interesting candidates for further explorations (Figure 6F): TCAP (also known as Telethonin) plays an essential role in the sarcomeric assembly by anchoring Titin in the Z-disk of the sarcomere (27) and is known to be involved in Limb-Girdle Muscular Dystrophy 2G (LGMD2G) (28); MYOM1 is a constituent of the sarcomeric M band and plays an important role in sarcomere assembly and integrity (29). Interestingly, MYOM1 is found mis-spliced in muscular biopsies from DM1 patients (30) and this splicing defect could be caused by the down-regulation of MBNL proteins (31).

Thus, we pursued our analysis by searching for cassette exons that demonstrate a significant difference in the level of inclusion between MBNL1(-/-), DKO and WT hiPSC-derived myotubes. A total of 547 cassette exons were found to be deregulated between DKO and WT hiPSC-derived myotubes, among which 296 and 251 exhibited a significant decrease and increase in inclusion, respectively (FDR < 0.05; Delta percent spliced-in (PSI) > 0.4) (Figure 7A and Table S5). We next compared these analyses with transcriptomic data generated from DM1 hiPSC-derived myotubes (Figure 7A). Of the 696 mis-splicing events (DSE) observed in DM1 hiPSC-derived myotubes, 224 (32%) appeared in MBNL1(-/-) hiPSC-derived myotubes. In the same way, 254 (36%) appeared in DKO hiPSC-derived myotubes and among these, 126 were commonly mis-spliced in DM1, MBNL1(-/-) and DKO hiPSC-derived myotubes (Figure 7B).

Functional annotation analysis of the DKO transcriptomic data revealed that the most relevant GO terms concerned skeletal muscle such as "sarcomere organization" (adjusted pvalue: 0.068) or "neuromuscular junction development" (adjusted pvalue: 0.06) (Figure 7C and Table S6). We highlighted 30 enriched DSEs involved in most of these muscle-related biological processes deregulated in DKO myotubes (Figure 7D), and found several already described alternative splicing defects associated with DM1 such as DMD exon 78, BIN1 exon 11 and BIN1 exon 7 (Figure 5C and Supplementary Figure 4B) (32, 33). Among these 30 DSEs,
21 were also mis-spliced in DM1 hiPSCs-derived myotubes in the same way they were in DKO myotubes (Supplementary Figure 6A). Only 3 DSEs (ABI2 ex10, MYL5 ex3 and TPM1 ex9) mismatched in the sense of inclusion/exclusion. Furthermore, only half of these 30 DSEs were also found deregulated in MBNL1(-/-) myotubes, which could explain why the biological processes deregulated in these cells are less related to skeletal muscle. In general, when DSEs concern both MBNL1(-/-) and DKO myotubes, the level of deregulation was worsened in DKO myotubes (Supplementary Figure 6A and 6B). In addition to DMD exon 78, BIN1 exon 11 and BIN1 exon 7, we validated 3 other DSEs in our MBNL knockouts hiPSC-derived myotubes (Supplementary Figure 6B), including the aberrant inclusion of MYOM1 exon 18 in DM1 hiPSC-derived myotubes compared to WT (Supplementary Figure 6A and Table S5) which was also validated in DKO myotubes (Supplementary Figure 6B).

Altogether, these data revealed sets of disrupted biological pathways associated to the loss of MBNL1 and MBNL2 that may be particularly relevant in the pathological context of DM1.
DISCUSSION

Cell-type-specific or developmental-stage-specific alternative splicing events are largely controlled by recruiting RNA-binding proteins (RBPs) that recognize specific regulatory sequences embedded in the pre-mRNA transcripts. In this study, we described the temporal requirement of the Muscleblind-like (MBNL) RBP family for defining human skeletal muscle maturation stages. Our results demonstrated that MBNL1 and MBNL2 were dispensable for the early myogenic commitment but played an important role for the late stages of myogenesis. When compared with hiPSC generated from DM1 patients, our results show that loss of MBNL1 and MBNL2 recapitulate the main cellular and molecular features associated to this neuromuscular disease. This study provides insights into the function of MBNL proteins during myogenesis. Given the technical limitations of studying embryonic and fetal human tissues, our study establishes new human cellular models to explore the role of these proteins in tissue development. Furthermore, the different hiPSC-based models developed in this study also pave the way towards the development of new therapeutic strategies capable to cope the loss of function of MBNL proteins.

While transcriptional regulation has been demonstrated to play an important role in myogenesis, there is a growing recognition of the contributions of alternative splicing to skeletal muscle development as well as the refinement of muscle function (34). Skeletal muscle was also one of the first tissues in which broad use of alternative splicing to increase cellular complexity was documented (35). A recent study has estimated the presence of hundreds of RBPs expressed in muscle of different species such as human, rodent, flies and worms, but only a small proportion (around 3%) have been studied (36). Specifically, the role of CELF (CUG-BP and ETR-3 like), Muscleblind-like (MBNL) and Fox families in controlling muscle-specific splice events has been largely documented in murine models. The Rbfox proteins are among the most heavily studied RBPs in the context of myogenesis. Gain- and loss-of-function
experiments have demonstrated that Rbfox1 and Rbfox2 coordinately regulate splicing of the transcription factors Mef2a and Mef2d to maintain the expression and proper activity of muscle-specific isoforms (34, 36). More recently, the generation of skeletal muscle-specific knockout of Rbfox1 and Rbfox2 in adult mice also revealed a role of these alternative splicing factors in the maintenance of skeletal muscle mass. This function could at least be attributable to the altered expression of calpain 3 splice isoforms that are not degraded rapidly leading to the accumulation of active calpain 3 and a defective proteostasis (37). As illustrated by this example, a major part of our knowledge has been gleaned from studies in animal models. Given the large number of RBPs expressed and with potential phenotypes in human muscle, this clearly illustrates the need for further studies examining the function of RBPs in human myogenesis.

In this context, human pluripotent stem cells represent an attractive cell source for producing myogenic cells and deciphering the function of different RBPs during this process. Until recently, the only efficient protocols allowing differentiation of human pluripotent stem cells into a reasonably mature muscle cells relied on the forced expression of transcription factors such as MyoD or Pax3/7 (38). These direct reprogramming approaches aim to bypass early developmental stages by overexpressing myogenic regulators and the cellular events occurring during differentiation remain largely unclear (39). More recently, stepwise induction protocols based on the use of small molecules and growth factors have been established and appeared as an alternative approach to better recapitulate the successive developmental stages involved in skeletal myogenesis (21, 40). By using such approach, our study adds a new dimension to the role of MBNL proteins in defining human skeletal muscle maturation stages. The exact mechanisms by which MBNL proteins affect these processes are however unclear. In contrast to Rbfox1/2 for which depletion has been shown to inhibit myoblast fusion in vitro through the mis-splicing of Mef2D (41, 42), our results indicate that depletion of MBNL
proteins leads to the aberrant expression of hundreds of genes. Since several of these genes encode factors involved in sarcomere organization, actin dynamics and vesicular trafficking events required for membrane fusion (43), further experiments will be required to evaluate the exact MBNL-based mechanisms required for myogenic differentiation.

Among the three MBNL paralogs, our results confirmed that MBNL1 is the most abundant MBNL protein during in vitro myogenesis (12, 14). We also observed a meticulous compensation of MBNL1 by MBNL2 since MBNL2 protein expression was increased in either MBNL1(-/-) and MBNL1(-/-); MBNL2(+/-) compared to WT and MBNL2(+/+) hiPSC-derived myotubes respectively. Moreover, transcriptomic analyses showed that several skeletal muscle-related biological processes were down-regulated in DKO myotubes compared to MBNL1(-/-) myotubes. Taking together, our results strengthen this compensation phenomenon between MBNL1 and MBNL2 proteins in myogenesis as suggested in prior studies in mice (15, 16). In contrast to MBNL1 and MBNL2, the expression of MBNL3 is found to be very low and limited in duration. The role of MBNL3 during myogenesis is not well understood. Whereas constitutive expression of Mbnl3 in murine muscle precursors cells appears to inhibit terminal differentiation (44), it has been recently shown that Mbnl3 depleted primary murine myoblasts displayed defective in vitro differentiation (24). Our results showed that triple knockouts hiPSC-derived skeletal muscle cells recapitulated all the phenotypes identified in double knockouts suggesting that MBNL3 might not play such important role during human myogenesis.

Sequestration of MBNL proteins is considered a major player in DM1 pathology. Comparative transcriptomic analysis between Mbnl1 knockout mice and transgenic mice expressing CUG expansion RNA have shown that the majority of changes induced by the abnormal expansions in skeletal muscle could be explained by reduced activity of Mbnl1 (23, 45, 46). However, it has been shown more recently that the combined loss of Mbnl1 and Mbnl2
was insufficient to reproduce the extensive gene expression changes detected in DM1, particularly in its congenital form which is the most severe form of the disease associated with large numbers of CTG expansion (24, 47, 48). Accordingly, we showed here that the loss of MBNL1 as well as its combined loss with MBNL2 do not fully recapitulate the gene expression changes that we observed in skeletal muscle cells derived from DM1 hiPSC carrying large number of CTG expansions. Altogether, these results strongly suggest that additional pathways may play non-negligible roles in the disease. Among the different hypotheses that can be formed, MBNL1 has been implicated in altered miRNA biogenesis (48), RNA localization (23) and in the misregulation of alternative polyadenylation (49) in DM1, and it would be relevant to investigate the effects of MBNL loss on these regulatory pathways. In addition, the stabilization and overexpression of CUG-BP1, due to its hyper-phosphorylation as a result of the alteration of the GSK3β pathway in DM1, have been shown to lead to several splicing defects (50, 51). Furthermore, the splicing factor hnRNP-H found to be deregulated in DM1 may also contribute to the disease spliceopathy (52).

Finally, to this day, only a couple of studies have described the generation of DM1 hiPSC-derived skeletal muscle cells and the capacity of these cellular systems to reproduce the main molecular features associated with DM1, namely the nuclear aggregation of mutated mRNAs and some alternative splicing defects. Despite promising results, all these attempts to generate skeletal muscle cells from DM1 hiPSCs were based on the forced expression of myogenic transcription factors MyoD or Pax7 which, as discussed above, are often undefined (53-55). The experimental system described here uses a transgene-free protocol to generate skeletal muscle cells from hiPSCs and opens new perspectives in term of disease modeling and drug screening. First, the different well-characterized steps of the protocol help to decipher the exact impact of the mutation on myogenic differentiation. Whereas most of the studies have focused on human primary skeletal muscle cells or immortalized skeletal muscle cells
originating from DM1 patients’ biopsies (26), our results indicate that early myogenic commitment is not affected by the presence of the DM1 mutation. In addition, the transgene-free protocol described here includes a freezing step allowing the possibility to generate large cell banks. Consequently, this represent an unlimited source of material that could be used for drug screening.

In summary, our results provide important new experimental support for the MBNL loss-of-function model for DM and introduce new human cellular models to examine the tissue-specific consequences of MBNL loss.
MATERIALS AND METHODS

Cell culture

Human induced pluripotent stem cells were derived by Phenocell® from human primary fibroblasts reprogrammed using sendai virus vectors carrying OCT4, KLF4, SOX2 and c-Myc (56). Two DM1 human induced pluripotent stem cell lines were generated from two independent patients’ fibroblasts containing around 2500 CTG repeats. DM1 hiPSC carry a mutant DMPK gene containing approximately 2700 CTG repeats. One control hiPSC line corresponding to the sibling matched control was also generated. Informed consents were obtained from all the patients included in this study, complying with the ethical guidelines of the institutions and with the legislation requirements of the country of origin. Experimental protocols were approved by the french minister of health (2019-A02599-48). All the hiPSC lines were used at passages around p10-p20 and were maintained in StemMACS iPS-Brew XF medium (Miltenyi Biotec®) in vitronectin (Gibco®) coated culture dishes. Culture medium was changed every two days and confluent cells were manually passaged every 5-7 days.

hiPSCs skeletal muscle differentiation experiments were performed using the commercially available Genea Biocells® skeletal muscle differentiation media following the protocol described previously (21). Briefly, hiPSCs were dissociated with StemPro Accutase Cell Dissociation Reagent (Gibco®) and plated at 3000 cells per cm² onto collagen I-coated plates (Gibco®) and maintained for 10 days in skeletal muscle induction medium (SKM01). At day 10, cells were dissociated with Trypsin-EDTA 0,05% (Gibco®), plated at 5000 cells per cm² onto collagen I-coated plates and maintained for 6 days in skeletal myoblast medium (SKM02). At day 16, myoblasts were dissociated and cryopreserved. For terminal differentiation, cells were thawed at 10,000 cells per cm² in SKM02 and after 2-3 days, when cells reached confluence, the medium was replaced by the myotube medium (SKM03). Cells were then maintained for 5 to 6 days in SKM03.
sgRNAs design

sgRNAs and primers used in this study are reported in Table S1. SpCas9 target sequences within the first coding exon of each MBNL paralog were determined by CRISPOR (http://crispor.tefor.net/). sgRNAs with the highest predicted efficiency, the lowest number of potential off-targets and having an enzymatic restriction site at the PAM sequence were selected. crRNA and tracrRNA were synthesized by Integrated DNA Technologies®.

RNP transfection of hiPSCs

Control hiPS cells were dissociated with StemPro Accutase Cell Dissociation Reagent (Gibco®), plated in 24-well plates at 40,000 cells per cm² and transfected the next day with a mixed of 30nM of sgRNA (by duplexing RNA oligos: crRNA and tracrRNA), 30nM of SpCas9 purified protein (gift from Jean-Paul Concordet, MNHN-CNRS UMR 7196 / INSERM U1154) and Lipofectamine Stem Transfection Reagent (ThermoFisher Scientific®), according to the manufacturer’s protocol.

Genomic DNA extraction and PCR

Genomic DNA was extracted from transfected hiPS cells either with QIAmp DNA Micro and Mini Kit (Qiagen®) or with QuickExtract DNA extraction solution (Lucigen®), according to the manufacturer’s instructions. PCR were performed using Q5 High-Fidelity DNA Polymerase (NEB®) with 30ng gDNA, 38 cycles of 10’’ at 98°C, 15’’ at 60°C and 45’’ at 72°C, with a final 10’ extension and with primers framing the target area according to each guide used (see Table S1).

RFLP assays to identify edited clones

Restriction Fragment Length Polymorphism (RFLP) presents the advantage to easily and rapidly estimate the presence of gene editing in a mixed population or in single clones. Briefly, 2µL PCR products were digested by 5 units of restriction enzyme (NEB®): BsrI for MBNL1.
and MBNL2 clones, and SmlI for MBNL3 clones. Cleaved fragments were then separated by gel electrophoresis. If CRISPR-mediated gene editing was successful, NHEJ-mediated INDELS abolished the restriction endonuclease recognition site. After restriction digest, clones that had lost their restriction site were selected and corresponding PCR products were sequenced by Sanger DNA sequencing (Eurofins®).

To ensure the purity of each clone selected, we used TOPO-TA cloning Kits for Sequencing (Invitrogen®) and One Shot TOP10 chemically Competent E. coli (Invitrogen®) according to manufacturer’s instructions. In order to determine the presence of subclones, at least 30 TA clones for each CRISPR cell line generated were analyzed by Sanger DNA sequencing (Eurofins®).

**Ion AmpliSeq transcriptome human gene expression kit**

For each sample, 50 ng of total RNA was reverse transcribed using the Ion AmpliSeq Transcriptome Human Gene Expression kit following the protocol of the manufacturer (Thermofisher Scientific®). The cDNA libraries were amplified and barcoded using Ion AmpliSeq Transcriptome Human Gene Expression core panel and Ion Xpress Barcode Adapter (Thermofisher Scientific®). The amplicons were quantified using Agilent High Sensitivity DNA kit before the samples were pooled in sets of eight. Emulsion PCR and Enrichment was performed on the Ion OT2 system Instrument using the Ion PI Hi-Q OT2 200 kit (Thermofisher Scientific®). Samples were loaded on an Ion PI v3 Chip and sequenced on the Ion Proton System using Ion PI Hi-Q sequencing 200 kit chemistry (200 bp read length; Thermofisher Scientific®). The quality control of the sequencing data was evaluated as described previously (57). The differential gene expression (DEG) between conditions was calculated with (v1.18.1 using R v3.4.1). Genes were considered differentially expressed when their adjusted pvalue was under 0.05 and their \(|\log2\text{FoldChange}|\) was upper than 0.4 (options: lfcThreshold=0.4, altHypothesis="greaterAbs"). Data are available on GEO database (GSE161029).
RNA-Seq library preparation and sequencing (TrueSeq polyA)

For each sample, 100ng of total RNA was used. mRNAs were purified using poly-T oligo from TruSeq Stranded kit (Illumina®) and were fragmented and reverse transcribed using the TruSeq Stranded kit following the protocol of the manufacturer. Final library was quantified using Agilent High Sensitivity DNA kit before the samples were pooled in sets of six for sequencing. 2nM of pooled libraries were denatured and the quantity of 1.8pM was used for cluster generation before paired-end sequencing on an Illumina Nextseq550 (Hight output 2*150pb run). Samples were sequenced with an average of 89 532 366 reads (paired-end) per sample.

The quality control of the sequencing data was evaluated using FastQC (v0.11.2). The reads were trimmed using Prinseq (v0.20.4) (--trim-right 20) and filtered by average quality score (--trim-qual 20) and cutadapt (v1.16) (58).

For the alternative splicing analysis, the raw reads were annotated on GRCh37.75 EnsEMBL reference annotation by Tophat2 (v2.0.13) and analyzed using only reads on exonic junctions with FaRLine pipeline and FasterDB database (59). Exons were considered differentially spliced when their adjusted pvalue was under 0.05 and their DeltaPSI (percent spliced-in) value was upper than 0.1. Data are available on GEO database (GSE161897).

RT-PCR and Agilent DNA chips analysis for Alternative Splicing

Total RNA was extracted using the RNeasy Micro/Mini kit (Qiagen®) and reverse transcribed using random hexamers and Superscript III Reverse Transcriptase kit (Invitrogen®) according to the manufacturer’s protocol. For splicing analysis, PCR amplification were carried out with recombinant Taq DNA polymerase (Invitrogen®) and primers listed in Table S1. The amplification was performed using a first step at 95°C for 3’ followed by 35 cycles of 45” at 95°C, 30” at 60°C, 1’ at 72°C, and a final 10’ extension at 72°C. The PCR products were analyzed using Agilent® DNA chips and quantified with the BioAnalyzer 2100.

Fluorescent In situ Hybridization
Cell preparation was performed as previously described. Briefly, cells were fixed with PBS buffer solution containing 4% PFA for 8’ at room temperature and incubated overnight at 4°C with 70% Ethanol (VWR®). After washes, cells were rehydrated with a solution of 5mM of MgCl2 (Sigma-Aldrich®) for 10’ at room temperature and then sequentially incubated with prehybridization buffer for 10’ and hybridization buffer containing the probe overnight at 37°C under agitation. Washing steps consisted in pre-warmed washing buffer for 1 hour at 37°C. Finally, cells were washed twice in PBS buffer and incubated 5’ in a Hoechst 33528 solution (Sigma-Aldrich®).

Nuclear foci detection was analyzed using the Cellomics CX7 high content imaging system (Thermofisher Scientific®) with the 20X objective and an automatically focus cell preparation. Images were acquired in High resolution camera mode on two channels and quantification was performed as previously described with the software attached. Each image was taken with z-stack and application of maximum projection.

**Immunocytochemistry**

After fixation with 4% PFA for 10’ at room temperature, cells were incubated over night at 4°C with the following primary antibodies: DESMIN (R&D System®, 1:100), PAX3 (DSHB®, 1:30), α-ACTININ (Sigma®, 1:250), MF20 (DSHB®, 1:200), MBNL1 - MB1a(4A8) (DSHB®, 1:100), MBNL2 - MB2a(3B4) (DSHB®, 1:100). Appropriated Alexa fluorescent secondary antibodies (Invitrogen®, 1:1000) and Hoechst (5μg/mL) were added for 1h. Image acquisitions were performed on the ImageXpress micro imaging system (Molecular Devices®) with the 20X objective and with three channels. Images were analyzed using MetaXpress software by quantifying the percentage of cells stained for PAX3 or DESMIN compared to the total number of cells.
Quantification of *in vitro* myoblasts fusion was performed counting the number of myotubes with at least 3 nuclei (manually counted) normalized to the total number of nuclei (at least 11,000 nuclei counted per experiment and per cell line).

**Protein extraction and Western blot analysis**

Western blots analyses were performed as previously described (59). Briefly, cells were lysed in RIPA 1X buffer (Sigma®) containing protease inhibitors (Sigma®) and phosphatase inhibitors (Roche®). Proteins were quantified by Pierce BCA Protein Assay kit (Pierce®). Protein extracts (15 to 20μg) were loaded on a 4-12% SDS-PAGE gradient (NuPage Bis–Tris gels, Invitrogen®) and transferred onto Gel Transfer Stacks Nitrocellulose membranes (Invitrogen®) using the iBlot2 Dry Blotting System (Invitrogen®). Membranes were then incubated overnight at 4°C with the following primary antibodies: MBNL1 - MB1a(4A8) (DSHB®, 1:1000), MBNL2 - MB2a(3B4) (DSHB®, 1:1000), MBNL3 – 5A11(sc-136168) (Santa Cruz, 1:100). After hybridization of the peroxidase-conjugated secondary antibody (1:10000), immunoreactive bands were revealed by using Amersham ECL Select Western Blotting Detection Reagents (GE Healthcare®). Equal protein loading was verified by the detection of β-Actin using the A3854 Monoclonal Anti-β-Actin–Peroxidase antibody (Sigma®, 1:10000).

**Statistical analysis**

All data were processed using Prism 8®. Values are represented as mean ± SD. For comparisons of more than two groups, statistical analyses were performed with by ordinary one-way analysis of variance (ANOVA) using Tukey’s multiple comparisons test. Values of p < 0.05 were considered significant (*p < 0.05; **p < 0.005; ***p < 0.0005; ****p < 0.0001).
ACKNOWLEDGEMENTS

I-Stem is part of the Biotherapies Institute for Rare Diseases (BIRD) supported by the Association Française contre les Myopathies (AFM-Téléthon). This project was supported by grants from Laboratoire d’Excellence Revive (Investissement d’Avenir; ANR-10-LABX-73), from the National Research Agency (ANR-16-CE17-0018) and from the doctoral school « Innovation Thérapeutique, du fondamental à l’appliqué » (ED 569) from Paris Saclay University. We thank Geneviève Gourdon’s team for the southern blot analyses of the CTG repeats length. We gratefully acknowledge support from the PSMN (Pôle Scientifique de Modélisation Numérique) of the ENS of Lyon for computing resource. We thank Lina El-Kassar and Karine Giraud-Triboult for karyotyping the cell lines.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST

The authors declare no conflicts of interest

AUTHOR CONTRIBUTIONS

A.M., M.C. and C.M.: conception and design. A.M., J.T. and C.M.: investigation and data collection. A.M., J.T., JB.D., H.P., M.J., J.P., C.L. and C.M.: formal analysis and interpretation. D.F. and C.M.: provision of different cellular resources. C.M.: financial support. A.M. and C.M.: manuscript writing.
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**FIGURE LEGENDS**
Figure 1: Generation of MBNL Knockout hiPSCs by CRISPR/Cas9
(A) Schematic representation of the different exons encoding Human MBNL1, MBNL2 and MBNL3. Alternative exons are indicated in blue and the targeted site of each guide-RNA is indicated by an orange arrow. (B) Edited clones were first identified by RFLP (Restriction Fragment Length Polymorphism). The on-target site is underlined and the PAM sequence is in red. The restriction endonuclease recognition site (BsrI in this case) is framed in purple. After locus-specific PCR and restriction digest, cleaved fragments are separated by gel electrophoresis into readily distinguishable profiles, allowing an efficient and rapid screening of single clones to further analyze by sequencing. (C) Examples of sequencing of the targeting site in Human MBNL1, MBNL2 and MBNL3 genes after NHEJ-mediated knockout using the CRISPR/Cas9 system in WT hiPSCs. The MBNL1(-/-) clone harbors a deletion of 14bp; MBNL2(-/-) a deletion of 4bp; MBNL1(-/-); MBNL2(-/-) (DKO) a deletion of 35bp and MBNL1(-/-); MBNL2(-/-); MBNL3(-/-) (TKO) a deletion of 7bp. The DKO clone was generated from MBNL2(-/-) clone and the TKO clone from the DKO clone. STOP codons induced by INDELS are framed in red. (D) Schematic representation of the different gene editing experiments performed to obtain MBNL knockout hiPSCs clones. Created with Biorender.com
Figure 2: Expression profile of the three MBNL paralogs during myogenic differentiation from hiPSCs
(A) Schematic representation of the differentiation protocol used to convert hiPSCs into skeletal muscle cells. Created with Biorender.com (B) Representative examples of western blot analysis for MBNL1 and MBNL2 at different time points of the myogenic differentiation. The detection of the two isoforms were highlighted by an arrowhead (for 40kDa) and an arrow (for 42kDa) as previously described (12). The graph represents the kinetics of expression of the three MBNL paralogs along the process of differentiation of WT hiPSCs. β-Actin was used as housekeeping protein for normalization. Data represent the mean ± SD values from 2 independent experiments. (C-D) Example of a western blot analysis for MBNL1 (C) and MBNL2 (D) proteins after 24 days of differentiation (myotube stage). The graphs on the right panel represent the quantification of MBNL proteins expression in the different conditions compared to WT. β-Actin was used as housekeeping protein for normalization. Data represent means ± SD values from 3 independent experiments (*, P < 0.05; **, P < 0.005; ***, P < 0.0005; ****, P < 0.0001 - Ordinary One-Way ANOVA, Tukey’s multiple comparisons test).
Figure 3: Presence of CUG foci in skeletal muscle cells derived from DM1 hiPSCs
(A) Fibroblasts from two DM1 patients and one healthy donor subject (WT; sibling of one DM1 patient) were used for reprogramming into hiPSCs. Southern blot analysis of the CTG repeats length present in the different fibroblasts and hiPSCs. (B) Characterization of WT and DM1 hiPSCs by flow cytometry for two markers of pluripotency (SSEA3 and TRA1-81). Genomic integrity of these hiPSC lines was validated by karyotyping analysis. All the MBNL KO clones were generated from the WT_1 cell line. (C) Southern blot analysis of the CTG repeat length in hiPSC-derived skeletal muscle cells (24 days after differentiation). (D) Representative images of double immunolabeling for intranuclear CUG repeat foci (red) and DAPI nuclear staining (blue) of WT and DM1 hiPSC-derived skeletal muscle cells (24 days of differentiation). The images below are higher magnifications of the inset to better visualize the presence of CUG RNA foci in DM1 nuclei. Scale bar 50 µM. (E) Quantification of the number of CUG repeat foci per cell as well as the percentage of cells with CUG RNA foci in skeletal muscle cells derived from WT and the two DM1 hiPSC lines (24 days of differentiation). The quantification of CUG repeat foci was performed as previously described (Maury et al., iScience 2019). Data represent the mean ± SD values from at least 3 independent experiments in quadruplicate (*, P < 0.05; **, P < 0.005; ***, P < 0.0005; ****, P < 0.0001 - Ordinary One-Way ANOVA, Tukey’s multiple comparisons test).
Figure 4: No impairment of the early myogenic commitment in the absence of the MBNL proteins
Representative images of immunolabeling for PAX3 (A) and DESMIN (B) on hiPSCs differentiated for 5 and 16 days respectively (PAX3 and DESMIN are in green, DAPI nuclear staining is in blue). Scale bar 10 µM. The graphs on the left represent the percentage of PAX3+ or DESMIN+ cells, after automated quantification. Data represent the mean ± SD values from at least 3 independent experiments in quadruplicate. (C) Alternative splicing analysis for DMD exon 78, INSR exon 11 and MBNL1 exon 5 in hiPSCs-derived myoblasts (16 days of differentiation). Data represent the mean ± SD values from 3 independent experiments in duplicate (*) P < 0.05; (**) P < 0.005; (***) P < 0.0005; (****) P < 0.0001 - Ordinary One-Way ANOVA, Tukey’s multiple comparisons test). Asterisks placed above individual bar correspond to the comparison with WT.
Figure 5: Depletion of MBNL1 and MBNL2 proteins impairs myogenic fusion and reproduces DM1-associated phenotypes
(A) Representative images of immunolabeling for DESMIN (green), Myosin Heavy Chain (MF20) (red) and DAPI nuclear staining (blue) in skeletal muscle cells generated from hiPSCs after 24 days of differentiation. Scale bar 5 µM. The graph on the right represents the number of myotubes containing at least 3 nuclei normalized to the total number of nuclei. A minimum of 11,000 nuclei has been quantified per experiment and per cell line. Data represent mean ± SD values after normalization on the value obtained with WT hiPSC-derived skeletal muscle cells (n = at least 3 independent experiments). Data were analyzed with an ordinary One-Way ANOVA, Tukey’s multiple comparisons test (*, P < 0.05; **, P < 0.005; ***, P < 0.0005; ****, P < 0.0001).

(B) Representative images of immunolabeling for MBNL1 and MBNL2 on hiPSC-derived skeletal muscle cells (24 days of differentiation). Scale bar 10 µM. Higher magnifications of the area in inset are provided to show the punctiform staining of MBNL1 and MBNL2 in DM1 hiPSC-derived skeletal muscle cells in comparison with WT hiPSC-derived skeletal muscle cells; Scale bar 5 µM.

(C) Alternative splicing analysis of DMD exon 78, INSR exon 11 and BIN1 exon 11 were performed by RT-PCR on hiPSC-derived skeletal muscle cells (24 days of differentiation). The mean ± SD values from 3 independent experiments in duplicate are shown (*, P < 0.05; **, P < 0.005; ***, P < 0.0005; ****, P < 0.0001 - Ordinary One-Way ANOVA, Tukey’s multiple comparisons test).
Figure 6: Depletion of MBNL1 and MBNL2 affects the expression of gene sets related to skeletal muscle
(A) PCA (Principal Component Analysis) mapping indicating distinct transcriptomic profiles between WT, MBNL1(-/-) and MBNL1(-/-); MBNL2(-/-) hiPSC-derived skeletal muscle cells (24 days of differentiation). (B) Schematic representation of the number of differentially expressed genes (DEGs) in MBNL1 KO hiPSCs-derived myotubes compared to WT (blue), DKO myotubes compared to WT (green) and DKO compared to MBNL1 KO myotubes (orange). The detailed list DEGs for each condition is provided in Table S3. (C) Venn diagram representation of the number of DEGs commonly deregulated between the three different conditions analyzed. The proportion of each subset of DEGs relative to the total number of DEGs (396 in total) is indicated in brackets. (D) The hierarchical clustering of the 122 DEGs detected in DKO hiPSC-derived skeletal muscle compared to WT hiPSC-derived skeletal muscle cells. This analysis indicates that MBNL1 KO are comparable to WT hiPSC-derived skeletal muscle cells regarding these DEGs. Gene expression is represented by a color code ranging from yellow for over-expressed genes to purple for under-expressed genes. (E) EnrichR based gene set enrichment analysis of the DEGs identified between MBNL1 KO hiPSCs and WT-derived skeletal muscle cells (left), and between DKO hiPSCs and WT-derived skeletal muscle cells (right). The top ten deregulated biological processes are classified according to their adjusted p-value. (F) Gene set enrichment analysis using EnrichR on the 82 down-regulated genes in DKO hiPSC-derived skeletal muscle cells when compared to WT hiPSC-derived skeletal muscle cells. The table on the right provides a list of specific genes down-expressed in DKO hiPSC-derived skeletal muscle cells (EnrichR analysis using GO Biological Process 2018 database).
Figure 7: Loss of MBNL1 and MBNL2 proteins leads to a large number of alternative splicing defects related to the development of skeletal muscle
(A) Number of differentially spliced exons (DSEs) in DM1, MBNL1 (-/-) and in DKO hiPSC-derived skeletal muscle cells compared to WT myotubes. (B) Venn diagram highlighting the percentage of common and different DSEs between the three conditions listed in A. All the DSEs are provided in Table S5. (C) EnrichR based analysis of gene set enrichments using the lists of DSEs observed in DM1, MBNL1(-/-) and DKO hiPSC-derived skeletal muscle cells when compared to WT hiPSC-derived skeletal muscle cells. The top ten most deregulated biological processes are classified according to their adjusted p-value. All the data about the enriched biological processes are detailed in Table S6. (D) List of 30 enriched DSEs covering most of the deregulated biological processes related to skeletal muscle and identified in DKO hiPSC-derived skeletal muscle cells compared to WT. For alternative splicing analysis, exons have been numbered according to FasterDB database. The numbers of the corresponding exons are indicated in brackets (EnrichR analysis using GO Biological Process 2018 database).

**ABBREVIATIONS**

Myotonic Dystrophy type 1 (DM1)

human induced pluripotent stem cells (hiPSC)

untranslated region (UTR)

Dystrophia myotoma protein kinase (DMPK)

Muscleblind-like RNA binding proteins (MBNL)

RNA-binding proteins (RBP)

CUGBP and ETR3-like proteins (CELF)

Knockout (KO)

double knockout (DKO)

Triple knockout (TKO)
Differentially expressed genes (DEGs)

Differentially splice events (DSEs)