Therapeutic Potential of AntagomiR-23b for Treating Myotonic Dystrophy

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Myotonic dystrophy type 1 (DM1) is a chronically debilitating, rare genetic disease that originates from an expansion of a non-coding CTG repeat in the dystrophia myotonica protein kinase (DMPK) gene. The expansion becomes pathogenic when DMPK transcripts contain 50 or more repetitions due to the sequestration of the muscleblind-like (MBNL) family of proteins. Depletion of MBNLs causes alterations in splicing patterns in transcripts that contribute to clinical symptoms such as myotonia and muscle weakness and wasting. We previously found that microRNA (miR)-23b directly regulates MBNL1 in DM1 myoblasts and mice and that antisense technology (“antagomiRs”) blocking this microRNA (miRNA) boosts MBNL1 protein levels. Here, we show the therapeutic effect over time in response to administration of antagomiR-23b as a treatment in human skeletal actin long repeat (HSALR) mice. Subcutaneous administration of antagomiR-23b upregulated the expression of MBNL1 protein and rescued splicing alterations, grip strength, and myotonia in a dose-dependent manner with long-lasting effects. Additionally, the effects of the treatment on grip strength and myotonia were still slightly notable after 45 days. The pharmacokinetic data obtained provide further evidence that miR-23b could be a valid therapeutic target for DM1.

INTRODUCTION
Myotonic dystrophy type 1 (DM1) is a rare genetic disease with no current effective treatment. DM1 is associated with a substantial disease burden resulting in impairment across many different patient systems and tissues. Muscle weakness and fatigue constitute the two most common disease manifestations, reported by 93% and 90% of patients, respectively, followed by muscle locking (73%). Other phenotypes include cardiac dysfunctions, cataracts, insulin resistance, and intellectual disability (OMIM; MIM: 160900). The disease is based on CTG repeat expansions occurring in the dystrophia myotonica protein kinase (DMPK) gene, which are transcribed into pathogenic mRNAs. The CUG repeats bind with high affinity to the muscleblind-like (MBNL) family of proteins, thereby inhibiting their normal function. In skeletal muscle and brain, MBNL1 and MBNL2, respectively, are preferentially expressed, whereas MBNL3 is expressed primarily during embryonic development and adult tissue regeneration. MBNL1 proteins are responsible for the regulation of splicing of several transcripts, specifically by causing a shift from fetal to adult splicing patterns, and act antagonistically to CUGBP Elav-like family member 1 (CELF) proteins in splice regulation, which are found upregulated in DM1. Consequently, transcripts, such as CLCN1, INSR, BIN1, and DMD, inappropriately take on a fetal splicing pattern, which leads to clinical symptoms. Indeed, the correction of erroneous Clcn1 alternative splicing in mouse models of DM eliminates chloride channelopathy and myotonia. MBNL proteins control RNA metabolism in additional ways, including fetal-to-adult polyadenylation patterns, stability, differential localization of mRNAs, and miRNA biogenesis.

The depletion of MBNL protein function has been shown to be a critical factor in the course of the disease. In loss-of-function experiments, Mbnl1 knockout mice and compound loss of Mbnl1 and Mbnl2 recapitulate several clinical symptoms for DM1, including myotonia, mis-splicing, reduced lifespan, and progressive skeletal muscle weakness. Mbnl1 knockout in mice also reflected the various cardiac dysfunctions and embryonic splice isoforms seen in human DM1 patients. Indeed, MBNL1 loss of function accounts for more than 80% of mis-splicing events and nearly 70% of expression defects. In complementary gain-of-function experiments, upregulation of Mbnl1, using a recombinant adeno-associated viral vector in a murine model that expresses 250 CTG repeats in the context of human skeletal actin long repeat (HSA13 mouse), rescued myotonia and mis-splicing of Clcn1, Zasp, Serca1, and Tnnt3. Likewise, overexpression of Muscleblind in the CUG repeats Drosofila model rescued eye phenotypes and heart and muscle histopathology, as well as reduced nuclear foci. Muscleblind overexpression also rescued muscle atrophy and excessive autophagy levels in an inducible fly model, and autophagy was reduced after treatment with Muscleblind-increasing chloroquine treatment in the same model. MBNL1 upregulation in DM1 mice and patient-derived fibroblasts is well tolerated and rescues several symptoms, such as myotonia and mis-splicing events, as well as the reduction of foci

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formation. Finally, genetic variations in the promoter of MBNL1 have been shown to correlate with disease severity, further implicating MBNL1 in DM1. Thus, DM1 is likely treatable because limited function of MBNL1 proteins can be compensated by enhanced expression of their normal endogenous genes.

The sequestration of MBNL1 in the DMPK repeats is not the only factor inhibiting its function. MicroRNAs (miRNAs) are regulators of the expression of mRNA transcripts and can have dozens of targets. By complementary binding to the 3’ UTR of mRNAs, miRNAs guide the RNA-induced silencing complex (RISC) and signal them for translational repression and decay. miRNAs are known to play a critical role in the mechanisms of DM1, especially once it was found that using miRNA sponges in DM1 blind and rescues muscle atrophy, improves lifespan, and reverses splicing events. In human cells, the expression of microRNA-23b-3p (hereafter referred to as miR-23b) is of importance in DM1, because it has been shown to regulate MBNL1 and -2 transcripts directly by luciferase reporter assay, and the silencing of this miRNA by use of antisense oligonucleotides, termed antagoniRs, induced a statistically significant improvement of pathological symptoms in DM1 cells and mice. In this study, we show the therapeutic response to the cholesterol-conjugated antagoniR-23b administration in dosage by subcutaneous and intravenous routes and washout time in an HSA1R mice model. By observing and quantifying the molecular response to drug administration, we conclude that miR-23b suppression by the use of antagoniRs is a viable treatment for DM1 in precise correspondence with miR-23b as a possible therapeutic target.

RESULTS
Therapeutic Effects of AntagomiRs Are Similar Using Subcutaneous or Intravenous Administration Routes
We have recently demonstrated that an antagoniR that reduces the activity of miR-23b in mammalian DM1 models, including patient cells and HSA1R mice, can rescue DM1-like phenotypes. In this previous study, we observed that miR-23b can bind directly to the 3’ UTR of MBNL1. We also observed that a single subcutaneous injection of the antagoniR against miR-23b at 12.5 mg/kg obtained significant rescue of Mbnl1 levels, mis-splicing, histological phenotypes, myotonia, and grip strength in HSA1R mice. In order to investigate whether the use of another route of administration could improve the efficacy of the antagoniR treatment in muscles, we compared the phenotypes of HSA1R mice treated with antagoniR-23b or antagoniR-scramble (SC) as a control for the chemistry of the antisense oligonucleotide at a concentration of 12.5 mg/kg by subcutaneous and tail-vein injection (intravenously [i.v.]). 4 days after injection, the animals were sacrificed, and the quadriceps (QD) and gastrocnemius (GT) muscles of the hind limbs were dissected to study the molecular effects of the treatments (Figure 1A).

We observed that antagoniR-23b, injected using intravenous administration, was only slightly more efficient in reducing miR-23b levels (Figure S1) and produced a significantly higher increase of Mbnl1 transcripts (Figures 1B and 1C), but in correlation with the miRNA levels, there was no significant difference between the Mbnl1 protein levels obtained with subcutaneous and the intravenous routes (Figures 1D–1G). Subsequent to this ELISA test, we verified the existence of both the nuclear and cytoplasmic fractions of Mbnl1 in the samples of both muscles extracted with PBS for the subcutaneous administration test. To do this, we carried out a western blot experiment in which we detected histone H3, a well-known nuclear protein; glyceraldehyde 3-phosphate dehydrogenase (Gapdh) for the cytoplasmic fraction; and Mbnl1, our study protein. As observed in Figure S2, in all of the western blot lanes, the presence of histone H3, Gapdh, and Mbnl1 is observed. This indicates that, despite extracting the protein with PBS, both nuclear and cytoplasmic phases of our protein are present in the extraction. The improvement of force, measured as percentage of normal force (PNF), and myotonia, was also similar using both routes (Figures 1H and 1I). Importantly, blood serum biochemistry, measured in total blood extracted before sacrifice, showed that the intravenous administration produced more alterations in comparison to the subcutaneous treatment and normalized to PBS (Table S1, tab A). Specifically, the amylase, alanine aminotransferase (ALT), and bile acid levels were significantly increased compared to the DM1 mice injected subcutaneously with antagoniR-23b, which could be biomarkers of pancreas and liver-related alterations. The SC also showed alterations in lipase after intravenous injection. However, although increased, the levels achieved were not enough to be considered clinically relevant, according to previous reports. Weight was evaluated before the treatment and again before sacrifice. Unaltered weight and visual necropsy confirmed that there were no relevant toxic effects in the treated mice. Analyzing together the white blood cell differential count of mice administered with antagoniR-23b with both types of administration, significant differences were found (*p = 0.0047), indicating that intravenous administration does produce changes, which supports our decision to administer subcutaneously. In the case of SC, no significant changes are seen. This is observed in the dendrograms of Table S1, tab B, where it is shown how mice treated with intravenously administered antagoniR-23b show a tendency to form a cluster, and those treated subcutaneously form another. This indicates that there is a significant difference between both routes of administration. Since there was not a significant difference between the delivery routes in Mbnl1 protein and functional improvement and also because of the biochemical blood and serum alterations seen in intravenous injection, the subcutaneous injection was chosen for the rest of drug administrations in this study, as it is less invasive and would be preferable in cases of administration to patients as a chronic treatment.

Effects of AntagomiR-23b after Subcutaneous Injection Depend on Dosage
In order to define the dose range in which antagoniR showed therapeutic effects, we studied the effects of subcutaneous injection of antagoniR-23b or -SC as a negative control at a dose approximately 4 times lower (3 mg/kg) and 4 times higher (40 mg/kg) than the previously used dose of 12.5 mg/kg. 4 days after injection, the animals were sacrificed, and the QD and GT muscles of the hindlimbs were dissected for molecular study (Figure 2A). The results showed a
Figure 1. Comparative Study between Subcutaneous and Intravenous Delivery of AntagomiRs in Mice

(A) Administration protocol for subcutaneous and intravenous injections of antagomiR-23b (23b) and antagomiR-SC (SC). All administrations were performed at a concentration of 12.5 mg/kg. (B and C) Mbnl1 relative transcript expression was quantified in gastrocnemius (B) and quadriceps (C) muscles using Gapdh as the endogenous control. (D and E) Mbnl1 relative protein expression was quantified by ELISA in gastrocnemius (D) and quadriceps (E) muscles and was normalized to total protein. (F and G) Western blots from pooled samples were also relatively quantified (RQ) in support of the ELISA results. Mbnl1 relative protein level was measured from gastrocnemius (F) and quadriceps (G) muscles and normalized to Gapdh. (H and I) Before sacrifice, mice were evaluated for grip strength (H), represented as percentage of normal force (PNF), and myotonia grade (I). Statistical comparisons shown were all performed against PBS-treated HSALR mice data (black dashed lines) via Student’s t-test. Additionally, results between mice treated with antagomiR-23b subcutaneous or intravenously were compared in all panels. p values: ns = not significant, *p < 0.05, **p < 0.01, and ***p < 0.001. Error bars = standard error of the mean (SEM).
dose-dependent decrease in miR-23b levels in both muscles (Figure 2B), only when antagomiR-23b and not SC was administered, which was concurrent to an increase of Mbnl1 at mRNA and protein levels (Figures 2C–2E). ELISA results for protein measurements were confirmed with western blot (Figure 2D). AntagomiR-23b at 3 mg/kg produced a slight but significant reduction of miR-23b in QD, which was not able to increase Mbnl1 protein or mRNA levels. Of note, reductions of more than 50% of miR-23b, obtained with the 12.5 mg/kg dose, produced a significant effect on Mbnl1 at both mRNA and protein levels. The 40-mg/kg dose achieved reductions in the miRNA levels of more than 90% and produced an important increase in the levels of Mbnl1 mRNA compared to the treatment with 12.5 mg/kg. This difference was not so prominent at the level of protein. Importantly, the administration of antagomiR-SC had no effect on the miRNAs or the Mbnl1 levels even at the highest concentration. In order to verify the direct effect of the antagomiR-23 on Mbnl1 and to rule out other possible effects of the silencing of the miRNA within the context of the disease, the levels of the HSA transgene that HSALR mice carry were measured by qRT-PCR. The qRT-PCR results revealed that treatment with antagomiRs-SC and -23b compared to PBS did not exert any effect on the HSA transgene in any of the study situations (Figure S3), ruling out a possible effect of these oligonucleotides at this level.

In parallel with the levels of Mbnl1 protein measured, Mbnl1-dependent splicing rescue was also observed in a dose-response manner for Nfx and Clcn1 transcripts, either in GT or QD muscles (Figures 3A–3D). There was a reduction in the inclusion of exon 7 and exon 7a, respectively. Representative electrophoresis gels for Nfx and Clcn1 can be visualized in Figure S4. Recovery of Clcn1 splicing was concomitant to an improvement of myotonia levels after injection with antagomiR-23b (Figure 3E), which is in agreement with previous studies. Grip strength increased in a dose-dependent manner, as well after treatment (Figure 3F). Of note, in both myotonia and grip-strength tests, we observed no difference between the results of mice treated at the concentration of 12.5 mg/kg or 40 mg/kg, which correlated with the levels of Mbnl1 protein detected.
The component’s analysis of serum biochemistry studies revealed that neither antagomiR-SC nor antagomiR-23b treatments caused significant alterations in comparison to PBS treatment, arguing against a specific effect of the miRNA reduction in the immune system activation (Table S1, tab C). Importantly, antagomiR-23b had no other relevant effects in the levels of the different tissue-damage biomarker...
analyzed and did not affect the weight of the animals or cause any other macroscopic alteration that could be detected in the necropsy at all of the dosages tested. Although the administration of the antagoniR at high doses has no effect on blood biochemistry and weight, the statistical analysis with ANOVA and Kruskal-Wallis revealed statistically significant changes between Friend leukemia virus B (FVB) and HSALR mice treated with PBS, SC, and antagoniR-23 for the levels of urea, lipase, and weight. This is novel data and indicates that these are changes that are normally present between healthy mice and mice modeling the disease. As for the white blood cell differential count, there were no significant differences between the groups (p = 0.0665), which means that there are no alterations in blood composition either by treatment or by dose. This can be observed in the dendrogram (Table S1, tab D), since the treated mice are not grouped into clusters. Overall, it seems that there are not relevant changes in blood biochemistry parameters depending on concentration.

One Single Injection of AntagomiR-23b Produces Long-Lasting Phenotypical Alterations

To define the period of effective treatment of DM1 phenotypes, we injected the antagoniR-23b or -SC at 12.5 mg/kg through subcutaneous injection and sacrificed the animals at different times postinjection (4, 15, 30, and 45 days) (Figure 4A). The GT and QD muscles from the hindlimbs were dissected after sacrifice, and total RNA and protein were quantified for levels of target miRNA, Mbnl1.
transcripts, and protein and the effect on Mbnl1-dependent splicing events. With regard to molecular changes, the highest therapeutic effects were observed 4 days after injection (Figure 4). However, there was a clear tendency to maintain reduced levels of miRNA and increased levels of Mbnl1 protein over time. A mild but significant overexpression of Mbnl1 protein levels was detected even 30 days in QD after a single injection and 45 days in GT after a single injection (Figures 4D and 4E). Of note, these levels of protein were enough to exert an effect on Mbnl1-dependent splicing events up to 15 days postinjection (Figure 5). Significant splice recovery was also seen for up to 15 days in transcripts for Clcn1 in GT and QD samples and in Nfix for GT samples. There was a reduction in the inclusion of exon 7 in Nfix and exon 7a in Clcn1, respectively. Representative electrophoresis gels of Nfix and Clcn1 splicing can be visualized in Figure S5.

The sustained effect of treatment with antagomiR-23b was also detected in the functional studies. Before sacrifice, mice were evaluated for grip strength and myotonia (Figures 5E and 5F). Although the greatest effect of rescue was again observed 4 days after injection, a reduction in myotonia and an increase in grip strength continued to be evident even 45 days after the injection. The component’s analysis of blood serum biochemistry studies and white blood cell differential counts showed no significant alterations caused by antagomiR-23b treatment at the different time-point studies in comparison to PBS treatment, except for amylase (Table S1, tab E). Visual necropsy and weight control showed no other significant alterations in the treated animals. There are significant differences in the weight in FVB compared to the PBS. For amylase, there is a significant difference between 23b and FVB in comparison to PBS (p = 0.027) compared to the rest of treatments, which means that we managed to reverse the levels of amylase from mice treated to those of FVB. As for the white blood differential count, there are no significant differences between the groups (p = 0.4856), which means that there are no alterations in blood composition either by treatment or by time. This can be observed in the dendrogram (Table S1, tab F), since the treated mice are not grouped into clusters. These results suggest that although the effects of antagomiRs on Mbnl1 levels and functional improvement are maintained in time, they do not have any delayed effect on blood biochemistry or tissue damage.

Effects of AntagomiR-23b after Subcutaneous Injection in the Mbnl1 Distribution and Foci in Muscle Fibers

From the previous experiments, we have been able to establish the subcutaneous route as a suitable administration mode for the antagomiR-23b. In the dose-response and time-response experiments, we see a plateau effect of Mbnl1 growth and miR-23b decline between the concentrations of 12.5 and 40 mg/kg and that the duration...
of this effect is between 4 and 15 days. Therefore, we selected subcuncaneous injection, a 12.5-mg/kg dose, and a time of 4 days to carry out two important tests to better understand the mechanism of action of the antagomiR-23b. In the first of the tests, we used immunofluorescence (IF) to analyze the location of the Mbnl1 protein in the muscle fibers of HSALR mice treated with PBS and antagomiR-SC and -23b. FVB mice were used as control of a normal Mbnl1 distribution. With support of the previous experiments, the increase in Mbnl1 protein detected in western blotting and ELISA with the antagomiR-23b was also detected by IF (Figure 6D). With the consideration that the Mbnl1 protein is sequestered in ribonuclear foci in HSALR mice, treatment with the antagomiR-23b compared to PBS or SC produced a significant increase in Mbnl1 expression and restored its distribution at the nucleus and cytoplasm level in muscle fibers. This distribution was similar to that observed in healthy FVB mice (Figures 6A–6E). This increase in Mbnl1 that we see in the muscle at the nuclear level is consistent with the splicing rescue shown previously at the 12.5-mg/kg dose. Since there is a complex balance among MBNL proteins, their sequence, and the formation of the characteristic ribonuclear foci of DM1 itself, a possible adverse effect to the increase in Mbnl1 could be a greater formation of foci. To test this hypothesis specifically, we quantified these foci in Q of mice treated with antagomiRs and PBS and discovered that they remained unchanged (Figures 6G–6J). In FVB, the existence of foci was practically nonexistent, as expected (Figure 6F).

DISCUSSION

A little-explored therapeutic strategy for DM1 is in the concept of therapeutic gene modulation, which seeks to increase or decrease the endogenous expression of a gene to alleviate a certain pathological state. Paradigmatic examples are the inhibition of estrogen receptors by antagonists in breast cancer or pharmacologically potentiating the expression of utrophin, a gene that is normally only expressed in the fetus, to compensate for the lack of dystrophin in Duchenne muscular dystrophy. For DM1, the strategy intends to enhance the endogenous expression of MBNLs, for which activity is limited in the disease by sequestration to expansions, which, as described above, cause symptoms such as atrophy, myotonia, and heart disease. In this work, we provide further evidence that miR-23b could be a valid therapeutic target for DM1, which we have previously shown to repress MBNL1 and -2 in natural conditions and in which blockage increases the levels of both proteins in model cells and DM1 mice.

We first show that there is a clear correlation between doses of antagomiR-23b and the degree of repression of miR-23b and an increase in the levels of Mbnl1 transcripts in mice, which does not occur in mice that are administered a control with the same chemistry but SC sequence. This dose-response is no longer observed in levels of Mbnl1 proteins and functional muscle measurements, such as myotonia and muscle strength, where the difference between doses of 12.5 mg/kg and 40 mg/kg is relatively small. This may be due to
a certain plateau effect where more repression of miR-23b results in discrete increases—which have only small, additional improvements at the functional level—at the level of the Mbnl1 protein. We consistently saw an increase in the level of Mbnl1 transcripts. Curiously, this increase was not proportional in Mbnl1 protein levels even at the highest dose. Perhaps low levels of miR-23b have little impact on Mbnl1 transcript stability but still manage to keep transcripts from being translated into protein, which can be expected from miRNAs that typically act within a precise margin. This may also be due to mRNA accumulation as P-bodies or stress granules, resulting in the transcript not being translated into protein.40,41 Another explanation can be due to alternative splicing, which can change the subcellular localization of miRNAs and potentially contribute to the lack of Mbnl1 protein being made. For example, in order for integrin α-3 transcripts to be translated, they must be located in focal adhesions, a process regulated by Mbnl2.22 That being said, the medium dosage of 12.5 mg/kg showed remarkable rescue of Mbnl1 transcripts. However, at the 40-mg/kg dose, the transcribed levels of Mbnl1 in mouse muscle exceeded 2 to 2.5 times the level of endogenous expression in FVB mice, according to the levels observed in other experiments where the level of expression of Mbnl1 mRNA between HSA41R mice treated with PBS and FVB is similar.28 Likewise, the 12.5-mg/kg dosage also shows a healthy rescue of Mbnl1 protein levels and distribution, mis-splicing, myotonia, and muscle weakness. These results emphasize that antagoniR-23b can be fine-tuned to an effective and nontoxic concentration.

The results from the time-response study were particularly promising after observing rescue effects well after 4 days of treatment. The antagoniR showed efficient entry into the mouse muscle cells evidenced by the significant biological and physiological effects (i.e., myotonia and grip-strength tests). Interestingly, these effects, although statistically non-significant, were still seen slightly, even after 45 days of the initial injection. It is true that the effect of the antagoniRs strongly decreases after 4 days, but over time, the decrease is much slower. Indeed, the effect seen at 45 days was similar to the amount of rescue observed at the minimal therapeutic dose of 3 mg/kg. It should be noted as well that the administration of this 12.5-mg/kg dose had no effect on the HSA transgene nor on the percentage of foci, thus ruling out the possible off-target effects that antagoniR-23b could have had at this level.

It is encouraging to see that reducing the activity of a miRNA by only a small amount can have relevant therapeutic effects. Indeed, a reduction of around 50% in miR-23b was sufficient for the increase in Mbnl1 levels, which led to significant splicing and muscle-function improvements downstream. In fact, a 50% reduction is equivalent to what could be described as a heterozygous individual and would rarely generate a dominant phenotype. Likewise, the 12.5-mg/kg dosage facilitated the restoration levels of Mbnl1. This strategy could be a tactic form of combatting DM1 by fine tuning the expression of Mbnl1 without directly disturbing the endogenous expression of DMPK or MBNL1.

Another advantage of the antagoniR-23b was its notable long-term effects and low toxicity. One of the most pressing issues in antisense oligonucleotide therapy, along with off-target effects, is the delivery to the muscle. However, we see a significant improvement in the physical mouse muscle in this experiment. There are precedents in the literature that affirm that anti-miRs with a cholesterol group, known as antagoniRs, are able to reach muscle, among other tissues, very well.42,43 therefore, we could say that we have managed to overcome this problem in mice despite the fact that exogenous administration of a chemically modified oligonucleotide can activate the body’s natural immune response.44 Additionally, the antagoniR has shown stability in vivo by its noted effects long term. Finally, the lack of toxic biochemistry profiles bodes well for the advancement of this particular chemistry.

miR-23b has been involved in several important developmental and cancer-related processes, in which it can either block tumorigenesis or enhance metastatic properties of cells depending on the biological context.45 Indeed, loss of function of miR-23b has shown an increase of metastasis and tumor growth pathways in breast46 and gastrointestinal47 cancers. Conversely, miR-23b has also been shown to promote proliferation in ovarian and prostate cancer through the downregulation of its target phosphatase and tensin homolog (PTEN), which is a known tumor suppressor.48,49 Whereas potential oncogenic effects cannot be assessed from 45 days of treatment, the association of miR-23b with the immune response can be partially addressed in our data. Specifically, neither the dose-response nor the time-response experiments provided any indication of proinflammatory activity, as indicated by healthy visual spleen size and monocyte counts in blood tests. It is true that decreasing miR-23b has been related to a strong increase in immune response50 and that high doses of antisense drugs administered in a short period of time give rise to an acute response that causes the spleen to double in size.51 Therefore, it should be noted that after 4 days with the antagoniR-23b and at the 40-mg/kg dose where miR-23b levels are practically nonexistent, no change in the spleen was visually observed. Taken together, low toxicity, high efficacy, and long-lasting biological effects at the molecular and functional levels highlight antagoniRs against miR-23b as a promising therapeutic strategy.

MATERIALS AND METHODS

Transgenic Mice and AntagoniR Administration

Mouse handling and experimental procedures conformed to the European law regarding laboratory animal care and experimentation (2003/65/CE) and were approved by Conselleria de Agricultura, Generalitat Valenciana (reference numbers A1529567788818 and A1458832800370). Homozygous transgenic HSA41R (line 20 b) mice52 were provided by Prof. C. Thornton (University of Rochester Medical Center, Rochester, NY, USA) and mice with the corresponding genetic background (FVB) were used as controls. AntagoniR against miR-23b and the SC control was purchased from Creative Biogene. The modified sequence and the preparation of the oligo for subcutaneous injections were performed as previously described.36 For intravenous injections, antagoniRs were reconstituted in PBS (KH2PO4 0.144 g/L, NaCl 9 g/L, Na2HPO4 0.795 g/L) at an adequate concentration to allow injection of the required
amount in only 100 μL. There are two PBS controls: one referring to
the control used for 12.5 mg/kg in previous experiments and another PBS for the 3-mg/kg and 40-mg/kg treatment concentrations. New mice injected with PBS have been introduced for the dose-response assay, because the mice used for this new assay are 1 month older than those previously injected at the dose of 12.5 mg/kg. The age of all of the mice used in this article ranges from 4.5 to 5.5 months, and mice were of the male sex.

RNA Extraction, RT-PCR, and qRT-PCR
Total RNA from murine GT and QD muscle was isolated using the miRNeasy Mini Kit (QIAGEN, Valencia, CA, USA), according to the manufacturer’s instructions. 1 μg of RNA was digested with DNase I (Invitrogen) and reverse transcribed with SuperScript II (Invitrogen) using random hexanucleotides. For subsequent PCR reactions, 20 ng of cDNA was used with the GoTaq polymerase (Promega). Specific primers were used to analyze the alternative splicing of Nfix and Clcn1 in mouse samples (QD and GT). Gapdh was used as the endogenous control using 0.2 ng of cDNA. PCR products were separated on a 2% agarose gel and quantified using ImageJ software (NIH). Percentage splice recovery (PSR) index was defined as $value_{%SI} \times %DSI$, divided by $value_{%DSI} \times %HSI$ (where SI is splicing inclusion of each sample, DSI is disease splicing inclusion, and HSI is healthy splicing inclusion). This ratio was calculated for Nfix and Clcn1. The primer sequences and exons analyzed are available in Cerro-Herreros et al. We used 1 ng of mouse tissue cDNA as a template for multiplex qRT-PCR using the Quantifast Probe PCR Kit reagent. Commercial TaqMan probes (QIAGEN) were used for mouse (Mbnl1; 6-carboxyfluorescein [FAM]-labeled probes) and reference (Gapdh; NHS ester [MAX]-labeled probe) genes. HSA transgene expression levels were determined by qRT-PCR, as described previously.53 Mouse results were normalized to Gapdh endogenous gene expression.

miRNA expression in muscle tissues was quantified using specific miRCURY-locked nucleic acid miRNA PCR primers (Exiqon), according to the manufacturer’s instructions. Relative gene expression was normalized to U1 and U6 small nuclear RNAs (snRNAs).

Expression levels were measured using an Applied Biosystems QuantStudio 5 Real-Time PCR System. Expression relative to the endogenous gene and control group was calculated using the 2$^{-\Delta\DeltaCT}$ method. Pairs of samples were compared using two-tailed t tests ($α = 0.05$), applying Welch’s correction when necessary.

ELISA
Mbnl1 protein was quantified by ELISA, according to the manufacturer’s instructions (MyBioSource), in GT and QD muscles from HSA$^{LR}$ mice. Briefly, 20–40 μg of muscle was homogenized in 200 μL of 1× PBS buffer (8 mM Na$_2$HPO$_4$, 150 mM NaCl, 2 mM KH$_2$PO$_4$, 3 mM KCl). All muscles used for the ELISA assay were processed at the same time using the same aliquot of PBS for all samples. Mbnl1 levels were expressed as nanogram of Mbnl1/mg of total protein. Total proteins were quantified with a bicinchoninic acid (BCA) protein assay kit (Pierce) using bovine serum albumin as a standard concentration range. The values were determined using a Tecan Infinite M200 PRO plate reader (Life Sciences).

Western Blotting
As confirmation of the results of ELISA, we performed a western blotting assay with sample pools. Each pool contained the protein extracted from samples of the same treatment. For total protein extraction, mouse muscles (GT and QD) were homogenized in radioimmunoprecipitation assay buffer (RIPA) buffer (150 mM NaCl, 1.0% IGEPAL, 0.5% sodium deoxycholate, 0.1% SDS, 50 mM Tris-HCl, pH 8.0), supplemented with protease and phosphatase inhibitor cocktails (Roche Applied Science). Total proteins were quantified with a BCA protein assay kit (Pierce) using bovine serum albumin as a standard concentration range. For the western blot assay, 20 μg of samples was denatured for 5 min at 100˚C, electrophoresed on 12% SDS-PAGE gels, transferred onto 0.45 μm nitrocellulose membranes (GE Healthcare), and blocked with 5% nonfat dried milk in PBS-Tween 20 (PBS-T; 8 mM Na$_2$HPO$_4$, 150 mM NaCl, 2 mM KH$_2$PO$_4$, 3 mM KCl, 0.05% Tween 20, pH 7.4). Membranes were incubated overnight (O/N) at 4˚C with primary mouse anti-MB1a (1:200, 4A8; Developmental Studies Hybridoma Bank) antibody. The anti-MBNL1 antibody was detected using horseradish peroxidase (HRP)-conjugated anti-mouse-immunoglobulin G (IgG) secondary antibody (1 h, 1:3,500; Sigma-Aldrich).

We determined the existence of the nuclear protein fraction in the muscle samples extracted with PBS. To do this, we loaded 20 μg of the protein onto polyacrylamide gels for SDS-PAGE. We transferred proteins to nitrocellulose membranes and carried out immunoblotting using anti-histone H3 (O/N; Millipore; 05-928, 1:1,000), followed by incubation with the appropriate secondary HRP-conjugated anti-rabbit-IgG secondary antibody (1 h, 1:3,500; Sigma-Aldrich). We diluted both antibodies in PBS containing 3% bovine serum albumin and 0.1% Tween 20.

Anti-GAPDH antibody (1 h, 1:3,500, clone G-9; Santa Cruz) was used as a loading control for mouse samples, followed by HRP-conjugated mouse-immunoglobulin G (IgG) secondary antibody (1 h, 1:5,000; Sigma-Aldrich). Immunoreactive bands were detected using an enhanced chemiluminescence (ECL) Western Blotting Substrate (Pierce), and images were acquired with an ImageQuant LAS 4000 (GE Healthcare). Quantification was performed using ImageJ software (NIH).

Fluorescent Methods
In situ detection and Mbnl1 IF were performed as previously described. Briefly, for foci detection, 10 mm frozen sections of the QD muscles were fixed in 3% paraformaldehyde (PFA)/PBS for 15 min at room temperature, washed with 1× PBS, and permeabilized with 0.5% Triton X-100/PBS for 5 min at room temperature. Fixed sections were incubated in the prehybridization buffer ($2 \times$ saline sodium citrate [SSC], 30% deionized formamide) for 10 min at room
temperature and hybridized with a Cy3-(CAG)7-Cy3-labeled probe, diluted 1:200 in hybridization buffer, 30% formamide, for 2 h at 37°C in the dark. After hybridization, we washed the muscle sections with a prehybridization buffer for 30 min at 42°C, washed twice with 1 × SSC for 15 min at room temperature, washed with 1 × PBS, incubated with fluorescein isothiocyanate (FITC)-labeled wheat germ agglutinin diluted 1:600 in PBS for 45 min at room temperature to stain cell membranes, washed with PBS, and mounted with 4',6-diamidino-2-phenylindole (DAPI) mounting media (Vector). A total of eight images per mice were taken using the LSM800 confocal microscope (Zeiss, Jena, Germany) at 400× magnification. To find the percentage of RNA nuclear foci, each image was quantified using the following formula:

\[
\text{% nuclear Foci} = \frac{\text{Nuclei with foci}}{\text{Total nuclei}} \times 100
\]

Localization of Mbnl1 by IF was carried out with 10 mm frozen sections of QD muscles, fixed in 3% PFA/PBS for 15 min, washed with PBS, permeabilized with 0.5% Triton X-100/PBS for 5 min, and blocked in 5% normal goat serum in PBS for 30 min, all at room temperature. Sections were incubated O/N at 4°C with primary mouse anti-MBNL1 (1:200 clone MB1a; The Wolfson Centre for Inherited Neuromuscular Disease; in blocking buffer), washed with PBS, incubated with the secondary antibody (goat anti-mouse-FITC labeled, 1:200 in blocking buffer) in the dark for 1 h at room temperature, washed with PBS, and mounted with DAPI mounting media (Vector). A total of eight images for mice were taken using LSM800 confocal microscope (Zeiss, Jena, Germany) at 400× magnification. The Mbnl1 signal was quantified, dividing green channel intensity by the muscle area. ImageJ software measures intensity and pixel-size area from the confocal images.

Electromyography Studies
Electromyography was performed before the treatment, at the halfway point, and at the time of sacrifice under general anesthesia, as previously described. Briefly, five needle insertions were performed in each QD muscle of both hind limbs, and myotonic discharges were graded on a scale from 1 to 3, where 1 = no myotonia; 2 = occasional myotonic discharge in <50% of the needle insertions; 3 = myotonic discharge in >50% of the insertions; and 4 = myotonic discharge in >50% of the insertions. The experiment was performed blindly to eliminate bias.

Forelimb Grip-Strength Test
The forelimb grip strength was measured with a Grip Strength Meter (BIO-GS3; Bioseb, USA). The peak pull force (measured in grams) was recorded on a digital force transducer when the mouse grasped the bar. The gauge of the force transducer was reset to 0 g after each measurement. The tension was recorded by the gauge at the time the mouse released its forepaws from the bar. We performed three consecutive measurements at 30-s intervals. The bodyweight measurement was performed in parallel. The experiment was performed blindly to eliminate bias.

The values of force are represented as the PNF, and it measures how close the weight-normalized strength values of treated HSA1LR mice are compared to force measured in FVB controls. This PNF index is obtained by normalizing the weight-relative force (WRF; =force/weight) of each mouse after treatment with its WRF before initiating the treatment and dividing this value for the force effect (FE), resulting in dividing the mean WRF of FVB mice between the mean WRF of HSA1LR treated with PBS at the same time point (PNF = WRF/FE).

Blood Assays
4, 15, 30, and 45 days following treatment, animals were sacrificed, and blood was collected by cardiac puncture exsanguination with K3-EDTA (SARSTEDT). The samples were analyzed by Laboratorios Montoro Botella (Valencia, Spain). White blood cell differential count (monocytes, stab cells, segmented cells, basophils, eosinophils, and lymphocytes) was measured with the Hematology Cell Counter ADVIA 120 (Siemens). The serum biochemistry profile (creatinine, urea, amylase, alkaline phosphatase, ALT, bilirubin, lipase, and bile acids) was analyzed with the cobas 600 CCE modular analyzer (Roche).

Statistical Analyses
In the molecular and functional studies, for comparison on mean data, we assumed that all parameters follow a normal distribution, and the samples were compared using two-tailed t-tests (\(p = 0.05\)), applying Welch’s correction when necessary. The statistical differences were estimated by the Student’s t-tests (\(p < 0.05\)) on normalized data. Sample size (\(n\)) can be seen in each figure. The graphs were generated using GraphPad Prism 6 software. The individual data for each mouse for the qPCR, ELISA, myotonia, force, splicing, Mbnl1, and foci IF can be found in Table S2.

Average values of blood parameters were compared by means of multivariate analysis of variance of compositional data using PBS-treated HSA1LR mice as a reference. This is done by transforming the white blood cell differential count parameters as a set of elements from a simplex under Aitchison geometry, applying ILR (isometric log ratio) transformation to the compositional data, and conducting a multivariate analysis of variance (MANOVA). Statistical analysis of the serum biochemistry profile was performed with a one-way ANOVA test or Kruskal-Wallis when application requirements did not hold for general linear modeling. In the cases in which the Kruskal-Wallis test gave a value lower than 0.05, a pairwise Wilcoxon rank sum test was performed with a p value correction for false discovery rate in order to see which of the treatments had different values from the rest. In all cases, all experiments (route, doses, or time) were analyzed together since it was not possible to analyze each group separately because there was no PBS control for each dose. Here, treatment type was also considered as the only independent factor using PBS-treated HSA1LR mice as a reference.

The cutoff for statistical significance was set as \(p = 0.05\). Compositional data were analyzed with R package “compositions,” whereas the serum biochemistry profile was analyzed with “R base.”
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.omtn.2020.07.021.

AUTHOR CONTRIBUTIONS

E.C.-H., M.P.-A., B.L., and R.A. designed the experiments. E.C.-H., I.G.-M., and N.M.-C. conducted all experiments. S.O. wrote the paper.

CONFLICTS OF INTEREST

B.L. is CEO and cofounder of Arthex Biotech. R.A. is advisor and cofounder of Arthex Biotech. The remaining authors declare no competing interests.

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Supplemental Information

Therapeutic Potential of AntagomiR-23b for Treating Myotonic Dystrophy

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Supplemental Material

Figure S1. Comparative study between subcutaneous and intravenous delivery of antagomiRs in mice. miR-23b relative expression was quantified from (A) gastrocnemius and (B) quadriceps muscles relative to U1 and U6 snRNA endogenous controls after subcutaneous or intravenous injection of PBS, antagomir-23b (23b), and antagomir-SC (SC). Statistical comparisons without brackets were performed against the data obtained in PBS-treated HSA LR mice (black dashed line) via Student’s t-test. p-value: ns=not significant, * p<0.05, ** p<0.01, *** p<0.001.

Figure S2. Both fractions cytoplasmic and nuclear of Mbnl1 protein are present in the samples extracted for ELISA with PBS. Western blot to detect Mbnl1, Histone H3 and Gapdh proteins in gastrocnemius (A) and quadriceps (B) treated with PBS, scrambled antagomir (SC) and antagomiR-23b (23b) for subcutaneous delivery of antagomiRs in mice. Histone H3 and Gapdh were used as the loading controls for the nuclear and cytoplasmic fractions, respectively.
Figure S3. Transgene quantification in dose-response study after treatment with antagomiRs. HSA relative expression was quantified from gastrocnemius (left) and quadriceps (right) muscles relative to Gapdh endogenous controls after subcutaneous injection of PBS, antagomir-23b (23b), and antagomir-SC (SC). The study was performed at 3 different concentrations: 3 mg/kg, 12.5 mg/kg, and 40 mg/kg by subcutaneous injection. Statistical comparisons without brackets were performed against the data obtained in PBS-treated HSA LR mice (black dashed line) via Student’s t-test.
Figure S4. Dose response study of Mbnl1-dependent splicings after treatment with antagomiRs. (A-B) Nfix and (C-D) Clcn1 splicings were quantified in (A, C) gastrocnemius and (B, D) quadriceps after treatment with PBS, antagomir-23b (23b), and antagomir-SC (SC). Control treatment PBS1 corresponds to 12.5 mg/kg treatment while PBS2 corresponds to 3 mg/kg and 40 mg/kg treatments. Levels of healthy control mice can be seen in FVB. Each treatment was statistically compared with its respective PBS control via Student's t-test. p-value: * p<0.05, ** p<0.01, *** p<0.001.
Figure S5. Time response study of Mbnl-dependent splicings after treatment with antagomiRs. (A-B) *Nfix* and (C-D) *Clcn1* splicings were quantified in (A, C) gastrocnemius and (B, D) quadriceps after treatment with PBS, antagomir-23b (23b), and antagomir-SC (SC) over 4, 15, 30, and 45 days. Levels of healthy control mice can be seen in FVB. Time interval was normalized to its respective PBS control via Student’s t-test. p-value: * p<0.05, ** p<0.01, *** p<0.001.
Table S1. Statistical analysis of blood parameters in the each experiment. (A. Biochemistry IV vs SUB) Here the biochemical profile for the intravenous and subcutaneous experiments is shown with the means and standard error of the mean (SEM) of each of the parameters in different groups so that the statistical test compares the two administrative routes, intravenous (IV) and subcutaneous (Sub.), between the scramble and antagomiR-23b. (B. WBCs IV vs SUB) Here the white blood cells (WBC) formula is shown for the intravenous vs subcutaneous experiment including the mean and the SEM of each of the parameters in different groups so that the statistical test compares the two administrative routes between the scramble and antagomiR-23b. It also includes dendograms for both IV and Sub. treatments. (C. Biochemistry Dose Response) Here the biochemical profile for the dose response experiments is shown with the means and standard error of the mean (SEM) of each of the parameters in different groups so that the statistical test compares the different dosages between the scramble and antagomiR-23b, along with FVB values. (D. WBCs Dose Response) Here the WBC formula is shown for the dosage experiment including the mean and the SEM of each of the parameters in different groups so that the statistical test compares the different dosages between the scramble and antagomiR-23b, along with FVB values. It also includes dendograms for each dosage. (1: 3 mg/kg, 2: 12.5 mg/kg, 3: 40 mg/kg.) (E. Biochemistry Time Response) Here the biochemical profile for the time response experiment is shown with the means and standard error of the mean (SEM) of each of the parameters in different groups so that the statistical test compares the different time treatments between the scramble and antagomiR-23b, along with FVB values. (F. WBCs Time Response) Here the WBC formula is shown for the time response experiment including the mean and the SEM of each of the parameters in different groups so that the statistical test compares the different dosages between the scramble and antagomiR-23b, along with FVB values. It also includes dendograms for all time points (1: 4 days, 2: 15 days, 3: 30 days, 4: 45 days). P-value: *p<0.05, **p<0.01, ***p<0.001.

Table S2. Individual data for mice in each experiment. (A. IV vs SUB) Individual mouse data is shown for the qPCR, ELISA, myotonia grade, and PNF experiments for the administration type assay. (B. Dose Response) Individual mouse data are shown for the qPCR, ELISA, myotonia grade, PNF, and Nfix and Clcn1 splicing experiments for the administered dose assay. (C. Time Response) Individual data per mouse is shown for the qPCR, ELISA, myotonia grade, PNF, and Nfix and Clcn1 splicing experiments for the administration time assay. (D. Mbnl1 distribution and Foci) The individual data per mouse for the immunofluorescence experiments of Mbnl1 and foci in the case of the optimal dose are shown.