Reversing established muscle atrophy following mechanical unloading is of great clinical challenge. Long noncoding RNAs (lncRNAs) have been demonstrated to play important roles in myogenesis. Here we identified a lncRNA (mechanical unloading-induced muscle atrophy-related lncRNA [IncMUMA]) enriched in muscle, which was the most downregulated lncRNA during muscle atrophy development in hindlimb suspension (HLS) mice. The in vitro and in vivo data demonstrated that the decreased expression levels of IncMUMA closely associated with a reduction of myogenesis during mechanical unloading. Mechanistically, IncMUMA promoted myogenic differentiation by functioning as a miR-762 sponge to regulate the core myogenic regulator MyoD in vitro. The enforced expression of IncMUMA relieved the decreases in MyoD protein and muscle mass in miR-762 knockin mice. Therapeutically, the enforced expression of IncMUMA improved the in vitro myogenic differentiation of myoblasts under microgravity simulation, prevented the muscle atrophy development, and reversed the established muscle atrophy in HLS mice. These findings identify IncMUMA as an anabolic regulator to reverse established muscle atrophy following mechanical unloading.

INTRODUCTION

Mechanical unloading induces obvious skeletal muscle atrophy and functional deficit, especially under microgravity environment during spaceflight. Exposure to microgravity for 3–6 months induces skeletal muscle atrophy with more than 40% impairment of functional properties in astronauts during spaceflight. Decreased muscle differentiation and protein synthesis are largely responsible for the mechanical unloading-induced skeletal muscle atrophy. However, to date there are no effective countermeasures to combat the mechanical unloading-induced skeletal muscle atrophy. Exercises are used to replace mechanical loads during spaceflight. Astronauts on the international space station spend 5 hr/week for aerobic exercise and 3–6 days/week for resistance exercise; the exercises are insufficient to prevent loss of muscle strength and endurance. For the pharmaceutical approaches, intake of essential amino acids has been proven to increase muscle protein synthesis in astronauts; but, there is concern that increasing amino acid intake during spaceflight may increase bone resorption. Other anabolic agents, such as insulin-like growth factor-I (IGF-I) treatment and anabolic androgenic steroids, have some concerns on increased risks (drug resistance and cardiac events), limiting their therapeutic use for muscle atrophy. Thus, it is desirable to investigate the molecular mechanisms of the compromised muscle anabolism during unloading-induced skeletal muscle atrophy to develop a promising muscle anabolic strategy.

Long noncoding RNAs (lncRNAs) (>200 nt) are implicated as important regulators in numerous physiological and pathological processes. lncRNAs also have been demonstrated to play important roles in myogenesis. The lncRNA H19 can positively regulate the progression of myogenic differentiation by repressing the expression of insulin–like growth factor 2 (IGF2) that negatively regulates the key myogenic regulator factor MyoD expression. Moreover, H19 also antagonizes microRNA let-7 that impairs insulin signaling and decreases glucose uptake in skeletal muscle, where H19 depletion results in premature muscle differentiation. The competing mode for microRNAs is also found in another lncRNA linc-MD1, which positively correlates with the muscle differentiation. The lncRNA Inc-mg promotes myogenesis by functioning as a competing endogenous RNA (ceRNA) that binds to microRNA-125b to elevate protein

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abundance of IGF2.\textsuperscript{15} The inhibition of lncRNA Malat1 by myostatin is associated with decreased myogenesis.\textsuperscript{16} The lncRNA Yam-1 is downregulated upon differentiation and acts as an inhibitor of myogenesis through in cis regulation of miR-715, which in turn targets Wnt7b to repress myogenesis.\textsuperscript{17} However, the specific roles of lncRNAs in muscle atrophy following mechanical unloading are still largely unknown. Thus, the aim of our study was to explore the role of lncRNAs in regulating muscle differentiation during the skeletal muscle atrophy following mechanical unloading.

In this study, we identified a lncRNA (mechanical unloading-induced muscle atrophy-related lncRNA [lncMUMA]) in mice skeletal muscle. The decreased expression level of lncMUMA was closely associated with a reduction of myogenesis during mechanical unloading in vitro and in vivo. Mechanistically, we demonstrate that lncMUMA functioned as a miR-762 sponge to regulate the core myogenic regulator MyoD in vitro and in vivo. Furthermore, we show that the enforced expression of lncMUMA improved the in vitro myogenic differentiation of myoblasts under microgravity simulation, prevented the muscle atrophy development, and reversed the established muscle atrophy in hindlimb suspension (HLS) mice. These findings could provide a novel strategy for the treatment of muscle atrophy following mechanical unloading.

RESULTS

lncMUMA Was the Most Downregulated lncRNA in Gastrocnemius Muscle from HLS Mice

We utilized lncRNA microarray to compare the differentially expressed lncRNAs in the gastrocnemius muscle between control and HLS mice (n = 3). The muscle mass was significantly decreased in HLS mice (Figure 1A). A 959-nt-long lncRNA (AK014246) was identified as the most downregulated lncRNA in the gastrocnemius muscle from HLS mice (Figure 1B; Figure S1; Table S1). Further real-time
PCR showed that the expression level of the lncRNA was much higher in the skeletal muscle than in other tissues and organs in mice (n = 10) (Figure 1C). Among different skeletal muscles, this lncRNA was the most abundant in gastrocnemius muscle (Figure S2A). Except in the skeletal muscle in hindlimb, the expression levels of the lncRNA in other tissues and organs or skeletal muscles in the forelimb and trunk were not significantly altered during HLS (Figure S2B). The lncRNA was named lncMUMA.

lncMUMA Expression Was Associated with Muscle Differentiation in Microgravity-Simulated C2C12 Myoblasts In Vitro and Muscle Mass in HLS Mice

We cultured C2C12 cells in a microgravity-simulated (MGS) environment. Interestingly, the IncMUMA expression was found dramatically decreased and reached a stable low level in the MGS group on day 5 of differentiation. The similar trend was also observed in the expression level of the myogenic marker MyoD protein in the MGS group. Consistently, simulated microgravity also suppressed myotube formation in C2C12 cells, evidenced by a much lower fusion index in the MGS group compared to the control group on day 7 of the differentiation (Figures 2A–2D). Consistent with the in vitro findings, we also found that the IncMUMA expression rapidly decreased in the gastrocnemius muscle of mice that underwent HLS, and it reached a stable low level on day 14 of HLS (n = 10) (Figure 2E). A similar trend was also observed in the change of muscle mass in HLS mice (Figures 2F and 2G).

IncMUMA Silencing Suppressed Muscle Differentiation in C2C12 Myoblasts and Decreased Muscle Mass and Strength in Adult Mice

To evaluate the biology function of IncMUMA in vitro and in vivo, a skeletal muscle-specific lentiviral vector system was constructed with either IncMUMA or IncMUMA short hairpin RNA (shRNA), and the lentivirus for cell and tissue transduction was subsequently prepared (Figure S3A). We transduced C2C12 cells with IncMUMA shRNA lentivirus under normal gravity. The MyoD protein level dramatically decreased in the IncMUMA shRNA group compared to that in the scrambled shRNA group. Consistently, skeletal muscle-specific IncMUMA silencing suppressed myotube formation in C2C12 cells, evidenced by a lower fusion index in the IncMUMA shRNA group compared to the scrambled shRNA group on day 7 of the differentiation (Figures S3B–S3D). The IncMUMA knockdown (KD) lentivirus was injected into the gastrocnemius muscle of adult mice (n = 10). The muscle mass and muscle fiber cross-sectional area (CSA) significantly decreased in the IncMUMA KD group compared to those in the scrambled shRNA group. Consistently, the muscle-specific force and the MyoD protein level were also reduced in the IncMUMA KD group compared to those in the scrambled shRNA group (Figures S3E–S3J).
lncMUMA Interacted with miR-762- and miR-762-Targeted MyoD in C2C12 Cells

To gain further insight into the mechanism by which lncMUMA regulates muscle differentiation, we predicted microRNA-762 (miR-762) was one of the target microRNAs (miRNAs) of lncMUMA by using RNAhybrid 2.12 (https://bibiserv.cbi. uni-bielefeld.de/rnahybrid/). The binding site of lncMUMA to miR-762 is located in 767–789 nt from the 5' end of lncMUMA (Figure 3A). We found that the expression level of miR-762 was unchanged, while MyoD protein expression significantly decreased in MGS C2C12 cells, which was closely associated with the decreased lncMUMA expression (Figures S4A–S4C). Moreover, consistent data were also found in the gastrocnemius muscle of HLS mice (n = 10) (Figures S4D–S4F).

To further validate the interaction between lncMUMA and miR-762, pull-down assay and luciferase reporter assay were performed. The biotin-labeled lncMUMA specifically pulled down miR-762 in a dose-dependent manner in miR-762-transfected C2C12 cells (Figure 3B). The miR-762 containing either wild-type binding site (miR-762-WT) or mutated binding site (miR-762-Mut) was cloned into the downstream of luciferase reporter gene (Figure 3C). It showed that lncMUMA transduction could reduce the luciferase activity of miR-762-WT, but not affect that of miR-762-Mut. Meanwhile, transduction of lncMUMA containing a mutated binding site could not reduce the luciferase activity of either miR-762-WT or miR-762-Mut (Figure 3D).

Moreover, MyoD was downregulated at the protein level by agomiR-762 (a miR-762 agonist) in C2C12 cells, whereas...
antagomiR-762 (a miR-762 inhibitor) elevated the MyoD protein expression (Figure 3E; Figure S5).

To validate the interaction between miR-762 and MyoD, the luciferase reporter gene containing either wild-type MyoD 3’ UTR-binding site (MyoD 3’ UTR) or mutated MyoD 3’ UTR-binding site for miR-762 (MyoD 3’ UTR-Mut) was constructed (Figure 3F). Luciferase reporter assay revealed that agomiR-762 (agomiR-762 containing wild-type-binding site for MyoD), but not agomiR-762-Mut (agomiR-762 containing mutated binding site for MyoD), could reduce the luciferase activity of MyoD 3’ UTR. Meanwhile, the luciferase activity of MyoD 3’ UTR-Mut was not repressed by agomiR-762 (Figure 3G). Furthermore, the luciferase activity of MyoD 3’ UTR was significantly increased after reducing the endogenous levels of miR-762 by treating C2C12 cells with antagomiR-762 (Figure 3H).

lncMUMA Vector, but Not Mutated lncMUMA Vector, Counteracted the Decreases in Expression of MyoD and Muscle Differentiation in miR-762-Overexpressing C2C12 Cells

To evaluate the interaction between lncMUMA and miR-762 on MyoD protein level and muscle differentiation, lncMUMA plasmid containing either wild-type or mutated binding site of lncMUMA with miR-762 was transfected into the C2C12 cells stably overexpressing miR-762. The expression of MyoD protein and myotube formation stayed at low levels in miR-762-overexpressing C2C12 cells (Figure 4A). Transduction of lncMUMA vector containing wild-type binding site, but not the mutated lncMUMA vector, elevated the MyoD protein level (Figure 4A). Moreover, the abundance of multinuclear myotubes was increased in the wild-type lncMUMA vector group, while the same phenomenon was not observed after mutated lncMUMA vector transduction (Figures 4B and 4C).

Skeletal Muscle-Specific Overexpression of lncMUMA Promoted MyoD, Muscle Mass, Structure, and Function in Muscle-Specific miR-762 Knockin Mice

To facilitate the mechanism study of the anabolic role of lncMUMA in vivo, muscle-specific miR-762 knockin mice were generated (Figure 5A). Compared to other tissues and organs, in the skeletal muscles of muscle-specific miR-762 knockin mice miR-762 was highly expressed (n = 10) (Figure S6A). The mice also showed lower muscle mass, lower muscle fiber CSA, and lower MyoD protein level in gastrocnemius muscle compared to the control mice (Figures S6B–S6D). lncMUMA overexpression increased the muscle mass, muscle fiber CSA, and muscle strength in muscle-specific miR-762 mice (n = 10) (Figures 5B–5E). In addition, lncMUMA overexpression elevated the expression level of MyoD protein in gastrocnemius muscle of muscle-specific miR-762 knockin mice (Figure 5F).
Enforced IncMUMA Expression Counteracted the Decreases in Muscle Differentiation in MGS C2C12 Myoblasts

To further explore the effect of IncMUMA in counteracting the suppressed muscle differentiation induced by MGS in vitro, we infected the C2C12 cells with IncMUMA lentivirus to overexpress the IncMUMA level and then cultured them in an MGS environment. The IncMUMA was stably overexpressed in the IncMUMA group, while IncMUMA expression was found dramatically decreased in the empty vector (EV) group (Figure 6A). There was no significant change in the MyoD protein level in the IncMUMA vector group, while it was significantly decreased in the EV group (Figure 6B). Moreover, much less myotube formed in the EV group compared to that in the IncMUMA group on day 7 of the differentiation in C2C12 cells (Figures 6C and 6D).

Skeletal Muscle-Specific Overexpression of IncMUMA Could Attenuate and Reverse the Decreases of MyoD, Muscle Mass, Structure, and Function following Mechanical Unloading

To evaluate the preventive effect of IncMUMA overexpression on muscle atrophy following mechanical unloading in vivo, the adult mice were locally injected with IncMUMA vector in gastrocnemius muscle and then underwent 42-day HLS (n = 10). IncMUMA overexpression attenuated the decreases in muscle histomorphometric (i.e., muscle mass and muscle fiber CSA) and functional (i.e., muscle strength) parameters in HLS mice (Figures S7A–S7D). In addition, the decreased MyoD protein in the gastrocnemius muscle of HLS mice was also successfully attenuated by IncMUMA overexpression (Figure S7E).

To further evaluate the effect of IncMUMA overexpression on reversing established muscle atrophy following mechanical unloading in vivo, the adult mice were locally injected with IncMUMA vector in gastrocnemius muscle and then underwent 42-day HLS. IncMUMA overexpression significantly attenuated the decreases in muscle histomorphometric (i.e., muscle mass and muscle fiber CSA) and functional (i.e., muscle strength) parameters in HLS mice (Figures S7A–S7D). In addition, the decreased MyoD protein in the gastrocnemius muscle of HLS mice was also successfully attenuated by IncMUMA overexpression (Figure S7E).
unloading, the adult mice with established muscle atrophy (on day 14 of HLS) were locally injected with either EV or lncMUMA lentivirus in the gastrocnemius muscle, and they were kept hindlimb suspended until day 42 (n = 10). The decreased muscle mass and muscle fiber CSA started to elevate after lncMUMA overexpression on day 14 of HLS (Figures 7A–7D). Moreover, MyoD protein level and muscle-specific force in the lncMUMA vector group were higher than those in the EV group on day 42 of HLS (Figure 7E).

**DISCUSSION**

In this study, the lncRNA lncMUMA in mice skeletal muscle that functions as a miR-762 sponge to promote skeletal muscle differentiation has been identified. Enforced expression of lncMUMA prevented the muscle atrophy development and reversed the established muscle atrophy following mechanical unloading in mice. lncMUMA could be a novel therapeutic target for treating muscle atrophy following mechanical unloading.

lncRNAs, including H19,12,13 linc-MD1,14 Yam-1,17 Malat1,16 and lnc-mg,15 have been demonstrated to regulate muscle differentiation, while little is known about their function during myogenesis in vivo. In this study, the newly identified lncRNA lncMUMA is highly expressed in skeletal muscle and associated with muscle atrophy following mechanical unloading in mice, implying its potential function in regulating muscle differentiation. It is further confirmed that...
lncMUMA promoted myogenesis in vitro and in vivo, evidenced by the reduced muscle mass and strength when silencing lncMUMA, which is consistent with the in vitro data. Furthermore, enforced lncMUMA expression in skeletal muscle prevents muscle atrophy and even reverses established muscle atrophy following mechanical unloading.

Recently, several studies have reported a regulatory mechanism of lncRNA, that lncRNA may function as ceRNA to sponge miRNA, thereby modulating the derepression of miRNA targets and imposing an additional level of post-transcriptional regulation. In our study, miR-762 is predicted to contain a binding site for lncMUMA by bioinformatics analysis. Pull-down assay and luciferase assay validate the direct binding of the miR-762 response elements on the lncMUMA transcript. Furthermore, the enhanced muscle differentiation in lncMUMA-infected C2C12 cells is attenuated by the wild-type miR-762, but not the mutated miR-762. It has been reported that miR-762 could promote the development of ovarian cancer by suppressing Menin. miR-762 has also been reported to regulate Interferon-induced transmembrane protein 5 (IFITM5), an osteoblast-specific membrane protein. This is the first study to reveal miR-762’s role in myogenesis.

MyoD is a key regulator during muscle differentiation. In our study, decreased MyoD protein expression is closely associated with the decreased lncMUMA expression in MGS C2C12 cells and HLS mouse muscle, while the expression level of miR-762 is unchanged. lncMUMA, but not mutated lncMUMA, counteracts the decreases in expression of MyoD and muscle differentiation in miR-762-overexpressed C2C12 cells. Furthermore, lncMUMA overexpression promotes MyoD, muscle mass, structure, and function in muscle-specific miR-762 knockin mice. These data indicate that lncMUMA interacts with miR-762 to post-transcriptionally regulate the MyoD protein.

To date, most studies of lncRNAs in skeletal muscle are investigated in the C2C12 cell line or animals under physiological condition; there’s no report to clarify the role of lncRNA in a mechanical unloading-induced muscle atrophy animal model. In this study, we overexpressed lncMUMA level in the skeletal muscle of HLS mice, which is an established animal model of muscle atrophy following mechanical unloading. Skeletal muscle-specific overexpression of lncMUMA before HLS prevents muscle atrophy. Furthermore, it could also reverse established muscle atrophy induced by unloading. The data show the therapeutic potential of lncMUMA in muscle atrophy.

We used mouse α-skeletal actin promoter to construct the lentiviral vectors to achieve skeletal muscle-specific gene expression with high efficiency in our study. The α-skeletal actin plays important roles in skeletal muscle contraction. Its promoter has been used for
skeletal muscle-specific gene delivery due to its high tissue specificity and expression efficiency.24,25 The lentivirus has been widely used in gene therapy research.26 It has also been used in the transduction for C2C12 myoblasts27–30 and skeletal muscle in mice.31,32 The intramuscular administration of lentiviral particles with z-skeletal actin promoter could generate high-level skeletal muscle-specific gene expression for 2 years.33 In our study, direct injection of lentiviral particles constructed with mouse z-skeletal actin promoter into the gastrocnemius muscle of mice resulted in the significantly higher expression level of target gene lncMUMA in gastrocnemius muscle from lncRNA lentivirus-treated mice than that in vector control mice.

In conclusion, the newly identified lncRNA lncMUMA acts as a miR-762 sponge to regulate MyoD protein, resulting in promoting muscle differentiation. lncMUMA could be a novel anabolic therapeutic agent to reverse established skeletal muscle atrophy following mechanical unloading.

MATERIALS AND METHODS

Animals

6-month-old C57BL/6J mice were used in the study. All the animals were maintained under standard animal housing conditions (12-hr-light and 12-hr-dark cycles and free access to food and water). All the experimental procedures were approved by the Committees of Animal Ethics and Experimental Safety of the Chinese University of Hong Kong.

HLS Procedure

The animals were subjected to HLS for 42 days following established procedure.40 Briefly, a strip of adhesive tape was applied to the animal’s tail, which was suspended by passing the tape through a fishline swivel that was attached to a metal bar on the top of the cage. The forelimbs were allowed to touch the grid floor and the animals could move around the cage for free access to food and water. The suspension height was adjusted to prevent the hindlimbs from touching any supporting surface while maintaining a suspension angle of approximately 30°. The distal tip of the tail was examined to ensure that the procedure did not occlude blood flow to the tail.

Cell Culture

The C2C12 mouse myoblast cell line was supplied by American Type Culture Collection (ATCC). Cells were cultured at subconfluent densities in growth medium made up of DMEM supplemented with 10% heat-inactivated fetal calf serum (FCS) and 1% penicillin-streptomycin. C2C12 myoblast cells were differentiated into myocytes or myotubes in differentiation medium, consisting of DMEM containing 2% heat-inactivated horse serum and 1% penicillin-streptomycin. All these cells were maintained in a humidified atmosphere containing 5% CO2 at 37°C.

Random Positioning Machine

For the cell culture under MGS environment, a random positioning machine (RPM) (Dutch Space, Leiden, the Netherlands) was used. The RPM is in essence a 3-dimensional clinostat in which the gravity vector is continually reoriented, allowing the simulation of microgravity conditions. The cells were plated into a T-25 flask or 35-mm cover-glass-bottom dish. The dish was sealed with gas-permeable parafilm to avoid the leakage of the medium, and the dishes were fixed near the center of 2 frames that simultaneously rotated independently of each other. The entire system was placed in an incubator set at 37°C. Control samples (1 g) were placed in the same incubator and near the RPM. Microgravity simulation lasted from 1 to 7 days. The medium was replaced with fresh medium every 72 hr.

Immunocytochemistry Evaluation and Fusion Index

Cells were fixed using 4% formaldehyde in PBS for 15 min and then washed with PBS twice for 5 min. Fixed cells were permeabilized and blocked with 1% BSA (BioShop) in PBST (PBS with 0.25% Triton X-100) for 30 min at room temperature. The cells were washed once with PBST and incubated in anti-myosin heavy chain (MyHC, 1:100; R&D Systems) supernatant for 1 hr at room temperature. The cells were washed with PBST three times and incubated with AlexaFluor488 secondary antibody (1:250; Cell Signaling Technology) in blocking solution for 1 hr in the dark at room temperature. Following extensive washes with PBS, cells were mounted in mounting medium with DAPI (Merck KGaA) and imaged using the Zeiss Observer Z1 microscope. The fusion index was determined by quantifying the number of nuclei within multinucleated MyHC-expressing myotubes divided by total nuclei using ImageJ software (NIH).

Isolation of Total RNA and Real-Time PCR Analysis

The total RNA from cell line and tissue samples was isolated by Trizol reagent (Invitrogen), following the manufacturer’s instructions. cDNA synthesis for mRNA and lncRNA detection was carried out using SuperScript III first strand synthesis system for RT-PCR (Invitrogen). The Fast start Universal SYBR Green Master (Roche) was applied for the qRT-PCR. Quantification of amplicons was
done using ABI 7300 software (Applied Biosystems). GAPDH and U6 were used as endogenous controls for normalization. All the primer sequences are listed in Table S2. The relative fold changes of candidate genes were analyzed by using the 2^(-ΔΔCt) method.  

**Microarray Analysis**

150 ng total RNA was used to generate amplified and biotinylated sense-strand cDNA with the GeneChip WT PLUS Reagent Kit (Affymetrix), according to the manufacturer’s instructions. cDNA was hybridized to GeneChip Mouse Transcriptome Array 1.0 (Affymetrix, Thermo Fisher Scientific) for 16 hr in a 45°C incubator, rotated at 60 rpm. After hybridization, the microarrays were washed, and then they were stained using the Fluidics Station 450 followed by scanning with the Affymetrix GeneChip Scanner 3000 7G. The data were analyzed in Expression Console software from Affymetrix using default analysis settings. Differentially expressed lncRNAs were identified when the paired t test p value was <0.05 and the fold change was greater than 2. The sample size was three mice per group.

**Lentiviral Vector Construction and Lentivirus Production**

For construction of the IncMUMA overexpression lentiviral vector, mouse α-skeletal actin (MSA) promoter (2.0 kb) and full-length IncMUMA were subcloned into the lentiviral GV112 vector, which was provided by Shanghai Genechem (Shanghai, China), according to the manufacturer’s instructions. For the IncMUMA-KD lentiviral vector, shRNA targeting IncMUMA or a negative control scramble sequence was subcloned into the GV112 vector, respectively. Three shRNA sequences were designed by Shanghai Genechem: shIncMUMA-1, 5'-GACAGTTGATACCTTCTCTTG-3'; shIncMUMA-2, 5'-ACCATGTAGGGTAGACTTTTG-3'; and shIncMUMA-3, 5'-GCCATGTAGGGTAGACTTTTG-3'.

For the production of lentivirus, the expression vectors were co-transfected with packaging plasmid helper 1.0 vector (Shanghai Genechem) and envelope plasmid helper 2.0 vector (Shanghai Genechem) into 293T cells using TransIT-LT1 (Mirus Bio). The medium volume was made up to 500 ml with DMEM (Thermo Fisher Scientific) and 10% FCS (Hyclone). The co-transfected DNA was lipofected into the 293T cells using Lipofectamine 2000 (Invitrogen). After 48 hr, the supernatant was collected and analyzed with real-time PCR.

**Cell Transduction**

Based on a previously reported protocol with modification, C2C12 mouse myoblasts were seeded in 6-well plates and cultured until 60% confluent. Medium was then removed and 1.5 × 10^6 IU viral particles was added together with 8 μg/mL hexadimethrine bromide (Sigma-Aldrich). The medium volume was made up to 500 μL with DMEM and then incubated at 37°C and 5% CO₂ for 8 hr, at which time the DMEM plus viral particles was changed to DMEM with 10% FCS and the cells were cultured for 1–7 days.

**Bioinformatics Analysis for Targeted Gene Prediction**

The corresponding targeted genes for IncMUMA and miR-762 were predicted using RNAhybrid 2.12 (https://bibiserv.citeseer. uni-bielefeld.de/rnahybrid/). RNAhybrid is a tool used for predicting biological targets of certain RNA by searching for the presence of conserved sites that match the seed region of an RNA.

**Luciferase Reporter Assays**

Following a previously established procedure, the putative sequences of the binding site in miR-762 and the mutated sequences were cloned into a pmirGLO Dual-luciferase miRNA Target Expression Vector (Promega, Madison, WI, USA) to form the reporter vector. The reporter vector was co-transfected with IncMUMA-WT or IncMUMA-Mut into C2C12 cells by using Lipofectamine 2000 (Invitrogen). After 48 hr, luciferase assay was performed using a Dual-Luciferase Reporter Assay System (Promega) according to the manufacturer’s protocol.

**Biotin-Labeled IncMUMA Pulled down**

Biotin-labeled IncMUMA was synthesized by Sangon Biotech (Shanghai, China). The C2C12 cells were transfected with miR-762 by using Lipofectamine 2000 (Invitrogen). Different doses of biotin-labeled IncMUMA (0.5, 5, and 50 mM) were incubated with cytoplasmic lysates from miR-762-transfected C2C12 cells for 30 min at room temperature, and complexes were isolated with streptavidin-coated magnetic Dynabeads (Dynal). After wash steps, the captured RNA was purified and analyzed with real-time PCR.

**Western Blotting**

According to previously established procedures, cells and muscle samples were harvested, washed with 1 × PBS, and lysed in NP40 lysis buffer (50 mM Tris-HCl, 150 mM NaCl, 0.1% NP-40, 5 mM EDTA, and 10% glycerol) with protease inhibitor cocktail (Sigma). Proteins were separated in SDS-PAGE, transferred, and immunoblotted with various antibodies. The antibodies used were anti-MyoD (1:1,000; Invitrogen) and anti-β-actin (dilution 1:3,000; Santa Cruz Biotechnology).

**Generation of Muscle-Specific miR-762 Knockin Mice**

For the generation of ROSA26-PCAG-STOPlox-miR-762-EGFP mice, targeting vector was constructed by inserting a STOPlox-miR-762-EGFP in the ROSA26 allele and then electroporating into embryonic stem cells from C57BL/6J mice. Then, these mice were intercrossed with Tg(ACTA1-cre/Esr1*)2Kesr/J (ACTA1-Cre) mice (The Jackson Laboratory), and the offspring were intraperitoneally administered with tamoxifen (500 mg in 200 μl PBS/day) for 5 days at the age of 2 months old to induce muscle-specific miR-762 knockin. The ACTA1 gene encodes skeletal muscle α-actin, which is the predominant actin isoform in the sarcomeric thin filaments of adult skeletal muscle. Thus, miR-762 was overexpressed in skeletal muscle under the control of the ACTA1 gene in the muscle-specific miR-762 knockin mice.

**In Vivo Lentiviral Particle Administration**

Following established protocols with minor modifications, the animals were anesthetized with an isophorone gas inhale system, and 15 μL viral preparation with a titer of 10^8 IU/mL was injected into the mid-portion of gastrocnemius muscles by using a Hamilton syringe (Sigma-Aldrich) with a 33G needle. The mice were sacrificed.
after 42-day HLS. The specific force generated by gastrocnemius muscle was evaluated by in situ muscle functional testing before sacrifice. The gastrocnemius muscles were collected and weighed. The muscles were cryosectioned for histological staining. The mean muscle fiber CSA was determined. IncMUMA level and mRNA were assessed by real-time PCR. The protein of MyoD was evaluated by western blot analysis.

In Situ Muscle Functional Testing

In situ muscle functional testing was performed following the established protocol.\(^2\) The animals were anesthetized with an isophorone gas inhalation system and then placed on a 37°C heat platform. The gastrocnemius muscle was exposed, and the distal tendon was attached to the lever arm of a position feedback motor. The knee joint was immobilized by clamping it to the platform (In Vivo Muscle Test system 1300A, Aurora Scientific). A needle electrode was inserted through the skin and positioned on the peroneal nerve to stimulate contraction of the gastrocnemius muscle. The peak isometric force at 100 Hz for 400 ms was recorded and calculated with the ASI Dynamic Muscle Control Software (DMC version v.5.1 beta, Aurora Scientific). Specific force was expressed as peak tetanic force normalized to the muscle physical CSA.

Histology Evaluation

The dissected gastrocnemius muscles were snap frozen in liquid nitrogen-cooled isopentane and then embedded in optimal cutting temperature (OCT) medium. Serial cross-sections (6-μm thickness) were cut from the mid-belly of the muscles on a cryostat at −20°C for histological staining. H&E staining was performed to examine muscle fiber CSA in slide cuts from mid-belly and to determine the CSA of the muscle fiber. Slides were visualized with an Axio Observer Z1 microscope (Carl Zeiss). The muscle area and the number of fibers were determined using ImageJ software (NIH). The mean muscle fiber CSA was obtained by dividing the area of over 300 muscle fibers in 6 different locations in one slide by the fiber number.\(^2\)

Statistical Analysis

Data are presented as mean ± SEM. Each separate sample was analyzed in triplicate to yield an average value, and "n" is the number of samples in each group (n = 3 for microarray analysis, n = 5 for in vitro evaluation, and n = 10 for in vivo evaluation). GraphPad Prism v.6.0 was used for statistical analysis of the data. All results between groups were analyzed by applying the one-way ANOVA\(^2\) and Student’s t test. A difference with p < 0.05 was regarded as statistical significance.

SUPPLEMENTAL INFORMATION

Supplemental Information includes eight figures and two tables and can be found with this article online at https://doi.org/10.1016/j.ymthe.2018.09.014.

AUTHOR CONTRIBUTIONS

B.-T.Z., G.Z., and A.L. supervised the whole project. Z.-K.Z. performed the animal study and wrote the manuscript. J. Li performed the in vitro study and interpreted the data. D.G. and C.L. performed the microarray data analysis and bioinformatics analysis. Z.Z. and J. Liu provided the technical support and their professional expertise.

CONFLICTS OF INTEREST

The authors have no conflicts of interest.

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