**FTO is required for myogenesis by positively regulating mTOR-PGC-1α pathway-mediated mitochondria biogenesis**

Xiaobo Wang¹,², Ning Huang¹,², Min Yang¹, Dandan Wei¹, Haoran Tai¹, Xiaojuan Han¹, Hui Gong¹, Jiao Zhou¹, Jianqiong Qin¹, Xiawei Wei¹, Honghan Chen¹, Tingting Fang¹ and Hengyi Xiao*¹

Global germ line loss of fat mass- and obesity-associated (FTO) gene results in both the reduction of fat mass and lean mass in mice. The role of FTO in adipogenesis has been proposed, however, that in myogenesis has not. Skeletal muscle is the main component of body lean mass, so its connection with FTO physiologic significance need to be clarified. Here, we assessed the impact of FTO on murine skeletal muscle differentiation by in vitro and in vivo experiments. We found that FTO expression increased during myoblasts differentiation, while the silence of FTO inhibited the differentiation; in addition, skeletal muscle development was impaired in skeletal muscle FTO-deficient mice. Significantly, FTO-promoted myogenic differentiation was dependent on its m6A demethylase activity. Mechanically, we found that FTO downregulation suppressed mitochondria biogenesis and energy production, showing as the decreased mitochondrial mass and mitochondrial DNA (mtDNA) content, the downregulated expression of mtDNA-encoding genes and peroxisome proliferator-activated receptor gamma coactivator 1 alpha (PGC-1α) gene, together with declined ATP level. Moreover, the involvement of mTOR-PGC-1α pathway in the connection between FTO and muscle differentiation is displayed, since the expression of FTO affected the activity of mTOR and rapamycin blocked FTO-induced PGC-1α transcription, along with the parallel alteration pattern of FTO expression and mTOR phosphorylation during myoblasts differentiation. Summarily, our findings provide the first evidence for the contribution of FTO for skeletal muscle differentiation and a new insight to study the physiologic significance of RNA methylation.

*Cell Death and Disease (2017) 8, e2702; doi:10.1038/cddis.2017.122; published online 23 March 2017*

Fat mass- and obesity-associated (FTO) gene originally attracted attentions as an obesity- and diabetes-related protein owing to the significant association between its genetic polymorphism and BMI of human beings.¹-⁴ The linkage of FTO to body development and metabolic homeostasis is further demonstrated by experimental data. For instance, FTO overexpression leads to weight increase and its knockout induces weight loss in mice.⁵⁻⁷ It is clear now that FTO works as a nucleic acid demethylase capable of removing methyl groups from single-strand DNA and RNA,⁸⁻¹⁰ and therefore regulating N⁶-methyladenosine (m⁶A) level of RNA in cells.¹¹ The cellular function of FTO remains largely unknown, as very limited biologic processes have been linked with the function of FTO. Given the predicted close association of FTO with energy homeostasis and body development, exploring the cellular processes connected with FTO function is a perspective research pursuit.

Studies have shown that FTO expression affects the hypothalamus-governed food intake in mice and the fat accumulation in different animals,⁷,¹²⁻¹⁴ leading to the impression that FTO targets neural and adipose tissues and functions on central nervous system secretion and adipose differentiation. However, these findings cannot fully explain the data obtained from other experiments. For example, the homologous knockout of the FTO gene caused >50% of mice death embryonically, and the reduction of body weight, not only the weight of fat mass but also that of lean mass.⁷,¹⁴ In addition, FTO expression in non-neural and non-fat cells influenced the cell proliferation.¹⁵ Combining these data with the fact that FTO expresses in many mammalian tissues,⁷ it is reasonable to persuade the biologic function of FTO is not limited at neural and adipose tissues.

Skeletal muscle is the biggest energy-producing and -consuming organ in human body and also releases various myokines that participate in the metabolism regulation of the whole body.¹⁶ Accordingly, the disorders of skeletal muscle are involved in metabolic diseases.¹⁷ Differentiated and functional muscle cells are called myotubes, and derived from myoblasts during embryo and post-natal periods. Muscle maintenance and repair in adulthood are basically dependent on the differentiation of satellite cells, which are quiescent muscle precursors locating along with muscle fibers.¹⁸ Upon activation, satellite cells reenter the cell cycle to proliferate and differentiate to myoblasts, followed by the maturation of multinucleated muscle fibers.¹⁹ The process of muscle differentiation or myogenesis is controlled by various...
transcription factors, particularly by myogenic regulatory factors (MRFs). Among MRFs, MYF5 and MYOD are crucial for myogenic determination, while myogenin and MRF4 mainly drive the terminal differentiation. Although increasing knowledge have been achieved, the exact mechanism regarding to skeletal muscle differentiation has not been fully established, that is not an ideal state for developing new approaches to govern metabolic homeostasis and confront muscle-related disorders.

Energy supply is another critical factor for muscle differentiation. A series of studies proposed the role of mTOR-PGC-1α-mitochondria axis in the regulation of myogenesis. The facts are that the impairment of mitochondrial function blocks myogenic differentiation and mTOR signaling pathway, a master regulator of cell metabolism and energy homeostasis, and positively regulates PGC-1α expression, a coactivator that controls mitochondrial biogenesis. Therefore, a situation capable of affecting the activity of mTOR pathway should be a potential regulator of mitochondria function and muscle differentiation.

The role of FTO in skeletal muscle is poorly understood, despite recently a couple of studies mentioned that the cells lacking FTO have decreased activity of the mTORC1 pathway, and the FTO-deficient mice have reduced lean mass. These observations suggest a potential connection among FTO, mTOR and myogenesis, raising a new

**Figure 1** Endogenous FTO expression elevates during myogenic differentiation. C2C12 cells and MPM cells were differentiated for indicated days in DM, phase-contrast microscopy of differentiated C2C12 cells (a) and MPM cells (d). Western blot analysis of whole-cell lysates from differentiated C2C12 cells (b) and MPM cells (e) with indicated antibodies. qRT-PCR analysis of differentiated C2C12 cells (c) and MPM cells (f) with FTO. Asterisks indicate statistical significance (*P < 0.05, **P < 0.01, ***P < 0.001)
hypothesis that FTO probably plays a role in muscle differentiation through mTOR-dependent manner.

Therefore, we asked if FTO can regulate myogenic differentiation in the present study and tried to elucidate the possible mechanism. For this sake, in vitro and in vivo experiments were performed using cultured myoblasts and skeletal muscle-specific FTO-deficient mice. Our results indicate the positive association between FTO protein and skeletal muscle differentiation. Furthermore, our results reveal that FTO regulates myogenic differentiation, at least in part through mTOR-PGC-1α-mitochondria axis. This is the first time to demonstrate the role of FTO in myogenic differentiation, and our study also provides a new line of evidence stressing the importance of mitochondria function on muscle differentiation.

Results

FTO expression influences myogenic differentiation in vitro. As the first step for understanding the correlation between FTO expression and myogenic differentiation, we measured the endogenous FTO expression during the differentiation process of two cell models: both go from myoblasts to myotubes under novel cultivation condition described in Materials and Methods section. As shown, along with the differentiation of C2C12 myoblasts (Figure 1a), the protein level (Figure 1b) and the mRNA level (Figure 1c) of FTO both increased, similar to the situation observed in mouse primary myoblasts (MPMs) (Figures 1d–f). These results indicate that the expression of endogenous FTO is upregulated during myogenic differentiation.

Then, we silenced or overexpressed fto to see if the alteration of fto can influence myogenic differentiation. Silencing the fto by siRNA in C2C12 myoblasts, decreased the myotubes formation during differentiation (Figures 2a and b). Consistently, the protein level of myogenin and myosin heavy chain (MHC), early and late differentiation markers, respectively, decreased in FTO silenced C2C12 cells (Figure 2c), together with an apparent decrease of myogenin and MHC mRNA (Figure 2d). To strengthen these findings, we established two MPM pools with fto stably silenced or overexpressed by lentivirus, refer to as MPM/shCtrl and MPM/shFTO myoblasts, respectively. We found the growth rate was around 70% and the protein level of cell cycle regulator Cyclin D1 was reduced in MPM/shFTO cells compared with MPM/shCtrl cells (Supplementary Figure 1). Therefore, we seeded more MPM/shFTO cells before differentiation induction to make the number of these cells equal with that of MPM/shCtrl cells when the differentiation was initiated. For MPM/shFTO myoblasts, the result was consistent with that seen in C2C12 cells with transfection of FTO siRNA, clearly showing that FTO silence suppressed myogenic differentiation. As to MPM/FTO myoblasts, however, the phenotype alteration was not observed, since the situation of myogenic differentiation was almost similar in FTO overexpressed cells and control cells (Figures 2e and f), resembled as observed when FTO overexpressed in C2C12 cells. Paralleling with the morphologic assessment, the influence of FTO expression on myogenic differentiation in these cells was confirmed by detecting myogenin and MHC expression (Figures 2g and h). These results suggest that FTO is required for myogenic differentiation.

Skeletal muscle development is impaired in skeletal muscle FTO deficiency mice. To gain in vivo evidence for the role of FTO in myogenesis, doxycycline-inducible skeletal muscle-specific FTO knockout mice were generated. The procedure includes three steps: (1) crossing FTOlox/lox mice (Figures 3a and b), (2) treating these mice with doxycycline (Figure 3c) to implement FTO knockout in skeletal muscle of adult mice and getting FTOlox/lox HSA-Cre mice (Figures 3a and b). (3) measuring in genome DNA demonstrates that exon 3 of the FTO gene was efficiently deleted in rectus femoris of FTOKO versus WT mice (Figure 3d), and examining the muscle specificity of FTO knockout by quantitative real-time PCR (qRT-PCR) and western blot assays (Figures 3e and f).

By administrating doxycycline to pregnant FTOlox/lox HSA-Cre mice crossed with FTOlox/lox mice, we evaluated the role of FTO expression in the skeletal muscle development of their newborn offspring (Figure 4a). Then, we confirmed that FTO was depleted in hindlimb muscle of these newborn mice (Figures 4b and c). The body and hindlimb weight and the length of hindlimb of WT and FTOKO offspring P1 mice were no different (Supplementary Figures 2a and b). In line with our in vitro experiments, MHC and α-actin expression in the muscle of FTOKO mice was lower than that of WT littermates (Figures 4b and c). Furthermore, based on H&E-stained tissue section of hindlimb muscles, we found that the size of myofibers obviously reduced, with enlarged interval gap and decreased number of myofibers (Figures 4d and e). In line with our in vivo experiments, MHC and α-actin expression in the muscle of FTOKO newborn mice was lower than that of WT littermates (Figure 4a). Then, we confirmed that FTO was depleted in hindlimb muscle of these newborn mice (Figures 4b and c). The body and hindlimb weight and the length of hindlimb of WT and FTOKO offspring P1 mice were no different (Supplementary Figures 2a and b). In line with our in vitro experiments, MHC and α-actin expression in the muscle of FTOKO mice was lower than that of WT littermates (Figures 4b and c). Furthermore, based on H&E-stained tissue section of hindlimb muscles, we found that the size of myofibers obviously reduced, with enlarged interval gap and decreased number of myofibers (Figures 4d and e, and Supplementary Figure 2c). By Image J program (NIH, Bethesda, MD, USA) analysis, it was found that, in comparison with the WT, the number of smaller sized myofibers <30 μm² in area were significantly more in FTOKO mice, whereas larger sized myofibers beyond 30 μm² in area were much less (Figure 4f). These results provided in vivo evidence for the influence of FTO expression on myogenesis.

The demethylase activity of FTO is required for myogenic differentiation. Given FTO is a demethylase of N6-methyladenosine (m^6A) in RNA,11 we next collected data to assess whether FTO-mediated m^6A demethylation in mRNA is required for myogenesis. First, we measured the m^6A level of mRNA by dot blot assay, and observed a severe decrease during myogenic differentiation and in MPM/FTO cells, whereas an obvious increase in MPM/shFTO cells and in the muscle of FTOKO newborn mice (Figures 5a and b). To ensure the responsibility of m^6A dot blot assay for FTO activity, we generated MPM cells stably overexpressing wild-type FTO protein or a point mutation of FTO protein that lacks the demethylase activity of FTO (R96Q),10 and conducted a measure under the condition where endogenous FTO expression was repressed by FTO mRNA 5′-UTR-specific siRNA. As shown in Figure 5c, the m^6A level of mRNA significantly increased in MPM/R96Q cells, compared with MPM/FTO cells. Parallel, R96Q overexpression impaired the
differentiation of MPM cells (Figure 5d), as the protein and mRNA levels of MHC declined (Figures 5e and f). The final set of evidence came from the use of pharmaceutical compound rhein, a verified inhibitor of FTO demethylase. 27 Similar with the effect of FTO silence, rhein treatment remarkably increased the m6A level of mRNA (Figure 5g) and impaired the differentiation of primary myoblasts (Figure 5h), with decreased MHC protein and mRNA level (Figures 5i and j). As expected, rhein did not affect the expression of FTO (Figures 5i and j). These results indicated that the m6A demethylation activity of FTO is required for its influence on myogenesis.

**FTO affects mitochondria biogenesis and function during myogenic differentiation.** To explore the mechanism underlying the role of FTO in myogenic differentiation, we measured the expression of several MRFs during the differentiation of myoblasts. Two well-known MRFs acting for the initiation of myogenesis, MyoD and Myf5, were not influenced upon FTO silence at mRNA level (Figure 6a). However, the mRNA of myogenin, encoding a MRF downstream of MyoD and Myf5, decreased in FTO silence cells on day 2 of differentiation (Figure 6a).

Considering mitochondria dysfunction is an important reason for the downregulation of myogenin gene but not the MyoD and Myf5 genes,28 we assumed that FTO may affect myogenin expression and consequential myogenesis through regulating mitochondrial functions. To verify this hypothesis, we evaluated the status of mitochondria function, particularly from the view of mitochondria biogenesis and ATP production. As to evaluate mitochondria biogenesis, we measured the mass of mitochondria by MitoTracker Green staining, the content of mitochondrial DNA (mtDNA) by qRT-PCR and the expression of the genes important for mitochondria...
biogenesis by qRT-PCR analysis. Our data show that the mass of mitochondria decreased in FTO silenced cells (Figures 6b and c) and the mtDNA content increased with differentiation but decreased in FTO silence cells (Figure 6d). In terms of the gene expression, the mRNA of PGC-1α, a master transcriptional coactivator for mitochondria biogenesis, was significantly declined upon FTO silence (Figures 6e and h), without alteration in mRNA stability (Figure 6f), and three downstream targets of PGC-1α, TFAM, cytochrome c and cox5a; all increased during myogenesis, while decreased upon FTO silence (Figures 6g and h). In respect of ATP production, we found that intracellular ATP level increased during myogenesis, but decreased upon FTO silence (Figures 6g and h). In respect of mTOR production, we found that intracellular ATP level increased during myogenesis, but decreased upon FTO silence (Figures 6i). Reversely, although FTO overexpression increased PGC-1α expression, the expression of cytochrome c and cox5a was not influenced (Figure 6i). These results together suggest that FTO silence inhibits myogenin expression during myogenic differentiation by downregulating the expression of PGC-1α gene, and then represses mitochondria biogenesis and function.

mTOR-PGC-1α pathway activation is crucial for the role of FTO in myogenic differentiation. Since mTOR-PCG-1α controls mitochondria biogenesis and FTO can activate mTORC1, we asked whether FTO regulated mitochondrial biogenesis through mTOR-PGC-1α pathway. Western blotting assays confirmed that mTORC1 pathway was activated during myogenic differentiation (Figure 7a), and mTORC1 pathway together with PGC-1α expression was downregulated upon FTO silence (Figure 7b). In addition, PGC-1α mRNA level was higher in undifferentiated MPM/FTO cells and this increase was abrogated when treated MPM/FTO cells with mTOR inhibitor rapamycin (Figure 7c). Because of its ability in influencing myogenic differentiation, we tested whether FTO activity affects the level of PGC-1α mRNA. Interestingly, R96Q mutation of FTO protein and the rhein treatment resulted in similar alteration in PGC1a expression as that caused by rapamycin treatment (Figure 7d). In contrast, the treatment of cells with insulin, the mTOR activator, restored the decrease of myotubes (Figure 7e) and the protein and mRNA levels of MHC and PGC-1α (Figures 7f and g) in FTO silence cells. Consistently, PGC-1α, TFAM, Cox5a and the phosphorylation of mTOR (S2448) were also decreased in mouse muscles lacking FTO (Figure 7h). Together, our findings suggest that FTO acts on the upstream of mTOR-PGC-1α pathway to regulate myogenic differentiation.

Discussion

In this study, we first demonstrated that FTO plays an important role in myogenic differentiation and skeletal muscle development, and its m6A demethylation activity is required for this role. We also found that the effect of FTO on muscle differentiation is mediated at least partially by mTOR-PGC-1α-mitochondria axis.

Several studies have shown FTO deficiency resulted in high perinatal lethality, and reduced body length, fat mass and lean
mass in mice,\textsuperscript{6,7} implying that FTO fundamentally impacts on the development and functions of body composition. Yun-gui Yang \textit{et al.} reported that FTO regulates adipogenesis, and thereby influences fat mass and body weight.\textsuperscript{29,30} We investigated the role of FTO in myogenesis, considering that skeletal muscle is a major component of lean mass and an essential insulin-sensitive organ similar as adipose.\textsuperscript{31} Our study not only revealed the contribution of FTO on myogenic differentiation in cultivated myoblasts, either established C2C12 myoblasts cell line or primary murine myoblasts, but also confirmed the impact of FTO on skeletal muscle development, using skeletal muscle-specific FTO-deficient mice. The \textit{in vivo} evidence is novel as it was produced from mice with muscle-specific FTO depletion. This line of mice provides us convenience for evaluating the role of local FTO in myogenesis and muscle differentiation. In terms of this point, it is a good model and tool for \textit{in vivo} study, because the complicated situation caused by the influence of FTO expressed in other tissues can be avoided. Given skeletal muscle is the key component of lean mass, our findings support the opinion that FTO influences lean mass development.

Many insights for the mechanism of muscle differentiation have been achieved, revealing that sequential expression of different MRFs is crucial.\textsuperscript{20} We particularly concerned the effect of FTO on myogenin, as myogenin is the key MRF that directly activates genes encoding myofiber proteins.\textsuperscript{32} Hinted by a previous study,\textsuperscript{28} we determined the connection of mitochondrial biogenesis and function with FTO-mediated myogenesis and myogenin expression. This result is understandable as the energy produced by mitochondria is indispensable for cell growth, differentiation and organic development,\textsuperscript{33} not to mention the differentiation and functioning of skeletal muscle requires particularly lots of energy supply.\textsuperscript{34} Then, how does FTO affect mitochondria biogenesis? Our data draft a cascade from FTO to mTOR and to PGC-1α, the master factor for mitochondria biogenesis. It has been reported that mTOR-YY1-PGC-1α-mitochondria axis exists in cells, where mTOR controls mitochondrial biogenesis and

\begin{figure}
\centering
\includegraphics[width=\textwidth]{Fig4}
\caption{Skeletal muscle development is impaired in skeletal muscle FTO deficiency mice. (a) Dox regimen assay scheme. Vertical lines indicate daily intervals. (b) qRT-PCR and (c) western blot analysis of whole-cell lysates of hindlimb muscles from WT and FTOKO offspring P1 mice with indicated genes and antibodies, respectively. (d) H&E analysis of hindlimb muscles from WT and FTOKO offspring P1 mice. Scale bars: 20 \(\mu\)m. (e) Quantification of H&E in (d). (f) Distributions of fiber sizes were analyzed. Asterisks indicate statistical significance (* \(P<0.05\), ** \(P<0.01\), *** \(P<0.001\)).}
\end{figure}
function through regulating the stability of YY1-PGC-1α transcripional complex. This axis is partially confirmed in our study, showing that FTO influences the expression of PGC-1α and mitochondrial biogenesis with mTOR dependency. Moreover, although just being the preliminary information, our data indicates that FTO is an upstream regulator of mTOR pathway, coinciding with a finding published previously standing on the fact that FTO affects mTORC1 activity.

FTO is a nucleic acid demethylase that removes methyl groups from both DNA and RNA. It is commonly accepted that its most important functional role is demethylation of N6-methyladenosine (m6A) in mRNA. For example, FTO controls exon splicing of adipogenic regulatory factor RUNX1T1 by regulating m6A levels around splice sites and thereby modulates differentiation. As FTO inhibitor rhein and FTO mutant (R96Q) that lacks demethylase activity inhibited myogenic differentiation, we consider the activity of FTO is required for myogenic differentiation. This finding provides new evidences and ideas for the mechanism study of skeletal muscle differentiation.

An interesting result in our study is that FTO depletion interferes with myogenic differentiation, while overexpression of FTO could not promote myogenic differentiation in vitro. The unmatched results from loss-of-function and strength-of-function experiments we got in vitro seem coincident with those reported previously. For example, according to the statement from Gao et al., FTO deficiency in mice results in an obvious reduction of lean mass. However, in the
demonstration of Church et al., overexpression of FTO does not increase lean mass, at least in male mice. Although more data are needed, we would like to propose a brief explanation for these unmatched resultants based on the line of our finding. It is that FTO deficiency can inhibit myogenic differentiation because of its significant role in suppressing mitochondria biogenesis, which can severely interfere muscle differentiation requiring ATP consuming; on the other hand, the overexpression of FTO does not invoke marked change in differentiation, which may be owing to the enough capacity of endogenous FTO expression for supporting muscle differentiation. This notion should be strengthened by our data that the expression of Cytochrome c and Cox5a genes, directly participating in mitochondria biogenesis, does not change in FTO overexpression cells. It also remains that although being an important transcriptional coactivator involving in mitochondria biogenesis, PGC-1α expression level is not the only determinant of mitochondria biogenesis, so its expression could not linearly affect myogenic differentiation. Actually, many studies have confirmed that the function of PGC-1α can be regulated at protein modification level, for instance, it can be acetylated by GCN5 acetyltransferase complex, and phosphorylated by p38 MAP kinase, AMP kinase and Akt/PKB.

Although we displayed that FTO could regulate myogenic differentiation, the precise mechanism of this relationship has not been revealed. It is still unclear how FTO affects the activity of mTOR pathway, which is the direct target gene of FTO as a
demethylase of m6A of RNA. Further studies focused on these issues are needed.

In summary, our findings demonstrate that the function of FTO is required for myogenic differentiation and suggest FTO-mediated mTOR-PGC-1α-mitochondrial axis involved in this regulation. This study is a basic line for further investigation of the molecular mechanisms in the role of FTO during myogenic differentiation, and will be informative for developing our understanding about muscle differentiation, RNA methylation and mTOR pathway regulation.

Materials and Methods

Isolation of MPMs. Primary myoblasts from about 10-day-old C57BL/6J were isolated and cultured following the protocol of Gharaibeh et al.39 In brief, hindlimb muscles were minced mechanically and digested with enzyme mixture: 0.2% collagenase II (Invitrogen, cat. 17101015, Carlsbad, CA, USA) and 0.05% trypsin in DMEM (Gibco, Carlsbad, CA, USA) for 45 min at 37 °C with slight agitation. The tissue was triturated vigorously using a 1 ml tip and passed through a 70 μm filter, and the cells were collected by centrifugation. Cells were suspended in primary myoblasts growth media (DMEM supplemented with 20% FBS and 1% penicillin/streptomycin) for 2 h at 37 °C; the non-adherent cells were then transferred to another plate. To get pure satellite cells, the following procedure was performed in strict accordance with the purifying method in the study by Gharaibeh et al.39 After about 1 week, the satellite cells proliferate as myoblasts naturally.

Cell culture. C2C12 and MPM were maintained at 37 °C and 5% CO2 in growth media (GM: DMEM medium containing 10% (v/v) FBS and 1% antibiotics). To induce myogenic differentiation, cells were grown to 70–80% confluence in GM and then switched to differentiation medium (DM: DMEM supplemented with 2% (v/v) horse serum and 1% antibiotics). A total of 293FT cells were also cultured in GM.

Lentivirus packaging and cell lines. The vectors containing cDNA of wild-type FTO and R96Q mutant of FTO were kindly gifted by Renbin Zhao.40 Primers bearing Xba1 and Not1 sites were used to generate PCR fragments that were subcloned into pLVX-IRES-ZsPuro lentiviral expression vector (Clontech, Shiga, Japan). Lentiviral shRNA construct for mouse FTO gene was purchased from...
GENECHEN (Shanghai, China). The target sequence was: AGAACCATAC-TATTGCC. Lentivirus was produced by co-transfection of lentivirus packing plasmids with psPAX2 and pMD2.G using Jet PRIME (PolyPlus, Illkirch, France) into 293FT cells following manufacturer's instruction. Medium was changed 24 h post transfection and the medium containing virus was collected after 72 h, followed by a centrifugation at 10 000 g for 10 min. The supernatant was used to infect MPM cells in the presence of 10 μg/ml polybrene (Sigma-Aldrich, cat. H9268, Carlsbad, CA, USA), or stored at –80 °C. Selection of resistant colonies was initiated 48 h later using 3 μg/ml puromycin (Life Technology, Carlsbad, CA, USA; cat. A1113803).

RNA interference. All siRNAs were purchased from Sheng Gong (Shanghai, China), the sequences used are provided in Supplementary Information. For knockdown experiments, cells were transfected with siRNAs using Jet PRIME (PolyPlus) following manufacturer's instruction. The transfection media were then replaced with DMEM supplemented with fetal bovine serum, and cells were ready for subsequent differentiation induction.

Western blots. Cells were lysed in RIPA buffer with a cocktail of protease inhibitors (Biotool, cat. B14002, Houston, TX, USA). Twenty micrograms of protein extracts were separated by SDS-PAGE and transferred to PVDF membranes (Millipore, cat. PVVP29232A, Billerica, MA, USA). Membranes were blocked with 5% non-fat milk, followed by overnight incubation with primary antibodies against FTO (Abcam, cat. ab92821, Cambridge, UK), myogenin (Zen Bioscience, cat. 501926), MHC (eBioscience, cat. 14-6503, Carlsbad, CA, USA), GAPDH (Abcam, cat. ab181602), α-actin (Beyotime Biotechnology, cat. AA132, Beijing, China), p-mTOR (Cell Signaling Technology, cat. 5536, Danvers, MA, USA), mTOR (Cell Signaling Technology, cat. 2972), p-p70S6k (Cell Signaling Technology, cat. 9705), p70S6k (Cell Signaling Technology cat. 9202), Cyclin D1 (Cell Signaling Technology, cat. 2922), PGC-1α (Sheng Gong, cat. D162041), TFAM (Zen Bioscience, cat. 600252) and Cox5a (Sheng Gong, cat. D261450). Detection was made with HRP-conjugated secondary antibody (Zen Bioscience, cat. 5013926) and signals were detected with ECL Plus Western Blotting Reagent Pack (Bio-Rad, Hercules, CA, USA).

qRT-PCR analysis. Cells and skeletal muscle were collected and washed twice with PBS, and total RNA was extracted using Trizol reagent (Takara, Shiga, Japan) following the manufacturer's instructions. Reverse transcription for mRNA was carried out using cDNA Synthesis Super Mix (Biotool, cat. B24403). qRT-PCR was carried out in an ABI cycler using SYBR Green qPCR Master Mix (Biotool, cat. B21203), and the relative amount of cDNA was calculated by the comparative CT method using the 18S ribosomal RNA sequences as control. The primer sequences used are provided in Supplementary Information.

Analysis of m6A levels in mRNA using dot blot. Purified mRNA was denatured at 95 °C for 5 min and cooled down on ice. Samples (150 ng) were spotted on Amersham Hybond-N+ membranes (GE Healthcare, cat. RPN303B, Chicago, IL, USA) and air dried for 5 min, then UV-crosslinked (2 × auto-crosslink, 1800 UV Stratalinker, STRATAGENE, La Jolla, CA, USA). Membranes were blocked with 5% non-fat milk, followed by overnight incubation with primary antibodies against RPN303B, Chicago, IL, USA) using Student t-test (*P<0.05, **P<0.01, ***P<0.001).

Conflict of Interest

The authors declare no conflict of interest.

Acknowledgements. This work was supported by National Natural Science Foundation of China (Grant Number 81273224) and National Key Research and Development Program (Grant Number 2016YFC1200203). We thank Dr. Caiguang Yang for helping the setup of m6A detection, Dr. Canhua Huang and Yuquan Wei for continuous supports, and Dr. Jie Zhang, Ping Lin, Xiujie Wang and Yi Chen for all technical assistance. We also thank Dr. Feng Li and Dr. Fang Fang for helping the setup of m6A detection. The authors thank Dr. Canhua Huang and Yuquan Wei for continuous supports, and Dr. Jie Zhang, Ping Lin, Xiujie Wang and Yi Chen for all technical assistance. We also thank Dr. Feng Li and Dr. Fang Fang for helping the setup of m6A detection.

Author contributions

XW and NH performed most of the experiments, data analyses and the manuscript preparation; HX guided the study planning, experiment processing and manuscript preparation; and MY, DW, HT, XH, HG, JQ, XW, HC and TF— all contributed to the mitochondrial mass. Cells were washed with PBS and incubated at 37 °C for 30 min with 100 nM MitoTracker Green FM (Molecular Probes, Carlsbad, CA, USA). Cells were collected by using trypsin/EDTA and resuspended in PBS. Fluorescence intensity was detected with excitation and emission wavelengths of 490 and 516 nm, respectively, and the values were corrected for total protein (mg/ml).

ATP level quantitation. Intracellular ATP level assay was performed using an ATP assay kit (MCE, cat. HY-17559, Princeton, NJ, USA) using Student t-test (*P<0.05, **P<0.01, ***P<0.001).

Conflict of Interest

The authors declare no conflict of interest.

Acknowledgements. This work was supported by National Natural Science Foundation of China (Grant Number 81273224) and National Key Research and Development Program (Grant Number 2016YFC1200203). We thank Dr. Caiguang Yang for helping the setup of m6A detection, Dr. Canhua Huang and Yuquan Wei for continuous supports, and Dr. Jie Zhang, Ping Lin, Xiujie Wang and Yi Chen for all around convenience.

Author contributions

XW and NH performed most of the experiments, data analyses and the manuscript preparation; HX guided the study planning, experiment processing and manuscript preparation; and MY, DW, HT, XH, HG, JQ, XW, HC and TF—all contributed to the
1. Frayling TM, Timpson NJ, Weedon MN, Zeggini E, Freathy RM, Lindgren CM et al. A common variant in the FTO gene is associated with body mass index and predisposes to childhood and adult obesity. Science 2007; 316: 889–894.  
2. Dina C, Meyre D, Gallina S, Durand E, Köttner A, Jacobsson P et al. Variation in FTO contributes to childhood obesity and severe adult obesity. Nat Genet 2007; 39: 724–726.  
3. Scott LJ, Mohike KL, Bonnycastle LL, Willer CJ, Li Y, Duren WL et al. A genome-wide association study of type 2 diabetes in Finns detects multiple susceptibility variants. Science 2007; 316: 1341–1345.  
4. Scuteri A, Samuels S, Chen WM, Uda M, Albiga G, Strait J et al. Genome-wide association scan shows genetic variants in the FTO gene are associated with obesity-related traits. PLoS Genet 2007; 3: e115.  
5. Church C, Moir L, McMurray F, Girard C, Banks GT, eboul L et al. FTO effect on energy demand versus food intake. Proc Natl Acad Sci USA 2007; 104: 967–972.  
6. Deeb S, Gullberg R, Franzen L, Ekelund U, Hallberg J, WC et al. Total physical activity and FTO genotype are associated with body fat distribution. PLoS ONE 2009; 4: e62561.

Supplementary Information accompanies this paper on Cell Death and Disease website (http://www.nature.com/ccddis)