Coupling of mitochondrial function and skeletal muscle fiber type by a miR-499/Fnip1/AMPK circuit

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Abstract

Upon adaption of skeletal muscle to physiological and pathophysiological stimuli, muscle fiber type and mitochondrial function are coordinately regulated. Recent studies have identified pathways involved in control of contractile proteins of oxidative-type fibers. However, the mechanism for coupling of mitochondrial function to the muscle contractile machinery during fiber type transition remains unknown. Here, we show that the expression of the genes encoding type I myosins, Myh7/Myh7b and their intronic miR-208b/miR-499, parallels mitochondrial function during fiber type transitions. Using in vivo approaches in mice, we found that miR-499 drives a PGC-1α-dependent mitochondrial oxidative metabolism program to match shifts in slow-twitch muscle fiber composition. Mechanistically, miR-499 directly targets Fnip1, an AMP-activated protein kinase (AMPK)-interacting protein that negatively regulates AMPK, a known activator of PGC-1α. Inhibition of Fnip1 reactivated AMPK/PGC-1α signaling and mitochondrial function in myocytes. Restoration of the expression of miR-499 in the mdx mouse model of Duchenne muscular dystrophy (DMD) reduced the severity of DMD. Thus, we have identified a miR-499/Fnip1/AMPK circuit that can serve as a mechanism to couple muscle fiber type and mitochondrial function.

Keywords contractile fiber type; gene regulation; microRNA; mitochondrial function; muscle

Subject Categories Metabolism; Musculoskeletal System

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Introduction

Contractile fiber type and energy production are two key determinants of skeletal muscle function. Skeletal muscle fibers exhibit significant physiological plasticity, reflected in different patterns of coordinated metabolic and structural gene expression (Pette & Staron, 2000; Zierath & Hawley, 2004; Schiaffino & Reggiani, 2011). Muscle fibers are classified into slow-twitch (type I) and fast-twitch (type II) based on the myosin heavy chain (MHC) gene expressed (Pette & Staron, 2000; Zierath & Hawley, 2004; Schiaffino & Reggiani, 2011; Liu et al., 2015). Type I myofibers are high endurance and are mitochondria-rich (red), relying largely on mitochondrial oxidative metabolism for ATP production (Pette & Staron, 2000; Zierath & Hawley, 2004; Schiaffino & Reggiani, 2011; Liu et al., 2015). In contrast, type II myofibers are low endurance and contain fewer mitochondria, and primarily rely on glycolytic burst metabolism for energy production (Pette & Staron, 2000; Zierath & Hawley, 2004; Schiaffino & Reggiani, 2011; Liu et al., 2015).

During development, and upon adaption to physiological stimuli and systemic disease, skeletal muscle fibers undergo extensive metabolic and structural remodeling (Holloszy & Coyle, 1984; Booth & Thomason, 1991; Baldwin & Haddad, 2001; Bassel-Duby & Olson, 2006; Rowe et al., 2014; Neuf et al., 2015). During fiber type transition, the contractile machinery and energy production system must be precisely coordinated to maintain muscle function (Schiaffino & Reggiani, 2011). Exercise training is effective in improving muscle fitness by increasing the proportion of slow-twitch muscle fibers and by promoting mitochondrial oxidative capacity (Holloszy & Coyle, 1984; Booth & Thomason, 1991; Baldwin & Haddad, 2001; Bassel-Duby & Olson, 2006; Rowe et al., 2014; Neuf et al., 2015). Conversely, skeletal muscle dysfunction, including reduced numbers of slow-twitch muscle fibers and diminished fuel-burning capacity, is linked to several human diseases, including metabolic disorders, heart failure, and muscular dystrophy (Holloszy & Coyle, 1984; Booth & Thomason, 1991; Baldwin & Haddad, 2001; Bassel-Duby & Olson, 2006; Neuf et al., 2015). The mechanism for precise coupling of mitochondrial function and muscle contractile machinery during fiber type transition remains unknown. Previous studies using genetically modified mice have provided important clues. The myosin heavy chain genes, Myh7/Myh7b, have been defined as key regulators of muscle fiber type switching by encoding a network of intronic miRNAs (van Rooij et al., 2009; Bell et al., 2010). Specifically, miR-208b, which is encoded within the type I MHC (Myh7) gene, and miR-499, which is
encoded within another type I MHC (Myh7b) gene, have been shown to drive a slow-twitch fiber program by downregulating transcriptional repressors that suppress a broad program of slow-twitch contractile protein gene expression (Hagiwara et al., 2005, 2007; Ji et al., 2007; van Rooij et al., 2009; Bell et al., 2010; Quiat et al., 2011). We have recently found that nuclear receptor ERRγ directly activates the promoters driving the expression of miR-499 and miR-208b in muscle (Gan et al., 2013). In addition, the expression of miR-499 positively correlates with slow-twitch fiber proportion and with metabolic parameters, such as VO2max and ATPmax, in humans (Gan et al., 2013). However, whether and how these miRNAs orchestrate mitochondrial function to match the slow-twitch muscle fiber is unclear.

In this study, we found that the expression of Myh7b/miR-499 and Myh7/miR-208b parallels the mitochondrial respiration capacity during muscle fiber type transition. We speculated that the intronic miR-499 and miR-208b might regulate the correlated changes in mitochondrial function during the transition phase. Using transgenic and loss-of-function mouse lines, and primary skeletal myotubes in culture, we found that a regulatory circuitry comprised of Fnip1/AMPK drives PGC-1α-dependent mitochondrial function downstream of miR-499. Thus, our results suggest a molecular mechanism for the adaptive orchestration of mitochondrial function muscle fiber type.

**Results**

**The expression of miR-499 and miR-208b parallels mitochondrial function during muscle fiber type transition**

As an initial step, to explore whether miRNAs in myosin genes orchestrate mitochondrial function to match muscle contractile fiber type, we examined the expression patterns of miR-499 and miR-208b during muscle fiber type transition. Consistent with previously published data (van Rooij et al., 2009; Bell et al., 2010), the induced expression of Myh7 and Myh7b genes paralleled the elevated expression of miR-208b and miR-499 during differentiation of skeletal myoblasts into mature myotubes (Fig 1A). A marked increase in mitochondrial respiration capacity correlated with upregulation of miR-499 and miR-208b during the transition phase (Fig 1A and B). Similar results were obtained when we compared different muscle types from adult wild-type mice. As shown in Fig 1C, miR-499 and miR-208b were expressed preferentially in slow-fiber-dominant soleus muscle. Isolated mitochondrial respiration studies demonstrated that pyruvate-driven mitochondrial respiration rates were significantly higher in soleus (Sol) muscle compared to fast-fiber-enriched white vastus (WV) muscle (Fig 1D). Together, these results showed dynamic expression of miR-499 and miR-208b that paralleled mitochondrial function during muscle fiber type transition.

**miR-499 drives a mitochondrial oxidative metabolism program in skeletal muscle**

To investigate the potential role of miR-499 and miR-208b in regulating mitochondrial function, we analyzed the muscle fuel utilization of transgenic mice overexpressing miR-499 in muscle (MCK-miR-499) given that miR-499 and miR-208b function redundantly in vivo (van Rooij et al., 2009; Bell et al., 2010). The real-time respiratory exchange ratio (RER) was measured during a run-to-exhaustion exercise protocol. The MCK-miR-499 mice had a lower RER than did their NTG littermate controls (Fig 1E), indicative of a tendency to preferentially oxidize fat over carbohydrate, consistent with enhanced exercise performance in these mice. Notably, levels of blood lactate following exercise were significantly lower in both younger and older MCK-miR-499 mice than in their NTG littermate controls (Fig 1F). Mitochondrial respiration rates were determined in the plantaris muscle of the MCK-miR-499 mice and corresponding NTG controls. Consistent with the increased aerobic metabolism in MCK-miR-499 mice, pyruvate-driven state 3 respiration rates were significantly higher in MCK-miR-499 muscle compared to the NTG controls (Fig 1G). Together, these results suggest that MCK-miR-499 muscle has increased mitochondrial function.

Comparative analysis of RNA isolated from gastrocnemius (GC) muscle of the MCK-miR-499 mice and NTG littermate controls, confirmed increased expression of many genes encoding slow contractile proteins in MCK-miR-499 muscle (Appendix Table S1). Notably, gene ontology (GO)-based analysis revealed that the primary upregulated genes were mitochondrial metabolic genes (Fig 2A). A broad array of genes encoding a variety of components of the mitochondria were induced in MCK-miR-499 GC muscle (Appendix Table S2). Real-time PCR confirmed that the expression of nuclear-encoded genes (Ndufa1, Ndufs4, Uqcr11, Cox7a1, Cox7b, Atp5e, and Atp5l) and mitochondrial-encoded genes (mt-Nd2, mt-Co1, mt-Atp6, and mt-Cytb) was induced in the GC muscle of MCK-miR-499 mice compared to NTG controls (Fig 2B). The expression of the oxidative biomarkers myoglobin and cytochrome c was also induced in the MCK-miR-499 GC muscles compared to NTG controls (Fig 2C). Moreover, we also found an increased expression of biomarker genes associated with fatty acid uptake (Cd36 and Lpl) in MCK-miR-499 muscle (Fig 2D). In addition, the expression of the glucose oxidation biomarker Ldhb (Gan et al., 2011) was increased, whereas Ldhα gene expression was decreased in MCK-miR-499 muscle (Fig 2D). The expected LDH isoenzyme activity shifts were confirmed by gel-activity studies (Fig 2E and F).

Consistent with the gene expression results, metachromatic myosin 1 ATPase staining confirmed a dramatic increase in type I muscle fibers in MCK-miR-499 GC muscle (NTG, 126 ± 42 per section versus MCK-miR-499, 551 ± 43 per section, n = 5 mice per group, P = 0.0006) (Fig 2G). To further evaluate the effect of miR-499 on muscle oxidative mitochondrial activity, we performed histochemical staining for succinate dehydrogenase (SDH) and α-glycerophosphate dehydrogenase (α-GPDH), which are hallmarks for oxidative and glycolytic metabolism, respectively (Schiappino & Reggiani, 2011). As expected, the SDH enzymatic activities were higher (NTG, 30.5 ± 2.0% versus MCK-miR-499, 58.7 ± 1.3%, n = 4 mice per group, P < 0.0001), whereas the α-GPDH activities were lower in the GC muscle of MCK-miR-499 mice compared to their NTG littermate controls (NTG, 74.8 ± 1.5% versus MCK-miR-499, 31.6 ± 1.4%, n = 5 mice per group, P < 0.0001) (Fig 2G). Together, these results demonstrate that miR-499 reprograms muscle for increased mitochondrial oxidative capacity, as well as fiber type.
Figure 1.

**A** Myh7b → miR-499 → Ex1 → Ex19 → Ex20

**B** OCR vs TIME

**C** Myh7b → miR-499 → Ex1 → Ex31 → Ex32

**D** OCR vs TIME

**E** Speed (m/min) vs Time (min)

**F** Blood lactate (mM) vs Time (months)

**G** Pyruvate/Malate

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miR-499 regulates muscle mitochondrial oxidative capacity through PGC-1α

We next sought to explore the targets and pathways downstream of miR-499 that are involved in regulation of muscle mitochondrial function. Using Ingenuity Pathways Analysis (http://www.ingenuity.com), we found that many of the genes upregulated in MCK-miR-499 muscle are known targets of PGC-1α (Fig EV1A), a master transcriptional co-regulator of mitochondrial function (Finck & Kelly, 2006; Handschin & Spiegelman, 2006). Real-time PCR confirmed that Ppargc1a (PGC-1α) mRNA levels were higher and Ppargc1b (PGC-1β) mRNA levels were lower in MCK-miR-499 muscle (Fig EV1B). Esrrb (ERRβ) mRNA was also modestly increased in MCK-miR-499 muscle, whereas Esrra (ERRα), Esrrg (ERRγ), Ppara (PPARγ), and Ppard (PPARδ) gene expressions were not induced in MCK-miR-499 muscle (Fig EV1B). Remarkably, PGC-1α protein levels were dramatically induced (14-fold) in the GC muscle of MCK-miR-499 mice compared to NTG controls (Fig 3A). Levels of PGC-1α protein were further compared across multiple muscle types in NTG and MCK-miR-499 mice. As shown in Fig EV1C, the protein levels of PGC-1α and myoglobin were also increased in WV muscle in MCK-miR-499 mice. Interestingly, there is no increase in mitochondrial DNA levels in MCK-miR-499 muscle compared with NTG controls (Fig EV1D), suggesting that PGC-1α affects mitochondrial respiration capacity without changes in mitochondrial content in MCK-miR-499 muscle.

To determine whether the muscle mitochondrial oxidative changes induced by miR-499 were due to the marked induction of PGC-1α protein, we bred MCK-miR-499 mice with muscle-specific PGC-1α KO (PGC-1α mKO) mice to generate 499Tg/PGC-1α KO mice, in which the PGC-1α gene is disrupted in muscle in a miR-499 transgenic background. Upon gross examination, disruption of PGC-1α markedly reduced the red coloration of MCK-miR-499 muscle (Fig EV2A and B). miR-499-mediated induction of oxidative biomarker genes (Ldhb, Cox5a, myoglobin, and cytochrome c) was significantly reduced in the absence of PGC-1α (Fig 3B and C). Furthermore, muscle-specific disruption of PGC-1α resulted in marked reduction of oxidative mitochondrial enzyme SDH activity in the GC muscle of MCK-miR-499 mice (499Tg, 56.9 ± 2.0% versus 499Tg/PGC-1α mKO, 35.4 ± 7.9%, n = 4 mice per group, P = 0.012) (Fig 3D). Interestingly, however, no change in GPDH staining and MHCI immunofluorescence and was observed in the GC muscle of 499Tg/PGC-1α mKO mice compared to MCK-miR-499 mice (GPDH: 499Tg, 34.3 ± 2.0% versus 499Tg/PGC-1α mKO, 41.0 ± 3.5%, n = 4 mice per group, P = 0.114) (MHCI: 499Tg, 347 ± 35 per section versus 499Tg/PGC-1α mKO, 463 ± 55 per section, n = 5 mice per group, P = 0.113) (Fig 3D). Similar observations were made in multiple muscle types of the 499Tg/PGC-1α mKO mice, in which the expression of slow-twitch muscle fiber genes was activated by miR-499 but not affected by disruption of PGC-1α in all soleus, GC, and WV muscle (Fig EV2C). Together, these results demonstrate that PGC-1α is required for the miR-499-mediated increase in mitochondrial oxidative capacity in muscle but not fiber type. It is possible that Sox6 inhibition (Fig EV2C) acts in parallel with PGC-1α signaling, thereby inducing slow myofiber formation in MCK-miR-499 mice.

We also assessed the physiological effect of PGC-1α gene disruption in MCK-miR-499 mice. As shown in Fig 3E, disruption of PGC-1α abolished the miR-499-mediated enhancement of exercise capacity and the RER-lowering effect during exercise. In addition, muscle-specific disruption of PGC-1α prevented the miR-499-mediated reduction of blood lactate levels following exercise (Fig 3F). Together, these results indicate that PGC-1α is required for the miR-499-mediated increase in oxidative metabolism in muscle.

miR-499 directly targets Fnin1, an AMPK interacting protein that negatively regulates AMPK-PGC-1α signaling and mitochondrial function

We next sought to determine the mechanism whereby PGC-1α-dependent mitochondrial function was induced in the MCK-miR-499 muscle. miRNAs inhibit mRNA translation or promote mRNA degradation through putative basepair interactions with target mRNAs (Bartel, 2004, 2009; Chen et al, 2009; Mendell & Olson, 2012). Thus, bioinformatics prediction combined with 3’UTR reporter assays were used to identify putative target mRNAs for miR-499 (Fig EV3). The TargetScan and MicroCosm programs were first used to identify putative target mRNAs for miR-499, and this gene list was next cross-matched for genes that were downregulated in MCK-miR-499 muscle (Fig EV3 and Appendix Table S3). Putative targets were then further selected for additional 3’UTR reporter validation assays.
Figure 2. MCK-miR-499 muscle is reprogrammed for increased capacity for mitochondrial oxidation.

A Gene ontology (GO) enrichment analysis of gene transcripts upregulated in MCK-miR-499 muscle identified a number of terms related to mitochondrial function and muscle contraction. Gene expression array data generated from the gastrocnemius muscle of 8-week-old male MCK-miR-499 mice were compared to littermate controls (NTG).

B Expression of the nuclear-encoded and mitochondrial-encoded genes (RT-qPCR) in the gastrocnemius muscle from the indicated genotypes (n = 5 mice per group). *P < 0.05.

C Representative Western blot analysis performed with gastrocnemius muscle total protein extracts prepared from the indicated mice using myoglobin, cytochrome c, and β-actin (control) antibodies (n = 8 mice per group).

D Expression of genes (RT-qPCR) involved in muscle glucose and fatty acid metabolism in the gastrocnemius muscle from the indicated genotypes (n = 5 mice per group). *P < 0.05.

E LDH isoenzymes were separated by polyacrylamide gel electrophoresis using whole cell extracts from NTG heart (Ht, control) and gastrocnemius muscle from the indicated mice. A representative gel showing 3–4 independent mice per group is shown.

F Quantification of LDH isoenzyme activity gel electrophoresis in (E). Values represent the mean % (± SEM) total LDH activity. *P < 0.05.

G Cross section of the gastrocnemius muscle from 3-month-old male NTG and MCK-miR-499 mice stained for myosin I ATPase activity (top), SDH (middle), and α-GPDH (bottom). Representative images are shown. Scale bar: 500 μm.

Data information: All values represent the mean ± SEM. P-value was determined using two-tailed unpaired Student’s t-test. Source data are available online for this figure.
Figure 3.

A. GC

B. Normalized AU

C. Cyt C/Tubulin

D. SDH

E. PGC-1α mKO

F. Blood lactate (mM)

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(Fig EV3). This analysis revealed that the transcript encoding folliculin-interacting protein-1 (Fnip1) is a potential direct target of miR-499. This finding was of interest because Fnip1 was originally identified as an AMPK-interacting protein (Baba et al, 2006), suggesting a role in the AMPK–PGC-1α regulatory pathway. In addition, mice with genetic deletion of Fnip1 have recently been shown to increase PGC-1α signaling and oxidative muscle fiber composition (Hasumi et al, 2015; Reyes et al, 2015), a phenotype that substantially recapitulates that of MCK-miR-499 mice. Thus, additional studies were focused on Fnip1.

As shown in Fig 4A, the miR-499 binding sequences in the Fnip1 3’UTR are highly conserved in mammals. miR-499 repressed an Fnip1 3’UTR reporter via the conserved miR-499 binding site in HEK293T cells (Fig 4B). In addition, expression of Fnip1, but not the related Fnip2, was decreased in MCK-miR-499 GC muscle compared to NTG controls (Fig 4C). Fnip1 protein levels were dramatically decreased in MCK-miR-499 GC muscle compared to NTG controls (Fig 4D). These results demonstrate that miR-499 directly targets Fnip1.

To directly determine the role of Fnip1 in miR-499-mediated induction of mitochondrial function, loss-of-function experiments were conducted in primary myotubes. siRNA-mediated knockdown of Fnip1 resulted in a marked increase in phosphorylated AMPKα (p-AMPKα) levels (Fig 5A). In addition, levels of PGC-1α transcript were increased in myotubes treated with Fnip1 siRNA (Fig 5B). To determine whether AMPK signaling is involved in this mechanism, the effect of AMPK inhibitor compound C was assessed together in the presence of Fnip1 siRNA-mediated knockdown. As shown in Fig 5C, AMPK inhibition abrogated the Fnip1 siRNA-mediated induction of Ppargc1a mRNA. These results further establish the importance of AMPK in this mechanism, which was consistent with increased AMPK activity (Fig 5D), and elevated PGC-1α levels (Fig 5A) in MCK-miR-499 muscle. Oxygen consumption rates (OCR) were measured in skeletal myotubes following knockdown of Fnip1 expression. Fnip1 inhibition significantly stimulated OCR in the presence of the uncoupler FCCP, a sign of enhanced mitochondrial function (Fig 5E). These results demonstrate that Fnip1 is a negative regulator of mitochondrial function in myocyte.

To assess the requisite role of miR-499 in the control of mitochondrial function in the absence of overexpression and determine whether Fnip1 is involved in this mechanism, miR-499 and Fnip1 loss-of-function studies were conducted in wild-type (WT) mouse primary skeletal myotubes. Because miR-499 and miR-208b function redundant, the impact of antisense-mediated inhibition of both miR-499 and miR-208b on Ppargc1a and Fnip1 gene expression was evaluated in WT myotubes. Consistent with miR-499 suppression of Fnip1, inhibition of both miRNAs modestly reduced Ppargc1a mRNA levels and, conversely, induced Fnip1 mRNA levels (Fig 5F). The effects of miR-499/miR-208b inhibition on myocyte OCR were next assessed alone and together with the presence of Fnip1 siRNA. As shown in Fig 5G, inhibition of both miRNAs resulted in significant decrease in maximal OCR in the presence of the uncoupler FCCP, and Fnip1 knockdown prevented the miR-499/miR-208b inhibition-mediated repression of myocyte OCR. These results further establish the relevance of miR-499-mediated regulation of mitochondrial function and demonstrate the importance of Fnip1 suppression in this mechanism. Together, our data support a model in which miR-499 directly targets Fnip1, activating AMPK–PGC-1α signaling, and thereby driving increased muscle mitochondrial capacity.

**Figure 3.** miR-499 regulates muscle mitochondrial oxidative capacity through PGC-1α.

A (Left) Representative Western blot analysis of PGC-1α protein expression in the gastrocnemius muscle from the indicated genotypes, heart (Ht) lysate from WT mice and PGC-1α mKO (MCK-Cre) mice as positive and negative controls, respectively. (Right) Quantification of the PGC-1α/tubulin signal ratios normalized (= 1.0) to the NTG control (n = 5 mice per group). *P < 0.001 versus NTG, †P < 0.001 (versus 499Tg).

B Expression of the Ldhb, Ldhα, Ms (myoglobin), and Cox5a genes (RT-qPCR) in the gastrocnemius muscle from the indicated genotypes (n = 5 mice per group). *P < 0.01 versus NTG, †P < 0.01 (versus 499Tg).

C (Left) Representative Western blot analysis performed with gastrocnemius muscle total protein extracts prepared from the indicated mice using cytochrome c, myoglobin, and α-tubulin (control) antibodies. (Right) Quantification of the Cyct/tubulin signal ratios normalized (= 1.0) to the NTG control. NTG, n = 5, 499Tg, n = 6, PGC-1α mKO, n = 5, 499Tg/PGC-1α mKO, n = 5. *P < 0.01 versus NTG, †P = 0.00018 (versus 499Tg).

D Cross section of gastrocnemius muscle from the indicated mice stained for SDH activity (top) as well as MHC immunofluorescence (IF; bottom); representative images are shown. Scale bar: 500 μm.

E Respiratory exchange ratio (RER) during a graded exercise regimen as described in Materials and Methods. Notably, no significant increase in exercise speed was observed in 499Tg/PGC-1α mKO mice compared to PGC-1α mKO mice. PGC-1α mKO, n = 5, 499Tg/PGC-1α mKO, n = 8. The bars represent the mean blood lactate levels from the indicated mice following a 25-min run on a motorized treadmill. NTG, n = 9, 499Tg, n = 7; PGC-1α mKO, n = 6, 499Tg/PGC-1α mKO, n = 9. *P = 0.0009 (NTG versus 499Tg), P = 0.316 (NS, not significant).

Data information: All values represent the mean ± SEM. P-value in (A and F) was determined using two-tailed unpaired Student’s t-test; P-value in (B and C) was determined using one-way ANOVA coupled to a Fisher’s least-significant difference (LSD) post hoc test.

Source data are available online for this figure.

Restoring the expression of miR-499 activates the slow-oxidative muscle fiber program and ameliorates muscular dystrophy in mdx mice

Fast-glycolytic myofibers are more sensitive to the dystrophic pathology in Duchenne muscular dystrophy (DMD) patients as well as in the preclinical mdx mouse model of DMD (Webster et al, 1988; Kuznetsov et al, 1998; Timmons et al, 2005; Ljubicic et al, 2014). Accordingly, promoting the slow-oxidative muscle fiber program was proposed as a strategy to protect against DMD (Ljubicic et al, 2014). Therefore, we examined the regulation of the miR-499 circuit in mdx mice, a disease model of DMD that is known to have less slow-twitch myofibers (Jeanson-Leh et al, 2014; Ljubicic et al, 2014). As expected, the levels of miR-499 and miR-208b were significantly downregulated in mdx muscle compared to WT controls (Fig 6A). The diminished mitochondrial respiration capacity in mdx muscle was associated with decreased expression of miR-499 and miR-208b (Fig 6A and B). These results are consistent with miR-499 regulating mitochondrial function.
We next determined whether transgenic restoration of the expression of miR-499 could reduce the severity of the dystrophic pathology in mdx mice. MCK-miR-499 mice were bred with the mdx mice to obtain mdx and mdx/499Tg mice. Whereas no difference in PGC-1α protein levels was observed in the GC muscle of mdx mice compared to WT controls, probably due to very low basal PGC-1α levels, activation of miR-499 in mdx muscle resulted in dramatic increase in expression of PGC-1α (Fig 6C). Fnip1 protein showed a trend toward increase in mdx muscle but significantly suppressed in mdx/499Tg muscle (Fig 6C). Interestingly, we found an increase in phosphorylated AMPKα (p-AMPKα) levels in mdx muscle and this is reversed by miR-499 activation (Fig EV4A). The activation of p-AMPKα is consistent with cellular sensing of relative energetic deficiency within mdx dystrophic myofibers, the correction of AMPK activity in mdx/miR-499Tg muscle could also reflect a restored energy production capacity. The expression of myoglobin that was repressed in mdx mice was completely restored by miR-499 overexpression (Fig 6D). In addition, miR-499 activation also reversed oxidative biomarkers such as Ldhb expression in mdx muscle. The expected LDH isoenzyme activity shifts were confirmed.

Figure 4. miR-499 directly targets Fnip1.
A The schematic shows the putative conserved miR-499 binding site within the 3’UTR of the Fnip1 gene.
B Luciferase reporters containing the wild-type Fnip1 3’UTR or Fnip1 3’UTR mutated in the predicted binding site of miR-499 were used in cotransfection studies in HEK293T cells in the presence or absence of plasmids expressing miR-499 (n = 3 independent experiments). *P < 0.0001 (Fnip1 3’UTR); *P = 0.023 (Mutated).
C RT-qPCR analysis of Fnip1 and Fnip2 mRNA levels in the gastrocnemius muscle of the indicated genotypes (n = 5 mice per group).
D (Top) Fnip1 protein expression in the gastrocnemius muscle from the indicated mice. (Bottom) Quantification of the Fnip1/tubulin signal ratio (n = 4 mice per group). *P = 0.014.

Data information: All values represent the mean ± SEM. P-value was determined using two-tailed unpaired Student’s t-test.
Source data are available online for this figure.
by activity gel studies (Fig EV4B and C). As predicted, mitochondrial respiration capacity was significantly increased in mdx/miR-499 muscle compared to mdx mice (Fig EV4D). Interestingly, the mitochondrial DNA levels remains unchanged in mdx/miR-499 muscle compared to mdx mice (Fig EV4E). MHC1 immunostaining demonstrated that miR-499 activation reversed the loss of type I
mice contain higher numbers of centralized nuclei. Transgenic induc- 
fibers compared to mdx alone (Fig EV5A). Muscle fibers from mdx 
muscle fiber program in mdx mice. 

We next characterized the dystrophic phenotype of the mdx/ 
499Tg mice. Mdx mice had a higher variation in fiber size and a 
higher number of small fibers compared to WT mice. The mdx/miR-
499Tg mice had a lower variation in fiber size, and had fewer small 
fibers compared to mdx alone (Fig EV5A). Muscle fibers from mdx 
mice contain higher numbers of centralized nuclei. Transgenic induc-
tion of miR-499 in mdx mice resulted in a significant reduction in the 
percentage of centrally located nuclei (mdx, 55.0 ± 3.4%; mdx/
499Tg, 42.1 ± 3.3%; n = 5 mice per group, P = 0.031) (Figs 6E and 
EV5B). Evans blue dye infiltration test also showed a decrease in 
damaged myofibers in tibialis anterior (TA) muscle of mdx/499Tg 
compared with mdx mice (Fig EV5C). Additionally, we also 
measured the levels of serum creatine kinase (CK), another hallmark 
of damaged muscles, in young WT, mdx, and mdx/499Tg mice. As 
eXpected, we observed a marked increase in CK levels in mdx mice 
compared to WT controls. Remarkably, transgenic induction of 
miR-499 in mdx mice resulted in a dramatic reduction of CK levels 
(mdx, 18,754 ± 4,330 U/l; mdx/499Tg, 5,832 ± 474 U/l; mdx, 
n = 10; mdx/499Tg, n = 14; P = 0.0003) (Fig 6F).

To assess muscle functionality and performance, we challenged 
mdx and mdx/499Tg mice with treadmill exercise. The mdx mice 
rain on average 475 m, and the mdx/499Tg mice ran signiﬁcantly 
further, 1,307 m, a distance indistinguishable from that run by WT 
controls (Fig 6G). We further conducted real-time RER measure-
ments to evaluate the muscle fuel utilization of 499Tg/mdx mice 
during exercise. In mdx mice, a rapid increase in RER values 
ocurred at a very low speed during the exercise challenge, which is 
consistent with a marked decrease in exercise capacity (Fig 6H). 
Conversely, 499Tg/mdx mice were able to exercise at much higher 
speeds before an increase in RER values was observed (Fig 6H), 
suggesting that 499Tg/mdx mice restored the capacity to utilize fatty 
acid oxidation during exercise. Together, these results strongly 
suggest that re-activation of miR-499 in mdx muscle signiﬁcantly 
Improves the pathophysiology of the muscular dystrophic 
phenotype.

Figure 5. Inhibition of Fnip1 reactivated AMPK-PGC-1α signaling and mitochondrial function in myocytes.

A (Left) Representative Western blot analysis performed on extracts of myotubes subjected to Fnip1 siRNA or control (Con) siRNA using phospho-AMPKα (Thr172) and AMPKα antibodies. (Right) Quantiﬁcation of the p-AMPKα/AMPKα signal ratios (n = 3 independent experiments). *P = 0.037.

B, C Results of RT-qPCR analysis performed on extracts of myotubes subjected to Fnip1 siRNA or scrambled Con siRNA as indicated. For (C), 48 h post-siRNA transfection, myotubes were treated for 24 h with DMSO or 10 μm compound C before harvest (n = 3 independent experiments). *P = 0.0001 (Fnip1 in B).

D (Top) Representative Western blot analysis performed on extracts of the gastrocnemius muscle isolated from NTG or MCK-miR-499 mice using phospho-AMPKα (Thr172), AMPKα, phospho-ACC (Ser79), or total ACC antibodies. (Bottom) Quantiﬁcation of the p-AMPKα/AMPKα and p-ACC/ACC signal ratios normalized (= 1.0) to the NTG control (n = 8 mice per group). *P < 0.01.

E Oxygen consumption rates (OCR) in primary mouse myotubes transfected with Fnip1 siRNA or Con siRNA. n = 7 separate experiments done with 5 biological replicates. *P < 0.05 (Pyruvate), *P < 0.01 (FCCP).

F Results of RT-qPCR analysis for WT primary mouse myotubes subjected to inhibition of both miR-499 and miR-208b (anti-miRs) (n = 4 independent experiments). *P = 0.037 (Fnip1).

Oxygen consumption rates (OCR) in primary mouse myotubes transfected with miR-499/miR-208b inhibitors alone and together with the presence of Fnip1 siRNA. n = 4 separate experiments done with 5 biological replicates. *P < 0.01 (anti-miRs versus Control).

Data information: All values represent the mean ± SEM. P-value in (A, B, D) was determined using two-tailed unpaired Student’s t-test; P-value in (C and G) was determined using one-way ANOVA coupled to a Fisher’s least-significant difference (LSD) post hoc test.

Source data are available online for this figure.

Figure 6. Restoring the expression of miR-499 activates the slow-oxidative muscle ﬁber program and ameliorates muscular dystrophy in mdx mice.

A Mean expression levels (RT-qPCR) in gastrocnemius muscle of 8-week-old male WT and mdx mice (n = 5 mice per group). *P = 0.009 (Myh7b), *P = 0.0007 (miR-499), *P = 0.0143 (Myh7), *P < 0.0011 (miR-208B).

B Mitochondrial respiration rates were determined from mitochondria isolated from the hindlimb muscle of the indicated mice using pyruvate as substrate. n = 3 separate experiments done with 7–8 biological replicates. *P < 0.01 (ADP).

C (Left) Representative Western blot analysis of PGC-1α (Top) and Fnip1 (Bottom) protein expression in the gastrocnemius muscle from the indicated genotypes with 

Figure 8. fnip1 transcript expression in skeletal muscle.

A (Top) Representative MHC1 immunofluorescence (IF) in the soleus of the indicated genotypes. Scale bar: 500 μm. (Bottom) Representative HE staining of the gastrocnemius muscle of the indicated genotypes. Scale bar: 100 μm.

B Five-week-old male WT, mdx, and mdx/MCK-miR-499 mice were euthanized, and serum creatine kinase activity was determined. WT, n = 10; mdx, n = 10; mdx/499Tg, n = 14; *P < 0.0001 (versus WT), *P = 0.0003 (versus mdx).

C The bars represent the mean running time and distance (± SEM) for 8-week-old male mice on a motorized treadmill. WT, n = 10; mdx, n = 13; mdx/499Tg, n = 8. *P = 0.01406 (Running time, versus WT), *P = 0.005756 (Running time, versus mdx). *P = 0.004558 (Running distance, versus WT), *P = 0.001668 (Running distance, versus mdx).

H Respiratory exchange ratio (RER) during a graded exercise regimen on a motorized treadmill. mdx, n = 8; mdx/499Tg, n = 6. *P < 0.05.

Data information: All values represent the mean ± SEM. P-value in (A, B and H) was determined using two-tailed unpaired Student’s t-test; P-value in (C, F and G) was determined using one-way ANOVA coupled to a Fisher’s least-significant difference (LSD) post hoc test.

Source data are available online for this figure.
Figure 6.
Discussion

Delineation of the mechanisms involved in the coordinate regulation of muscle metabolic and structural programs during fiber type transition has implications for new therapeutic approaches for many human diseases, including metabolic disorders and muscular dystrophy. Previous studies have established that nuclear receptors such as PPARs and ERRs, along with their co-regulators PGC-1α and AMPK are key regulators of myocytes energy metabolism (Lin et al., 2002; Wang et al., 2004; Arany et al., 2005; Schuler et al., 2006; Narkar et al., 2008, 2011; Rangwala et al., 2010; Zechner et al., 2010; Gan et al., 2011, 2013). Herein, we discover a novel mechanism for muscle contractile property tightly coupled to its metabolic capacity during fiber type transition. Specifically, the myosin Myh7b gene encodes miR-499, which directly inhibits Fnip1, leading to activation of AMPK-PGC-1α signaling and thereby triggering a muscle mitochondrial oxidative metabolism program. We therefore propose a model for the adaptive mitochondrial function during muscle fiber type transition via the miR-499/Fnip1/AMPK circuit (Fig 7). This mechanism likely represents a general paradigm for efficiently couple cellular energy consumption with ATP production under an array of diverse physiological and pathophysiological circumstances.

Nuclear receptor ERRγ was recently shown to drive a slow-oxidative muscle fiber program in skeletal muscle, including increased fatty acid oxidation, mitochondrial oxidative phosphorylation, and slow-twitch muscle fiber composition (Rangwala et al., 2010; Narkar et al., 2011; Gan et al., 2013). We have recently found that ERRγ directly activates the promoters driving the expression of miR-499 and miR-208b in muscle (Gan et al., 2013). In the present study, our data suggest that miR-499 is a regulator of mitochondrial function that acts downstream of the ERRγ signaling. Exercise training induces the expression of ERRγ as well as miR-499 in muscle (Gan et al., 2013). It is likely that the miR-499/Fnip1/AMPK circuit serves as a major nexus between slow myofiber stimuli and the adaptive mitochondrial function, such as exercise training in which the transcription changes in contractile myosin genes parallels the changes in muscle metabolic properties (Kirschbaum et al., 1990; Schiaffino & Reggiani, 2011). Conversely, chronic immobilization and disuse is predicted to have the opposite effect.

We found that miR-499 regulates muscle metabolism by directly targeting Fnip1. The observed role of Fnip1 is of interest given its role in regulating muscle metabolism (Baba et al., 2006; Hasumi et al., 2015; Reyes et al., 2015). Notably, knockout of the gene encoding folliculin, an Fnip1 interacting protein, has also been shown to induce slow-oxidative muscle transformation (Hasumi et al., 2012), which further emphasizes the importance of Fnip1-related signaling in muscle metabolism. Our data strongly suggest that the major effect of miR-499 on muscle energy metabolism might be mediated through inhibition of Fnip1 protein. First, we found that Fnip1 mRNA and protein expression in muscle were markedly suppressed in MCK-miR-499 mice. Second, Fnip1 null mice recapitulated many of the

![Figure 7. A model for the adaptive mitochondrial function during muscle fiber type transition.](https://example.com/figure7.png)

The schematic depicts a proposed model for the miR-499/Fnip1/AMPK circuit that orchestrates mitochondrial function to match muscle contractile machinery during fiber type transition.
metabolic phenotypes observed in the MCK-miR-499 mice, including enhanced red coloration, increased capacity for SDH activity and mitochondrial respiration. Third, key regulators of aerobic metabolism such as AMPK/PGC-1α and oxidative biomarkers induced in MCK-miR-499 muscle were similarly activated in Fnip1 knockout mouse. Fourth, direct manipulation of Fnip1 in primary myotubes mimicked the effects of miR-499 on AMPK/PGC-1α levels and mitochondrial respiration in muscle. However, unlike Fnip1 (Reyes et al., 2015), an interesting finding of our study is that miR-499 drives an increase in slow-twitch muscle fiber proportion in a PGC-1α-independent manner. It is possible that Sox6, another direct target of miR-499, acts in parallel with Fnip1/PGC-1α signaling to induce slow myofiber formation in MCK-miR-499 muscle. Consistent with this notion, a recent study has concluded that Sox6 acts independently of PGC-1α on muscle fiber type switching in mice (Quiat et al., 2011). Interestingly, we also found evidence that miR-499 activates glucose oxidation in muscle. The Ldhb/Ldha gene expression ratio is increased in MCK-miR-499 muscle. These results were of interest because the Ldhb isoenzyme favors the reaction that converts lactate to pyruvate which, in turn, provides substrate for the mitochondrial TCA cycle, whereas Ldha favors the reverse reaction to produce lactate from pyruvate. Therefore, miR-499 drives an LDH isoenzyme shift that diverts pyruvate into the mitochondrion to fully catabolize glucose for maximal ATP production, contributing to enhanced exercise capacity and mitochondrial activity, and decreased post-exercise blood lactate in MCK-miR-499 mice.

Fast-glycolytic myofibers are more sensitive to the dystrophic pathology in Duchenne muscular dystrophy (DMD) patients as well as in mdx mice (Webster et al., 1988; Kuznetsov et al., 1998; Timmons et al., 2005; Lubicic et al., 2014). In this study, we found that downregulating the miR-499 circuit in the mdx muscle diminished the mitochondrial function in parallel. Based on our data, it is tempting to speculate that downregulated miR-499 signaling contributes to a vicious cycle, further promoting the progression of muscular dystrophy. Direct manipulation of miR-499 is able to restore the slow-oxidative muscle fiber program and prevent the hallmarks of DMD, including serum CK and exercise capacity. Interestingly, a recent study has concluded that Sox6 acts independently of PGC-1α on muscle fiber type switching in mice (Quiat et al., 2011). Interestingly, we also found evidence that miR-499 activates glucose oxidation in muscle. The Ldhb/Ldha gene expression ratio is increased in MCK-miR-499 muscle. These results were of interest because the Ldhb isoenzyme favors the reaction that converts lactate to pyruvate which, in turn, provides substrate for the mitochondrial TCA cycle, whereas Ldha favors the reverse reaction to produce lactate from pyruvate. Therefore, miR-499 drives an LDH isoenzyme shift that diverts pyruvate into the mitochondrion to fully catabolize glucose for maximal ATP production, contributing to enhanced exercise capacity and mitochondrial activity, and decreased post-exercise blood lactate in MCK-miR-499 mice.

Materials and Methods

Animal studies

All animal studies were conducted in strict accordance with the institutional guidelines for the humane treatment of animals and were approved by the IACUC committees at the Model Animal Research Center (MARC) of Nanjing University. MCK-miR-499 mice (van Rooij et al., 2009) originally generated at UT Southwestern were back-crossed to the C57BL/6J background for more than six generations. To generate mice with a muscle-specific disruption of the PGC-1α allele in MCK-miR-499 mice (herein named 499Tg/PGC-1α mKO), PGC-1α f/f mice (Jackson Laboratory, stock no. 009666) were crossed with mice expressing Cre under control of the muscle creatine kinase (MCK) promoter (Jackson Laboratory, stock no. 006475) to achieve muscle-specific deletion of PGC-1α, and these mice were then crossed with MCK-miR-499 mice to obtain 499Tg/PGC-1α mKO mice. The mdx mice were purchased from the Jackson Laboratory (stock no. 000476), and male MCK-miR-499 mice were bred with homozygous female mdx mice to generate 499Tg/mdx mice. Male offspring were genotyped and categorized in mdx and 499Tg/mdx mice, and mice at the age of 5–9 weeks were used. Littermate controls were used in all cases. Mice were randomly assigned to various analyses. Investigators involved in the histological analysis were blinded. Investigators performing animal handling, sampling, and raw data collection were not blinded.

Mitochondrial respiration studies

Mitochondrial respiration rates were measured in saponin-permeabilized plantaris or extensor digital longus muscle fibers with pyruvate/malate as substrates as described previously (Saks et al., 1998; Coen et al., 2013; Lai et al., 2014). In brief, the muscle fibers were separated and transferred to BIOPS buffer (7.23 mM K2EGTA, 2.77 mM CaK2EGTA, 20 mM imidazole, 20 mM taurine, 50 mM potassium 2-[N-morpholino]-ethanesulfonic acid, 0.5 mM dithiothreitol, 6.56 mM MgCl2, 5.7 mM ATP, and 14.3 mM phosphocreatine [PCr], pH 7.1). The muscle fiber bundles were then permeabilized with 50 μg/ml saponin in BIOPS solution. Measurement of oxygen consumption in permeabilized muscle fibers was performed in buffer Z (105 mM potassium 2-[N-morpholino]-ethanesulfonic acid, 30 mM KCl, 10 mM KH2PO4, 5 mM MgCl2, 5 mg/ml BSA, 1 mM EGTA, pH7.4) at 37°C and in the oxygen concentration range 220–150 nmol O2/ml in the respiration chambers of an Oxygraph 2K (Oroboros Inc., Innsbruck, Austria). Following measurement of basal, pyruvate (10 mM)/malate (5 mM) respiration, maximal (ADP-stimulated) respiration was determined by exposing the mitochondria to 4 mM ADP. Uncoupled respiration was evaluated following addition of oligomycin (1 μg/ml). Respiration rates were determined and normalized to fiber bundle dry weight or wet weight using Datlab 3 software (Oroboros Inc., Innsbruck, Austria), and the data were expressed as “pmol O2/s/mg dry weight” or “pmol O2/s/mg wet weight”.

Skeletal muscle mitochondria were isolated from hindlimbs of WT and mdx mice as previously described (Zechner et al., 2010). Total protein was quantified by a BCA assay (Thermo Scientific),
and respiration rates of the isolated mitochondria containing 4 \( \mu \)g protein/well were measured using a XF24 analyzer (Seahorse Bioscience Inc.) per the manufacturer’s protocol. Respiration was determined using pyruvate as substrate. Following measurement of the basal respiration, maximal ADP-stimulated respiration was determined by exposing mitochondria to 4 mM ADP. Uncoupled respiration was evaluated following the addition of oligomycin (1 \( \mu \)M) to inhibit ATP synthase; this was followed by addition of the uncoupler FCCP (2 \( \mu \)M), and then the addition of antimycin A (0.2 \( \mu \)M). Respiration rates were expressed as “OCR pmoles/min 4 \( \mu \)g protein”.

Exercise studies

Mice were acclimated (run for 9 min at 10 m/min followed by 1 min at 20 m/min) to the treadmill for 2 consecutive days prior to the experimental protocol. Endurance exercise was determined as described previously (Gan et al, 2013). In brief, fed mice were run for 10 min at 10 m/min followed by a constant speed of 20 m/min until exhaustion. Tail blood was taken after exercise and measured for lactate (Lactate Scout, Senelab, Germany) according to manufacturer’s instruction.

Respiratory exchange ratios (RER) during exercise were determined as described previously (Gan et al, 2011). Briefly, mice were placed in an enclosed treadmill attached to the Comprehensive Laboratory Animal Monitoring System (CLAMS) (Columbus Instruments) for 15 min at a 0° incline and 0 m/min. The mice were then challenged with two-min intervals of increasing speed at a 0° incline. The increasing speeds used in the protocol were 10, 14, 18, 22, 26, 28, 30, 32, 34, 36, 38, 40, 42, 44, and 46 m/min. Measurements were collected before the exercise challenge, throughout the challenge, and following failure. For the mdx and mdx/miR-499Tg mice running capacity test, the increasing speeds used in the protocol were 10, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, and 25 m/min.

Histological analyses

Muscle tissue was frozen in isopentane that had been cooled in liquid nitrogen. MHC ATPase and immunofluorescence (IF) stains were conducted as previously described (Zechner et al, 2010; Gan et al, 2013). MHC ATPase stains were performed at pH 4.31. Under these acidic conditions, MHC2 isoforms are inactivated whereas MHC1 is still functional, resulting in the addition of black dye to MHC1-positive muscle fibers. For IF stains, the muscle fibers were stained with antibodies directed against MHC1 (BA-D5) (MHC1, green; All fast MHC2, black [unstained]). H&E, SDH, and \( \alpha \)-GPDH staining were performed as previously described (Zechner et al, 2010). Quantification of histological staining was performed in a blinded manner with Image-Pro Plus software.

Serum creatine kinase assay

Mouse blood was collected, and the serum was isolated using heparin-coated collection tubes. Serum creatine kinase activity was then determined in a blinded fashion with a Beckman Coulter AU5400 using the Creatine Kinase Assay Kit (Beckman Coulter) according to the manufacturer’s protocol.

Gene expression array studies

Total RNA isolated from the gastrocnemius muscle of 8-week-old male MCK-miR-499 and NTG littermate mice was used for gene expression array studies performed as previously described using an Affymetrix GeneChip Mouse Gene 1.0 ST array (Lai et al, 2014). The Affymetrix probe level data were processed using Robust Multi-Array Analysis (RMA algorithm using the Expression Console software Version 1.1 default settings) to obtain normalized summary scores of expression for each probe set on each array. Partek analysis using the ANOVA was used to identify genes differentially expressed in two conditions. Three independent samples per group were analyzed. The criteria for a regulated gene were a fold change greater than 1.3 (either direction) and a significant P-value (\(< 0.05\)) versus NTG. For pathway analysis, the filtered data sets (a total of 1,002 upregulated genes) were uploaded into DAVID Bioinformatics Resources 6.7 to review the bio pathways using the Gene Ontology database. The functional annotation analysis was used to interpret data, and the pathways were reviewed using the cellular compartments defined by Gene Ontology, which is ranked by P-value of a regulated term in Fig 2A. The gene expression arrays have been deposited in the NCBI Gene Expression Omnibus and are accessible through GEO Series accession number GSE72581.

RNA analyses

Quantitative RT–PCR was performed as described previously, with modifications (Gan et al, 2011, 2013). Briefly, total RNA was extracted from mouse muscle or primary myotubes using RNAiso Plus (Takara Bio). Isolated total RNA integrity was electrothermally verified by ethidium bromide staining. About 1 \( \mu \)g total RNA samples were then reverse transcribed with the PrimeScript RT Reagent Kit with gDNA Eraser (Takara Bio) using random hexamer primers according to the manufacturer’s instructions. Real-time quantitative qPCR was performed using the ABI Prism Step-One system with SYBR\textsuperscript{®} Premix Ex Taq\textsuperscript{TM} (Takara Bio). A four-step experimental run protocol was used: (i) denaturation program (30 s at 95°C); (ii) amplification and quantification program repeated 40 times (5 s at 95°C; 34 s at 60°C); (iii) melting curve program (60–95°C with a heating rate of 0.3°C per 15 s and a continuous fluorescence measurement); (iv) and cooling program down to 40°C. Specific oligonucleotide primers for target gene sequences are listed in Appendix Table S4. Arbitrary units of target mRNA were corrected to the expression of 36b4.

Mitochondrial DNA analyses

Genomic/mitochondrial DNA was isolated using the RNAiso Plus (Takara Bio), followed by back extraction with 4 M guanidine thiocyanate, 50 mM sodium citrate, and 1 M Tris and an alcohol precipitation. Mitochondrial DNA content was determined by SYBR Green analysis (Takara Bio). The levels of NADH dehydrogenase subunit 1 (mitochondrial DNA) were normalized to the levels of lipoprotein
lipase (genomic DNA). The primer sequences are noted in Appendix Table S4.

**TaqMan miRNA**

TaqMan methods were used to assess miRNA expression as previously described (Gan et al., 2013). The TaqMan MicroRNA Assays were specific for each miRNA (Assay ID: hsa-miR-208b, 2290; mmu-miR-499, 1352) or for the internal controls (Assay ID: snoRNA202, 1232) (Applied Biosystems). Thermal cycling was performed on an ABI Prism Step-One system. The relative miRNA level was corrected to snoRNA202.

**Antibodies and immunoblotting studies**

Antibodies directed against MHC1 (BA-D5) (6 μg/ml) were purchased from the Developmental Studies Hybridoma Bank; antibodies directed against p-AMPK (Thr172) (2535) (1:1,000), AMPKα (5831) (1:1,000), p-ACC (11818) (1:1,000), and ACC (3676) (1:1,000) were all from Cell Signaling; anti-Fnip1 antibody (ab61395) (1:1,000) was purchased from Abcam, anti-cytochrome c (5831) (1:1,000), p-ACC (11818) (1:1,000), and ACC (3676) (1:1,000) were all from Cell Signaling; anti-Fnip1 antibody (ab61395) (1:1,000) was purchased from Abcam, anti-cytochrome c (bs1089) (1:1000) and anti-alpha-tubulin (bs1699) (1:5,000) antibody were from BioWorld, anti-myoglobin (sc-25607) (1:1,000) and anti-beta-actin (sc-47778) (1:5,000) were from Santa Cruz, and anti-PGC-1alpha (sc-80184) (1:1,000) was developed in the laboratory of Daniel Kelly as previously described (Leone et al., 2005). Western blotting studies were performed as previously described (Gan et al., 2011, 2013).

**LDH isoenzyme analysis**

LDH isoenzyme patterns were determined as previously described (Gan et al., 2011). Protein extracted from mouse hearts served as the positive control.

**Cell culture and RNAi experiments**

Primary muscle cells were isolated from skeletal muscles as previously described (Gan et al., 2011, 2013). For differentiation, cells were washed with PBS and re-fed with 2% horse-serum/DMEM differentiation medium and re-fed daily. siRNAs (Genepharma) targeting mouse Fnip1 (siRNA pool, #1: 5’-GGAGAUAGUUCUUCUCUCUUTT; #2 5’-GCAGGUACAGCAACCCAATT; #3 5’-GGUGGCUA CUGCAUCUUUTT) and microRNA inhibitors (Genepharma, anti-miR-499: AAACAUACUCUGAAGCUUUA; anti-miR-208b: ACAAC CCUUUUGUUCUCUAU) were transfected into primary myoblasts at a final concentration of 50 nM using Lipofectamine 2000 transfection reagent (Invitrogen) according to the manufacturer’s instructions. Cells were then differentiated for 3 days prior to harvest. For AMPK inhibition study, 48 h post-siRNA transfection, myotubes were treated for 24 h with DMSO or 10 μM compound C (Selleckchem).

**Oxygen consumption measurements**

Cellular oxygen consumption rates (OCR) were measured using the XF24 analyzer (Seahorse Bioscience Inc.) per the manufacturer’s protocol. The basal OCR was first measured in XF Assay Media without sodium pyruvate, followed by administration of 10 mM sodium pyruvate. Uncoupled respiration was evaluated following the addition of oligomycin (2 μM) to inhibit ATP synthase, by addition of the uncoupler FCCP (2 μM), and then followed by the addition of rotenone/antimycin (1 μM). Immediately after measurement, total protein levels were measured with the Micro BCA Protein Assay Kit (Thermo Scientific) for data correction.

**Cell transfection and luciferase reporter assays**

The luciferase assay was performed as previously described, with modification (Gan et al., 2011). The pCMV6-miR-499 vector was kindly provided by Dr. Eric Olson (UT Southwestern). To generate the 3’UTR luciferase reporters of miR-499 putative targets, the 3’UTRs containing the predicted binding site of miR-499 were amplified by PCR and inserted into the pGL3-promoter (Promega). Site-directed mutagenesis was performed using complementary oligonucleotides as follows (with mutated nucleotides shown in lowercase): 5’-GTATCTTGCATTAAATAATTGATATTGTTAATTTGAAACCATG (Fnip1 3’UTRmut). HEK293T cells were obtained from the American Type Culture Collection, and were cultured at 37°C and 5% CO₂ in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum, 1,000 U/ml penicillin, and 100 μg/ml streptomycin. Transient transfections in HEK293T cells were performed using PEI Transfection Reagent (Polysciences) following the manufacturer’s protocol. Briefly, 350 ng of reporter was cotransfected with 100 ng of pCMV6-miR-499 and 25 ng of CMV promoter-driven Renilla luciferase to control for transfection efficiency. Cells were harvested...
24 h after transfection. The luciferase assay was performed using Dual-Glo (Promega) according to the manufacturer’s recommendations. All transfection data are presented as the mean ± standard error of the mean (SEM) for at least three separate transfection experiments.

**Statistical analyses**

All mouse and cell studies were analyzed by Student’s t-test (two-tailed) or one-way ANOVA coupled to a Fisher’s least-significant difference (LSD) post hoc test when more than two groups were compared. No statistical methods were used to predetermine sample sizes. Data represent the mean ± SEM, with a statistically significant difference defined as a value of P < 0.05.

**Expanded View** for this article is available online.

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**Author contributions**

JL and XL contributed equally to this work and performed most of the experiments with assistance from DZ, Lla, TF, Lli, LX, and YK, and QZ, RBV, M-SZ, DPK, and XG contributed reagents and provided scientific insight and discussion. ZG provided oversight of the study including experimental design and data interpretation and wrote the manuscript. All authors reviewed and contributed to the manuscript.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**


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