Lnc-ORA interacts with microRNA-532-3p and IGF2BP2 to inhibit skeletal muscle myogenesis

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Skeletal muscle is one of the most important organs of the animal body. Long noncoding RNAs play a crucial role in the regulation of skeletal muscle development via several mechanisms. We recently identified obesity-related IncRNA (Inc-ORA) in a search for long noncoding RNAs that influence adipogenesis, finding it impacted adipocyte differentiation by regulating the PI3K/protein kinase B/mammalian target of rapamycin pathway. However, whether Inc-ORA has additional roles, specifically in skeletal muscle myogenesis, is not known. Here, we found that Inc-ORA was significantly differentially expressed with age in mouse skeletal muscle tissue and predominantly located in the cytoplasm. Overexpression of Inc-ORA promoted C2C12 myoblast proliferation and inhibited myoblast differentiation. In contrast, Inc-ORA knockdown repressed myoblast proliferation and facilitated myoblast differentiation. Interestingly, silencing of Inc-ORA rescued dexamethasone-induced muscle atrophy in vitro. Furthermore, adeno-associated virus 9–mediated overexpression of Inc-ORA decreased muscle mass and the cross-sectional area of muscle fiber by upregulating the levels of muscle atrophy–related genes and downregulating the levels of myogenic differentiation–related genes in vivo. Mechanistically, Inc-ORA inhibited skeletal muscle myogenesis by acting as a sponge of miR-532-3p, which targets the phosphatase and tensin homolog gene; the resultant changes in phosphatase and tensin homolog suppressed the PI3K/protein kinase B signaling pathway. In addition, Inc-ORA interacted with insulin-like growth factor 2 mRNA-binding protein 2 and reduced the stability of myogenesis genes, such as myogenic differentiation 1 and myosin heavy chain. Collectively, these findings indicate that Inc-ORA could be a novel underlying regulator of skeletal muscle development.

Skeletal muscle development is regulated by a series of myogenic regulatory factors (MRFs) (1). The MRF family plays a crucial positive role in skeletal muscle myogenic determination and differentiation during embryogenesis and postnatal myogenesis and includes myogenic differentiation 1 (MyoD), myogenic factor 5, myogenin (MyoG), and myogenic regulatory factor 4 (2). Once MRFs have been activated, many myogenic transcription factors form obligate heterodimers with their coregulators to activate the myoblast differentiation program by regulating the transcription of many genes, including coding and noncoding genes. In addition, myoblast proliferation leads to an increase in the number of nuclei, contributing to muscle growth in some forms of muscle hypertrophy in adults.

Skeletal muscle atrophy is controlled by the balance between the protein degradation rate and protein synthesis rate and induced by various stressors, including starvation, denervation, mechanical unloading, inflammation, and aging (3). This balance reflects the physiological condition of the muscle fiber, and breakage results in muscular dystrophy (4). Two critical protein degradation pathways, the autophagic–lysosomal and ubiquitin–proteasome systems, are activated during skeletal muscle atrophy (5, 6). These pathways include many atrophy-related genes, which are regulated by specific transcription factors. Muscle RING finger 1 (MuRF1) and muscle atrophy F-box (MAFbx) are atrophy markers, which represent two of many E3 ubiquitin ligases that are mostly expressed in skeletal muscle. Knockdown of MAFbx prevents skeletal muscle loss during fasting, and the absence of MuRF1 alleviates dexamethasone (Dex)-induced muscle atrophy in mice (7). The PI3K/protein kinase B (AKT) signaling pathway is one of the signaling pathways that regulate mammalian skeletal muscle atrophy (8).

The animal genome contains abundant noncoding RNAs, which serve as regulators of gene expression at the transcriptional, translational, and epigenetic levels. Noncoding RNAs regulate various muscle biological processes (9–11). Recent studies have confirmed that functional long noncoding RNAs (lncRNAs) are involved in skeletal muscle development, including skeletal muscle cell proliferation, differentiation, injury, atrophy, and regeneration by chromatin remodeling, transcription regulation, and microRNA sponge absorption (12–18). Although the effects of these lncRNAs in skeletal muscle myogenesis have been partially characterized, the function and regulatory mechanism of obesity-related IncRNA (Inc-ORA) in this process remains elusive.

In the present study, we found that Inc-ORA was significantly differentially expressed in skeletal muscle between two

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important developmental stages. Furthermore, the results indicated that lnc-ORA promoted myoblast proliferation, inhibited myoblast differentiation, and induced muscle atrophy in vitro. Moreover, overexpression of lnc-ORA reduced muscle mass and the cross-sectional area of muscle fibers by upregulating the levels of muscle atrophy–related genes and downregulating the levels of myoblast differentiation–related genes in vivo. Mechanistic investigations showed that lnc-ORA functioned as a sponge for miR-532-3p and insulin-like growth factor 2 mRNA-binding protein 2 (IGF2BP2), which activated phosphatase and tensin homolog (PTEN) and attenuated PI3K/AKT signaling, a critical pathway of myogenesis and muscle atrophy. Based on the aforementioned results, our findings provide a novel strategy for the regulation of skeletal muscle development.

**Results**

*Lnc-ORA is a potential regulator of skeletal muscle development*

To confirm whether lnc-ORA is associated with skeletal muscle development, the absolute expression of lnc-ORA in the tibialis anterior, gastrocnemius (GAS), and extensor digitorum longus muscles from 8-week-old and 52-week-old mice was examined. The results showed that the levels of lnc-ORA were much higher in 52-week-old mice than in 8-week-old mice.
mice (Fig. 1, A–C). Furthermore, the level of Inc-ORA was the highest at 12 h and then gradually decreased from 24 to 48 h during myoblast proliferation (Fig. 1D). The level of Inc-ORA gradually increased during the early stage and then decreased in the late stage of C2C12 cell differentiation (Fig. 1E), indicating that Inc-ORA was involved in myoblast proliferation and differentiation during muscle formation. To confirm the stability of Inc-ORA in C2C12 cells, a half-life experiment was performed. The results showed that the half-life of Inc-ORA was approximately 12 h, indicating that Inc-ORA is stably expressed in C2C12 cells (Fig. 1F). In addition, based on Inc-ORA–targeted genes, Gene Ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses were performed. GO term analysis revealed that Inc-ORA participated in the regulation of skeletal muscle development, muscle atrophy, cellular metabolic process, muscle hypertrophy, especially myoblast proliferation and differentiation (Fig. 1G). KEGG analysis indicated that Inc-ORA could regulate biological processes through the PI3K/AKT, peroxisome proliferator–activated receptors, mammalian target of rapamycin, and forkhead box O signaling pathway (Fig. 1H). Collectively, these results imply that Inc-ORA is a potential regulator of skeletal muscle development, partly through the PI3K/AKT signaling pathway.

**Figure 2.** Overexpression of Inc-ORA promotes myoblast proliferation. A, EdU and DAPI (nuclei) staining analysis 24 h after transfection of pclnc-ORA and pcDNA3.1 empty plasmid in proliferating myoblasts (n = 3). The scale bar represents 200 μm. B, percentage of EdU-positive cells/total cells (n = 3). C, flow cytometry analysis 24 h after transfection of pclnc-ORA and pcDNA3.1 empty plasmid (n = 3). D, statistical results of flow cytometry (n = 3). E, mRNA levels of cyclin E, cyclin D1, and PCNA 24 h after overexpression of Inc-ORA (n = 3). F, Western blot detection of cyclin E, cyclin D1, and PCNA (n = 3). G, quantitation of the protein level in F (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. DAPI, 4′,6-diamidino-2-phenylindole; EdU, 5-ethyl-20-deoxyuridine; Inc, long noncoding; Inc-ORA, obesity-related IncRNA; pclnc-ORA, pcDNA3.1-lnc-ORA; PCNA, proliferating cell nuclear antigen.
Overexpression of lnc-ORA promotes myoblast proliferation, whereas knockdown of lnc-ORA inhibits myoblast proliferation

To investigate the role of lnc-ORA in myoblast proliferation, lnc-ORA overexpression and knockdown experiments were carried out in proliferating C2C12 cells. The results showed that overexpression and knockdown of lnc-ORA worked well (Fig. S1, A and B). Overexpression of lnc-ORA increased the number of 5-ethynyl-2-deoxyuridine (EdU)-positive cells (Fig. 2, A and B). A flow cytometry analysis also indicated that overexpression of lnc-ORA increased the number of cells that progressed to S phase (Fig. 2, C and D). In addition, cell count assay showed that overexpression of lnc-ORA increased the total cell number (Fig. S1C). Moreover, overexpression of lnc-ORA increased the mRNA and protein levels of proliferation-related genes, including cyclin E, cyclin D1, and proliferating cell nuclear antigen (Fig. 2, E–G). Furthermore, knockdown of lnc-ORA decreased the number of EdU-positive cells (Fig. 3, A and B), the total cell number (Fig. S1D), the number of cells in S phase (Fig. 3, C and D), and the mRNA and protein levels of proliferation-related genes (Fig. 3, E–G). Taken together, these results indicate that lnc-ORA promotes myoblast proliferation.

Figure 3. Knockdown of lnc-ORA inhibits myoblast proliferation. A, EdU and DAPI (nuclei) staining analysis 24 h after transfection of silnc-ORA and si-NC in proliferating myoblasts (n = 3). The scale bar represents 200 μm. B, percentage of EdU-positive cells/total cells (n = 3). C, flow cytometry analysis 24 h after transfection of silnc-ORA and si-NC in proliferating myoblasts (n = 3). D, statistical results of flow cytometry (n = 3). E, mRNA levels of cyclin E, cyclin D1, and PCNA 24 h after knockdown of lnc-ORA (n = 3). F, Western blot detection of cyclin E, cyclin D1, and PCNA (n = 3). G, quantitation of protein level in F (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. DAPI, 4’,6-diamidino-2-phenylindole; EdU, 5-ethynyl-2-deoxyuridine; Inc, long noncoding; lnc-ORA, obesity-related IncRNA; PCNA, proliferating cell nuclear antigen; silnc-ORA, siRNA-lnc-ORA.
Overexpression of Inc-ORA inhibits myogenic differentiation, whereas knockdown of Inc-ORA promotes myogenic differentiation

To verify the effect of Inc-ORA on myogenic differentiation, overexpression and knockdown experiments were performed during C2C12 cell differentiation. C2C12 cells were induced by differentiation culture medium (Fig. S2A). After myoblasts were transfected with pcDNA3.1-Inc-ORA or siRNA-Inc-ORA (silnc-ORA) vector, the levels of Inc-ORA markedly changed after myogenic induction (Fig. S2, B and C), showing that overexpression and knockdown of Inc-ORA worked well. Overexpression of Inc-ORA decreased the number of myosin heavy chain (MyHC)-positive cells, the differentiation index, and the multinuclear fusion index (Fig. 4, A–C). Moreover, overexpression of Inc-ORA downregulated the mRNA and protein levels of myogenic markers, including MyoD, MyoG, and MyHC (Fig. 4, D–F). Conversely, knockdown of Inc-ORA increased the number of MyHC-positive cells and the differentiation index, as well as the multinuclear fusion index at day 4 after myogenic induction (Fig. 5, A–C). Knockdown of Inc-ORA upregulated the mRNA and protein levels of these markers.

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**Figure 4. Overexpression of Inc-ORA inhibits myogenic differentiation.** A, immunofluorescent staining of MyHC and DAPI (nuclei) on day 4 of differentiation after overexpression of Inc-ORA (n = 3). The scale bar represents 100 μm. B, statistical analysis of the differentiation index in A (n = 3). C, statistical analysis of the fusion index in A (n = 3). D, mRNA levels of myogenic genes on day 4 of differentiation after overexpression of Inc-ORA (n = 3). E, Western blot detection of MyoD, MyoG, and MyHC (n = 3). F, quantitation of protein levels in E (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. DAPI, 4',6-diamidino-2-phenylindole; Inc, long noncoding; Inc-ORA, obesity-related IncRNA; MyHC, myosin heavy chain; MyoD, myogenic differentiation 1; MyoG, myogenin.
Taken together, these results demonstrate that lnc-ORA inhibits myogenic differentiation of myoblasts.

**Knockdown of lnc-ORA rescues Dex-induced muscle atrophy in vitro**

To examine whether lnc-ORA could influence muscle atrophy, a Dex-induced muscle atrophy model was used in C2C12 cells (Fig. 6A). The level of lnc-ORA was significantly upregulated by the Dex treatment (Fig. 6B), and an increase in the protein levels of MAFbx and MuRF1 also occurred (Fig. 6, C and D). Furthermore, knockdown of lnc-ORA markedly rescued Dex-induced muscle atrophy, as shown by MyHC staining (Fig. 6E). Knockdown of lnc-ORA decreased the levels of MAFbx and MuRF1 and increased the levels of myogenic differentiation factors (Fig. 6, F and G). Therefore, knockdown of lnc-ORA rescued Dex-induced muscle atrophy in vitro, suggesting that lnc-ORA could be a potential therapeutic target for treating muscle atrophy.
Overexpression of lnc-ORA decreases muscle mass and induces muscle atrophy in vivo

To investigate the role of lnc-ORA in the regulation of muscle development in vivo, 10-week-old mice were injected with adeno-associated virus-GFP (AAV-GFP) or AAV-lnc-ORA overexpression virus and sacrificed after 8 weeks. Compared with the AAV-GFP control group, the AAV treatment group showed decreased size and mass of the tibialis anterior and GAS muscles (Fig. 7, A and B). Meanwhile, the muscle percentage of AAV-lnc-ORA–injected mice was lower (Fig. 7, C). H&E staining showed that the cross-sectional areas of GAS fibers were dramatically smaller in the AAV-lnc-ORA group than in the AAV-GFP group (Fig. 7, D and E). Overexpression of lnc-ORA significantly decreased the expression levels of MyHC and MyoD but increased the expression levels of MAFbx and MuRF1 (Fig. 7, F and G). The PI3K/AKT signaling pathway was inhibited in the GAS muscles of AAV-lnc-ORA mice (Fig. 7, F and H). Together, these results suggest that overexpression of lnc-ORA decreases muscle mass and induces muscle atrophy in vivo.

Lnc-ORA sponges miR-532-3p to control myogenesis through the PTEN/PI3K/AKT signaling pathway

To investigate the molecular mechanism of lnc-ORA in myogenesis, FISH experiments were performed to confirm the subcellular locations of lnc-ORA in myoblasts and myotubes. The results showed that lnc-ORA was predominantly expressed in the cytoplasm of both myoblasts and myotubes (Fig. 8, A and B). This finding indicates that lnc-ORA plays a role in the post-transcriptional regulation mechanism in the...
cytoplasm. Moreover, we predicted that miR-532-3p with a conserved seed sequence could be adsorbed by lnc-ORA (Fig. S3, A and B) and that PTEN acts as a target of miR-532-3p (Fig. S3 C). To confirm this hypothesis, we constructed wildtype and mutant dual luciferase reporters, psi-CHECK 2.0-lnc-ORA or psi-CHECK 2.0-PTEN 3'UTR (Fig. 9, A and B). The dual-luciferase reporter assay revealed that miR-532-3p bound to lnc-ORA transcripts and the PTEN 3'UTR (Fig. 9, C and D). Furthermore, significantly enriched miR-532-3p and lnc-ORA were measured through argonaute 2 RNA immunoprecipitation (RIP) assay (Fig. S3 D). Moreover, lnc-ORA was significantly enriched by biotin-labeled miR-532-3p compared with the control or mutated miR-532-3p (Fig. S3 E). We have also detected the absolute copy number

Figure 7. Overexpression of Inc-ORA decreases muscle mass and induces muscle atrophy in vivo. A, images of mouse skeletal muscles. B, statistical analysis of muscles mass (n = 5). C, percentage of skeletal muscle (TA and GAS) (n = 5). D, H&E staining images of GAS cross section. The scale bar represents 200 μm. E, statistics of fiber area in D (n = 5). F, Western blot detection of MyHC, MyoD, MAFbx, MuRF, p-Pi3K, Pi3K, p-AKT, and AKT in the GAS (n = 3). G and H, quantitation of protein level in F (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. AKT, protein kinase B; GAS, gastrocnemius; Inc, long noncoding; Inc-ORA, obesity-related lncRNA; MAFbx, muscle atrophy F-box; MuRF, Muscle RING finger; MyHC, myosin heavy chain; MyoD, myogenic differentiation 1; p-AKT, phosphorylated AKT; p-Pi3K, phosphorylated Pi3K; TA, tibialis anterior.
of miR-532-3p under the different conditions. The results showed that the absolute copy number of miR-532-3p exhibited opposite trends as that of lnc-ORA under different conditions (Fig. S4, A–E). Based on the aforementioned results, we speculated that lnc-ORA acted as a competing endogenous RNA to sponge miR-532-3p. Furthermore, miR-532-3p mimics promoted myogenic differentiation, as shown by immunofluorescence staining of MyHC-positive myotubes (Fig. S5, A and B). Besides, rescue experiments indicated that miR-532-3p attenuated the positive effect of lnc-ORA on myoblast proliferation (Fig. 9, E and F), whereas it rescued the repressive effect of lnc-ORA on myoblast differentiation.

Figure 8. Subcellular localization of lnc-ORA in proliferating and differentiating myoblasts. A, subcellular localization of lnc-ORA by FISH in myoblasts. Special FISH probes against lnc-ORA, U6, and 18S were modified by Cy3 in red. 18S is a cytoplasmic marker, and U6 is a nuclear marker. The nucleus was stained by DAPI in blue. The scale bar represents 50 μm. B, subcellular localization of lnc-ORA by FISH in myotubes. Special FISH probes against lnc-ORA, U6, and 18S were modified by Cy3 in red. 18S is a cytoplasmic marker, and U6 is a nuclear marker. The nucleus was stained by DAPI in blue. The scale bar represents 50 μm. DAPI, 4′,6-diamidino-2-phenylindole; Inc, long noncoding; Inc-ORA, obesity-related IncRNA.

Figure 9. Lnc-ORA functions as a ceRNA for miR-532-3p to control myogenesis. A and B, schematic of the double luciferase assay vector. C and D, analysis of the luciferase reporter assay (n = 3). E, EdU and DAPI (nuclei) staining analysis. The scale bar represents 200 μm. F, statistical analysis in E (n = 3). G, immunofluorescent staining of MyHC analysis. The scale bar represents 200 μm. H, statistical analysis in G (n = 3). I, Western blot detection of PTEN and myogenic factors (n = 3). J, statistical analysis in I (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. ceRNA, competing endogenous RNA; DAPI, 4′,6-diamidino-2-phenylindole; EdU, 5-ethynyl-2′-deoxyuridine; Inc, long noncoding; Inc-ORA, obesity-related IncRNA; MyHC, myosin heavy chain; PTEN, phosphatase and tensin homolog.
Moreover, miR-532-3p showed the opposite effect against lnc-ORA on the protein levels of PTEN, phosphorylated PI3K (p-PI3K), and phosphorylated AKT (p-AKT) (Fig. 9, I and J). In contrast, lnc-ORA knockdown reduced the protein level of PTEN and increased the protein levels of p-PI3K and p-AKT (Fig. S6). Therefore, lnc-ORA inhibited myogenic differentiation and promoted muscle atrophy through absorption of miR-532-3p via PTEN/PI3K/AKT signaling pathway.

**Lnc-ORA contributes to myogenesis by competitively binding IGF2BP2**

To further explore the mechanism by which lnc-ORA regulates myogenesis, we used a biotinylated lnc-ORA probe to perform an RNA pull-down assay, followed by mass spectrometry. Through this analysis, we identified IGF2BP2, an RNA-binding protein, as being bound to lnc-ORA, and we confirmed this interaction by Western blotting (Fig. 10A). Next, RIP was performed in C2C12 cells using an IGF2BP2 antibody. Lnc-ORA was present in the IGF2BP2 RIP sample at a much higher level than in the control IgG RIP sample (Fig. 10B). It has been demonstrated that IGF2BP2 could promote RNA stability. We then used this half-life assay to explore the RNA stability of lnc-ORA. The results showed that the interaction of lnc-ORA and IGFBP2 increased lnc-ORA stability and the ability to sponge miR-532-3p (Fig. 10, C and D). Moreover, lnc-ORA directly bound to IGF2BP2 and negatively regulated the IGF2BP2-mediated stability of myogenesis genes such as MyoD and MyHC (Fig. 10, E and F).

Together, we used RNA pull-down experiments to identify IGF2BP2 as an important binding partner of lnc-ORA in the myogenesis process. Based on the aforementioned results, we suggest that lnc-ORA, acting as a miR-532-3p sponge and interacting with IGF2BP2 to control PTEN-mediated PI3K/AKT signaling, inhibits skeletal muscle myogenesis and induces muscle atrophy (Fig. 11).

**Discussion**

Skeletal muscle is closely associated with physiological function, muscle strength, and metabolic performance, which influence human chronic disease, quality of life, and animal meat production. Recently, several studies have indicated that lncRNAs are implicated in skeletal muscle formation. However, the effect and molecular mechanism of lnc-ORA on myoblast proliferation, differentiation, and muscle atrophy are unknown. In the present study, we found that lnc-ORA was differentially expressed in mouse skeletal muscle with age, functioning as a miR-532-3p sponge and interacting with IGF2BP2 to inhibit myogenesis and induce muscle atrophy, and it could be a novel target for the regulation of skeletal muscle development.

LncRNAs have been demonstrated to regulate myoblast proliferation and differentiation, such as Myoparr (19), Linc-smad7 (20), Lnc-31 (21), and Gm26917 (22). Our previous study revealed that knockdown of lnc-ORA inhibited pre-adipocyte proliferation and differentiation (23). In the current study, we found that lnc-ORA was differentially expressed in mouse skeletal muscle with age, functioning as a miR-532-3p sponge and interacting with IGF2BP2 to inhibit myogenesis and induce muscle atrophy, and it could be a novel target for the regulation of skeletal muscle development.

Figure 10. Lnc-ORA contributes to myogenesis by competitively binding IGF2BP2. A, Western blotting identified IGF2BP2 bound with lnc-ORA. B, RIP assay revealed that IGF2BP2 was bound with lnc-ORA (n = 3). C, lnc-ORA RNA stability assays in C2C12 myoblasts (n = 3). D, expression level of miR-532-3p when IGF2BP2 was overexpressed (n = 3). E and F, MyoD and MyHC RNA stability assays in C2C12 cells (n = 3). Data represent the mean ± SD. *p < 0.05; **p < 0.01. IGF2BP2, insulin-like growth factor 2 mRNA-binding protein 2; lnc, long noncoding; lnc-ORA, obesity-related lncRNA; MyHC, myosin heavy chain; MyoD, myogenic differentiation 1; RIP, RNA immunoprecipitation.
study, we found that overexpression of Inc-ORA promoted myoblast proliferation but inhibited myotube formation. Furthermore, its knockdown had the opposite effect, implying that Inc-ORA is a vital negative regulatory factor of skeletal muscle myogenesis.

An increasing number of studies have also indicated that lncRNAs, including IncIRS1 [24], IncMUMA [25], Atrolnc-1 [26], Pctl1 [27], and SMN-AS1 [28], play a significant role in the regulation of muscle atrophy. In the current study, the level of Inc-ORA markedly increased in a Dex-induced muscle atrophy model, and knockdown of Inc-ORA significantly rescued muscle atrophy by inhibiting MAFbx and MuRF1 and promoting myogenic differentiation factor expression. In addition, skeletal muscle atrophy is defined as a decline in skeletal muscle mass because of muscle cellular shrinkage [29, 30]. Interestingly, we found that overexpression of Inc-ORA significantly decreased skeletal muscle mass and the cross-sectional area of muscle fibers in vivo. Therefore, it is suggested that Inc-ORA could be a promising therapeutic target for muscle atrophy.

LncRNAs not only regulate target genes at the transcriptional level in the nucleus but also participate at the post-transcriptional level in the cytoplasm, suggesting that the subcellular localization of lncRNAs is related to their regulatory mechanism [31–33]. Here, we found that Inc-ORA is distributed in both the nucleus and cytoplasm of either myoblasts or myotubes, but the proportion in the cytoplasm is much greater than that in the nucleus, indicating that Inc-ORA may play an important role in post-transcriptional regulation. Generally, lncRNAs function by absorbing miRNAs to regulate the expression of target genes [34–36]. Bioinformatics prediction showed that Inc-ORA acts as a competing endogenous RNA to absorb miRNA-532-3p, which could target PTEN. As an important factor of myogenic genes, PTEN could negatively regulate the PI3K signaling pathway, which has well-known anabolic effects to promote myoblast differentiation in skeletal muscles [37]. Furthermore, inhibition of the PI3K signaling pathway could activate the transcription factor forkhead box O family [38, 39], which induces the expression of the MuRF-1 and MAFbx genes implicated in the induction of muscle atrophy [40, 41]. Based on our results, we confirmed that Inc-ORA inhibited myogenic differentiation and aggravated muscle atrophy by sequestering miRNA-532-3p from inhibition of PTEN. In general, a single miRNA is able to recognize and inhibits a large number of target genes. Conversely, a single gene may be targeted by numerous miRNAs. Of course, Inc-ORA could be predicted to sequester other miRNAs in some biological processes, such as miR-149-5p, miR-320-5p, and miR-760-5p. Here, we confirmed that Inc-ORA acts as a miR-532-3p sponge in skeletal muscle myogenesis. IGF2BP2 could promote the stability and translation of RNAs [42]. Moreover, IGF2BP targets 76 genes that are associated with muscle development [43]. Consistent with the molecular mechanism of IncMYOD [44], we found that Inc-ORA competitively bound to IGF2BP2 and negatively regulated the IGF2BP2-mediated stability of myogenic genes such as MyoD and MyHC.

In conclusion, we found that Inc-ORA acts as a miR-532-3p sponge and interacts with IGF2BP2 to suppress the PI3K/AKT signaling pathway, resulting in the inhibition of myogenesis and induction of skeletal muscle atrophy. Based on our findings, we hypothesize that Inc-ORA could be a novel underlying regulator of skeletal muscle development.

Experimental procedures

Animals

C57/BL6J male mice were purchased from the Medical Laboratory Animal Center of Xi’an Jiaotong University (Xi’an, China). All animal experiments were approved by ethics committee of animal welfare and health of Northwest A&F University (NWAFU-314020038).

Cell culture, transfection, and RNA stability assay

The C2C12 mouse myoblast cell line was acquired from American Type Culture Collection. Cells were cultured with growth medium, which was composed of high-glucose Dulbecco’s modified Eagle’s medium and 10% fetal bovine serum (Gibco). After the cells reached 80 to 90% confluence, C2C12 cell differentiation was induced by differentiation medium consisting of Dulbecco’s modified Eagle’s medium and 2% horse serum (Gibco). To study myoblast proliferation, transfection experiments were performed when cells reached 40 to 50% confluence. Cells were transfected for studying myogenic differentiation when cells reached 70 to 80% density. All transfection procedures were performed with Xtreme-GENE Transfection Reagent (Roche) in accordance with the manufacturer’s instructions. The miR-532-3p mimics and inhibitor, silnc-ORA and siNC, were synthesized by Ribobio. The overexpression vectors of Inc-ORA and Inc-ORA mutant were constructed with the pcDNA3.1 plasmid. The empty pcDNA3.1 vector was used as a control plasmid. To construct Dex-induced muscle atrophy in vitro, well-differentiated C2C12 cells were treated with either 50 μM Dex (D4902; Sigma–Aldrich) or PBS (control), and then siRNA targeting Inc-ORA was transfected 36 h after treatment. The half-life of Inc-ORA was detected according to our previous report [45].
Role of lnc-ORA in skeletal muscle myogenesis

**RNA-fluorescent in situ hybridization**

In situ hybridization of lnc-ORA in myoblasts and myotubes was performed with a FISH kit (Ribobio). After washing three times with PBS, C2C12 cells were fixed with 4% paraformaldehyde for 30 min. Next, the cells were prehybridized for 30 min with prehybridization solution after permeabilization with 0.5% Triton X-100 buffer for 5 min. Then, the specific oligodeoxynucleotide probes of anti-18S RNA, anti-U6, and anti-lnc-ORA were hybridized at 37 °C overnight. Finally, the cells were stained with 4′,6-diamidino-2-phenylindole dye and photographed using a confocal laser-scanning microscope.

**RNA isolation and relative quantitative real-time PCR**

Total RNA from different cell and tissue samples was extracted by TRizol reagent (TaKaRa) in accordance with the manufacturer’s protocol. Complementary DNA synthesis was carried out by reverse transcription kits (TaKaRa). Quantitative real-time RT-PCR (qRT-PCR) was performed with a Bio-Rad iQTM5 system (Bio-Rad). For the miR-532-3p expression level, the specific primers of miR-532-3p and reference gene U6 (RiboBio) were used for reverse transcription. The 2−ΔΔCt method was used to analyze qRT-PCR data. Mouse GAPDH was used as a reference gene. The primer sequences of genes detected in this study are listed in Table S1.

**Western blotting**

Total protein of the cell or tissue samples was harvested using radioimmunoprecipitation lysis buffer after washing three times with PBS. Immunoblotting was carried out according to our previous method (10). The primary antibodies targeted the following proteins: MyHC (1:500; no. MAB4470; R and D Systems), MyoD (1:500; no. NB100-56511; Novus Biologicals), AKT (1:1000; no. 9272S; CST), p-AKT (Ser473) (1:1000; no. 9272S; CST), β-catenin (1:200; no. sc-58880; Santa Cruz Biotechnology), MyoG (1:1000; no. NB100-56510SS; Novus Biologicals), MuRF1 (1:200; no. sc-398608; Santa Cruz Biotechnology), MAFbx (1:200; no. sc-166806; Santa Cruz Biotechnology), cyclin E (1:100; no. sc-377100; Santa Cruz Biotechnology), cyclin D1 (1:100; no. sc-450; Santa Cruz Biotechnology), PTEN (1:500; no. sc-7974; Santa Cruz Biotechnology), IGF2BP2 (1:100; no. sc-377014; Santa Cruz Biotechnology), and proliferating cell nuclear antigen (1:100; no. sc-56; Santa Cruz Biotechnology). The secondary antibodies included goat anti-rabbit IgG (1:3000; no. BA1054; BosterBio) and goat anti-mouse IgG (1:3000; no. BA1050; BosterBio).

**Overexpression of lnc-ORA by AAV infection in vivo**

The recombinant AAV viruses (AAV9 serotype) expressing lnc-ORA or GFP were acquired by Hanbio (46). Ten-week-old C57BL/6J male mice with eight mice in each group were injected with 150 μl AAV virus including full-length lnc-ORA (AAV-lnc-ORA) or control (AAV-GFP) at 5 × 1012 vg/ml titers.

**Frozen section and H&E staining**

Frozen sectioning and H&E staining of GAS samples were performed according to our published method (47).

**Flow cytometry, EdU, and cell counting kit-8 assays**

Flow cytometry, EdU, and cell counting kit-8 experiments were performed according to our previous report (48). Briefly, after the C2C12 cell density reached 40% confluence, the cells were treated with pcDNA3.1-lnc-ORA or silnc-ORA as well as miR-532-3p mimics or miR-532-3p inhibitor. After treatment for 24 h, the cells were used to detect proliferation-related analysis.

**Immunofluorescence assay**

The immunofluorescence assay was performed as described in our previous procedure (49). Briefly, cells were fixed with 4% paraformaldehyde and blocked with 5% bovine serum albumin for 30 min. Then, the cells were incubated with MyHC primary antibody overnight at 4 °C. Next, the samples were incubated with secondary antibody for 1 h at 37 °C and 4′,6-diamidino-2-phenylindole for 10 min. The differentiation index was determined as the percentage of MyHC-positive nuclei among the total nuclei, and the myotube fusion index was determined as the distribution of the nucleus number among the total myotubes according to a previous report (50).

**Bioinformatics analysis**

miRbase (http://www.mirbase.org/) and TargetScan (http://www.targetscan.org/vert_71/) were used to predict target genes of miR-532-3p. GO analysis is a functional analysis for neighboring genes and coexpressed genes of lnc-ORA with GO categories. The GO categories are derived from GO (http://www.geneontology.org), which consists of three structured networks of defined terms that describe gene product attributes. The KEGG pathway enrichment analysis for neighboring genes and coexpressed genes of lnc-ORA is based on the latest KEGG database (https://www.genome.jp/kegg).

**Dual-luciferase activity assay**

The putative binding sites (wildtype and mutated) of miR-532-3p within Inc-ORA and the 3′-UTR of PTEN were synthesized using General Biosystems and inserted into psiCHECK-2 vector (Promega). Dual-luciferase activity was analyzed according to our previous report (48). Briefly, human embryonic kidney 293T cells were transfected with miR-532-3p mimics or negative control mimics as well as psiCHECK2 reporter wildtype or mutant vector. After transfection for 48 h, the relative luciferase activity was analyzed following the manufacturer’s protocol (Promega).

**Absolute quantitative real-time PCR**

Absolute qRT-PCR was used to analyze copy number of Inc-ORA and miR-532-3p under different conditions. Briefly, T-vectors containing a fragment of Inc-ORA and miR-532-3p
were constructed and diluted via titration to generate the standard curve. Total RNA (200 ng) from each sample was reverse transcribed, and complementary DNA was used as a template in qRT-PCR. The Ct values were used to calculate the copy numbers of lnc-ORA and miR-532-3p in each sample, according to the standard curve.

**RNA pull-down assay**

Linearizing DNA was biotin labeled and transcribed in vitro using Biotin RNA Labeling Mix and T7/SP6 RNA polymerase (Roche) and purified with the RNase Mini Kit (QIAugen). One milligram of protein was incubated with 3 μg of biotinylated RNA for 1 h at room temperature. After that, 40 μl streptavidin-coupled beads were added to each reaction and incubated for 1 h at room temperature. Finally, the beads were washed in RIP buffer for five times, and the pulled-down proteins were used for Western blotting. For mass spectrometry, the pulled-down proteins in C2C12 cells were separated by 10% SDS-PAGE and then subjected to silver staining. The differentially expressed bands were excised and analyzed by mass spectrometry (Novogene).

**miRNA pull-down assay**

A miRNA pull-down assay was performed as previously reported (51). C2C12 cells were transfected with biotinylated miR-532-3p: wildtype miR-532-3p (miR-532-3p-Bio) or mutated miR-532-3p (G–C mutation in the lnc-ORA—binding sites, miR-532-3p-MutBio) and biotinylated control (negative control-Bio) when 70 to 80% of confluency was reached and induced into myogenic differentiation at full confluence. Six days after induction, cell lysates were collected and incubated with Dyna M-280 streptavidin magnetic beads (#11205D; Thermo Fisher Scientific). The product was then treated with RNase-free DNase I (Roche), and RNA was purified using an RNase Mini Kit (QIAGEN). The enrichment of lnc-ORA was detected through qRT-PCR.

**RNA immunoprecipitation assay**

Well-differentiated C2C12 cells were subjected to an RIP assay after 6 days of induction. Briefly, cells were collected, lysed in complete RIP buffer provided in the EZ-Magna RIP kit (Millipore), and then incubated with RIP buffer containing magnetic beads conjugated to anti-arginoula 2 or anti-IGF2BP2 antibody. Subsequently, the samples were digested with proteinase K, and then the RNA was purified from the precipitates. The concentration of RNA was determined using a NanoDrop system (Thermo Fisher Scientific). Finally, qRT-PCR was performed to detect the existence of miR-532-3p and lnc-ORA.

**Statistical analysis**

All replicate experiments (including cell-based and mouse-based experiments) were biological replicates that were repeated at least three times. All analyses involved the use of SPSS, version 23 (SPSS, Inc). All data are represented as the mean ± SD. Comparisons of two groups were determined by Student’s t test, and comparisons of multiple groups were determined by one-way ANOVA with Tukey’s post hoc test. The assumption of normality was tested by the Shapiro–Wilk test. All statistical tests were two-tailed, and p < 0.05 was considered statistically significant.

**Data availability**

The data used to support the findings of this study are available from the corresponding author upon request.

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**Supporting information**—This article contains supporting information.

**Author contributions**—R. C. and Q. Z. data curation; R. C., Y. W., and W. Y. investigation; R. Z. methodology; R. C. writing original draft; R. C. and W. P. writing - review and editing; W. P. project administration; R. Z. conceptualization; W. P. funding acquisition; and W. P. supervised the project.

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**Conflict of interest**—The authors declare that they have no conflicts of interest with the contents of this article.

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