The remarkable regeneration capability of skeletal muscle depends on the coordinated proliferation and differentiation of satellite cells (SCs). The self-renewal of SCs is critical for long-term maintenance of muscle regeneration potential. Hypoxia profoundly affects the proliferation, differentiation, and self-renewal of cultured myoblasts. However, the physiological relevance of hypoxia and hypoxia signaling in SCs in vivo remains largely unknown. Here, we demonstrate that SCs are in an intrinsic hypoxic state in vivo and express hypoxia-inducible factor 2A (HIF2A). HIF2A promotes the stemness and long-term homeostatic maintenance of SCs by maintaining their quiescence, increasing their self-renewal, and blocking their myogenic differentiation. HIF2A stabilization in SCs cultured under normoxia augments their engraftment potential in regenerative muscle. Conversely, HIF2A ablation leads to the depletion of SCs and their consequent regenerative failure in the long-term. In contrast, transient pharmacological inhibition of HIF2A accelerates muscle regeneration by increasing SC proliferation and differentiation. Mechanistically, HIF2A induces the quiescence and self-renewal of SCs by binding the promoter of the Spry1 gene and activating Spry1 expression. These findings suggest that HIF2A is a pivotal mediator of hypoxia signaling in SCs and may be therapeutically targeted to improve muscle regeneration.
Transient HIF2A inhibition promotes satellite cell proliferation and muscle regeneration

Liwei Xie,1,2 Amelia Yin,1,2 Anna S. Nichenko,2 Aaron M. Beedle,4 Jarrod A. Call,3,5 and Hang Yin1,2
1Department of Biochemistry and Molecular Biology, 2Center for Molecular Medicine, and 3Department of Kinesiology, The University of Georgia, Athens, Georgia, USA. 4Department of Pharmaceutical Sciences, Binghamton University-SUNY, Binghamton, New York, USA. 5Regenerative Bioscience Center, The University of Georgia, Athens, Georgia, USA.

The remarkable regeneration capability of skeletal muscle depends on the coordinated proliferation and differentiation of satellite cells (SCs). The self-renewal of SCs is critical for long-term maintenance of muscle regeneration potential. Hypoxia profoundly affects the proliferation, differentiation, and self-renewal of cultured myoblasts. However, the physiological relevance of hypoxia and hypoxia signaling in SCs in vivo remains largely unknown. Here, we demonstrate that SCs are in an intrinsic hypoxic state in vivo and express hypoxia-inducible factor 2A (HIF2A). HIF2A promotes the stemness and long-term homeostatic maintenance of SCs by maintaining their quiescence, increasing their self-renewal, and blocking their myogenic differentiation. HIF2A stabilization in SCs cultured under normoxia augments their engraftment potential in regenerative muscle. Conversely, HIF2A ablation leads to the depletion of SCs and their consequent regenerative failure in the long-term. In contrast, transient pharmacological inhibition of HIF2A accelerates muscle regeneration by increasing SC proliferation and differentiation. Mechanistically, HIF2A induces the quiescence and self-renewal of SCs by binding the promoter of the Spry1 gene and activating Spry1 expression. These findings suggest that HIF2A is a pivotal mediator of hypoxia signaling in SCs and may be therapeutically targeted to improve muscle regeneration.

Introduction
The coordinated proliferation and differentiation of adult stem cells provide sustainable cell sources for tissue repair and regeneration. In support of long-term tissue homeostasis, many adult stem cell populations fine-tune the balance between proliferative and quiescent states so as to both meet the need for tissue repair and lessen cell cycle–associated stresses. Ineffective reversal of proliferation and quiescence in adult stem cells has been implicated in aging and many diseases (1).

Adult skeletal muscle stem cells, also called satellite cells (SCs), are essential for skeletal muscle regeneration (2). SCs and their undifferentiated progeny universally express paired box 7 (Pax7), a transcription factor that has been utilized to identify both quiescent and proliferative SCs or manipulate gene expression specifically in SCs (3, 4). In uninjured muscle, SCs reside closely juxtaposed with contractile myofibers beneath the basal lamina and are mitotically quiescent. SC quiescence is jointly maintained by, but not limited to, multiple mechanisms including TTP-mediated RNA decay, SIRT1-dependent histone deacetylation, and restraint of myogenic and proliferation-associated gene expression by microRNAs (5–8). Upon muscle injury, SCs transit through a G1(Meta) phase and enter the cell cycle in response to the activation of HGF/FGF-mediated RTK signaling and p38/MAPK signaling (9–11). After limited rounds of proliferation, a subset of SCs returns to quiescence and the niche location adjacent to myofibers (self-renewal), whereas many SCs undergo myogenic differentiation and eventually repair damaged myofibers. Angiopoietin-1/Tie2 RTK signaling, the RTK negative regulator Spry1, and inactivation of p38/MAPK signaling promote SC self-renewal (11–13). Besides these intrinsic mechanisms, dynamic remodeling of the muscle microenvironment also plays a critical role in directing SC behavior during muscle repair (2). However, how SCs fine-tune the scale of reversible quiescence and balance proliferation versus differentiation remains incompletely understood.

Oxygen (O2) is essential for mitochondrial bioenergetics and energy homeostasis; low O2 tension (hypoxia) underlies many disease conditions (14). The cellular responses to hypoxia are mainly mediated by hypoxia-inducible factors (HIFs), which form heterodimeric complexes between an O2-sensitive α-subunit (HIF1A/HIF2A/HIF3A) and a stable β-subunit (HIF1B) (15). Under normoxia, the α-subunit of HIF is hydroxylated and targeted for proteasome degradation. Under hypoxia, limited O2 stabilizes the HIF complex, which binds hypoxia-responsive elements (HREs) (5′-RCGTG-3′) and activates target gene expression. Many types of adult stem cells prefer hypoxic niches to maintain their quiescent and/or undifferentiated states (16). In vitro hypoxic conditions also profoundly impact the proliferation and differentiation of cultured SCs (myoblasts) (17). In addition, HIF1A KO or compound HIF1A/HIF2A double-KO in adult SCs distinctly affects SC numbers and muscle regeneration (18, 19). Yet, it remains elusive whether hypoxia is of any physiological relevance to SC in vivo and, if so, whether HIF-mediated hypoxia signaling regulates SC quiescence or can be potentially targeted to improve muscle regeneration.

Here, we demonstrated that SCs are inherently hypoxic within their in vivo niche. Quiescent SCs (QSCs) expressed HIF2A, but not...
HIF1A. We found that HIF2A was essential for the maintenance of SCs in a quiescent and undifferentiated state and for vif

Results

QSCs are hypoxic in the niche and express HIF2A, but not HIF1A. The partial oxygen tension (pO2) of most adult tissues ranges from approximately 2% to 9% (20). To assess the intracellular pO2 of SCs in their niche, a hypoxia probe, pimonidazole, was administered in vivo. This probe forms protein adducts in hypoxic cells in situ (where pO2 is <1.3%) (21). Because the antibodies against the SC marker Pax7 and pimonidazole are of the same isotype, pimonidazole was i.p. injected into tamoxifen-treated 3-month-old adult Pax7cre/ERT2 R26R CAG-Sun1/sfGFP mice (hereafter referred to as SC-INTACT mice), within which greater than 99% of Pax7+ QSCs were genetically labeled with nuclear membrane–located GFP (nmGFP) in the muscle (Supplemental Figure 1, A–C; supplemental material available online with this article; https://doi.org/10.1172/JCI96208DS1) (22). On myofibers isolated from extensor digitorum longus (EDL) muscles, we detected abundant pimonidazole adducts in over 99% of nmGFP+ QSCs (particularly within the cytoplasm), indicating that QSCs were in a hypoxic state in vivo (Figure 1, A and B). We further confirmed the hypoxic state of QSCs using another hypoxia probe, CCI-103F. In adult WT C57BL/6 mice, we detected CCI-103F adducts in the cytoplasm of approximately 97% of Pax7+ QSCs (Figure 1, C and D). In contrast to SCs, the sarcoplasm of myofibers or myonuclei had little staining for pimonidazole/CCI-103F. Given this clear contrast, we investigated whether pimonidazole is capable of detecting hypoxia within myofibers. EDL myofibers from SC-INTACT mice were cultured ex vivo under hypoxic conditions (1% pO2) and stained for pimonidazole/CCI-103F. The nonhypoxic state of myofibers is consistent with the sarcoplasm of myofibers, and at the periphery of myonuclei, pimonidazole adducts in both the cytoplasm and nuclei of SCs, indicating that QSCs were in a hypoxic state in vivo (Figure 1, A and B). We further confirmed the hypoxic state of QSCs using another hypoxia probe, CCI-103F. In adult WT C57BL/6 mice, we detected CCI-103F adducts in the cytoplasm of approximately 97% of Pax7+ QSCs (Figure 1, C and D). In contrast to SCs, the sarcoplasm of myofibers or myonuclei had little staining for pimonidazole/CCI-103F. Given this clear contrast, we investigated whether pimonidazole is capable of detecting hypoxia within myofibers. EDL myofibers from SC-INTACT mice were cultured ex vivo under hypoxic conditions (1% pO2) and labeled with pimonidazole in the culture media (Supplemental Figure 1D). Under this defined hypoxic condition, we detected pimonidazole adducts in both the cytoplasm and nuclei of SCs, in the sarcoplasm of myofibers, and at the periphery of myonuclei, but not within myonuclei (Supplemental Figure 1E). Thus, QSCs were hypoxic in vivo, whereas the adjacent myofibers were likely not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The nonhypoxic state of myofibers is consistent with the notion that the physiological pO2 of resting skeletal muscle is not hypoxic. The sustained hypoxia in SCs might be permissive for HIF1A stabilization. Indeed, we detected discernible yet weak HIF1A staining in approximately 7% of Pax7+ SCs at 3 dpi, in approximately 43% of SCs at 5 dpi, and in approximately 20% of SCs at 7 dpi, but was undetectable at 9 dpi (Figure 2, C and G). The dynamics of HIF1A expression in SCs appeared to be positively correlated with SC proliferation (Pearson’s correlation coefficient = +0.42), corroborating the reported function of HIF1A in promoting SC proliferation under hypoxia (26).

Genetic ablation of HIF2A in QSCs leads to transient activation, proliferation, and differentiation of SCs. Upon muscle injury, the robust SC proliferation following diminished HIF2A expression implicates a role for HIF2A in the maintenance of SC quiescence. To investigate this potential function of HIF2A, we administered tamoxifen to 3-month-old adult Pax7cre/ERT2 HIF2Afl/fl mice (hereafter referred to as SC-HIF2AKO mice), which resulted in genetic ablation of HIF2A specifically in SCs (Figure 3, A and B). At 10 days post tamoxifen-induced recombination (10 dpr), we found that HIF2A expression was abolished in approximately 80% of SCs in uninjured EDL myofibers (Figure 3, B and C). The myogenic transcription factor MyoD is absent in QSCs yet expressed in activated/proliferative SCs (27). We found that HIF2A ablation led to a remarkable increase in MyoD+ SCs in uninjured myofibers (Figure 3, B and D), suggesting that HIF2A-depleted SCs were spontaneously activated in uninjured muscles.

To confirm that SCs proliferate after HIF2A ablation, we administered EdU to SC-HIF2AKO mice at 11 to 15 dpr (Figure 3E). At 16 dpr, uninjured TA muscles from SC-HIF2AKO mice contained increased numbers of Pax7+ Ki67+ SCs and Pax7+ EdU+ SCs compared with those from control mice (Figure 3, F–H).
Figure 1. QSCs are hypoxic in the niche and express HIF2A, but not HIF1A. (A) Timeline of in vivo pimonidazole labeling in SC-INTACT mice and representative confocal images of uninjured/resting EDL myofibers (n > 50 myofibers from n = 3 mice) showing that nmGFP+ QSCs were pimonidazole+. Scale bars: 50 μm and 10 μm (insets). Inset images show that pimonidazole signals were relatively enriched in the cytoplasm of QSCs. Arrowheads indicate a QSC; asterisks indicate a myonucleus. (B) Percentage of pimonidazole+ QSCs. (C) Timeline of in vivo CCI-103F labeling in C57BL/6 mice and representative images of uninjured/resting EDL myofibers (n > 50 myofibers from 3 mice) showing that nmGFP+ QSCs were CCI-103F+. Arrowheads indicate a QSC; asterisks indicate a myonucleus. Scale bar: 20 μm. (D) Percentage of CCI-103F+ QSCs. (E and F) Representative images of uninjured/resting EDL myofibers from C57BL/6 mice (n > 50 myofibers from 6 mice/group) showing that most QSCs were HIF2A+, but HIF1A+. Scale bars: 10 μm. (G) Percentage of HIF1A+ and HIF2A+ QSCs. Data represent the mean ± SEM.

detected the proliferation markers Ki67 and Edu only in Pax7+ SCs, consistent with the specificity of HIF2A ablation in the SC compartment (Figure 3F).

To investigate whether SCs differentiate following proliferation, we performed SC lineage tracing in Pax7creERT2 HIF2Afl/fl; R26RAG-CamKII-Sun1/sfGFP mice (hereafter referred to as SC-HIF2AKO-INTACT mice) and control SC-INTACT mice, in which nmGFP-labeled SCs may remain as Pax7+ SCs or differentiate into Pax7- myonuclei (Figure 3, I and J). At 16 dpr, we observed that nmGFP-Pax7+ SCs increased in uninjured TA muscles from SC-HIF2AKO-INTACT mice, confirming SC proliferation upon HIF2A ablation; meanwhile, these uninjured, resting muscles also contained increased nmGFP-Pax7+ myonuclei, indicating massive myogenic differentiation of HIF2A-ablated SCs (Figure 3, J and K). The myogenic transcription factor myogenin is specifically expressed in differentiating myogenic cells (2). Uninjured EDL myofibers from SC-HIF2AKO-INTACT mice (16 dpr) contained an increased number of nmGFP+ myogenin+ cells compared with myofibers from control mice (Figure 3L and Supplemental Figure 3A), further confirming that HIF2A ablation in SCs induces myogenic differentiation.

To investigate whether SC apoptosis follows HIF2A ablation, we performed TUNEL labeling of TA muscle sections from SC-HIF2AKO-INTACT and control SC-INTACT mice. For both types of mice, TUNEL+ apoptotic nuclei counted for less than 0.5% of nmGFP+ nuclei (of SCs and SC progeny) at 16 dpr (Supplemental Figure 3, B and C). In contrast, TUNEL+ nuclei were readily detected in nuclease-treated muscle sections (technical positive control; Supplemental Figure 3D). Thus, SCs undergo activation, proliferation, and differentiation, but not apoptosis, in uninjured, resting muscles after HIF2A ablation.

It has been previously reported that the depletion of 1 HIF α-subunit provokes an increase of the other (28). At 16 dpr, we detected weak HIF1A expression in approximately 15% of Pax7+ SCs on uninjured EDL myofibers from SC-HIF2AKO mice, but not in SCs from control EDL myofibers (Supplemental Figure 3, E and F). As HIF2A expression was abolished in more than 90% of SCs at 16 dpr (data not shown), HIF2A depletion appeared not to induce HIF1A compensation in the majority of SCs.

Long-term ablation of HIF2A results in the loss of SC homeostatic self-renewal. The long-lasting regeneration capability of skeletal muscle depends on the self-renewal and homeostatic maintenance of SCs. We next investigated the impact of HIF2A ablation on SC self-renewal. Upon tamoxifen-induced HIF2A ablation, the number of Pax7+ SCs in TA muscles from SC-HIF2AKO mice increased to approximately 110% by day 16, returned to normal levels at 1 month, and fell to approximately 20% of normal levels at 6 months (Figure 4, A and B), revealing that HIF2A ablation impaired SC self-renewal and homeostatic maintenance. Six months after tamoxifen treatment, SC-depleted muscles in SC-HIF2AKO mice showed a discernable increase in myofiber cali-
Figure 2. Muscle repair following eccentric contraction–induced injury is concomitant with dynamic alterations of HIF2A and HIF1A expression in SCs. (A–C) Representative images of EDL myofibers from injured muscles at various time points (n >50 myofibers from 3 mice/group/time point) and stained for Pax7, DAPI, and EdU (A), HIF2A (B), or HIF1A (C). Scale bars: 20 μm. Arrowheads indicate SCs. (D) Number of Pax7+ SCs per myofiber. (E) Percentage of EdU+ SCs in total SCs. (F) Percentage of HIF2A+ SCs in total SCs. (G) Percentage of HIF1A+ SCs in total SCs. Data represent the mean ± SEM.
HIF2A stabilization under normoxia promotes quiescence, self-renewal, and stemness of SCs yet impedes myogenic differentiation. Hypoxia treatment increases the engraftment efficiency of cultured SCs (myoblasts) yet impairs their proliferation and differentiation (29, 30). The above phenotypes of HIF2A-ablated SCs led us to investigate whether HIF2A stabilization under normoxia is sufficient to elicit effects similar to those of hypoxia, which would be beneficial to SC transplantation-based therapy. After 72 hours in normoxic culture (21% pO2), SC clusters formed on myofibers as a result of the activation and proliferation of individual SCs compared with muscles from age-matched controls (Figure 4, C and D), yet with comparable maximal torques (Supplemental Figure 3G). Therefore, HIF2A is essential for the quiescence and long-term homeostatic maintenance of SCs in adult uninjured muscle.

Figure 3. Genetic ablation of HIF2A in QSCs leads to transient activation, proliferation, and differentiation of SCs. (A) Timeline of genetic ablation of HIF2A in QSCs. (B) Representative images of myofibers from SC-HIF2AKO mice and control littermates (n >50 myofibers from 5 mice/group; 10 dpr). Immunofluorescence of Pax7 (red), HIF2A (green), MyoD (purple), and DAPI (blue) staining revealed HIF2A+ MyoD+ and HIF2A+MyoD− SCs (arrowheads) in SC-HIF2AKO and control mice, respectively. Scale bar: 10 μm. (C) Number of HIF2A+ and HIF2A− SCs per myofiber (10 dpr). (D) Number of MyoD+ and MyoD− SCs per myofiber (10 dpr). (E) Timeline characterizing SC proliferation after HIF2A ablation in QSCs. (F) Representative cross-sectional images of TA muscles from SC-HIF2AKO mice and control littermates (n = 6 mice/group; 16 dpr). Immunofluorescence of Pax7 (red), Ki67 (green), EdU (purple), and DAPI (blue) staining revealed an increase in Ki67−EdU+ SCs (arrowheads) in SC-HIF2AKO mice. Scale bar: 20 μm. (G) Number of Ki67− and Ki67+ SCs per TA section. (H) Number of EdU− and EdU+ SCs per TA section. (I) Timeline for tracing SC fates after HIF2A ablation in QSCs. (J) Representative images of TA muscles from SC-HIF2AKO-INTACT and control SC-INTACT mice (n = 6 mice/group; 16 dpr). Immunofluorescence of nmGFP, Pax7, laminin B2, and DAPI revealed increased nmGFP+Pax7+ SCs (arrowheads) and nmGFP+Pax7+ myonuclei (asterisks) in SC-HIF2AKO-INTACT mice. Scale bar: 20 μm and 5 μm (insets). Inset images show that both nmGFP+Pax7+ SCs and nmGFP+Pax7+ myonuclei are adjacent to the basal lamina. (K) Number of nmGFP+Pax7−SCs and nmGFP-Pax7+ myonuclei per TA section. (L) Number of nmGFP+myogenin+ differentiating SCs per EDL myofiber (16 dpr). **P < 0.01 and ***P < 0.005, by 2-sided Student’s t test. Data represent the mean ± SEM.
Unlike the nuclear localization of Pax7 and MyoD, we detected HIF2A in the cytoplasm of SCs, suggestive of its loss of function under normoxia (Supplemental Figure 4A). The cytoplasmic staining was specific to HIF2A, as evidenced by its absence in SC clusters derived from HIF2A-ablated SCs (16 dpr; Supplemental Figure 4B). Like cultured SCs, C2C12 myoblasts cultured under normoxia also expressed a low but detectable level of HIF2A that was mainly present in the cytoplasmic fraction, which is in line with the notion that normoxia destabilizes HIF2A (Supplemental Figure 4C).

To stabilize HIF2A in SCs under normoxia, we transfected cultured SCs with a plasmid expressing GFP (to trace transfection) and a triple mutated form of HIF2A (HIF2ATM, carrying P405A, P530V, and N851A mutations of murine HIF2A), which is resistant to O2-induced hydroxylation and proteasome degradation (Figure 5A) (31). After 72 hours of normoxic culture, control myofibers that were transfected only with GFP had 4.6 GFP+ SC clusters and 4.3 GFP+ SCs per cluster on average (Figure 5, B and C). In contrast, myofibers that were transfected with HIF2ATM-GFP had a markedly reduced number and size of GFP+ SC clusters (~1.9 clusters/myofiber and ~2.5 SCs per cluster on average; Figure 5, B and C). Thus, HIF2A stabilization elicited effects opposite those of HIF2A ablation, supporting the idea that HIF2A promotes SC quiescence and impedes SC proliferation. In comparison, the transfection of a WT form of HIF2A (HIF2AWT) under normoxia (1% pO2), and HIF2ATM transfection did not further repress the proliferation (Supplemental Figure 4E). This observation corroborates with previous findings that hypoxia impairs myoblasts proliferation (17, 29) and implicates a role of HIF2A stabilization in this hypoxic effect.

To assess the impact of HIF2A stabilization on SC self-renewal and differentiation, we stained the normoxia-cultured SC clusters/myofibers for Pax7 and MyoD (Figure 5, A–C). Compared with the control myofibers, HIF2ATM-transfected myofibers had increased numbers of Pax7+MyoD– self-renewing SCs (HIF2ATM, 0.6 vs. control, 0.35 cells per cluster) yet decreased numbers of Pax7+MyoD+ committed myogenic SCs (HIF2ATM, 0.4 vs. control, 2.6 SCs per cluster) (Figure 5C). No such difference was observed after HIF2AWT transfection (Supplemental Figure 4D). Thus, HIF2A stabilization promotes the self-renewal but impairs the myogenic differentiation of SCs.

To confirm that the above effects are SC autonomous, control and HIF2ATM-transfected C2C12 myoblasts were differentiated in 2% horse serum for 5 days (21% pO2) and stained for Pax7 and myosin heavy chain (MyHC) (Supplemental Figure 4G). Myoblasts transfected with the control plasmid (GFP+) were either Pax7+MyHC– (self-renewed reserve cells) or Pax7–MyHC+ (differentiated myocytes). In contrast, cells transfected with HIF2ATM (GFP+) were almost exclusively Pax7+MyHC+ (Supplemental Figure 4G), confirming that HIF2A intrinsically impedes myogenic differentiation but augments self-renewal.

The above observations suggest that HIF2A stabilization may facilitate the engraftment efficiency of SCs after transplantation (remaining as stem cells). To assess this potential effect, we isolated myofibers (and residing SCs as donor cells) from SC-INTACT mice. The specific presence of nmGFP on the SC nuclear membrane allowed us to trace the location and fates of transplanted SCs. We transplanted the myofibers with control and HIF2ATM plasmids (no GFP expression) and transplanted them into cardiotoxin-injured (CTX-injured) TA muscles (1 dpi) of cognate 3-month-old C57BL/6 host mice (Figure 5D). As massive SC transplantation may alter muscle regeneration progression and hence indirectly
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Figure 5. HIF2A stabilization under normoxia promotes quiescence, self-renewal, and stemness of SCs yet impedes myogenic differentiation. (A) Diagram depicting the timeline and plasmids used for HIF2A stabilization in normoxic SC culture. Pound signs denote the number and locations of point mutations in HIF2ATM. (B) Representative images of transfected (GFP) SC clusters on myofibers from C57BL/6 mice (n > 50 myofibers from 7 mice/group). Scale bar: 5 μm. (C) Number of SC clusters, Pax7+ SCs per SC cluster (72 h), and Pax7+MyoD+SCs per SC cluster (72 h). (D) Diagram of the experimental scheme for transplantation of HIF2A-stabilized SCs and tracing of their cell fates in vivo. (E) Cross-sectional images of TA muscles that were transplanted with HIF2ATM- or control plasmid–transfected SCs (n = 5 mice/group; 21 days after the first injury). Immunofluorescence of Pax7 and nmGFP revealed 2 fates of transplanted SCs: engrafted SCs that retained stemness (nmGFP+Pax7+; circles, bottom) and engrafted SCs that differentiated into myonuclei (nmGFP+Pax7–; arrowheads). Scale bars: 20 μm. (F and H) Percentage of engrafted and self-renewed nmGFP+Pax7+ SCs in the total SC pool after the first round (21 dpi; n = 5 mice/group in F) and second round (30 dpi; n = 6 mice/group in H) of regeneration. (G and I) Number of nmGFP+Pax7+ differentiated myonuclei per TA muscle section after the first round (G) and second round (I) of regeneration. *P < 0.05, **P < 0.01, and ***P < 0.005, by 2-sided Student’s t test. Data represent the mean ± SEM.

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SCs per section) (Supplemental Figure 4H) and the percentage of engrafted SCs in the total Pax7+ SC pool (control, 2.5% vs. HIF2A, 4.8%) (Figure 5F), indicating that HIF2A improves SC engraftment efficiency. In contrast, HIF2ATM transfection decreased the number of nmGFP+Pax7+ myonuclei (derived from nmGFP+ donor SCs; control, 43 nmGFP+ myonuclei vs. HIF2A, 32 nmGFP+ myonuclei) (Figure 5G), which echoes the negative effect of HIF2A stabilization on myogenic differentiation in cultured SCs and myoblasts.

The long-term outcome of SC transplantation depends on a fine balance between the self-renewal and differentiation of engrafted SCs; either excessive self-renewal or unrestrained differentiation expectedly leads to a failure to support repetitive muscle repair. Thus, we further investigated whether the skewed self-renewal and differentiation potential of HIF2A-stabilized SCs should eventually enhance the long-term engraftment of these stem cells and their contribution to muscle repair. To this aim, we injured the regenerated TA muscles that received SC engraftment (either control or HIF2ATM-transfected SCs) for the second time (Figure 5D). At 30 dpi, the TA muscles from both groups completed regeneration and contained regenerated myofibers of comparable calibers (data not shown). Compared with control TA muscles, the TA muscles that originally engrafted HIF2ATM-transfected SCs...
Figure 7. Transient pharmacological inhibition of HIF2A in CTX-injured muscle promotes SC proliferation and accelerates muscle regeneration. (A) Timeline of pharmacological inhibition of HIF2A after CTX-induced muscle injury in C57BL/6 mice. (B) Representative images of HIF-c2- or 1% DMSO–treated TA muscles (3 dpi) from C57BL/6 mice (n = 6 mice/group). Immunofluorescence revealed increased Pax7+EdU+ proliferative SCs (arrowheads) and decreased Pax7+EdU– quiescent SCs (circles). Scale bar: 20 μm. (C) Number of Pax7+ EdU+ SCs per TA muscle section 3 dpi (n = 6). (D) Number of Pax7+ SCs per TA muscle section 7 dpi, 10 dpi, and 30 dpi (n = 6 or 7). (E) Representative eMyHC and laminin B2 immunofluorescence in HIF-c2- or DMSO-treated TA muscles 7 dpi and 30 dpi (n = 3). Scale bars: 20 μm. (F) Immunoblots showing the expression levels of HIF2A, eMyHC, MyoD, and tubulin in HIF-c2- or DMSO-treated TA muscles 7 dpi (n = 3). (G) Number of myofibers per TA muscle section 30 dpi. (H) Maximal torques of uninjured TA muscles and HIF-c2- or DMSO-treated TA muscles 7 dpi, 10 dpi, and 30 dpi (n = 6 or 7). (I) Timeline of pharmacological inhibition of HIF2A after CTX-induced muscle injury in C57BL/6 mice and SC-HIF2AKO mice. (J) Representative Pax7 immunofluorescence images of HIF-c2- or DMSO-treated TA muscles from SC-HIF2AKO mice (n = 6 mice/group) 10 dpi and number of Pax7+ SCs per TA muscle section 7 dpi. Scale bar: 20 μm. (K) Representative eMyHC and laminin B2 immunofluorescence images of HIF-c2- or DMSO-treated TA muscles from SC-HIF2AKO (n = 6 mice/group) 7 dpi. Scale bar: 20 μm. *P < 0.05, **P < 0.01, and ***P < 0.005, by 2-sided Student’s t test. Data represent the mean ± SEM.
all enhancement of the contribution to muscle regeneration and self-renewal potential suggests that HIF2A stabilization improves the long-term outcome of SC transplantation.

Genetic ablation of HIF2A transiently improves muscle regeneration but impairs long-term muscle regeneration potential. As HIF2A ablation in QSCs led to transient SC proliferation and myogenic differentiation, we sought to understand whether these effects may improve muscle repair. Stretch-induced injury is of physiological relevance, yet the diminished HIF2A expression in SCs after injury negates the necessity for genetic HIF2A ablation in this injury model. Unlike stretch-induced injury, HIF2A expression dampened but was still present in many SCs after CTX-induced injury (3 dpi; 3-month-old C57BL/6 mice) (Supplemental Figure 5A); meanwhile, most SCs (along with many Pax7– cells) expressed HIF1A, which is in line with reported massive blood vessel damage in CTX-injured muscles (Supplemental Figure 5A) (32). Thus, we examined TA muscle regeneration following CTX-induced injury in SC-HIF2AKO mice and their control littermates, which were administered tamoxifen 8 to 10 days before CTX injury (to deplete HIF2A in SCs from SC-HIF2AKO mice). SC-specific HIF2A ablation increased the number of Pax7+ SCs at 10 dpi in SC-HIF2AKO mice compared with that detected in control mice (control, ~320 SCs vs. SC-HIF2AKO; ~420 SCs per TA section) (Figure 8).
HIF2A, the proliferative SC marker MyoD, and the early regenerator, HIF-c2–treated muscles had reduced expression levels of 7-B-D). By 7 dpi, HIF-c2–treated muscles contained regenerated muscle promotes SC proliferation and accelerates muscle regeneration. The above observations suggest that a transient HIF2A inhibition outcomes of muscle regeneration in SC-HIF2AKO mice echo the transient and long-term effects of HIF2A ablation in SCs, namely HIF2A ablation leads to transient increases in SC proliferation and myogenic differentiation, whereas HIF2A ablation in the long term results in loss of the SC population and regeneration failure. Transient pharmacological inhibition of HIF2A in CTX-injured muscle promotes SC proliferation and accelerates muscle regeneration. The above observations suggest that a transient HIF2A inhibition following hypoxia-burdened muscle injury may benefit regeneration by promoting SC proliferation yet lessening the side effect of SC loss. To transiently inhibit HIF2A in muscle, we intramuscularly administered a HIF2A inhibitor (HIF-c2, CAS no. 1422955-31-4; 5 mg/injection/day over 3 consecutive days), which reduced HIF2A expression in muscle (including SCs) (Supplemental Figure 6A) as previously reported (33). We injected HIF-c2 or the carrier solution (1% DMSO) into CTX-injured TA muscles of adult C57BL/6 mice (daily injections, 1-3 dpi) (Figure 7A). HIF-c2 treatment increased the number of Pax7+EdU+ proliferative myofibers by 3 dp and the number of Pax7+ SCs by 7 dp and 10 dpi (Figure 7, B-D). By 7 dp, HIF-c2-treated muscles contained regenerated myofibers of noticeably increased calibers and decreased expression of eMyHC (Figure 7E and Supplemental Figure 6B). Consistently, HIF-c2-treated muscles had reduced expression levels of HIF2A, the proliferative SC marker MyoD, and the early regeneration marker eMyHC (Figure 7F). Additionally, HIF-c2-treated muscles recovered approximately 53% of the maximal torque measured before injury, whereas the control muscles only recovered approximately 32% of the maximal torque by 7 dpi (Figure 7H), indicating accelerated muscle regeneration. At 30 dpi, when muscle regeneration was completed, both HIF-c2-treated and control muscles contained comparable numbers of Pax7+ SCs and recovered approximately 90% of their maximal torque (Figure 7, D and H), suggesting that the HIF-c2 treatment regimen does not affect SC self-renewal or the future contractile capacity of regenerated muscle. However, regenerated myofibers in HIF-c2–treated muscles had increased myofiber calibers (Figure 7, E and G). HIF-c2–treated muscles had no obvious sign of fibrosis and had a myofiber type composition similar to that of control muscles (Supplemental Figure 6, C-E). Therefore, transient inhibition of HIF2A after CTX-induced muscle injury accelerates the progression of muscle regeneration.

As HIF-c2 treatment is not specific to SCs, we asked whether the beneficial effects are attributable to HIF2A inhibition in SCs. Thus, we repeated the same HIF-c2 treatment regimen (with 1% DMSO as a control) in CTX-injured TA muscles from SC-HIF2AKO mice, in which HIF2A had been genetically ablated in SCs (CTX injury at 10 dp) (Figure 7I). By 7 dp, DMSO-treated muscles from SC-HIF2AKO mice had increased numbers of Pax7+ SCs compared with DMSO-treated muscles from WT mice (Figure 7J). However, in SC-HIF2AKO mice, HIF-c2 treatment did not further increase the number of Pax7+ SCs by 7 dp (compared with DMSO-treated SC-HIF2AKO mice) (Figure 7J). In addition, eMyHC levels and the morphology of regenerating myofibers were comparable in HIF-c2– and DMSO-treated muscles of SC-HIF2AKO mice (Figure 7K). Thus, the beneficial effects of HIF-c2 on muscle regeneration in WT mice were probably attributable to HIF2A inhibition in SCs.

Spryl is a target of HIF2A in QSCs. Spryl1 is highly expressed in QSCs, but not in proliferative SCs, and is essential for the self-renewal and maintenance of SCs (13, 34). Bioinformatic analysis identified 3 conserved HREs within the proximal promoter of the Spryl gene (Supplemental Figure 7A), suggesting that HIF2A promotes SC self-renewal by activating Spryl. In luciferase assays, the HRE-containing Spryl promoter was specifically responsive to stabilized HIF2A, but not stabilized HIF1A, in C2C12 myoblasts (Figure 8A). Arp7a and Pdk3 are specific target genes of HIF2A and HIF1A, respectively (15, 35, 36). Consistently, an Arp7a promoter was specifically responsive to stabilized HIF2A, whereas a Pdk3 promoter was only responsive to stabilized HIF1A.

To confirm that HIF2A at its native expression level directly binds to the Spryl promoter under hypoxia, we performed HIF2A ChIP using 2 × 10^6 mmGFP+ QSC nuclei that were fixed in vivo and isolated from SC-INTACT mice (Supplemental Figure 7B and see Methods). In support of the efficacy and specificity of HIF2A ChIP in QSCs, ChIP-qPCR revealed that a Cav1 promoter, which was previously shown to be associated with HIF2A in hypoxic colorectal epithelial cells (37), was also enriched by 3.4 fold in QSC HIF2A ChIP samples, whereas no enrichment was detected for the Pdk3 promoter (Figure 8B). ChIP-qPCR also revealed an approximately 13-fold enrichment of the Spryl promoter in HIF2A ChIP samples, indicating that HIF2A directly binds to the Spryl promoter in QSCs (Figure 8B).

To investigate whether HIF2A activates Spryl1 in QSCs, we isolated mmGFP+ SCs from SC-INTACT and SC-INTACT-HIF2AKO mice (at 10 dpi) by FACS and performed reverse transcription
quantitative PCR (RT-qPCR). SCs from SC-INTACT-HIF2AKO mice had reduced expression of Hif2a and the QSC markers Calcrl and Cd36 (Figure 8C) (8,38), confirming that HIF2A ablation leads to SC activation. Importantly, HIF2A-ablated SCs expressed decreased levels of Spryl1 and the HIF2A targets Atp7a and Cxcl12 (Figure 8O) (39), indicating that HIF2A transactivates Spryl1 (and Atp7a and Cxcl12) in SCs.

We further investigated whether Spryl1 expression is responsive to HIF2A expression levels in vitro. Under normoxia, 2 HIF2A-targeting shRNAs independently knocked down Hif2a (but not Hif1a) in C2C12 myoblasts, which also decreased the expression of Spryl1, Atp7a, and Cxcl12 (Figure 8D and Supplemental Figure 7C). Likewise, HIF-c2, although not affecting Hif2a mRNA levels, reduced HIF2A protein levels in C2C12 myoblasts as well as decreased the mRNA levels of Spryl1, Atp7a, and Cxcl12 (Figure 8E and Supplemental Figure 7D). Instead, HIF-c2 increased the levels of Hif1a mRNA and HIF1A protein and mRNA levels of the known HIF1A targets Pdk3 and Slc2a1 (also known as Glut1) (Supplemental Figure 7, D and E) (40). Conversely, overexpression of HIF2ATM in C2C12 myoblasts increased the mRNA levels of Spryl1, Atp7a, and Cxcl12 (Figure 8F and Supplemental Figure 7F). Thus, Spryl1 is an HIF2A target gene, and HIF2A transactivates Spryl1 in SCs and myoblasts.

Discussion
This study uncovers a previously unappreciated hypoxic state of SCs in their native niche. Pimonidazole and CCI-103F form non-specific protein adducts when pO2 is less than 1.3% (or 10 mmHg) and have been widely used to determine hypoxic states of stem cells and tumor cells in vivo (41-43). QSCs in healthy resting muscle stained positive for pimonidazole and CCI-103F, indicating that SCs are intracellularly hypoxic. This finding echoes a previous observation that SCs isolated from postmortem muscle retain their viability and regenerative potential (44) and implies that SCs adapt to hypoxia. The pO2 in healthy resting muscle ranges from 3% to 4% (25-34 mmHg) (24, 45, 46). Consistently, we found that the sarcoplasm of myofibers had much less evident staining for pimonidazole and CCI-103F. The distinct pO2 in SCs and myofibers is intriguing, as these 2 types of cells are juxtaposed. It has been reported that SCs are preferentially localized adjacent to capillaries (47). Thus, the hypoxic state of SCs is probably not the result of poor oxygenation in their extracellular milieu, but rather is due to an unknown intrinsic mechanism. Notably, hematopoietic stem cells in the bone marrow are also pimonidazole-, irrespective of the variable distance from the vasculature or local pO2 (1%-4%) and are distinct from adjacent pimonidazole- cells (48, 49). Thus, a hypoxic state seems to be physiologically inherent to some adult stem cell populations, which is suggestive of an obligatory role of hypoxia in stem cell biology.

Mammalian cells sense and cope with hypoxia by stabilizing HIFs (HIF1A, HIF2A, and HIF3A). In line with the hypoxic state, QSCs express HIF2A, but not HIF1A. This finding is novel but not surprising (50). Recent studies demonstrated that HIF1A and HIF2A, although sharing many common targets, often have disparate or even opposite functions (50). In particular, HIF1A promotes glucose uptake, augments glycolysis flux, and reduces glucose oxidation in the TCA cycle under hypoxia or pseudohypoxia, all of which favor biosynthesis and cell proliferation in many types of fast-cycling cells, such as hematopoietic stem cells and many types of cancer cells (50-52). In contrast, HIF2A promotes stem cell characteristics in multiple types of cancer stem cells, and HIF2A reexpression suppresses the growth of soft tissue sarcoma (53-55). In addition, HIF2A enhances the stemness of human embryonic stem cells and the reprogramming efficiency of human induced pluripotent stem cells (15, 41). Thus, it is conceivable that the selective stabilization of HIF2A (but not HIF1A) in SCs ensures their quiescence and stemness. Indeed, HIF2A-ablated SCs proceed to proliferate, differentiate, and eventually become depleted in uninjured muscles, reflecting a pivotal role of HIF2A in the maintenance of this adult stem cell population.

This study also revealed intriguing expression patterns of HIF2A and HIF1A in SCs during muscle repair: after stretch muscle injury, HIF2A was diminished in SCs, followed by SC proliferation and transient HIF1A expression in a subset of SCs; in contrast, after CTX-induced injury, most SCs expressed both HIF2A and HIF1A. The emerging HIF1A expression in SCs apparently correlates with vascular damage in muscle and local hypoxia and ischemia. In support of this view, it has been shown that HIF2A controls chronic hypoxia signaling under medium-level hypoxia, whereas HIF1A mediates acute hypoxia responses under transient extreme hypoxia (53, 54). However, it is notable that the above-mentioned HIF1A/HIF2A dynamics cannot be the sole result of HIF-α stabilization/destabilization in response to tissue oxygenation, as evidenced by the findings that (a) following HIF2A depletion, HIF1A emerged in a small percentage of SCs in uninjured, resting muscles, wherein local pO2 probably remained unchanged; and (b) after stretch-induced muscle injury, most SCs remained pimonidazole-, yet HIF2A expression had diminished. Instead, these observations strongly implicate O2-independent mechanisms in HIF1A and HIF2A regulation in SCs. Notably, the global translation profile of QSCs markedly differs from that of activated SCs (56), and the translation of Hif2a mRNA (but not Hif1a mRNA) is regulated by the insulin receptor/PI3K/mTORC2 pathway and IREBP1 (57, 58). Thus, it is conceivable that HIF1A and HIF2A expression in SCs may be regulated at the translational level during muscle repair, an idea that warrants further investigation in the future.

This study shows consistent evidence of SC-specific gain and loss of function of HIF2A and supports the notion that HIF2A promotes SC quiescence or a return to quiescence (self-renewal), while it impedes the myogenic differentiation of these cells. Given that self-renewal, proliferation, and differentiation divergently affect SC numbers, we found that HIF2A ablation led to distinct consequences in terms of the size of the SC pool and muscle regeneration capacity. Shortly after HIF2A ablation, SCs transiently expanded in number and proceeded to myogenic differentiation, which benefits regeneration, with increased input from the stem cell compartment. In contrast, chronic HIF2A loss in SCs skewed the self-renewal versus differentiation propensities and exhausted the stem cell pool, which was destructive to muscle regeneration.

Reduced stemness and the consequent poor long-term engraftment potential of in vitro-expanded myoblasts presents one of the major hurdles for SC-based therapies for degenera-
five muscle diseases (59). Myoblasts cultured under hypoxia have increased quiescence, self-renewal, and engraftment efficiency (17). In this study, HIF2A stabilization in normoxic culture elicited the same series effects as hypoxia. Intriguingly, HIF2A-stabilized SCs outperformed the control transplanted SCs and the endogenous SCs in terms of their self-renewal capability and overall contribution to the myonuclear compartment after repetitive regeneration. It seems paradoxical that HIF2A promotes SC quiescence and also increases the number of engrafted SCs. It is also apparently conflicting that HIF2A impeded myogenic differentiation, yet HIF2A-stabilized SCs gave rise to more myonuclei. However, it is notable that a small percentage of SCs preferentially undergo asymmetric cell divisions, divide slowly, and withdraw early from the cell cycle (11, 60–62). When transplanted, these SCs have long-term self-renewal ability and can efficiently regenerate injured muscles, which are characteristics of true stem cells (60). In this perspective, the above apparent conflicting observations may actually reflect the increased stemness of HIF2A-stabilized SCs, characterized by the improved ability to maintain a quiescent and undifferentiated stem cell state as well as to sustain a myogenic progenitor pool and support muscle regeneration in the long term. Thus, compared with in vitro hypoxia treatment, HIF2A stabilization could afford lasting stemness enhancement and long-term engraftment of myoblasts in transplantation-based therapies for degenerative muscle diseases.

This study also revealed that Spry1 is a target of HIF2A in SCs. Like Hif2a, Spry1 mRNA is abundant in QSCs and down-regulated in proliferative SCs, and Spry1 ablation in SCs results in a loss of SC quiescence in uninjured muscle and impaired SC self-renewal during muscle regeneration (13, 34). These similar expression patterns and phenotypes suggest a genetic interaction between Spry1 and HIF2A. In this study revealed strong evidence that HIF2A binds to the Spry1 promoter and transactivates Spry1. Interestingly, HIF1A appeared not to transactivate Spry1, which is in line with its absence in SCs. Previous studies showed that HIF1A and HIF2A require distinct coactivators for transcriptional activation (63, 64). Further investigation of specific targets and coactivators of HIF1A and HIF2A in SCs would be instrumental in elucidating their distinct functions in SCs.

Various types of muscle injury involve different levels of vascular damage and eventually have distinct effects on HIF expression in SCs (32). We postulate that the vasculature in regenerative muscle coordinates with SCs via hypoxia signaling to meet distinct needs for SC proliferation upon different types of muscle injury. Understanding the hypoxia signaling in regenerative muscle may shed light on specific therapeutic strategies for muscle injuries complicated with different levels of hypoxia and/or ischemia. In this study, we demonstrate that HIF2A loss of function benefited muscle regeneration after CTX-induced injury, which involved vascular damage and remodeling and heterogeneous HIF2A reduction in activated SCs. HIF2A inhibition augmented SC proliferation and accelerated myogenic differentiation, apparently alleviating a negative impact of hypoxia/ischemia on muscle regeneration capacity. The beneficial effects of HIF2A inhibition are probably not the result of a compensatory expression of HIF1A, given that SC-specific HIF1A ablation has been shown to increase myogenic differentiation without an effect on SC expansion following ischemic muscle injury (18). Additionally, SC-specific ablation of both HIF1A and HIF2A impairs muscle regeneration, with reduced SC expansion and self-renewal (19). A previous study reported that myogenin Cre-mediated HIF2A ablation results in myofiber switching from slow to fast types (65). In addition, muscle regeneration with HIF1A ablation in SCs ends in muscle fibrosis (18). In this study, we did not observe these side effects following transient HIF2A inhibition in regenerative muscle. Given the accelerated recovery of muscle contractile function, transient HIF2A inhibition represents an appealing strategy for improving muscle regeneration after hypoxia/ischemia-involved muscle injury.

Methods

Animal experimental procedures. All animals were housed at 23°C under 12-hour light/12-hour dark cycles with ad libitum access to food and water. All mouse strains were purchased from The Jackson Laboratory (Pax7 CreERT2, no. 017763; R26R<cre>fl<floxed> (R26R<cre>fl<floxed>) no. 021039; and Hif2afl<floxed>, no. 008407).

To induce Cre activity, tamoxifen (20 mg/ml in corn oil) was administered i.p. for 3 consecutive days (100 mg/kg/d). To assess myofiber damage, 1% Evans blue (4 ml/kg in saline) was administered i.p. To assess proliferation, EdU (50 mg/kg in DMSO) was administered i.p. for induction of muscle injury, CTX (0.5 nmol, 100 μl) was injected into TA muscles.

Plasmid construction. For HIF1ATM/HIF2ATM expression, pcDNA3_mHIF2A_MYC and pcDNA3_mHIF1A_MYC plasmids were purchased from Addgene (nos. 44027 and 44028). HIF ORFs were PCR amplified from a mouse cDNA library using the primers HIF2A ORF_S (5’-CACAGCTGACAAAGGAGAAAAGGAG-3’) and HIF2A ORF_AS (5’-GTTGGCCTGGTCAGACAGCAGC-3’) and cloned into the pMEGIP plasmid (pMEGIP; System Biosciences). For HIF2AWT expression, the mHIF2A ORF was PCR amplified from a mouse cDNA library using the primers HIF2A ORF_S (5’-ACAAGCTGACCAAGGAGAAAAGGAG-3’) and HIF2A ORF_AS (5’-GTTGGCCTGGTCAGACAGCAGC-3’) and cloned into the pMEGIP plasmid.

For transplantation of HIF2ATM-transfected SCs and myofibers, the mHIF2ATM ORF was subcloned into the pcDH-CMV-MCS-EF1-Puro plasmid (pCMEP; System Biosciences), which does not express GFP.

For HIF2A knockdown, HIF2A shRNA oligonucleotide no. 1 (5’-TGCATTGGTTACAGTGACCAGATCTGATGGAAAGC-3′) and HIF2A shRNA oligonucleotide no. 2 (5’-TGCATTGGTTACAGTGACCAGATCTGATGGAAAGC-3′) were cloned into pMSCV-P2GM plasmids (Addgene; no. 19750).

In situ detection of hypoxic SCs by hypoxyporphers. Hypoxyporphers-1 (pimonidazole; Hypoxyprobe Inc.; 60 mg/kg in PBS) and Hypoxyporphers-F6 (CCI-103F; Hypoxyprobe Inc.; 60 mg/kg in DMSO) were administered i.p. 1.5 and 3 hours before euthanasia, respectively. To assess Hypoxyporphers-1 in staining hypoxic myonuclei, myofibers from tamoxifen-induced Pax7 CreERT2 R26R<cre>fl<floxed> mice were cultured in a hypoxia chamber (STEMCELL Technologies) supplied with 1% O2 and 5% CO2 for 20 hours and in the presence of Hypoxyporphers-1 (100 μM) for 2 hours.

Maximal contraction force measurement and eccentric contraction-induced injury. The peak isometric torque of the ankle dorsiflexors was assessed as previously described (66). Briefly, the left foot of anesthe-
tized mice was placed on a foot plate attached to a servomotor (model 300C-LR; Aurora Scientific). Two Pt-Ir electrode needles (model E2-12; Grass Technologies) were inserted percutaneously into either side of the peroneal nerve. The ankle joint was secured at a 90° angle. The peak isometric torque was achieved by varying the current delivered to the peroneal nerve at a frequency of 200 Hz and a 0.1-ms square wave pulse. Torque (N-mm) was normalized by the body mass (kg) to account for differences in body size.

To induce injury, the dorsiflexors were subject to 100 electronically stimulated eccentric contractions, during which the foot was passively moved from the 0° position (perpendicular to the tibia) to 20° of dorsiflexion. The dorsiflexors were stimulated at 200 Hz for a 100-ms isometric contraction followed by an additional 50-ms stimulation while moving from 20° dorsiflexion to 20° plantarflexion at an angular velocity of 800°/s. Eccentric contractions were repeated every 10 seconds.

Myofiber isolation, culture, and transfection. EDL myofibers were isolated as described before (67). Briefly, EDL was dissected and digested in 0.2% Type I Collagenase ( Worthington) in DMEM at 37°C for 1.5 hours. Single myofibers were isolated by triturating the digested EDL muscle with polished Pasteur pipettes. Myofibers were cultured in horse serum-coated 24-well plates in DMEM (4.5 g/l glucose) supplemented with 20% FBS, 1% sodium pyruvate, and 1% chicken embryo extract and 1% penicillin-streptomycin. For myofiber transfection, myofibers were cultured for 12 hours and transfected with 2 μg plasmids with Lipofectamine 3000 (Thermo Fisher Scientific) for 6 hours.

SC and myofiber transplantation. Myofibers were isolated from tamoxifen-induced SC-INTACT mice, cultured for 12 hours, transfected with plasmids for 6 hours, and briefly washed. Ten transfected myofibers were loaded into a twenty-eight–gauge insulin syringe and injected into each TA muscle that was damaged by CTX twenty-four hours beforehand.

Myoblast culture, transfection, and differentiation. Primary myoblasts were isolated from 1-week-old C57BL/6 male mice as described previously (68). Primary myoblasts were cultured on collagen-coated dishes in Ham’s F-10 medium supplemented with 20% FBS, 5 μg/l basic FGF, and 1% penicillin-streptomycin. C2C12 myoblasts were purchased from ATCC and cultured in DMEM (4.5 g/l glucose) medium supplemented with 10% FBS and 1% penicillin-streptomycin. For hypoxia treatment, primary myoblasts and C2C12 myoblasts were cultured in a hypoxia chamber supplied with 1% O₂ and 5% CO₂. C2C12 myoblasts were transfected with plasmids using TransIT-X2 (Mirus Bio). C2C12 myoblasts (100% confluent) were differentiated in DMEM (4.5 g/l glucose) medium supplemented with 2% horse serum for 5 days.

Cell-cycle analysis of primary myoblasts. Primary myoblasts were infected with lentivirus packaged from a pCMV5G-HIF2aTM plasmid or a control pCMV5G empty plasmid, selected with puromycin (0.5 μg/ml) for 1 week, fixed with 70% ethanol for 10 minutes, and stained with Hoechst 33342 (10 μg/ml). Myoblasts were run into a HyperCyAn Cytometer (Beckman Coulter), and profiles were analyzed with FlowJo, version 10.

FACS. SC sorting was performed on a MoFlo XDP Cell Sorter (Beckman Coulter), mmGFP-SCs (~2 × 10⁵) from 3-month-old SC-INTACT and SC-HIF2AKO-INTACT mice (7 dpi) were FACS sorted as previously described (68).

Immunofluorescence staining and imaging. For staining of muscle sections, muscle sections were fixed with paraformaldehyde (PFA)/PBS (1%, 10 min), quenched with glycine (50 mM, 10 min), permeabilized with Triton X-100 (0.5%, 10 min), blocked with Mouse On Mouse (M.O.M.) Blocking Reagent (Vector Laboratories) and 5% BSA/5% normal goat serum/PBS, and incubated with the following primary antibodies: anti-Pax7 (1:5; Developmental Studies Hybridoma Bank [DSHB] Pax7); anti-HIF2A (1:250; Novus Biologicals; NB100-122); anti-HIF1A (1:250; Novus Biologicals; NB100-105); anti-Ki67 (1:1,000; Abcam; ab15580); anti-laminin B2 (1:1,000; MilliporeSigma; 05-206); anti–eMyHC (1:50; DSHB; FL652); and anti–MyHC type I, IIA, or IIB (1:50; DSHB; BA-D5; SC-71; BF-F3) overnight at 4°C. For EdU staining, a Click-IT Edu Fluorescence Kit (Life Technologies, Thermo Fisher Scientific) was used according to the manufacturer’s instructions before primary antibody application. Sections were washed in PBS/0.1% Tween-20, incubated with Alexa Fluor-labeled secondary antibodies (1:200, 1 h), and then washed and mounted with DAPI-containing mounting medium (Life Technologies, Thermo Fisher Scientific).

For staining of myofibers and myoblasts, the myofibers and myoblasts were fixed in PFA (4%, 10 min), blocked in 5% BSA/5% normal goat serum/PBS, and incubated with the following primary antibodies: anti-Pax7 (1:50); anti-MyoD (1:250; MilliporeSigma; M6190); anti-HIF2A (1:250); anti-HIF1A (1:250); and anti-GFP (1:1,000; 101Bio; P601). Mounted slides were imaged on a Zeiss LSM 710 confocal microscope.

TUNEL labeling. Muscle sections were stained using a FlowTACS Apoptosis Detection Kit (Thermo Fisher Scientific) following the manufacturer’s instructions. Briefly, sections were fixed by formaldehyde (3.7%, 10 min), washed with PBS, incubated with 20 μl Cytotnin solution (Thermo Fisher Scientific) for 30 minutes, and washed with PBS. At this step, sections assigned for positive controls were further incubated with 25 μl TACS-Nuclease (Thermo Fisher Scientific) solution at room temperature for 30 minutes. All sections were washed with 1× labeling buffer, incubated with the labeling reaction mix (1 μl 50× MnCl₂, 1 μl TdT dNTP mix, 1 μl TdT enzyme, and 50 μl 1× labeling buffer) at room temperature for 1 hour, incubated with 1× stop buffer, and then washed in PBS. All sections were incubated with DAPI and streptavidin–Alexa Fluor 647 (1:200) for 10 minutes in the dark and then washed and mounted for confocal microscopy.

Picrosirius red staining. Muscle sections were stained with Picrosirius red solution (0.1% in picric acid, 15 min), washed with acetic acid (1%, 3 times), and mounted with Cytoseal 60 Mounting Medium (Thermo Fisher Scientific).

SC nuclei isolation. SC nuclei were isolated by isolation of nuclei tagged in specific cell types (INTACT). SC-INTACT mice were administered tamoxifen 1 week before nuclei isolation to induce sfGFP- and Myc-tagged Sun1 localization on SC nuclear membranes. Anti-GFP (2.5 μg; 101Bio; no. P601) and anti-Myc (2.5 μg; MilliporeSigma; no. 05-419) antibodies were incubated with Protein A Dynabeads (50 μl 50% slurry; Thermo Fisher Scientific) in PBS on a rotating platform at 4°C overnight. Cross-link HIF2A in situ on SC chromatin in vivo, SC-INTACT mice were transcardially perfused with 30 ml 0.5% PFA and 30 ml of 125 mM glycine as previously described (69). Limb and trunk muscles were dissected, ground in a mortar with liquid nitrogen, and homogenized (15 ml Wheaton dounce, loose pestle) in homogenization buffer (250 mM sucrose, 2 mM MgCl₂, 25 mM KCl, 1% NP40) for 30 strokes. The homogenate was filtered using 100-μm and 40-μm strainers and centrifuged at 1,000 g at 4°C for 5 minutes to collect crude nuclei. Nuclei were resuspended in 10 ml homogenization buffer and loaded

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on top of a 2-step buffer cushion (lower buffer: 500 mM sucrose, 2 mM MgCl₂, 25 mM KCl and 40% glycerol; upper buffer: 340 mM sucrose, 2 mM MgCl₂, 25 mM KCl, and 40% glycerol). The nuclei were centrifuged in a swing bucket rotor at 1,000 g at 4°C for 10 minutes. Nuclei enriched at the interface were collected with long-neck Pasteur pipettes and checked under a fluorescence microscope. The above homogenization and centrifugation procedure was repeated until most nmGFP⁺ SC nuclei were physically separated from tissue debris or other nmGFP⁻ nuclei. Nuclei were incubated with antibody-coated Dynabeads in INTACT IP buffer (340 mM sucrose, 2 mM MgCl₂, 25 mM KCl, and 5% glycerol) on a rotating platform (8 rpm/min) at 4°C overnight. Nuclei tethered on Dynabeads were isolated on magnetic stands and washed (~6 times) with PBS to remove the nonbound nmGFP⁺ nuclei. After each round of washing, the purity and yield of nmGFP⁺ SC nuclei were assessed under a microscope.

ChIP and ChIP-qPCR. INTACT-purified SC nuclei (with Dynabeads) were resuspended in 100 μl lysis buffer (1% SDS, 10 mM EDTA, 50 mM Tris HCl, pH 8.1, 1× protease inhibitor cocktail) and sonicated at 4°C in a Bioruptor Pico Sonicator (Diagenode) using 15 on/off cycles of 30:30 seconds. The sheared chromatin (~300 bp) was diluted in IP dilution buffer (50 mM HEPES-KOH, pH 7.5, 140 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 0.1% SDS) and centrifuged at 20,000 g at 4°C for 10 minutes. The supernatant was transferred to siliconized tubes and incubated with 2 μg HIF2A antibody (Novus Biologicals) or rabbit IgG (Santa Cruz Biotechnology) on a rotating platform at 4°C overnight. PBS-washed Protein A Dynabeads (30 μl 50% slurry) were added to the chromatin-antibody mixture and incubated on a rotating platform at 4°C for 4 hours. All ChIP beads were sequentially washed with low salt, high salt, lithium, and TE buffer 2 times each and reverse cross-linked with 1 M NaCl for 6 hours at 65°C. Samples were digested with 10 μg RNase A (37°C, 0.5 h) and 20 μg proteinase K (55°C, overnight). Genomic DNA (gDNA) for ChIP analysis was purified by phenol/chloroform/isoamyl alcohol extraction followed by precipitation with 100% isopropanol, 0.3 M sodium acetate, and GlycolBlue (Thermo Fisher Scientific) at –20°C overnight. The protein concentration was quantified by BCA Protein Assays (Thermo Fisher Scientific). Membranes were blocked with 5% nonfat milk/TBST, probed with the primary antibodies anti-HIF2A (1:1,000), anti–HIF1A (1:1,000), anti–HA tag (1:10,000; Thermo Fisher Scientific; 26183), and anti–α-tubulin (1:5,000; MilliporeSigma; T6199), incubated with ECL reagents (Santa Cruz Biotechnology), and exposed to x-ray films.

Luciferase assay. A Spry1 promoter region (~341 to +39 bp relative to the transcription start site (TSS)) and a Pdk3 promoter region (~107 to +101 bp relative to the TSS) were PCR amplified from genomic DNA (C57BL/6 male) and subcloned into pGL4.18 (Promega). pGL4.18_Atp7a has been reported previously (35). pGL4.18-Spry1, pGL4.18-Atp7a, and pGL4.18-Pdk3 (200 ng each) were cotransfected with pcDNA3-HIF1ATM, pcDNA3-HIF2ATM, or a control empty vector into C2C12 myoblasts. pcDNA-LacZ (100 ng) expressing β-gal was cotransfected as an internal transfection control. Forty-eight hours after transfection, myoblasts were lysed with Luciferase Cell Lysis Buffer (NEB), followed by a firefly luciferase/β-gal assay with homemade luciferase and β-gal solutions. The luciferase assay solution consisted of 0.1 M Trizma, pH 7.8, 0.5 M MgCl₂, 0.1 M ATP, and 10 mM luciferin. The β-gal solution consisted of 4 mg/ml O-nitrophenyl-β-D-galactoside, 60 mM Na₂HPO₄, 40 mM NaH₂PO₄, 10 mM KCl, and 1 mM MgSO₄.

Statistics. All values in the figures represent the mean ± SEM and reflect 3 or more biological replicates. Statistical significance was determined with GraphPad Prism 6 (GraphPad Software) using a 2-sided Student’s t test. A P value of less than 0.05 was considered statistically significant.

Study approval. All animal studies were approved by the IACUC of the University of Georgia and performed in strict accordance with institutional guidelines.

See complete unedited blots in the supplemental material.

Author contributions. HY conceived of the study. LX, AY, AMB, JAC, and HY designed the study methodology. LX, AY, ASN, AMB, JAC, and HY performed the experiments. HY wrote the original draft of the manuscript.
script, and LX, AY, AMB, JAC, and HY wrote, reviewed, and edited the manuscript. AMB, JAC, and HY were responsible for resources. HY supervised the study.

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Address correspondence to: Hang Yin, 316 Riverbend Road, Center for Molecular Medicine, Room 2228, the University of Georgia, Athens, Georgia 30602, USA. Phone: 706.583.0655; Email: hyin@uga.edu.

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