RESEARCH ARTICLE

Myosin heavy chain-embryonic regulates skeletal muscle differentiation during mammalian development

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ABSTRACT

Myosin heavy chain-embryonic (MyHC-emb) is a skeletal musclespecific contractile protein expressed during muscle development. Mutations in MYH3, the gene encoding MyHC-emb, lead to Freeman-Sheldon and Sheldon-Hall congenital contracture syndromes. Here, we characterize the role of MyHC-emb during mammalian development using targeted mouse alleles. Germline loss of MyHC-emb leads to neonatal and postnatal alterations in muscle fiber size, fiber number, fiber type and misregulation of genes involved in muscle differentiation. Deletion of Myh3 during embryonic myogenesis leads to the depletion of the myogenic progenitor cell pool and an increase in the myoblast pool, whereas fetal myogenesis-specific deletion of Myh3 causes the depletion of both myogenic progenitor and myoblast pools. We reveal that the non-cell-autonomous effect of MyHC-emb on myogenic progenitors and myoblasts is mediated by the fibroblast growth factor (FGF) signaling pathway, and exogenous FGF rescues the myogenic differentiation defects upon loss of MyHC-emb function in vitro. Adult Myh3 null mice exhibit scoliosis, a characteristic phenotype exhibited by individuals with Freeman-Sheldon and Sheldon-Hall congenital contracture syndrome. Thus, we have identified MyHC-emb as a crucial myogenic regulator during development, performing dual cell-autonomous and non-cell-autonomous functions.

This article has an associated 'The people behind the papers' interview.

KEY WORDS: Skeletal muscle, Myosin heavy chain-embryonic, Myogenesis, Development, Mice, Muscle progenitors, FGF, Signaling

INTRODUCTION

The vertebrate skeletal muscle develops through a complex series of steps involving cell fate determination, cell migration and differentiation. The early myogenic progenitors originate in the somites, expressing Pax3 followed by Pax7 paired box transcription factors, and migrate to their target tissues such as limbs, the diaphragm or the tongue. Commitment to the myogenic lineage occurs as a result of four basic helix-loop-helix transcription factors called myogenic regulatory factors (MRFs) – Myf5, MyoD, MRF4

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and myogenin. Once committed, the progenitors become myoblasts that fuse with each other and with progenitors to generate myofibers, which are tubular multinucleate fibers that contain the contractile machinery essential for muscle contraction. Myofibers are composed of sarcomeres, the functional contractile unit of the skeletal muscle, made up of alternating arrays of thick and thin filaments. MRFs are crucial for the transcriptional regulation of the structural and contractile proteins that form the mature muscle fibers.

During mammalian development, myofibers are generated in two distinct stages, namely the embryonic and fetal phases of myogenesis. During the embryonic phase [embryonic day (E) 9.5-13.5 in mice], a small proportion of the embryonic myogenic progenitors fuse with each other to give rise to primary myofibers, whereas in the fetal phase (E14.5-17.5), fetal myogenic progenitors fuse with each other and with primary myofibers to give rise to secondary myofibers (Biressi et al., 2007a,b; Zammit et al., 2008). Although embryonic and fetal myogenic progenitors differ from each other with respect to their morphology, proliferation rates and response to growth factors, the myofibers derived from them, namely primary and secondary myofibers, respectively, also exhibit distinct characteristics (Biressi et al., 2007b; Stockdale, 1992; Zammit et al., 2008).

The sarcomeric thick filament is primarily composed of myosin, a heterohexamer comprising a pair each of myosin heavy chains (MyHCs), regulatory light chains and essential light chains. MyHCs have an N-terminal globular head region with actin-binding and ATPase activity, a neck region where the light chains bind, and a Cterminal coiled-coil tail region (Schiaffino and Reggiani, 1996). The contractile velocity of muscles is directly correlated with their myosin ATPase activity, varying from muscle to muscle and classified broadly as fast or slow (Barany, 1967). Of the seven mammalian skeletal muscle MyHCs, five are expressed during adult life: MyHC-IIa (encoded by *Myh2*), MyHC-IIx (encoded by *Myh1*) and MyHC-IIb (encoded by Myh4) are adult 'fast' isoforms; MyHC-slow (encoded by Myh7) is the 'slow' isoform; and MyHC-extraocular (encoded by Myh13) is a unique 'fast' isoform expressed in the extraocular and laryngeal muscles (Narusawa et al., 1987; Parker-Thornburg et al., 1992; Weydert et al., 1983; Wieczorek et al., 1985). Mice lacking MyHC-IIb or -IIx are significantly smaller, weigh less than the controls, and exhibit reduced grip strength, muscle weakness and increased interstitial fibrosis (Acakpo-Satchivi et al., 1997). In addition, two developmental isoforms, MyHC-embryonic (encoded by Myh3) and MyHC-perinatal (encoded by Myh8), are expressed during embryonic, fetal and neonatal development (Condon et al., 1990; Periasamy et al., 1984, 1985; Schiaffino et al., 2015; Whalen et al., 1981). Although MyHC-slow is expressed in the adult muscle, it is also expressed during developmental stages in the myofibers. MyHC-embryonic and -perinatal (MyHC-emb and -peri) are not normally expressed in adult muscle, except during skeletal muscle regeneration following injury or disease when they are transiently reexpressed (Sartore et al., 1982; Schiaffino et al., 1986).



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Although several transcriptional regulators and their functions in the distinct phases of myogenesis have been well studied, little is known about the roles of downstream effectors such as MyHCs in myogenic differentiation. This is especially interesting with respect to developmental MyHCs because their expression correlates with specific developmental time points and myogenic phases, and mutations in these MyHCs lead to congenital syndromes. Mutations in the MyHC-emb encoding *MYH3* gene lead to Freeman–Sheldon (FSS), Sheldon–Hall (SHS) and Multiple-Pterygium congenital contracture syndromes, indicating that developmental MyHCs have vital, yet unidentified, roles during development (Toydemir et al., 2006b,c).

In this study, we characterize the role of MyHC-emb in myogenic differentiation during embryonic, fetal, neonatal and postnatal stages of development using targeted mouse alleles. We find that MyHC-emb performs novel cell-autonomous and non-cell-autonomous functions during myogenesis. At the cell-autonomous level, MyHC-emb regulates muscle fiber size, fiber number and fiber type, whereas at the non-cell-autonomous level it regulates myogenic progenitor and myoblast differentiation. MyHC-emb expressed in

myofibers, mediates non-cell-autonomous effects on myogenic progenitors and myoblasts via secreted fibroblast growth factor (FGF) signals. Supplementation with FGF in the culture media rescues the myogenic differentiation defects caused by loss of MyHC-emb function *in vitro*. Adult *Myh3* null mice exhibit scoliosis, one of the phenotypes exhibited by individuals with FSS and SHS. Our results demonstrate that MyHC-emb is a crucial regulator of mammalian myogenesis.

RESULTS

Loss of MyHC-embryonic leads to alterations in myofiber number, area and fiber type

MyHC-emb is expressed during the embryonic and fetal stages of muscle development, and mutations in the *MYH3* gene coding for MyHC-emb protein cause congenital contracture syndromes (Chong et al., 2015; Tajsharghi et al., 2008; Toydemir et al., 2006c). Because few studies have been undertaken to investigate the molecular functions of MyHC-emb, we generated *Myh3* targeted mouse alleles. We generated a conditional $Myh3^{/13-7}$ allele, where exons 3-7 of Myh3 were flanked by LoxP sites (Fig. 1A), and a



Fig. 1. Loss of MyHC-embryonic leads to neonatal myogenic differentiation defects. (A,B) Schematics depicting the $Myh3^{3/3-7}$ allele, where LoxP sites (red arrowheads) flank exons 3-7 (A) and the $Myh3^{4}$ allele lacking exons 3-7 of Myh3 (B). (C-D") Cross-sections through the hind limbs of P0 $Myh3^{+/+}$ (C,C') and $Myh3^{4/4}$ (D,D') mice labeled by immunofluorescence for MyHC-slow (red), laminin (green) and DAPI (blue). 'E' and 'S' denote EDL and soleus muscles, respectively, and 'Tib' and 'Fib' denote tibia and fibula bones, respectively, in C' and D'; C" and D" are magnifications of boxed areas of the flexor digitorum longus (FDL) muscle from C' and D', respectively. (E,F) Laminin (green) labeling on sections through the soleus muscles of P0 $Myh3^{+/+}$ (E) and $Myh3^{4/4}$ (F) mice. (G,H) Quantification of myofiber number through the EDL (G) and soleus (H) muscles of P0 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice. (I,J) Quantification of myofiber area through the EDL (I) and soleus (J) muscles of P0 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice, grouped into 0-50 µm², 50-150 µm² and above 150 µm². (K) Quantification of MyHC-slow, MyHC-lla and β-actin from P0 $Myh3^{+/+}$ and $Myh3^{4/4}$ shank whole cross-section normalized to total area. (L-N) Western blots for MyHC-slow, MyHC-lla and β-actin from P0 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice hind limb protein lysates (L) and densitometric quantification (M,N). Data are mean±s.e.m. of a minimum of three independent experiments. Scale bars: 200 µm (D'); 20 µm (F).

 $Myh3^A$ allele, where exons 3-7 were deleted (Fig. 1B). Both $Myh3^{fl3-7}$ and $Myh3^A$ alleles were verified by PCR (Fig. S1A-C). In crosses between $Myh3^{A/+}$ heterozygous mice, $Myh3^{A/A}$ homozygous pups were not obtained at the expected 25% frequency but at about 14% (17 out of 126), suggesting that approximately half of $Myh3^{A/A}$ genotype animals die *in utero* during embryonic or fetal stages, and that MyHC-emb might have crucial functions during developmental stages. We find that MyHC-emb is expressed in the wild-type fetal heart and is absent in the $Myh3^{A/A}$ heart at E16.5, where it might have essential functions (Fig. S4D) (Rutland et al., 2011). We validated that $Myh3^{A/A}$ animals are null for MyHC-emb using immunofluorescence and qPCR (Fig. S2A-C).

Using $Myh3^{\Delta/\Delta}$ animals that survive, we investigated whether loss of MyHC-emb causes myofiber differentiation defects. We started by quantifying the number of myofibers per unit area in two representative muscles, the extensor digitorum longus (EDL), which is rich in fast fibers, and the soleus, which is rich in slow fibers, at postnatal day 0 (P0). There was a significant increase in the total fiber number in both the EDL and soleus upon loss of MyHCemb (Fig. 1G,H). We also observed that myofiber size was reduced in the soleus in $Mvh3^{\Delta/\Delta}$ animals, whereas it was not affected in the EDL (Fig. 1E,F; Fig. S2D,E). Upon quantification of the myofiber area of representative slow (soleus) and fast (EDL) muscles by grouping myofibers in 0-50 μ m², 50-150 μ m² and above 150 μ m² categories, we observed a significant increase in myofibers with the smallest area (0-50 μ m²) in the soleus upon loss of MyHC-emb, whereas there was no effect on the EDL (Fig. 1I.J). Thus, MyHCemb regulates two crucial muscle fiber properties in neonatal mice: myofiber number in fast and slow muscles, and myofiber crosssectional area in slow muscles such as the soleus.

We next tested whether the loss of MyHC-emb affects muscle fiber type, by labeling $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ P0 mouse hind limb cross-sections with MyHC-slow and laminin (basal lamina marker) (Fig. 1C-D"). We found a striking, almost twofold increase in the number of MyHC-slow⁺ fibers over the whole shank cross-section per unit area in $Myh3^{\Delta/\Delta}$ animals (Fig. 1C-D",K). We validated the increase in MyHC-slow protein levels using western blot and found a dramatic sixfold increase in MyHC-slow protein levels in $Myh3^{\Delta/\Delta}$ animals (Fig. 1L,M). In addition, a fivefold increase in MyHC-IIa protein levels also occurred upon the loss of MyHC-emb, signifying that loss of MyHC-emb leads to changes in levels of other MyHC isoforms and fiber types (Fig. 1L,N). Thus, our findings indicate that MyHC-emb function is crucial for regulating myofiber number, size and type at neonatal stages.

At birth, $Myh3^{\Delta/\Delta}$ pups weighed significantly less than $Myh3^{+/+}$ siblings (Fig. 2A). To test whether the myofiber defects observed at P0 persisted into postnatal stages, we documented the whole-muscle weights of the tibialis anterior (TA), gastrocnemius and quadriceps muscles of $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ mice at P15 and P30 postnatal time points, and found that all three muscles weighed significantly less in $Mvh3^{\Delta/\Delta}$ mice at both time points (Fig. 2B,C). Upon quantification of the myofiber area of P15 (Fig. 2H) and P30 (Fig. 2L) TA muscles, we found a significant increase in the proportion of myofibers with the smallest area (100-500 μ m²) and a significant decrease in the proportion of myofibers with a larger area $(1000-1500 \,\mu\text{m}^2, 1500-2000 \,\mu\text{m}^2 \text{ at P15 and } 1500-2000 \,\mu\text{m}^2,$ 2000-2500 μ m² at P30) in *Mvh3^{Δ/Δ}* mice. In the fast fiber-rich EDL muscle, we found a significant increase in the proportion of myofibers with the smallest area (100-500 μ m²) and a significant decrease in the proportion of myofibers with a larger area (1000-1500 µm²) at P15 (Fig. 2I), whereas no significant differences were observed at P30 (Fig. 2M) in $Myh3^{\Delta/\Delta}$ mice. In the slow fiber-rich

soleus muscle, we found a significant increase in the proportion of myofibers with the smallest area (100-500 μ m²) and a significant decrease in the proportion of myofibers with a comparatively larger area (500-1000 μ m²) at P15 (Fig. 2D,E,J) in *Myh3^{Δ/Δ}* mice. The significant increase in the proportion of myofibers in the soleus with the smallest area (100-500 μ m²) was maintained, although the decrease in the proportion of myofibers with a larger area was not significant at P30 (Fig. 2F,G,N) in *Myh3^{Δ/Δ}* mice. Interestingly, the number of MyHC-slow⁺ fibers per unit area was significantly increased in the TA and soleus but not the EDL at P15 (Fig. 2D,E,K), whereas this increase was not observed at P30 in *Myh3^{Δ/Δ}* mice (Fig. 2F,G,O).

In summary, the observed reduced whole-body weight at P0 persists through the postnatal stages where individual muscles weigh significantly less in $Myh3^{\Delta/\Delta}$ mice. However, although the increase in the number of smaller myofibers and decrease in larger myofibers is apparent in all three muscles at P15 in $Myh3^{\Delta/\Delta}$ mice, the differences are seen in only two muscles at P30. Similarly, although the number of MyHC-slow⁺ fibers increases significantly in two muscles at P15 in $Myh3^{\Delta/\Delta}$ mice, it is not observed in any muscle tested at P30. Overall, these results indicate that several myofiber defects seen at neonatal stages in $Myh3^{\Delta/\Delta}$ mice persist into the postnatal stages, which are at least partially compensated for by P30.

Distinct muscles respond differently to loss of MyHCembryonic

As loss of MyHC-emb led to alterations in crucial myofiber characteristics, we performed a whole-transcriptome RNA-Seq experiment to compare the gene expression profiles of P0 $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ animals, in order to identify specific affected genes and pathways. For this, we used four muscles, the TA, quadriceps, gastrocnemius and diaphragm muscles, which express varying amounts of different MyHC isoforms (Fig. 3A-D). We found 218 genes that were significantly misregulated in Myh3 null muscles (Table S3). Only two genes, *Nfil3* and *Btg2*, were misregulated across any three muscles, and 42 genes were misregulated across two muscles (Fig. 3H; Table S3).

We found that the genes misregulated in two or more muscles were mostly involved in myogenic differentiation. For example, *Btg2*, a transcriptional coregulator misregulated in all four muscles, is involved in muscle development and differentiation (Feng et al., 2007), and Adamts15, a metalloproteinase misregulated in three muscles, is essential for myoblast fusion (Stupka et al., 2013). Transcripts for genes that encode proteins essential for muscle contractility, such as Tpm3, Mvl3 and Mvl2, were misregulated in multiple muscles (Table S3). Interestingly, 12 genes misregulated upon loss of Myh3 were identified in a study to discover transcriptional regulators involved in early muscle differentiation using C2C12 myoblasts (Rajan et al., 2012) (Table S4). Of these 12 candidates, eight genes were misregulated in more than one muscle in our study (Table S4). Six of these candidate genes were tested for their role in early C2C12 differentiation by knockdown assays and were found to be vital for differentiation (Rajan et al., 2012) (Table S4). All of this confirms that MyHC-emb is necessary for normal myogenic differentiation and its absence leads to the misregulation of muscle differentiation-related genes.

The diaphragm has the fewest misregulated genes in common with the three limb muscles, possibly because the diaphragm is a specialized muscle that is anatomically and functionally distinct from the limb (Fig. 3H). Another difference between these muscles



Fig. 2. Loss of MyHC-embryonic leads to postnatal myogenic differentiation defects. (A) Graph showing body weight of P0 $Myh3^{+/+}$ and $Myh3^{4/4}$ pups. (B,C) Graphs showing TA, gastrocnemius and quadriceps muscle weights of $Myh3^{+/+}$ and $Myh3^{4/4}$ animals at P15 (B) and P30 (C). (D-G) Representative images from cross-sections through the soleus of P15 and P30 $Myh3^{+/+}$ mice (D,F), and $Myh3^{4/4}$ (E,G) mice labeled by immunofluorescence for MyHC-slow (red), laminin (green) and DAPI (blue). (H-J) Quantification of myofiber area through the TA (H), EDL (I) and soleus (J) muscles of P15 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice, grouped into 100-500 µm², 500-1000 µm², 1000-1500 µm², dove 1500 µm² (for EDL and soleus), 1500-2000 µm² (for TA), 2000-2500 µm² (for TA) and above 2500 µm² (for TA). (K) Quantification of MyHC-slow⁺ fibers in cross-sections of the TA, EDL and soleus muscles of P15 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice, grouped into 100-500 µm², 1000-1500 µm², above 1500 µm² (for EDL and soleus), 1500-2000 µm² (for TA), 2000-2500 µm² (for TA) and above 2500 µm² (for TA). (L) Quantification of myofiber area through the TA (L), EDL (M) and soleus (N) muscles of P15 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice, grouped into 100-500 µm², 1000-1500 µm², above 1500 µm² (for EDL and soleus), 1500-2000 µm² (for TA), 2000-2500 µm² (for TA) and above 2500 µm² (for TA). (O) Quantification of MyHC-slow⁺ fibers in cross-sections of the TA, EDL and soleus (N) muscles of P30 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice, grouped into 100-500 µm², 500-1000 µm², 1000-1500 µm², dove 1500 µm² (for EDL and soleus), 1500-2000 µm² (for TA), 2000-2500 µm² (for TA) and above 2500 µm² (for TA). (O) Quantification of MyHC-slow⁺ fibers in cross-sections of the TA, EDL and soleus muscles of P30 $Myh3^{+/+}$ and $Myh3^{4/4}$ mice normalized to total area. Data are mean±s.e.m. of a minimum of three independent experiments. Scale bars: 25 µm (E); 50 µm (G).

is that the diaphragm is more mature at birth, as it starts working immediately with the onset of respiration, unlike limb muscles, which start working gradually (Kelly et al., 1991). One hundred and fifty-six genes were uniquely misregulated in a single muscle type, which could be genes that have distinctive roles in the specific muscles, depending on fiber type characteristics, functional properties and anatomical position (Fig. 3H). Next, we validated misregulated candidate genes in the quadriceps and the diaphragm using qPCR (Fig. 3E,F). Certain candidate genes identified by the RNA-Seq experiment were also misregulated upon *Myh3* knockdown during C2C12 myogenic differentiation, indicating that MyHC-emb affects similar processes and pathways *in vivo* and *in vitro* (Fig. 3G).

In summary, genes related to differentiation were misregulated upon loss of MyHC-emb function, indicating that MyHC-emb is crucial for proper myogenic differentiation. Although MyHC-emb is expressed uniformly across muscles, the misregulated genes are not the same in different muscles, possibly because of inherent differences between muscles, which reflects their fiber type and metabolic diversity.

Loss of MyHC-embryonic leads to the misregulation of other MyHC isoforms

Myh3 is tightly linked with other fast *Myh* isoforms, and, therefore, we tested whether loss of MyHC-emb affects the expression of other MyHCs (Allen and Leinwand, 2001; Sartorius et al., 1998) (Fig. S2F). We found that *Myh2* transcript levels were significantly upregulated in *Myh3*^{Δ/Δ} animals (Fig. 3I,J; Fig. S2G,H). *Myh1* transcript levels were significantly upregulated in the quadriceps and diaphragm of *Myh3*^{Δ/Δ} mice (Fig. 3I,J), whereas *Myh8* was significantly upregulated in the quadriceps and gastrocnemius (Fig. 3I; Fig. S2H). Remarkably, *Myh2*, which is upregulated in all of the muscles, is located adjacent to *Myh3* in the *Myh* gene cluster on chromosome 11, whereas *Myh1* and *Myh8* are located downstream of *Myh2* (Vikstrom et al., 1997; Weiss et al., 1999) (Fig. S2F). Thus, loss of MyHC-emb leads to compensatory



Fig. 3. Loss of MyHC-embryonic leads to global misregulation of genes involved in myogenic differentiation. (A-D) Volcano plots depicting results from the RNA-Seq experiment comparing P0 $Myh3^{+/+}$ and $Myh3^{4//4}$ samples for quadriceps (A), TA (B), gastrocnemius (C) and diaphragm (D) muscles. The adjusted *P*-values are on a log10 scale, and significantly up- or down regulated candidates are marked as dark spots on the volcano plot. (E,F) Selected candidate genes from the RNA-Seq were validated by qPCR for quadriceps (E) and diaphragm (F). (G) Candidates from the RNA-Seq were tested on *Myh3* or control siRNA-treated C2C12 cells at day 5. (H) Venn diagram depicting the number of candidate genes obtained and the degree of overlap in the RNA-Seq results comparing P0 $Myh3^{+/+}$ and $Myh3^{4//4}$ muscles. (I,J) Quantification of *Myh* isoform transcript levels by qPCR on P0 quadriceps (I) and diaphragm (J) muscles of $Myh3^{+/+}$ and $Myh3^{4//4}$ mice. The graphical data represent the mean±s.e.m. of a minimum of three independent experiments.

upregulation of other Myh genes in the fast *Myh* gene cluster, with the genes located physically closest to *Myh3* upregulated in more muscles. Transcript levels of *Myh7*, which is not part of the MyHC chromosome 11 cluster, were significantly downregulated in $Myh3^{A/A}$ animals in the TA and quadriceps (Fig. S2G and Fig. 31).

MyHC-embryonic regulates myogenic progenitor differentiation in a non-cell-autonomous manner during development

As loss of MyHC-embryonic led to changes in myofiber size, number and type, as well as the misregulation of genes involved in muscle differentiation at neonatal stages, we hypothesized that the differentiation defects are most likely a result of embryonic- or fetal-specific requirements of MyHC-emb. To decipher the embryonic- and fetal-specific roles played by MyHC-emb, we made use of two Cre drivers, $Pax3^{CreKI/+}$, which causes Cremediated recombination in embryonic and fetal myogenic lineages, and $Pax7^{iCre/+}$, which recombines only in the fetal myogenic lineage (Engleka et al., 2005; Hutcheson et al., 2009; Keller et al., 2004). At E13.5, during embryonic myogenesis, deletion of *Myh3* in *Pax3^{CreKI/+};Myh3^{Δ//I3-7}* led to a significant reduction in the protein levels of the myogenic progenitor marker Pax7 to half that of the control (Fig. 4A,B), whereas levels of the committed myoblast markers MyoD and myogenin were significantly increased (Fig. 4A,B). This indicates that loss of MyHC-emb accelerates the differentiation of myogenic progenitors leading to the depletion of the progenitors, as seen by a reduction in Pax7 levels, and a concomitant increase in differentiated myoblasts, as indicated by the increased levels of the myoblast markers MyoD and myogenin (Fig. 4A,B). MyHC-slow protein levels were upregulated in *Pax3^{CreKI/+};Myh3^{Δ//I3-7}* E13.5 embryos (Fig. 4A).

At E16.5, during fetal myogenesis, the deletion of *Myh3* in $Pax7^{iCre/+};Myh3^{\Delta/fl3-7}$ also resulted in a significant reduction in Pax7 protein levels (Fig. 4A,C). Interestingly, unlike E13.5, the levels of the committed myoblast markers MyoD and myogenin were significantly decreased in $Pax7^{iCre/+};Myh3^{\Delta/fl3-7}$ E16.5 embryos



Fig. 4. MyHC-embryonic non-cell-autonomously regulates myogenic progenitor differentiation during embryonic and fetal myogenesis. (A-G) Western blots for Pax7, MyoD, myogenin, MyHC-slow, caspase 3 and β-actin on protein lysates from hind limbs of E13.5 $Pax3^{Cree/k/l}$; $Myh3^{\Delta/lf3-7}$, E16.5 $Pax7^{iCre/+}$; $Myh3^{\Delta/lf3-7}$ and P0 $Myh3^{\Delta/lA}$, and controls (A); and their densitometric quantification (B-G). (H-I") Immunofluorescence for Pax7 (green) and DAPI (blue) on cross-sections from E16.5 control (H-H") and $Pax7^{iCre/+}$; $Myh3^{\Delta/lf3-7}$ (I-I") embryo hind limbs; H" and I" are magnifications of boxed areas from H' and I', respectively. (J,K) Immunofluorescence for MyoD (green) and DAPI (blue) on cross-sections from E16.5 control (J) and $Pax7^{iCre/+}$; $Myh3^{\Delta/lf3-7}$ (K) embryo hind limbs. (L,M) Quantification of Pax7⁺ myogenic progenitors (L) and MyoD⁺ myoblast numbers (M) normalized to total area from E16.5 $Pax7^{iCre/+}$; $Myh3^{\Delta/lf3-7}$ and the control embryo. Data are mean±s.e.m. of a minimum of three independent experiments. Scale bars: 200 µm (I'); 20 µm (I" and K).

(Fig. 4A,C). MyHC-slow protein levels were upregulated approximately sevenfold in $Pax7^{iCre/+};Myh3^{\Delta/fl3-7}$ E16.5 embryos (Fig. 4A,E), which suggests that the elevated MyHC-slow protein levels and MyHC-slow⁺ fibers seen at P0 (Fig. 1C-D",K; Fig. 4A) are due to increased MyHC-slow protein levels during embryonic and fetal myogenesis. We investigated this further at the neonatal stage by comparing Pax7 protein levels in $Myh3^{+/+}$ and $Myh3^{\Delta/d}$ P0 mouse muscles. This comparison indicated that there was no difference in Pax7, MyoD or myogenin levels between the control and MyHC-emb null animals (Fig. 4A,D). Levels of cleaved caspase 3, a cell death

marker, did not change in $Pax7^{iCre'+}$; $Myh3^{\Delta/fl3-7}$ E16.5 embryos, indicating that decreased levels of the myogenic progenitor and myoblast markers were not due to elevated cell death (Fig. 4A,F). However, increased cell death was evident at P0 in $Myh3^{\Delta/d}$ muscles (Fig. 4A,G and Fig. S5A-B'), which fits well with the increased *Trim63 (MuRF1)* transcript levels, a muscle-specific ubiquitin ligase that targets proteins for degradation, in P0 $Myh3^{\Delta/d}$ muscles by RNA-Seq (Witt et al., 2005).

As we observed reduced Pax7 levels during embryonic and fetal myogenesis, we next examined whether the number of $Pax7^+$

muscle progenitors varied between control and $Pax7^{iCre/+}$; $Myh3^{\Delta/fl3-7}$ embryos (Fig. 4H-I"). We found a drastic \sim 50% reduction in the number of Pax7⁺ myogenic progenitors in Pax7^{iCre/+}:Mvh3^{Δ/fl3-7} embryos (Fig. 4H'',I'',L). Similarly, we also observed a ~60% decline in MyoD⁺ myoblasts in Pax7^{iCre/+};Myh3^{Δ/fl3-7} embryos (Fig. 4J,K,M). To validate this, we used germline Myh3 null E16.5 embryo ($Myh3^{\Delta/\Delta}$) hind limb muscles. We found that both Pax7⁺ progenitors and levels of MyoD were significantly reduced in $Myh3^{\Delta/\Delta}$ embryos, corroborating the fetal-specific loss of MyHCemb results (Fig. S5F,G,H). There were no changes observed in the number of dividing, phospho-histone H3⁺ (PHH3) nuclei between control and $Pax7^{iCre/+}$; $Myh3^{\Delta/fl3-7}$ embryos, indicating that the rate of cell division is not responsible for the observed decrease in Pax7⁺ progenitors and MyoD⁺ myoblasts (Fig. S3A). This confirms that loss of MyHC-emb accelerates the differentiation of myogenic progenitors during fetal myogenesis, and causes the depletion of Pax7⁺ progenitors and MyoD⁺ myoblasts. We did not observe any difference in the number of Pax7⁺ muscle progenitors in $Myh3^{\Delta/\Delta}$ mice at P0, suggesting that the effect of MyHC-emb on Pax7⁺ myogenic progenitors is restricted to embryonic and fetal stages of development (Fig. S3B-D). We carried out semi-quantitative RT-PCR for *Myh3*, comparing cDNA derived from neonatal muscle progenitors (isolated by fluorescence-activated cell sorting) with whole-muscle cDNA, and found that *Myh3* is not expressed in muscle progenitors (Fig. S4B). As MyHC-emb is not expressed in myogenic progenitors or myoblasts and its expression is limited to myofibers (Yoshida et al., 1998), the depletion of progenitor and myoblast pools that we observed must be mediated by secreted signals arising from the myofibers. Thus, MyHC-emb is essential for embryonic and fetal myogenesis by non-cell-autonomously regulating the differentiation rate of myogenic progenitors and myoblasts.

MyHC-embryonic knockdown causes reserve cell depletion and decreased fusion index

To confirm our *in vivo* results, we carried out siRNA-mediated knockdown of *Myh3* during C2C12 myogenic differentiation *in vitro*. We observed a knockdown efficiency of \sim 80% or higher (Fig. 5A,B). As observed *in vivo*, MyHC-slow protein levels exhibited an approximate sevenfold increase upon *Myh3*



Fig. 5. *Myh3* depletion causes reduction in reserve cell number and decreased fusion index. (A-C) Western blots for MyHC-emb, MyHC-slow and β-actin on control and *Myh3* siRNA-treated C2C12 cells over 9 days of differentiation (A), and densitometric quantification (B,C); 'C' and 'S' denote control and *Myh3* siRNA along with the specific day of differentiation (A). (D-E') MyHC (red), phalloidin (green) and DAPI (blue) immunofluorescence on control (D,D') and *Myh3* (E,E') siRNA-treated C2C12 cells at day 5 of differentiation (D' and E' are magnifications from D and E, respectively, with white arrows marking reserve cells). (F) Quantification of reserve cell number per unit area (mm²) from control and *Myh3* siRNA-treated C2C12 cells. (G-J) Western blots for MyoD, myogenin, caspase 3 and β-actin on control and *Myh3* siRNA-treated C2C12 cells during differentiation (G) and densitometric quantification (H-J); 'C' and 'S' denote control and *Myh3* siRNA along with the specific day of differentiation (G). (K) Fusion index of myofibers formed by differentiating myoblasts for 7 days, isolated from *Myh3^{+/+}* and *Myh3^{+/+}* (E').

knockdown at days 5 and 7 of differentiation (Fig. 5A,C). Next, we tested whether the depletion of Myh3 leads to an altered rate of differentiation of myogenic progenitors. To achieve this, we labeled the reserve cell pool, i.e. the population of cycling undifferentiated cells in the differentiated C2C12 culture, using F-actin (Burattini et al., 2004) (Fig. 5D-E'). We found that the number of reserve cells was dramatically reduced by ~85% upon Myh3 knockdown (Fig. 5D-F). We hypothesized that any of three possibilities could explain the reduction in reserve cells: reduced cell proliferation, increased cell death or increased rate of differentiation. We did not observe any change in the number of proliferative PHH3⁺ cells between the control and Myh3 siRNA-treated cells (Fig. S3E,F). Although cell death, as measured by levels of cleaved caspase 3 protein, was elevated on day 1 of Myh3 knockdown, it was significantly less compared with control samples on days 5, 7 and 9 of differentiation (Fig. 5G,J). Thus, we ruled out decreased rate of cell division or increased cell death as being responsible for the reduction in reserve cell number, which must therefore occur as a result of the alteration in the rate of myogenic differentiation. To test this, we analyzed the protein levels of MyoD and myogenin following Myh3 knockdown, and observed an initial upregulation on days 3-5 (except for myogenin at day 3, where we observed a downregulation), followed by a reduction in expression of both at later time points (days 7-9) of differentiation (Fig. 5G-I). This indicates an initial increased rate of differentiation, which depletes the reserve cell pool, which in turn causes a decline in differentiation rate. Again, this effect must be non-cellautonomous as MyHC-emb expression is restricted to myofibers and is not expressed by reserve cells or myoblasts. We found that differentiation was compromised upon Myh3 knockdown, with a significantly lower fusion index in Myh3 siRNA-treated cells, which could be caused by cell-autonomous and non-cellautonomous signals (Fig. S5E). A significant reduction in the fusion index was also observed with differentiating primary myoblasts from $Myh3^{\Delta/\Delta}$ neonatal mice compared with controls (Fig. 5K; Fig. S5C,D). The results from these experiments have striking similarities to the developmental loss of MyHC-emb, with similar effects on the myogenic progenitor and myoblast pool, fiber type, and differentiation.

The non-cell-autonomous effects of MyHC-embryonic on myogenic differentiation are mediated by FGF signaling

To validate whether non-cell-autonomous signals mediated by MyHC-emb are crucial for normal myogenic differentiation, we differentiated C2C12 cells in conditioned media derived from Mvh3 or control siRNA-treated C2C12 cells. Interestingly, we observed that C2C12 cells treated with Myh3 siRNA-conditioned media had larger and increased numbers of myofibers (Fig. 6A,B). The fusion index was significantly increased in cells differentiated in the presence of Myh3 siRNA-conditioned media (Fig. 6C), confirming that non-cell-autonomous, secreted signals mediated by MyHC-emb (expressed in myofibers) are crucial for regulating myogenic differentiation. The distinction between cell-autonomous and non-cell-autonomous effects are apparent in the fusion index experiments where cells were allowed to differentiate following Mvh3 knockdown (both cell-autonomous and non-cell-autonomous effects) (Fig. S5E), compared with differentiating cells in conditioned media from Myh3-silenced cells (only non-cellautonomous effects) (Fig. 6C).

Next, to identify signaling pathways that are misregulated upon loss of MyHC-emb, we carried out a pathway analysis using the RNA-Seq results comparing neonatal $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ muscles. Among the top five misregulated pathways, four, namely IGF1 (insulin-like growth factor 1), ILK (integrin-like kinase), PTEN (phosphatase and tensin homolog) and STAT3 (signal transducer and activator of transcription 3), were related to mitogen-activated protein kinase (MAP kinase) signaling (Table 1).

The FGF pathway, in which signaling is mediated through MAP kinase activation, has been reported to mediate differentiation and



Fig. 6. The non-cell-autonomous effect of MyHC-embryonic on myogenesis is mediated by FGF signaling. (A-C) Immunofluorescence labeling for MyHC (red) and DAPI (blue) on C2C12 cells differentiated for 4 days in conditioned media from control and Myh3 siRNA-treated cells (A,B) and quantification of the fusion index (C). (D) Western blots for p-FGFR4, Spry2, p-Akt, p-Stat3 and β-actin on protein lysates from E16.5 Pax7^{(Cre/+};Myh3^{Δ/ff3-7} and P0 Myh3^{Δ/Δ} compared with controls. (E) Mass spectrometric analysis of secretome showing an abundance of FGF ligands in control and Myh3 siRNA-treated C2C12 cells. The graphical data represent the mean±s.e.m. of a minimum of three independent experiments. Scale bar: 25 µm.

Table 1. RNA-Seq data analysis

Sample number	Canonical pathway	p-value	Percentage of genes*
1	IGF-1 signaling	0.00002	18.6% (18/97)
2	Valine degradation I	0.00005	38.9% (7/18)
3	ILK signaling	0.00006	14.0% (26/186)
4	PTEN signaling	0.00009	16.1% (19/118)
5	STAT3 pathway	0.0001	19.2% (14/73)

Pathway analysis of the RNA-Seq data from P0 $Myh3^{+/+}$ and $Myh3^{4/d}$ muscle samples identifying the major pathways altered, their *P*-value and the percentage of genes affected in the respective pathway upon loss of MyHC-emb.

maintenance of the stem cell pool (Goetz and Mohammadi, 2013; Pawlikowski et al., 2017; Tsang and Dawid, 2004). FGFR4, a FGF family receptor, is central to myogenic differentiation during development, playing a key role in regulating the rate of differentiation of myogenic progenitors and myoblasts (Lagha et al., 2008; Marics et al., 2002). Interestingly, we found that levels of activated forms of known FGF pathway members, p-FGFR4, p-Akt and p-Stat3, were decreased during fetal myogenesis-specific (E16.5) loss of MyHC-emb (Fig. 6D). A similar decrease in the FGF target Spry2 was also observed, indicating that FGF pathway activation is compromised upon loss of MyHC-emb during fetal myogenesis (Fig. 6D). By P0, p-FGFR4 and Spry2 levels were similar to the control, whereas p-Akt and p-Stat3 levels continued to be reduced upon loss of MyHC-emb (Fig. 6D), possibly as a result of the compensatory effect of adult MyHCs, which are expressed by P0. To further confirm the effect of MyHC-emb on the FGF pathway, we carried out a mass spectrometric analysis to identify secreted FGF levels comparing the secretome from control and Myh3 siRNAtreated C2C12 cells. We found that levels of FGFs classically known to activate FGFR4 (especially FGF4, FGF6, FGF8 and FGF17) (Ornitz and Itoh, 2015) were reduced upon Mvh3 knockdown (Fig. 6E). Furthermore, we quantified the transcript levels of FGFs known to bind FGFR4 (FGF1, FGF2, FGF4, FGF5, FGF6 and FGF8) using qPCR in E16.5 limb muscles of Pax7^{iCre/+}; $Myh3^{+/+}$ and $Pax7^{iCre/+}$; $Myh3^{\Delta/fl}$ embryos to identify the specific FGFs that are misregulated. We found that FGF1 and FGF2 are significantly downregulated, and FGF4 is significantly upregulated upon loss of MyHC-emb function (Fig. S4A). These results suggest that FGF signaling mediates the effect of MyHC-emb on myogenic differentiation.

Next, we tested whether FGF modulates the myogenic differentiation defects observed upon Myh3 knockdown. Knockdown of Myh3 led to a drastic reduction in the reserve cell number (Fig. 5D-F) and, therefore, we quantified the reserve cell number in Myh3 siRNA-treated cells grown in FGF-supplemented media. We found reserve cell numbers were restored to wild-type levels upon FGF supplementation, indicating that exogenous FGF can rescue the effect of loss of MyHC-emb function (compare Fig. 7A-C with Fig. 5D-F). A significant increase in the fusion index was observed upon FGF supplementation of Myh3 siRNAtreated C2C12 cells (Fig. 7D). To validate the effect of FGF signaling upon Myh3 knockdown, we also analyzed the levels of the myogenic differentiation markers MyoD and MyoG at days 3, 5 and 7 of culture (Fig. 7E). Interestingly, we found that both MyoD and MyoG levels decreased at days 3 and 5, whereas they increased at day 7, in the presence of exogenous FGF (Fig. 7E-G). This is in contrast to MyoD and MyoG levels increasing initially and decreasing at day 7 upon Myh3 knockdown (Fig. 5G-I). Levels of MyHC-slow, a marker of myogenic differentiation, increased upon FGF supplementation at days 5 and 7 (Fig. S4C). Thus, our results

clearly demonstrate that FGF rescues the differentiation defects observed as a result of loss of MyHC-emb function.

Although surviving $Myh3^{\Delta/\Delta}$ mice looked relatively normal at birth, other than decreased body weight, we observed that by 4-6 weeks of age, all $Myh3^{\Delta/\Delta}$ mice exhibit severe scoliosis, one of the phenotypes seen in individuals with FSS and SHS who have mutations in *MYH3* (Fig. 7H,I).

DISCUSSION

We have characterized the role of MyHC-emb, one of two MyHCs expressed in the skeletal muscle during mammalian development. Although not absolutely required for muscle differentiation, we find that MyHC-emb underpins novel cell-autonomous and non-cell-autonomous functions during myogenesis.

Loss of MyHC-embryonic leads to cell-autonomous effects during muscle development

Loss of MyHC-emb function caused diverse cell-autonomous effects on neonatal muscle fibers, including alterations in fiber number, size and type. Intriguingly, although the effects on fiber number, type and size were not uniform across all muscles, they persisted into postnatal stages and were partially compensated for by ~4 weeks of age, emphasizing the importance of MyHC-emb in myogenesis during development and neonatal stages. Although MyHC-emb is uniformly expressed across muscles, it might have distinct functions in different muscles based on muscle contractile and metabolic properties. Previous studies on adult MyHCs indicate that loss of Myh1 leads to an increase in the proportion of very small and large fibers (Acakpo-Satchivi et al., 1997), and specific adult muscles were affected to varying degrees with respect to fiber size and interstitial fibrosis in Myh4 and Myh1 null mice (Allen et al., 2000, 2001). The RNA-Seq experiment identified genes that are misregulated upon loss of *Myh3*, where 156 genes were uniquely misregulated in any of four muscles, confirming that loss of Myh3 has distinct effects on different muscles. Broadly, the misregulated genes were related to myogenic differentiation or muscle structure, indicating that MyHC-emb has essential roles during development.

MyHC-embryonic regulates embryonic and fetal myogenesis through non-cell-autonomous signals mediated by the FGF pathway

In addition to its cell-autonomous effects, loss of MyHC-emb function led to non-cell-autonomous effects on C2C12 reserve cells, the pool of undifferentiated cells that exhibit characteristics of satellite cells (Alli et al., 2013; Stuelsatz et al., 2010; Yoshida et al., 1998). We also find that MyHC-emb mediates non-cellautonomous effects on myogenic progenitors and myoblasts during embryonic and fetal myogenesis in vivo. Previous studies show that the Pax7⁺ muscle stem cell pool and their differentiation are regulated by signals from differentiated myofibers, including the mechanical force of muscle contraction (Esteves de Lima et al., 2016). Although the myogenic progenitor pool depletion upon loss of MyHC-emb was consistent across embryonic and fetal myogenesis, the myoblast pool behaved differently, possibly as a result of inherent differences between embryonic and fetal myogenic progenitors, as previously described (Biressi et al., 2007a,b; Stockdale, 1992).

Both positive and negative signals regulate the rate of differentiation of myogenic progenitors and myoblasts. We and others have previously shown that extrinsic signals from the



Fig. 7. Supplementation of FGF rescues the effect of loss of MyHC-embryonic on myogenesis. (A-B^{*m*}) Immunofluorescence labeling for MyHC (red), phalloidin (green) and DAPI (blue) on C2C12 cells at day 5, where *Myh3* has been knocked down, treated with control media without FGF (A-A^{*m*}) or with FGF-supplemented media (B-B^{*m*}); A^{*m*} and B^{*m*} are magnifications from A^{*n*} and B^{*n*}, respectively. (C,D) Quantification of reserve cell number per unit area (mm²) (C) and fusion index (D) from *Myh3* siRNA-treated C2C12 cells at day 5, treated with FGF-supplemented media compared with the control. (E-G) Western blots for MyoD, myogenin and β-actin on *Myh3* siRNA-treated C2C12 cells at days 3, 5 and 7 of differentiation, grown in the presence or absence of FGF (E), and densitometric quantification (F-G). The symbols, '-' and '+' denote absence or presence of FGF in the media (E). (H,I) Six-week old *Myh3^{4/A}* mice exhibit scoliosis (I), compared with control *Myh3^{4/A}* animals (H). (J) Model summarizing the cell-autonomous and non-cell-autonomous roles of MyHC-embryonic during embryonic, fetal and neonatal myogenesis, where it regulates FGF levels, which control the rate of myogenic differentiation. Data are mean±s.e.m. of a minimum of three independent experiments. Scale bars: 100 µm (B^{*m*}); 33 µm (B^{*m*}).

muscle connective tissue fibroblasts regulate muscle differentiation and fiber type (Joe et al., 2010; Mathew et al., 2011). The FGF pathway is crucial to myogenesis. FGF signaling has been reported to regulate Pax3, Pax7, MyoD and myogenin expression, as well as terminal differentiation (Edom-Vovard et al., 2001; Groves et al., 2005; Lagha et al., 2008). FGF signaling is also known to regulate the choice between proliferation and differentiation of myogenic precursors (Ben-Yair and Kalcheim,

2005; Marics et al., 2002). Our results indicate that MyHC-emb regulates the differentiation of myogenic progenitors and myoblasts via the FGF signaling pathway, potentially by regulating FGF secretion from the myofibers. Exogenous FGF was able to rescue the myogenic differentiation defects caused by Myh3 knockdown in C2C12 cells, demonstrating that FGF signaling mediates the non-cell-autonomous effects of MyHC-emb on myogenesis.

Recently, we and others described the first animal models for FSS using Drosophila, which has only one skeletal muscle MyHC isoform (Das et al., 2019; Rao et al., 2019). In the current work, we have generated the first mouse model for studying FSS. Individuals with FSS exhibit several clinical symptoms including muscle contractures, dental crowding, scoliosis, cryptorchidism, strabismus and hearing loss (Stevenson et al., 2006). Adult mice null for Myh3 exhibit scoliosis (Fig. 7H,I), a phenotype seen in most individuals with FSS (Stevenson et al., 2006). Detailed characterization of the *Myh3* null mouse is required to verify whether they exhibit other symptoms seen in FSS patients. Whereas individuals with FSS have missense mutations in the MYH3 gene, which are thought to impede the ATPase activity of the protein, our mouse model is a loss of function of Myh3. Strikingly, missense mutations in human FGFR3 have been reported to cause camptodactyly-tall stature-scoliosishearing loss (CATSHL) syndrome, where individuals exhibit camptodactyly and scoliosis, phenotypes that are also observed in FSS and SHS caused by MYH3 mutations, indicating that MyHCemb and FGF signaling might have common roles during development (Toydemir et al., 2006a,c).

Model and conclusions

Based on our findings, we propose that MyHC-embryonic has crucial cell-autonomous and non-cell-autonomous functions during embryonic, fetal, neonatal and postnatal stages of myogenesis. MyHC-emb within myofibers during embryonic and fetal stages is required to regulate FGF levels, which control the rate of differentiation of myogenic progenitors and myoblasts into myofibers (Fig. 7J). Reductions in FGF levels cause accelerated differentiation of progenitors and myoblasts, which depletes the pool of progenitors and myoblasts (a non-cell-autonomous effect of MyHC-emb) (Fig. 7J). In addition, MyHC-emb also regulates levels of other MyHCs, as well as fiber type, number and size (cellautonomous effects of MyHC-emb) (Fig. 7J).

Thus, our study shows for the first time that developmental MyHCs are important regulators that play crucial cell-autonomous and non-cell-autonomous roles during skeletal muscle development. Future work on how the observed phenotypic changes contribute to scoliosis should shed light on the role of MyHC-embryonic in congenital contracture syndromes.

MATERIALS AND METHODS

Mice

Myh3^{fl3-7/+} mice were generated by flanking exons 3-7 of Myh3 with LoxP sites, according to published protocols (Wu et al., 2008). Briefly, an 11.3 kb genomic fragment of mouse Myh3 was recombineered into the pStartK vector from bacterial artificial chromosome clone RP23-67L23 (Children's Hospital Oakland Research Institute - CHORI, California, USA). By a series of recombineering and cloning steps, LoxP sites, FRT-PGKNeo-FRT and restriction enzyme sites were added to the construct and validated (Wu et al., 2008). The targeting construct was transferred to the pWS-TK2 vector, linearized and used for integration into embryonic stem cells (ESC) using positive and negative selection (Wu et al., 2008). Genomic DNA from ~200 ESC clones was screened for 5' and 3' LoxP integration by PCR, and 24 clones were identified as positive for both 5' and 3' targeting events. Two clones were chosen for microinjection into blastocysts. The resulting chimeric animals from these were crossed with C57Bl/6J wild-type mice and the offspring tested for targeting. Once targeted animals were identified, they were bred for 5 or 6 generations to bring them into the C57Bl/6J background, and the neo cassette was removed by crossing with the R26R^{Flpe} mice (Farley et al., 2000). $Myh3^{\Delta/+}$ mice were generated by crossing the $Myh3^{fl3-7/+}$ mice with the ubiquitous Cre-expressing HprtCre mice (Tang et al., 2002). The Myh3^{fl3-7/+} mice were generated at the Transgenic and Gene Targeting Core (University of Utah, Salt Lake City, UT, USA).

Other Cre-drivers used were $Pax3^{Cre}$ (Engleka et al., 2005) and $Pax7^{iCre}$ (Keller et al., 2004). *C57Bl/6J* wild-type mice were used in this study. All of the animal maintenance and experiments were performed according to Institutional Animal Care and Use Committee (IACUC) approved protocols of the University of Utah and the Regional Centre for Biotechnology Institutional Animal Ethics Committee.

Genotyping

Genotyping was carried out by PCR using genomic DNA extracts prepared from mouse ear clips. Primer sequences used for genotyping Myh3 wildtype, floxed and delta alleles are listed in Table S1, under 'Primers used for genotyping'. Briefly, mouse genomic DNA preparation was carried out using the HotSHOT lysis method (Truett et al., 2000). A PCR mastermix was prepared as follows: 7.5 μ l GoTaq G2 Hot Start Green Master Mix (Promega, M7423); 8 μ l of a 5 μ M solution of primers for Myh3 (one forward and two reverse) (0.26 μ M final concentration); 2.1 μ l DNAse free water; and 3 μ l HotSHOT DNA. The PCR cycling conditions were as follows: 95°C for 5 min; 34 cycles of 95°C for 30 s, 60°C for 20 s and 72°C for 40 s; 72°C for 10 min; and 10°C for 10 min. The PCR products were separated on 2% agarose gel, and Myh3 genotypes were determined based on band sizes (delta, 179 bp; wild type, 231 bp; floxed, 279 bp).

Cell culture

C2C12 mouse myoblasts (ATCC, CRL-1722) were cultured and maintained according to ATCC guidelines in growth medium containing DMEM-Dulbecco's Modified Eagle Medium (Gibco, 11995065) supplemented with 10% (v/v) fetal bovine serum (Sigma-Aldrich, F2442) and 2% penicillin-streptomycin (Gibco, 15140122). C2C12 cells were differentiated in differentiation media containing DMEM, 2% (v/v) horse serum (BioAbChem, 72-0460) and 2% penicillin-streptomycin.

For knocking down *Myh3* expression in C2C12 myoblasts, reverse transfection was performed. Briefly, ~30,000 cells were seeded on gelatincoated coverslips (Neuvitro, GG-12) in each well of a 24-well plate (Nunc, 142485) layered with the transfection mix. The transfection mix comprised 100 μ l Opti-MEM (Gibco, 31985070), 50 nM of *Myh3* or control siRNA (Ambion, s70258 and 4390847, respectively) and 2 μ l Lipofectamine RNAiMAX (Invitrogen, 13778150). Cells were cultured in growth medium for 48 h to allow growth to ~80% confluence and efficient transfection. Subsequently, differentiation was induced by replacing the growth medium with differentiation medium. For RNA and protein lysate preparation, the same protocol was used, except that cells were seeded directly in the wells. All treatments were carried out uniformly between control and *Myh3* siRNA-transfected cells.

For the conditioned media experiments, C2C12 cells were cultured in a 24-well dish after treatment with control and *Myh3* siRNA, respectively, and the conditioned media collected at day 5 of plating. Next, \sim 30,000 C2C12 cells were plated on gelatin-coated coverslips in growth media. After 48 h, the media were replaced with conditioned media from control and *Myh3* siRNA-treated cells, and cultured for 4 days, following which coverslips were processed for immunofluorescence analysis.

For the FGF rescue experiments, C2C12 cells cultured in 24-well dishes were treated with *Myh3* siRNA with (for immunofluorescence) or without (for protein lysates) gelatin-coated coverslips. For immunofluorescence analysis, 10 µg/ml FGF2 (Sigma-Aldrich, SRP4038) was added 48 h after plating to half of the wells and the other half were used as untreated controls. For protein lysate preparation, 10 µg/ml FGF2 was added to half of the wells at 48, 72 and 96 h after plating for 3-, 5- and 7-day samples, respectively. The other half were used as untreated controls for each time point.

For primary myoblast culture, hind limb muscles were isolated from $Myh3^{4/4}$ and $Myh3^{4/4}$ mice at postnatal day 0 (P0), minced with surgical blades and then processed for pre-plating as described previously (Goetsch et al., 2015). After 3 h of pre-plating, the nonadherent cells in the supernatant (enriched for myoblasts) were transferred to gelatin-coated coverslips in individual wells of a 24-well dish, where they were grown in growth medium. After 3 days in growth medium, the cells were shifted to differentiation medium and grown for an additional 7 days, following which the coverslips were processed for immunofluorescence.

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RNA isolation, cDNA synthesis and quantitative PCR (qPCR)

RNA was isolated from the TA, quadriceps, gastrocnemius and diaphragm muscles of $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ mice at postnatal day 0 (P0) using the RNeasy Lipid Tissue Mini Kit (Qiagen, 74804). RNA was isolated from C2C12 cells using the RNeasy Mini Kit (Qiagen, 74106), cDNA was prepared using SuperScript III Reverse Transcriptase (Invitrogen, 18080-044) and oligo (dT) (Invitrogen, 58862) according to the manufacturer's protocol. Quantitative PCR was performed using SYBR Green (Applied Biosystems, 4367659) on the ABI 7500 Fast Real Time PCR System (Applied Biosystems) and normalized to Gapdh transcript levels. The genes studied and primers used for qPCR are listed in Table S1. Fgf qPCR primers are from a previous study (Du et al., 2016). The expression of target genes in the mutant muscles and Myh3 siRNA-transfected cells were normalized to that of wild-type muscles and control siRNA-transfected cells, respectively (Livak and Schmittgen, 2001). A minimum of four biological replicates of mutant and knockout mouse muscle samples and three biological replicates of Myh3 and control siRNA transfected samples were used for the expression analysis.

RNA-sequencing

Quadriceps, TA, gastrocnemius and diaphragm muscles from six Myh3+/+ and $Myh3^{\Delta/\Delta}$ animals, respectively, were harvested at postnatal day 0 (P0) and RNA extracted using the RNeasy Lipid Tissue Mini Kit (Oiagen, 74804) according to the manufacturer's protocol. The integrity and concentration of the isolated RNA was verified using the 2200 TapeStation (Agilent). Library preparation was performed using the TruSeq Stranded mRNA sample preparation kit (Illumina) with oligo dT selection according to the manufacturer's protocol and single-end 50 bp reads were generated using a HiSeq 2000 instrument (Illumina). Transcript annotations for mm10 (M_musculus, Dec 2011) were used from Ensembl. Reads were aligned using Pysano and annotated splice junctions generated using USeq (www.sourceforge.net/projects/USeq). Splice junction reads were mapped to genomic coordinates using the SamTranscriptomeParser application in USeq. Differential gene expression was identified using the Defined Region Differential Seq (DRDS) application in USeq, following which paired-sample differential gene expression analysis was performed using DESeq2 (Love et al., 2014). RNA-Seq metrics were generated using Picard's CollectRnaSeqMetrics, samples clustered using custom R scripts (Analysis/ Plots), and significant genes were run in IPA to generate pathway analyses. A Venn diagram was generated to represent the differentially expressed genes identified by RNA-Seq, common between muscles or unique to specific muscles in $Myh3^{\Delta/\Delta}$ knockout animals compared with $Myh3^{+/+}$ animals.

Western blots

For protein isolation, cells were lysed in ice-cold radioimmunoprecipitation assay (RIPA) buffer (Sigma-Aldrich, R0278-500 ml) containing protease inhibitor (Sigma-Aldrich, P8340-5 ml). Protein was isolated from E13.5 $Pax3^{Cre/+};Myh3^{+/+}$ and $Pax3^{Cre/+};Myh3^{\Delta/fl3-7}$, and E16.5 $Pax7^{iCre/+};Myh3^{+/-}$ and $Pax7^{iCre/+}$; $Myh3^{\Delta/fl3-7}$ embryo hind limbs using Qproteome FFPE Tissue Kit (Qiagen, 37623). Total protein was isolated from $Myh3^{+/+}$ and $Myh3^{\Delta/\Delta}$ E16.5 embryo hearts or P0 hind limb muscles by homogenization using the Precellys 24 homogenizer (Bertin Technologies). Quantification of the protein samples was performed using a Pierce BCA Protein Assay Kit (Thermo Scientific, 23225) as per the manufacturer's protocol. Protein samples were separated on a 10% SDS-PAGE and transferred onto a PVDF membrane (Millipore, iPVH00010) at 4°C for 2 h. Western blot was performed using standard procedures: blocking in 5% skimmed milk (Himedia, RM1254-500GM) for 3 h; washes in PBS containing 0.1% Tween 20 (Sigma-Aldrich, P7949-100ML); incubation overnight with primary antibody at 4°C; 2 h incubation with HRP-conjugated secondary antibody at room temperature; and signal detection using the HRP substrate (Millipore, WBLUF0100). Blots were imaged using ImageQuant LAS 4000 (GE). Blots were stripped using standard procedures and reprobed for β-actin. Densitometry was performed to quantify the amount of protein normalized to beta-actin levels using the ImageQuant software. Antibodies used in these experiments are listed in Table S2.

Immunofluorescence and microscopy

Embryos at the appropriate stages were harvested from timed matings (E13.5 $Pax3^{Cre/+};Myh3^{+/+}$ and $Pax3^{Cre/+};Myh3^{\Delta/fl3-7}$ embryos, and E16.5

Pax7iCre/+;Myh3+/+ and Pax7iCre/+;Myh3^(13-7), fixed and embedded in Optimal Cutting Temperature (OCT) (Tissue-Tek). Hind limbs from $Myh3^{\Delta/\Delta}$ and $Myh3^{+/+}$ neonates at P0 were embedded in OCT and flash frozen in 2-methyl butane cooled in liquid nitrogen. Specific hind limb muscles from $Myh3^{\Delta/\Delta}$ and $Myh3^{+/+}$ mice at P15 and P30 were also embedded in OCT and flash frozen in 2-methyl butane cooled in liquid nitrogen. Samples were sectioned at 10 µm using a cryomicrotome (Thermo Scientific; Microm HM 550) and adjacent sections collected on coated glass slides (VWR, VWRU48311-703). Sections from the middle of the limb (P0) or the muscle (P15 and P30) were processed as described below with adjacent sections used for detection of MyHC-emb, MyHC-slow, Pax7, MyoD, Laminin and PHH3. For immunofluorescence, tissue sections on slides or C2C12/primary cells on coverslips were fixed in 4% paraformaldehyde (PFA) for 20 min and washed with PBS. For tissue sections, the antigen was retrieved wherever required (specified in Table S2) by heating samples to 120°C for 5 min in citrate buffer (1.8 mM citric acid and 8.2 mM sodium citrate in water) in a 2100 PickCell Retriever (Aptum Biologics). Tissue sections and cells were blocked with 5% goat serum (BioAbChem, 72-0480) in PBS containing 0.1% Triton-X-100 (MP Biochemicals, 194854) for 1 h at room temperature, incubated overnight at 4°C in an appropriate concentration of primary antibody (Table S2), rinsed three times with PBS, incubated with secondary antibody for 2 h at RT, and rinsed with PBS. Where amplification was required, the samples were incubated with a biotin-conjugated secondary antibody for 2 h at room temperature and then with Strep-coupled fluorophore for 1 h at room temperature, and washed three times with PBS. Samples were postfixed in 4% PFA, rinsed in distilled water and mounted using DAPI Fluoromount-G (Southern Biotech, 0100-20). Fluorescence microscopy was performed using a Leica TCS SP5 II or Nikon A1R confocal microscope.

Preparation of secretome proteins and mass spectrometric analysis

Cells were cultured for 48 h in complete medium initially, following which it was replaced with serum-free medium. All traces of serum were removed by rinsing the cells three times with PBS before adding the serum-free medium. Medium for the secretome analysis was collected after 48 h of culture. Samples were centrifuged at 150 g for 5 min at 4°C to remove intact cells and debris. Protease inhibitor was added and the media were concentrated by ultrafiltration using the Amicon Ultra-15 Centrifugal Filter Unit (Millipore, UF C905024). Proteins were precipitated with cold acetone at -20° C overnight, and collected by centrifugation at 19,000 g for 1 h at 4°C. Pellets were air-dried at 37°C, solubilized in 4× Laemmli buffer and boiled for 10 min to denature proteins. Proteins were separated by 10% SDS-PAGE, stained with Coomassie Brilliant Blue R-250 and protein bands excised using sterile blades. The gel slices were destained, dehydrated and digested with mass spectrometry grade trypsin (Pierce Trypsin protease; Thermo Scientific, 90057) at 37°C overnight. Peptides were extracted by ultrasonication in extraction solution (50% Acetonitrile/5% trifluoroacetic acid) and the extract was dried in a vacuum concentrator at RT. The samples were desalted with modified Pierce C18 Zip-Tips (Thermo Scientific, 87 782) and loaded on a LC-MS/MS mass spectrometer (Triple TOF 5600 Sciex). The raw data of MS-MS (wiff and mgf files) were processed and subjected to database searches using Search GUI. Search results were processed, combined and interpreted in the form of emPAI (exponentially modified Protein Abundance Index) using Peptide Shaker, run with a false discovery rate of 1%.

Cell counts, fiber counts and statistics

For the fusion index analysis, five randomly selected nonoverlapping fields of view (2×3 tiles) per coverslip were imaged using identical settings. Myotubes were labeled for myosin heavy chain using a mixture of My32 and MyHC-slow antibodies and nuclei by DAPI. Counts for total nuclei, number of myotubes and nuclei within the myotubes per unit area, were performed using ImageJ (Schindelin et al., 2012; Schneider et al., 2012). The fusion index was calculated as the percentage of nuclei within myotubes (myotubes with at least two nuclei), compared with the total number of nuclei. For reserve cell counts, *Myh3* and control siRNA-treated cells were cultured, stained using phalloidin, myosin heavy chain antibodies (mixture

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of My32 and MyHC-slow antibodies) and DAPI (to identify nuclei), after which images were captured. The total number of DAPI⁺ nuclei and phalloidin⁺ reserve cells were counted using spot and annotation functions in Imaris software (www.bitplane.com/), and normalized to unit area. PHH3⁺ nuclei were counted manually and the total DAPI⁺ nuclei were counted using the particle analyzer function in ImageJ (Schindelin et al., 2012; Schneider et al., 2012).

Pax7⁺, MyoD⁺ and PHH3⁺ cell counts from the entire P0 or E16.5 hind limb cross-section was performed using the spot function, and total P0 or E16.5 hind limb muscle cross-sectional area was quantified using the surface function in Imaris software. MyHC-slow fiber count was performed using the fiber typing function, and fiber area using the fiber properties function with the semiautomatic muscle analysis using segmentation of histology software (SMASH) (Smith and Barton, 2014).

Data from all of the experiments were analyzed with parametric unpaired *t*-tests using the GraphPad Prism software. Data are mean \pm s.e.m. The *P*-value is indicated on each graph and **P*-value ≤ 0.05 is considered significant.

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Competing interests

The authors declare no competing or financial interests.

Author contributions metadata

Conceptualization: M.A., G.K., S.J.M.; Methodology: M.A., A.S., P.K., A.K., A.B., M.S., G.K., S.J.M.; Validation: M.A., A.S., P.K., A.K., A.B., M.S., S.J.M.; Formal analysis: M.A., A.S., P.K., A.K., A.B., M.S., S.J.M.; Investigation: M.A., A.S., P.K., A.K., A.B., M.S., G.K., S.J.M.; Resources: S.J.M.; Data curation: M.A., A.S., P.K., A.K., A.B., M.S., S.J.M.; Writing - original draft: M.A., A.S., P.K., A.K., A.B., M.S., S.J.M.; Writing - review & editing: M.A., P.K., M.S., S.J.M.; Supervision: G.K., S.J.M.; Project administration: G.K., S.J.M.; Funding acquisition: G.K., S.J.M.

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Data availability

RNA-Seq datasets have been deposited in GEO under accession number GSE100331.

Supplementary information

Supplementary information available online at http://dev.biologists.org/lookup/doi/10.1242/dev.184507.supplemental

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Supplementary Data





Fig. S1: Verification of *Myh3* **targeting and deletion.** (A) 5' LoxP targeting in *Myh3*_{fl3-7/+} mice was verified by PCR across the LoxP site; animal numbers 1, 3 and 4 show two bands, with the larger molecular weight band resulting from incorporation of the LoxP site and the smaller molecular weight band from the wild type homologous chromosome, with animals 2 and 5 being wild type. (B) 3' LoxP targeting in *Myh3*_{fl3-7/+} mice was verified by PCR for Neomycin cassette, which is part of the 3' targeted locus, in the same 5 animals tested in A for 5' LoxP targeting; animal numbers 1, 3 and 4 are positive for Neomycin while 2 and 5 are negative. (C) Deletion of *Myh3* exons 3-7 in the *Myh3*_{d/+} animals was verified by PCR with 3 primers, 2 amplifying across the deleted region to give a product and the third primer giving a product in cases where there is no deletion. Results show that animals 1 and 5 have the deletion (smaller molecular weight deleted band and larger molecular weight wild type band as these are heterozygotes), while animals 2, 3 and 4 are wild type (single larger molecular weight band).





Fig. S2: Validation of *Myh3* **knockout.** (A-B') Cross sections through hind limbs of P0 $Myh3_{+/+}$ (A, A') and $Myh3_{4/4}$ (B, B') mice labeled by immunofluorescence for Laminin (red), MyHC-emb (green), and

DAPI (Blue). (C) Quantitation of *Myh3* transcript levels by qPCR in P0 gastrocnemius, tibialis anterior, quadriceps and diaphragm muscles of $Myh3_{+/+}$ and $Myh3_{4/4}$ mice. (D-E) Laminin (green) labeling through the EDL muscles of P0 $Myh3_{+/+}$ (D) and $Myh3_{4/4}$ (E) mice. (F) Schematic showing the *Myh* gene cluster on mouse chromosome 11 which is ~293 kb in size, where Myh3 is followed by Myh2, Myh1, Myh4, Myh8 and Myh13. (G-H) Quantitation of transcript levels of other Myh isoforms by qPCR in the P0 TA (G) and gastrocnemius (H) muscles of $Myh3_{+/+}$ and $Myh3_{4/4}$ mice. The graphical data represent the mean ±SEM of a minimum of 3 independent experiments. (Scale bar in B' is 100 microns and E is 20 microns).



Fig. S3: Loss of MyHC-emb function does not affect cell proliferation. (A) Quantification of phosphohistone H3+ cells at E16.5 in $Pax7_{iCre/+};Myh3_{\Delta/fl3-7}$ and control hind limbs normalized to total area. (B-C') Pax7 (red), Laminin (green) and DAPI (blue) immunofluorescence on $Myh3_{+/+}$ (B-B') and $Myh3_{\Delta/\Delta}$ (C-C') knockout P0 hind limb sections. (D) Quantification of Pax7+ cell number normalized to total area in P0 $Myh3_{+/+}$ and $Myh3_{\Delta/\Delta}$ hind limb sections. (E-E') Representative images of control siRNA (E) and Myh3siRNA (E') treated C2C12 cells labeled for PHH3 (green) and DAPI (blue) at day 3 of differentiation. (F)

Quantification of PHH3₊ cells in control and *Myh3* siRNA treated C2C12 cells at days 3, 5 and 7 of differentiation. The graphical data represent the mean \pm SEM of a minimum of 3 independent experiments. (Scale bar in C' is 40 microns and E' is 50 microns).



Fig. S4: FGF levels are misregulated upon loss of MyHC-emb function and FGF supplementation promotes differentiation. (A) Quantitation of transcript levels of FGFs known to bind FGFR4 by qPCR in E16.5 limb muscles of $Pax7_{iCre/+};Myh3_{+/+}$ and $Pax7_{iCre/+};Myh3_{4/f}$ embryos. (B) MyHC-emb transcripts are not detectable in muscle stem cell cDNA as opposed to muscle cDNA by semi-quantitative RT-PCR; 1 denotes positive control (GAPDH) and 2 denotes *Myh3* PCR. (C) Western blots for MyHC-slow and beta-actin on *Myh3* siRNA treated C2C12 cells at days 5 and 7 of differentiation, grown in the presence or absence of FGF, with '-' and '+' denoting absence or presence of FGF in the media. (D) Western blots for MyHC-emb, and beta-actin from E16.5 *Myh3*+/+ and *Myh3*_A/_A embryo heart protein lysates. The graphical data represent the mean ±SEM of a minimum of 3 independent experiments.





Fig. S5: *Myh3* null mice exhibit increased cell death, reduced fusion index and reduction in muscle progenitors. (A-B') Immunofluorescence for My32 (red), Caspase 3 (green), and DAPI (blue) on P0 $Myh_{3+/+}$ (A-A') and $Myh_{3\pm/-}$ (B-B') embryo hind limb cross sections; the left panels show Caspase 3 (A, B) and the right panels are the merge (A', B'). (C-D) Representative immunofluorescence images for MyHC (red) and DAPI (blue) on primary myoblasts, isolated from $Myh_{3+/+}$ (C) and $Myh_{3\pm/-}$ (D) P0 mice, differentiated for 7 days, for which fusion index is shown in Figure 5K. (E) Fusion index of control and Myh_3 siRNA treated C2C12 cells at days 3, 5 and 7 of differentiation. (F) Western blots for MyoD, and beta-actin from E16.5 $Myh_{3+/+}$ and $Myh_{3\pm/-}$ embryo hind limb protein lysates. (G-G') Immunofluorescence for Pax7 (green), and DAPI (blue) on cross sections through E16.5 $Myh_{3+/+}$ (G) and $Myh_{3\pm/-}$ (G') embryo hind limbs. (H) Quantification of Pax7+ cell number normalized to total area in E16.5 $Myh_{3+/+}$ and $Myh_{3\pm/-}$ hind limb sections. The graphical data represent the mean ±SEM of a minimum of 3 independent experiments. (Scale bar in B' is 20 microns, D is 100 microns and G' is 25 microns respectively).

Table S1: List of primers

Gene	Direction	Primer sequence 5' to 3'	location	Product size
		-		(bp) using
				cDNA
				template
	1	Primers used for qPCR analysis		1
<i>Myh1</i> (MyHCIIx)	Forward	CGGTGGTGGAAAGAAAGG	Exon17	154bp
	Reverse	CAGGAGTCTTGGTTTCATT	Exon18	
<i>Myh2</i> (MyHCIIa)	Forward	CCAAGAAAGGTGCCAAGAAG	Exon17	147bp
	Reverse	CGGGAGTCTTGGTTTCATTG	Exon18	
Myh3 (MyHCemb)	Forward	ATGAGTAGCGACACCGAGATG	Exon3	117bp
	Reverse	AAAGCAGTAGGTTTTGGCAT	Exon3	
Myh4 (MyHCIIb)	Forward	GCTTGAAAACGAGGTGGAAA	Exon40	190bp
	Reverse	CCTCCTCAGCCTGTCTCTTG	Exon41	
<i>Myh7</i> (MyHCslow)	Forward	AGGGCGACCTCAACGAGAT	Exon32	92bp
	Reverse	CAGCAGACTCTGGAGGCTCTT	Exon32	
Myh8 (MyHC-peri)	Forward	AACAGAAACGCAATGCTGAGG	Exon38	135bp
	Reverse	TCGCCTGTAATTTGTCCACCA	Exon39	
MyoD	Forward	GCTGCCTTCTACGCACCTG	Exon1	119bp
	Reverse	GCCGCTGTAATCCATCATGC	Exon2	
Myogenin	Forward	CAGTACATTGAGCGCCTACAG	Exon1	163bp
	Reverse	GGACCGAACTCCAGTGCAT	Exon2	_
Myf5	Forward	CCTGTCTGGTCCCGAAAGAAC	Exon2	130bp
	Reverse	GACGTGATCCGATCCACAATG	Exon3	_
Mrf4	Forward	ATCAGCTACATTGAGCGTCTACA	Exon1	173bp
	Reverse	CCTGGAATGATCCGAAACACTTG	Exon2	_
Pax7	Forward	TGTTGGGCTCTTCAAGGTCT	Exon9	130bp
	Reverse	GGAATGTGGAGGAGGATGC	Exon9	_
GAPDH	Forward	GACTTCAACAGCAACTCCCACT	Exon6	169bp
	Reverse	GGTCCAGGGTTTCTTACTCC	Exon7	-
Fgf1	Forward	GTAGTTTCCTAGAGGCAGGTTG		
	Reverse	TGATAAAGTGGAGTGAAGAGAGC		
Fgf2	Forward	GAAACACTCTTCTGTAACACACTT		
	Reverse	GTCAAACTACAACTCCAAGCAG		
Fgf4	Forward	ACTCGTCGGTAAAGAAAGGC		
	Reverse	GACACGAGGGACAGTCTTC		
Fgf5	Forward	AACTCCTCGTATTCCTACAATCC		
	Reverse	CGGATGGCAAAGTCAATGG		
Fgf6	Forward	CTGTACACAACGCCCAGCTT		
	Reverse	TTGTTTGGAAGGAGGGTTTCTC		
Fgf8	Forward	CATGGCAGAAGACGGAGAC		
	Reverse	ACTCGGACTCTGCTTCCAAA		
	•	Primers used for RNAseq validation		•
Tpm3	Forward	CCCTGAGCCCAAACTTAT	Exon9	173bp
· ·	Reverse	GGCGATGAGATGATGTTC	Exon9	±
Ppargc1a	Forward	TCACACCAAACCCACAGAAA	Exon5	127bp
	Reverse	GGTCAGAGGAAGAGATAAAG	Exon6	±

Table S1: List of primers

Lpin1	Forward	CAAACAAGCCAGTGACAACG	Exon10	132bp			
	Reverse	AGGGAGATGGCGATGGAT	Exon10				
Ankrd2	Forward	CATTTTCTTTCCCTGGGCTTGG	Exon7	230bp			
	Reverse	TTCTGCTCTGATTCTGGCTCGG	Exon9				
Myl2	Forward	GTCCACATCATTACCCACGG	Exon7	118bp			
	Reverse	AGAGCCAAGACTTCCTG	Exon7				
Adamts15	Forward	GAGACACAACCCAAACAAGT	Exon4	152bp			
	Reverse	CCTCGCAGTATTTTCCACCG	Exon5				
Cebpb	Forward	CGGGTTTCGGGACTTGATGC	Exon1	127bp			
	Reverse	ACCCCGCAGGAACATCTTTA	Exon1				
Csrp3	Forward	ACCACAAGCAACCCTTCCAAAT	Exon4	255bp			
	Reverse	GTGTAAGCCCTCCAAACCCAAT	Exon6				
Btg2	Forward	GCTGCTTTGTATGGGTGGAT	Exon2	231bp			
	Reverse	AAAATGGGGAAGGTTGCTCT	Exon2				
Myl3	Forward	CGGGAAGGAGTGGTTCGGAC	Exon7	104bp			
	Reverse	AAAGGCAAGCACAGGTAGGT	Exon7				
Trim63	Forward	GGTGCCTACTTGCTCCTTGT	Exon3	190bp			
	Reverse	ATTCTCCTTGGTCACTCTGC	Exon5				
Pde4d	Forward	GCTTTGGAGGCTGTGTTCA	Exon13	121bp			
	Reverse	CGAGTTCCGAGTTTGTATTG	Exon14				
	Primers used for Genotyping						
Myh3	Forward	CGTCTGAGAGGCTTCCATTC	Wildtype	231bp			
	Reverse	TAGGTTTTGGCATCAAAGGG	Floxed	279bp			
	Reverse	TATCCTTCACGCTCTCCCAC	Delta	179bp			

Table S2. Antibodies used for immunofluorescence and western blots

Antibody	Туре	Source	Product No.	Working Concentration (ug/ml)	Antigen retrieval for tissue sections
Primary antibodies		·			
Pax7	Mouse IgG1	Developmental Studies Hybridoma Bank	PAX7	2.4	Yes
MyHCemb	Mouse IgG1	Developmental Studies Hybridoma Bank	F1.652	3	Yes
MyHCslow	Mouse IgG1	Sigma	M8421 (NOQ7.5.4D)	1.5 (IF), 6 (western)	Yes
MyHCfast	Mouse IgG1	Sigma	M4276 (MY-32)	10	Yes
MyoD	Mouse IgG1	Santa Cruz Biotechnology	sc-32758 (5.8A)	4 (IF), 2 (western)	Yes
MyoG	Mouse IgG1	Santa Cruz Biotechnology	sc-12732 (F5D)	2	Western only
Myf5	Rabbit polyclonal	Santa Cruz Biotechnology	sc-302 (C-20)	1	Western only
Caspase 3	Rabbit IgG	Cell Signaling Technology	9664 (Asp175) (5A1E)	0.02	No
Phospho-Histone H3	Rabbit IgG	Thermo Fisher Scientific	PA5-17869 (Ser10)	0.29	Yes
β-Actin	Mouse IgG2b	Cell Signaling Technology	3700 (810D10)	1.25	Western only
Laminin	Rabbit polyclonal	Sigma	L9393	2.5	Yes
Phospho-p42/44 MAPK (Thr202/Tyr204)	Rabbit IgG	Cell Signaling Technology	4370	0.01	Western only
Phospho-p38 MAPK (Thr180/Tyr182)	Rabbit IgG	Cell Signaling Technology	4511	0.02	Western only
Phospho-Akt (Ser473)	Rabbit IgG	Cell Signaling Technology	4060	0.02	Western only
Phospho-Stat3 (Tyr705)	Mouse IgG1	Cell Signaling Technology	4113	0.01	Western only

Secondary antibodies, fluorescent coupled conjugates							
Cy2/Cy3 conjugated	Goat	Jackson ImmunoResearch Laboratories	115-225-146/ 115-165-	7.5	IF only		
Goat anti-mouse			146				
Cy2/Cy3 conjugated	Goat	Jackson ImmunoResearch Laboratories	111-225-144/ 111-165-	7.5	IF only		
Goat anti-rabbit			144				
Biotin conjugated Goat	Goat	Jackson ImmunoResearch Laboratories	115-065-020	2.8	IF only		
anti-mouse							
Biotin conjugated Goat	Goat	Jackson ImmunoResearch Laboratories	111-065-144	2.8	IF only		
anti-rabbit							
Cy2 conjugated	-	Jackson ImmunoResearch Laboratories	016-220-084	3.6	IF only		
streptavidin							
Cy3 conjugated	-	Jackson ImmunoResearch Laboratories	016-160-084	3.6	IF only		
streptavidin							
Oregon Green 488	-	Life Technologies	O7466	0.6 Units/ml	IF only		
Phalloidin							
Peroxidase-AffiniPure	Goat	Jackson ImmunoResearch Laboratories	111-035-144	0.08	Western only		
Goat anti-rabbit							
Peroxidase-AffiniPure	Goat	Jackson ImmunoResearch Laboratories	111-035-003	0.08	Western only		
Goat anti-mouse							

Table S3: List of RNAseq selected genes and their log2fold change values in Diaphragm, Gastrocnemius, Quadriceps and TA muscles. Only genes with statistically significant log2 fold change values are shown.

Genes	Gene Symbol	Diaphragm	Gastrocnemius	Quadriceps	Tibialis anterior
B cell translocation gene 2, anti- proliferative	Btg2	0.77	0.47	0.63	0.41
Nuclear factor, interleukin 3, regulated	Nfil3	0.51	0.38	0.48	0.47
Cysteine and glycine-rich protein 3	Csrp3	0.76	0.67	0.71	-
Tribbles homolog 1 (Drosophila)	Trib1	0.56	0.50	0.47	-
Phosphodiesterase 4D, cAMP specific	Pde4d	0.56	-	0.55	0.69
CCR4 carbon catabolite repression 4- like (S. cerevisiae)	Ccrn4l	0.53	-	0.52	0.45
Cell adhesion molecule 3	Cadm3	-	-0.55	-0.43	-0.39
CCAAT/enhancer binding protein (C/EBP), beta	Cebpb	-	0.63	0.56	0.50
Myosin, light polypeptide 3	Myl3	-	-0.46	-0.68	-0.66
Calmodulin binding transcription activator 1	Camta1	-	0.44	0.38	0.38
Early B cell factor 2	Ebf2	-	-0.30	-0.31	-0.36
Solute carrier family 2 (facilitated glucose transporter), member 1	Slc2a1	-	0.53	0.57	0.66
RIKEN cDNA C130074G19 gene	C130074G 19Rik	-	-0.31	-0.33	-0.26
Peroxisome proliferative activated receptor, gamma, coactivator 1 alpha	Ppargc1a	-	0.42	0.58	0.52
Tripartite motif-containing 63	Trim63	-	0.45	0.69	0.55
Tropomyosin 3, gamma	Tpm3	-	-0.34	-0.47	-0.41
Family with sequence similarity 107, member A	Fam107a	-	0.48	0.52	0.69
a disintegrin-like and metallopeptidase (reprolysin type) with thrombospondin type 1 motif, 15	Adamts15	-	0.34	0.55	0.43
Lipin 1	Lpin1	-	0.46	0.59	0.75
a disintegrin-like and metallopeptidase (reprolysin type) with thrombospondin type 1 motif, 1	Adamts1	-	0.31	0.40	0.29
Ankyrin repeat domain 2 (stretch responsive muscle)	Ankrd2	1.01	0.61	-	-

ST3 beta-galactoside alpha-2,3- sialyltransferase 5	St3gal5	0.42	0.54	-	-
Uridine-cytidine kinase 2	Uck2	0.31	0.35	-	-
Ankyrin repeat domain 1 (cardiac muscle)	Ankrd1	0.54	0.54	-	-
v-maf musculoaponeurotic fibrosarcoma oncogene family, protein F (avian)	Maff	0.63	-	0.48	-
salt inducible kinase 1	Sik1	0.54	-	0.39	-
xin actin-binding repeat containing 2	Xirp2	-	0.87	0.48	-
shisa family member 3	Shisa3	-	0.74	0.74	-
collagen, type XXII, alpha 1	Col22a1	-	0.53	0.35	-
regulator of cell cycle	Rgcc	-	0.67	0.46	-
heat shock protein family, member 7 (cardiovascular)	Hspb7	-	0.63	0.45	-
major facilitator superfamily domain containing 2A	Mfsd2a	-	0.72	0.48	-
peptidase inhibitor 16	Pi16	-	-0.42	-0.46	-
family with sequence similarity 214, member B	Fam214b	-	0.48	0.40	-
stathmin-like 4	Stmn4	-	0.59	0.46	-
ATP-binding cassette, sub-family B (MDR/TAP), member 4	Abcb4	-	0.56	0.46	-
BarH-like homeobox 2	Barx2	-	-0.49	-0.54	-
progestin and adipoQ receptor family member VIII	Paqr8	-	0.53	0.55	-
dual specificity phosphatase 5	Dusp5	-	0.53	0.59	-
musculoskeletal, embryonic nuclear protein 1	Mustn1	-	0.42	0.47	-
solute carrier family 35, member F5	Slc35f5	-	0.39	0.41	-
BCL2-associated athanogene 3	Bag3	-	0.29	0.31	-
G protein-coupled receptor 133	Gpr133	-	-0.37	-0.45	-
meningioma 1	Mn1	-	0.48	-	0.67
solute carrier family 43, member 2	Slc43a2	-	0.44	-	0.40
hypoxia inducible factor 3, alpha subunit	Hif3a	-	0.49	-	0.49

maestro heat-like repeat family member 1	Mroh1	-	0.33	-	0.43
ankyrin repeat and SOCS box- containing 2	Asb2	-	0.29	-	0.32
nuclear receptor subfamily 4, group A, member 1	Nr4a1	-	0.49	-	0.46
ankyrin repeat and zinc finger domain containing 1	Ankzf1	-	0.37	-	0.43
ubiquitin specific peptidase 2	Usp2	-	-	0.28	0.34
solute carrier family 15, member 4	Slc15a4	-	-	0.39	0.38
transmembrane protein 37	Tmem37	-	-	0.46	0.46
solute carrier family 7 (cationic amino acid transporter, y+ system), member 8	Slc7a8	-	-	0.39	0.37
cytochrome b-561 domain containing 1	Cyb561d1	-	-	-0.46	-0.58
kelch-like 38	Klhl38	-	-	0.53	0.74
carbonic anhydrase 3	Car3	-	-	-0.38	-0.57
galactosidase, beta 1-like 2	Glb1l2	-	-	0.54	0.47
solute carrier family 25 (mitochondrial carrier, phosphate carrier), member 25	Slc25a25	-	-	0.51	0.54
deiodinase, iodothyronine, type II	Dio2	-	-	0.56	0.54
RIKEN cDNA 2410131K14 gene	2410131K 14Rik	-	-	0.45	0.34
myosin, light polypeptide 2, regulatory, cardiac, slow	Myl2	-	-	-0.77	-1.21
pyridoxal (pyridoxine, vitamin B6) kinase	Pdxk	0.47	-	-	-
heat shock 105kDa/110kDa protein 1	Hsph1	0.37	-	-	-
ATPase, class II, type 9A	Atp9a	0.42	-	-	-
TCDD-inducible poly(ADP-ribose) polymerase	Tiparp	0.54	-	-	-
chemokine (C-C motif) ligand 21A (serine)	Ccl21a	0.54	-	-	-
ADAMTS-like 2	Adamtsl2	0.53	-	-	-
serum/glucocorticoid regulated kinase 1	Sgk1	0.45	-	-	-
ADP-ribosylation factor-like 4D	Arl4d	0.59	-	-	-

sodium channel, voltage-gated, type V, alpha	Scn5a	0.42	-	-	-
family with sequence similarity 220, member A	Fam220a	0.33	-	-	-
keratocan	Kera	0.51	-	-	-
OTU deubiquitinase with linear linkage specificity	Otulin	0.36	-	-	-
atonal homolog 8 (Drosophila)	Atoh8	0.49	-	-	-
RIKEN cDNA 6030419C18 gene	6030419C 18Rik	0.44	-	-	-
phosphodiesterase 4B, cAMP specific	Pde4b	0.49	-	-	-
NA	NA	0.35	-	-	-
myosin binding protein C, cardiac	Mybpc3	-	0.66	-	-
microtubule-associated protein tau	Mapt	-	0.74	-	-
carboxylesterase 5A	Ces5a	-	0.80	-	-
epidermal growth factor-containing fibulin-like extracellular matrix protein 1	Efemp1	-	-0.58	-	-
myosin, heavy polypeptide 2, skeletal muscle, adult	Myh2	-	0.76	-	-
tripartite motif-containing 9	Trim9	-	0.72	-	-
T cell lymphoma invasion and metastasis 2	Tiam2	-	0.59	-	-
ubiquitin carboxy-terminal hydrolase L1	Uchl1	-	0.45	-	-
calmodulin-like 3	Calml3	-	0.59	-	-
microtubule associated tumor suppressor candidate 2	Mtus2	-	0.60	-	-
runt related transcription factor 2	Runx2	-	0.60	-	-
methyltransferase like 21C	Mettl21c	-	0.58	-	-
family with sequence similarity 81, member A	Fam81a	-	0.56	-	-
MAP/microtubule affinity-regulating kinase 1	Mark1	-	-0.30	-	-
proline synthetase co-transcribed	Prosc	-	0.26	-	-

Sfrp4	-	-0.57	-	-
Tnn	-	0.57	-	-
Cnksr1	-	0.56	-	-
Dnajb2	-	0.42	-	-
Mb	-	0.40	-	-
Igsf21	-	0.47	-	-
Dach1	-	-0.38	-	-
Tmem125	-	0.45	-	-
Cspg4	-	0.43	-	-
Camk1g	-	0.52	-	-
Slc6a18	-	0.52	-	-
Мус	-	0.32	-	-
Tub	-	-0.49	-	-
Plat	-	-0.27	-	-
Ttc9	-	0.38	-	-
Spi1	-	-0.43	-	-
Lamc2	-	0.46	-	-
Spsb2	-	0.44	-	-
Rgs5	-	-0.31	-	-
Tspan18	-	-0.29	-	-
Mir17hg	-	0.36	-	-
Ecm1	-	-0.37	-	-
Ntn1	-	-0.37	-	-
Wdr95	-	0.43	-	-
	Sfrp4TnnCnksr1Onajb2MbIgsf21Dach1Tmem125Cspg4Camk1gSlc6a18MycTubPlatTtc9Spi1Lamc2Spsb2Rgs5Tspan18Mir17hgEcm1Ntn1Wdr95	Sfrp4-Tnn-Cnksr1-Dnajb2-Mb-Igsf21-Dach1-Tmem125-Cspg4-Camk1g-Slc6a18-Myc-Tub-Tub-Plat-Spi1-Spi2-Spsb2-Rgs5-Tspan18-Ntn1-Wdr95-	Sfrp4 - -0.57 Tnn - 0.57 Cnksr1 - 0.56 Dnajb2 - 0.42 Mb - 0.40 Igsf21 - 0.47 Dach1 - -0.38 Tmem125 - 0.43 Camk1g - 0.43 Camk1g - 0.52 Skc6a18 - 0.52 Myc - 0.32 Tub - -0.49 Plat - 0.38 Spi1 - 0.43 Lamc2 - 0.44 Rgs5 - - Tspan18 - - Mir17hg - 0.36 Ecm1 - - Wdr95 - 0.43	Sfrp4 - -0.57 - Tmn - 0.57 - Cnksr1 - 0.56 - Dnajb2 - 0.42 - Mb - 0.40 - Igsf21 - 0.47 - Dach1 - -0.38 - Tmem125 - 0.43 - Cspg4 - 0.43 - Camk1g - 0.52 - Slc6a18 - 0.52 - Myc - 0.32 - Tub - -0.27 - Plat - 0.38 - Spi1 - 0.46 - Spi1 - 0.44 - Rgs5 - 0.31 - Tspan18 - -0.29 - Mir17hg - 0.36 - Ecm1 - -0.37 - Mir195 - 0.33 -

coagulation factor II (thrombin) receptor-like 3	F2rl3	-	-0.45	-	-
wingless-type MMTV integration site family, member 10B	Wnt10b	-	-0.46	-	-
melanophilin	Mlph	-	0.42	-	-
Fras1 related extracellular matrix protein 1	Frem1	-	0.31	-	-
RNA, 7SK, nuclear	Rn7sk	-	0.42	-	-
vacuolar protein sorting 37B (yeast)	Vps37b	-	-0.26	-	-
sodium channel, voltage-gated, type III, alpha	Scn3a	-	0.40	-	-
C1q and tumor necrosis factor related protein 2	C1qtnf2	-	-0.39	-	-
CD34 antigen	Cd34	-	-0.15	-	-
family with sequence similarity 174, member B	Fam174b	-	-0.36	-	-
FAD-dependent oxidoreductase domain containing 2	Foxred2	-	-	0.61	-
ring finger protein 144B	Rnf144b	-	-	0.38	-
gap junction protein, delta 4	Gjd4	-	-	0.39	-
angiotensinogen (serpin peptidase inhibitor, clade A, member 8)	Agt	-	-	0.47	-
dynamin 3, opposite strand	Dnm3os	-	-	0.23	-
solute carrier family 3 (activators of dibasic and neutral amino acid transport), member 2	Slc3a2	-	-	0.26	-
TSC22 domain family, member 1	Tsc22d1	-	-	0.39	-
homeobox C8	Hoxc8	-	-	-0.39	-
NA	NA	-	-	0.46	-
Solute carrier family 35, member E1	Slc35e1	-	-	0.38	-
NCK associated protein 1 like	Nckap11	-	-	0.39	-

pleckstrin homology domain containing, family F (with FYVE domain) member 1	Plekhf1	-	-	0.38	-
sema domain, transmembrane domain (TM), and cytoplasmic domain, (semaphorin) 6A	Sema6a	-	-	-0.19	-
growth associated protein 43	Gap43	-	-	-0.41	-
kinesin family member 26A	Kif26a	-	-	-0.29	-
carbonic anhydrase 14	Car14	-	-	0.46	-
low density lipoprotein receptor- related protein 11	Lrp11	-	-	-0.46	-
R-spondin 3 homolog (Xenopus laevis)	Rspo3	-	-	-0.35	-
RIKEN cDNA 2500002B13 gene	2500002B 13Rik	-	-	0.43	-
twinfilin, actin-binding protein, homolog 2 (Drosophila)	Twf2	-	-	0.25	-
sestrin 2	Sesn2	-	-	0.43	-
troponin C, cardiac/slow skeletal	Tnnc1	-	-	-	-0.55
melanin-concentrating hormone receptor 1	Mchr1	-	-	-	0.81
phosphoserine aminotransferase 1	Psat1	-	-	-	-0.45
myosin, light polypeptide kinase 2, skeletal muscle	Mylk2	-	-	-	0.57
serine hydroxymethyltransferase 1 (soluble)	Shmt1	-	-	-	0.41
sema domain, immunoglobulin domain (Ig), short basic domain, secreted, (semaphorin) 3C	Sema3c	-	-	-	-0.36
SH2B adaptor protein 2	Sh2b2	-	-	-	0.48
ankyrin repeat and SOCS box- containing 11	Asb11	-	-	-	0.53
myosin, heavy polypeptide 7, cardiac muscle, beta	Myh7	-	-	-	-0.43
solute carrier family 2 (facilitated glucose transporter), member 4	Slc2a4	-	-	-	0.42
troponin T1, skeletal, slow	Tnnt1	-	-	-	-0.32
family with sequence similarity 214, member A	Fam214a	-	-	-	0.43
PHD finger protein 10	Phf10	-	-	-	-0.28

asparagine synthetase	Asns	-	-	-	-0.50
F-box protein 31	Fbxo31	-	-	-	0.39
cysteinyl-tRNA synthetase	Cars	-	-	-	-0.42
tetraspanin 13	Tspan13	-	-	-	-0.29
adenosine monophosphate deaminase 3	Ampd3	-	-	-	0.56
lysine (K)-specific demethylase 4B	Kdm4b	-	-	-	0.28
patatin-like phospholipase domain containing 7	Pnpla7	-	-	-	0.47
myosin VC	Myo5c	-	-	-	0.55
pantothenate kinase 1	Pank1	-	-	-	0.53
dihydrolipoamide branched chain transacylase E2	Dbt	-	-	-	0.47
nicotinamide phosphoribosyltransferase	Nampt	-	-	-	0.35
DDB1 and CUL4 associated factor 11	Dcaf11	-	-	-	0.24
potassium voltage gated channel, Shaw-related subfamily, member 4	Kcnc4	-	-	-	-0.48
glycyl-tRNA synthetase	Gars	-	-	-	-0.36
inscuteable homolog (Drosophila)	Insc	-	-	-	-0.53
aldehyde dehydrogenase 18 family, member A1	Aldh18a1	-	-	-	-0.42
nitric oxide synthase 1, neuronal	Nos1	-	-	-	0.50
proteolipid protein (myelin) 1	Plp1	-	-	-	0.32
pyruvate dehyrogenase phosphatase catalytic subunit 2	Pdp2	-	-	-	0.52
myosin, light polypeptide 6B	Myl6b	-	-	-	-0.34
ADP-ribosylhydrolase like 1	Adprhl1	-	-	-	-0.32
transport and golgi organization 2	Tango2	-	-	-	0.48
ankyrin repeat and SOCS box- containing 14	Asb14	-	-	-	0.49
glycoprotein galactosyltransferase alpha 1, 3	Ggta1	-	-	-	0.34
protein phosphatase 1K (PP2C domain containing)	Ppm1k	-	-	-	0.51
RIKEN cDNA 1110018N20 gene	1110018N 20Rik	-	-	-	-0.47

corneodesmosin	Cdsn	-	-	-	-0.49
WD repeat domain 1	Wdr1	-	-	-	-0.30
Smith-Magenis syndrome chromosome region, candidate 8 homolog (human)	Smcr8	-	-	-	0.42
Kruppel-like factor 15	Klf15	-	-	-	0.50
tyrosyl-tRNA synthetase	Yars	-	-	-	-0.32
NA	NA	-	-	-	0.49
glutamate-ammonia ligase (glutamine synthetase)	Glul	-	-	-	0.45
epithelial membrane protein 1	Emp1	-	-	-	-0.27
methionine-tRNA synthetase	Mars	-	-	-	-0.34
Rho GTPase activating protein 36	Arhgap36	-	-	-	-0.20
mitochondrially encoded 12S rRNA	mt-Rnr1	-	-	-	0.44
ankyrin repeat and SOCS box- containing 10	Asb10	-	-	-	0.47
transmembrane protein 52	Tmem52	-	-	-	0.41
exportin, tRNA (nuclear export receptor for tRNAs)	Xpot	-	-	-	0.29
ankyrin repeat domain 9	Ankrd9	-	-	-	-0.26
cystathionase (cystathionine gamma- lyase)	Cth	-	-	-	0.43
host cell factor C1 regulator 1 (XPO1- dependent)	Hcfc1r1	-	-	-	-0.23
solute carrier family 44, member 2	Slc44a2	-	-	-	0.22
F-box protein 32	Fbxo32	-	-	-	0.44
myosin, heavy polypeptide 3, skeletal muscle, embryonic	Myh3	-	-	-	-0.32
tetratricopeptide repeat domain 38	Ttc38	-	-	-	0.45
myosin, light polypeptide 9, regulatory	Myl9	-	-	-	-0.33
zinc finger, FYVE domain containing 21	Zfyve21	-	-	-	0.35
solute carrier family 25, member 38	Slc25a38	-	-	-	0.38
RIKEN cDNA 4933431E20 gene	4933431E 20Rik	-	-	-	0.37

SLIT and NTRK-like family, member 4	Slitrk4	-	-	-	-0.43
peroxisome proliferator activated receptor alpha	Ppara	-	-	-	0.46
paired related homeobox 1	Prrx1	-	-	-	-0.22
transmembrane protein 131	Tmem131	-	-	-	0.21
phosphodiesterase 7A	Pde7a	-	-	-	0.33
methyltransferase like 21E	Mettl21e	-	-	-	0.46

Rajan S et al (2012)				MyHC-emb ^{Δ/Δ} RNA-seq		
Genes	Log Fold change	Expression stage and effect	Validated	Log2 fold change	Muscles	
Nr4a1	Max 3.29	Early	VOG	0.492792(G)	Gastrocnemius	
	Min -0.08	upregulated	yes	0.460666(TA)	and TA	
	Max 0.94	Early upregulated	yes	0.774867(D)	Diaphragm,	
Btg2				0.473973 (G)	Gastrocnemius,	
	Min 0.00			0.631396(Q)	Quadriceps and	
				0.412594(TA)	TA	
Ccrn41	Max 1.52 Min 0.00	Early upregulated	No	0.537682(D)	Diaphragm,	
				0.525946(Q)	Quadriceps and	
	101111 0.000	uprogutated		0.451526(TA)	TA	
Ankrd1	Max 1.22	Early	Noc	0.542368 (D)	Diaphragm and	
	Min -0.14	upregulated	yes	0.541384(G)	Gastrocnemius	
Maff	Max 0.83	Early	yes	0.631282(D)	Diaphragm and	
	Min -0.06	upregulated		0.484519(Q)	Quadriceps	
Ankrd2	Ankrd2 Max 4.12 Late		Ves	1.016406(D)	Diaphragm and	
/ IIKI u2	Min	upregulated	yes	0.616889(G)	Gastrocnemius	
Cebpb	Max 0 Min-0.09	Late downregulated	No	0.634893(G)	Gastrocnemius,	
				0.56011(Q)	Quadriceps and	
				0.509648(TA)	TA	
Asb2	_	-	ves	0.298487(G)	Gastrocnemius	
	M 0.00			0.326305(TA)	and TA	
Myc	Max 0.99	Early	yes	0.321544(G)	Gastrocnemius	
	Min -0.22	Upregulated	-			
Prrx1	$\begin{array}{c} \text{Max } 0.00 \\ \text{Min } 1.23 \end{array}$	Early and late	No	-0.22888(TA)	TA	
Hoxc8	Max 0.00	Early				
	Min -1 13	downregulated	No	-0.39671 (Q)	Quadriceps	
	Max 0.03	Late				
Runx2	Min -1.24	downregulated	No	0.60068(G)	Gastrocnemius	

Table S4: Table showing details of candidate genes common between MyHC-emb $^{\Delta/\Delta}$ RNA-seq analysis and transcription factors regulating myogenesis found by Rajan et al (2012).