

PhD Thesis

**Coordinating growth arrest and myogenesis in muscle stem cells:
A molecular and cellular analysis**



A Dissertation

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LIST OF ABBREVIATIONS

bHLH	basic Helix-Loop-Helix
BMP	Bone Morphogenetic Protein
BWS	Beckwith-Wiedemann Syndrome
CAK	Cdk-Activating Kinase
CDK	Cyclin-Dependent Kinase
CDKI	Cyclin-Dependent Kinase Inhibitor
DKO	Double Knock-Out
DTA	Diphtheria Toxin Fragment
DP	Differentiation-regulated transcription factor-1 Polypeptide
EMT	Epithelial–Mesenchymal Transition
ECM	ExtraCellular Matrix
FAP	Fibro/Adipogenic Progenitor
Fucci	Fluorescent Ubiquitynation-based Cell Cycle Indicator
HGF	Hepatocyte Growth Factor
KID	Kinase Inhibitory Domain
LIMK	LIM domain Kinase
MRF	Myogenic Regulatory Factor
MyHC	Myosin Heavy Chain
NICD	Notch IntraCellular Domain
NLS	Nuclear Localization Signal
PDGFR	Platelet-Derived Growth Factor Receptor
PP	Pocket Protein
PIC	PW1+ interstitial cell
Rb	RetinoBlastoma protein
SF	Scatter Factor
SP	Side Population
VEGF	Vascular Endothelial Growth Factor

ABSTRACT

Tightly controlled growth arrest coordinates the equilibrium between cell proliferation and cell differentiation during embryonic tissue formation as well as in adulthood during stem cell-mediated tissue regeneration. Myogenic differentiation requires a coordinated course of tissue-specific gene expression and irreversible cell cycle exit. However, I contributed to showing -using genetic manipulation in the mouse embryo- that these processes can be uncoupled. During development, growth arrest in myogenic cells is mediated by the cyclin-dependent kinase inhibitors (CDKIs) p21 and p57, which act redundantly to promote cell cycle exit. We demonstrated that skeletal muscle progenitors require a direct interaction with the differentiating myoblasts via the Notch signaling pathway to maintain their pool. We also identified a muscle-specific regulatory element of p57 that directly receives the input of Myogenic Regulatory Factors (MRFs) and Notch downstream targets. During my Ph.D. I examined whether this regulatory mechanism is also involved in postnatal myogenesis.

Adult skeletal muscle has a remarkable regenerative capacity, involving a stem cell population, called satellite cells (SCs), located on close contact to the myofibers under the basal lamina. At the transition from juvenile to adult state (around 3-4 week during postnatal growth in mice) they enter a non-cycling, quiescent state. Upon injury the SCs rapidly get activated, expand by proliferation and provide differentiated progeny for muscle repair, while a subpopulation self-renews and re-enters quiescence, allowing the support of additional rounds of muscle damage. To understand the mechanisms regulating acquisition of quiescence, we explored the role of the aforementioned CDKIs in adult muscle. Although absent from quiescent SCs, they become rapidly expressed upon activation (even in proliferating myoblasts) and their levels remain high in the differentiating muscle cells. Strikingly, during the course of differentiation p57 translocated from the cytoplasm of activated myoblasts to the nuclei of differentiating cells. Since p57 deficient mice die at birth, we generated a conditional knock-out allele to perform functional studies at the postnatal stages. This new allele, in which the coding region can be removed by the loxP/Cre recombination system, also contains a β -galactosidase reporter allowing the identification of p57-expressing cells. We generated a complete

loss of function allele using a ubiquitously expressed Cre, and observed developmental and perinatal phenotypes similar to previously described germline knock-outs. Furthermore, we showed that the reporter inserted in the p57 conditional allele faithfully recapitulates p57 expression profile at embryonic and adult tissues. Conditional ablation of p57 in adult SCs resulted in reduced myogenic differentiation in primary myoblast culture. Similarly, p21-null myoblasts exhibited proliferation and differentiation defects in single myofiber cultures. In vivo regeneration studies with p21 mutants showed an early impact on the SC pool, while both SCs and muscle structure were re-established by the end of the regeneration process. My Ph.D. work suggests that p21 and p57 are at play during adult myogenesis and cell cycle exit, although via different mechanisms compared to the developmental scenario. They both work at the early steps of satellite cell activation but do not compensate for each other's loss. Future studies will elucidate whether they lie genetically downstream of the MRFs and Notch targets during postnatal myogenesis.

RÉSUMÉ

Au cours du développement embryonnaire, tout comme chez l'adulte, la formation ainsi que la régénération des tissus nécessitent une régulation fine du cycle cellulaire afin de maintenir l'équilibre entre la prolifération et l'entrée en différenciation des cellules. La différenciation myogénique nécessite une coordination parfaite entre l'expression des gènes spécifiques au développement musculaire et la sortie irréversible du cycle cellulaire des cellules constitutives du tissu. Cependant, j'ai contribué à montrer que ces processus peuvent être découplés chez l'embryon via la génération de modèles murins génétiquement modifiés.

Au cours de la différenciation myogénique chez l'embryon, l'arrêt du cycle cellulaire est contrôlé par les inhibiteurs de kinases cyclines-dépendantes (CDKI) p21 et p57 qui présentent une activité redondante. Nous avons démontré que les cellules progénitrices du muscle squelettique requièrent une communication directe avec les myoblastes en différenciation via la voie de signalisation cellulaire Notch afin de maintenir cette population dans un état indifférenciée. De plus, nous avons mise en évidence la présence d'un élément de régulation spécifique du muscle dans la séquence du

gène p57 répondant aux facteurs myogéniques (Myogenic Regulatory Factors, MRFs) et aux gènes cibles de la voie de Notch. Dans la poursuite de ces travaux, j'ai ensuite étudié l'implication de ce mécanisme de régulation au cours de la myogenèse postnatale pendant mon travail de thèse.

Les cellules souches du muscle, ou cellules satellites, sont localisées sous la lame basale au contact des myofibres et confèrent une capacité de régénération remarquable au tissu musculaire. Chez la souris, au cours de la transition entre le stade juvénile et le stade adulte (c'est à dire 3-4 semaines après la naissance) les cellules satellites rentrent en état de quiescence. Sous l'action d'un stimuli externe ou lors d'une blessure musculaire, les cellules satellites s'activent et prolifèrent. Au cours de la régénération du muscle, une sous-population va se différencier pour réparer le muscle lésé alors qu'une partie de cette population va s'auto-renouveler afin de maintenir le stock de cellules souches quiescentes.

Dans le but de comprendre les mécanismes qui régissent l'entrée en quiescence de ces cellules, nous avons étudié le rôle de p21 et p57 et dans le muscle adulte. Bien que l'expression des gènes codants pour ces CDKs ne soit pas détectée dans les cellules satellites quiescentes, ils sont rapidement détectés dès leur activation mais également dans les myoblastes et les cellules musculaires en différenciation où ils sont fortement exprimés. Au cours de la différenciation myogénique p57 est transloqué depuis le cytoplasme des myoblastes jusqu'au noyau des cellules en différenciation.

Chez la souris, l'ablation du gène p57 est létale à la naissance. Afin de pouvoir étudier le rôle fonctionnel de cette protéine après la naissance, nous avons généré un modèle murin présentant une modification génique conditionnelle permettant de muter le gène p57. Cette construction conditionnelle utilise le système de recombinaison Cre/LoxP qui permet d'exciser la séquence codante du gène p57. Elle contient également le gène rapporteur β -galactosidase afin de pouvoir identifier les cellules qui expriment p57. La perte totale de fonction générée par l'utilisation d'une Cre recombinase exprimée de manière ubiquitaire a permis de caractériser les phénotypes observés au cours du développement embryonnaire et au cours de la période périnatale. Les phénotypes observés sont identiques aux phénotypes décrits précédemment chez les souris présentant une perte

de fonction du gène p57. De plus, nous avons pu caractériser le profile d'expression de ce gène au cours du développement embryonnaire et dans les différents tissus chez l'adulte grâce au gène rapporteur.

Dans les cultures primaires de cellules satellites adultes, la délétion de p57 conduit à une diminution de la différenciation myogénique. De même, les cultures de fibres isolées issues des myoblastes mutants pour le gène p21, présentent des défauts de prolifération et de différenciation. In vivo, l'étude de régénération chez les mutants p21 montre une réduction précoce de la population satellitaire. Paradoxalement, la population des cellules souches du muscle ainsi que la structure du tissu musculaire sont entièrement reconstituées à la fin du processus de régénération.

Mon travail de thèse suggère que p21 et p57 jouent un rôle important dans la myogenèse et la régulation du cycle cellulaire. Ces protéines ont une action similaire qui est déterminante sur l'activation des cellules satellites et agissent de manière précoce sur ces dernières. Cependant, chez l'adulte leur activité semble distincte lorsqu'un des deux CDKIs est manquant contrairement au stade embryonnaire. Mes travaux ouvrent des perspectives nouvelles sur le rôle de p21 et p57 en aval de la voie de signalisation de Notch et des MRFs au cours de la myogenèse post-natale.

ABSTRAKT

Das Gleichgewicht zwischen Zellproliferation und Zelldifferenzierung wird während der embryonalen Gewebekonstruktion sowie während der stammzellvermittelten Geweberegeneration im Adultstadium durch einen streng kontrollierten Wachstumsarrest koordiniert. Die myogene Differenzierung erfordert sowohl eine koordinierte Abfolge von gewebespezifischer Genexpression als auch einen irreversiblen Zellzyklusaustritt. Jedoch konnte ich durch genetische Manipulation von Mausembryos dazu beitragen, aufzuzeigen, dass diese beiden Prozesse voneinander entkoppelt werden können. Ein Wachstumsarrest myogener Zellen wird während der Entwicklung durch die Cyclin-abhängigen Kinaseinhibitoren (CKIs) p21 und p57 vermittelt. Letztere wirken dabei redundant, um einen Zellzyklusaustritt zu fördern. Wir konnten bereits nachweisen, dass skelettale Muskelvorläuferzellen

eine direkte Interaktion mit differenzierenden Myoblasten über den Notch-Signalweg benötigen, um ihren Pool aufrecht zu erhalten. Des Weiteren haben wir ein muskelspezifisches regulatorisches Element von p57 identifiziert, das direkten Input von myogenen Regulationsfaktoren (MRFs) und Notch Stromabwärts-Zielen erhält. Im Rahmen meiner Doktorarbeit habe ich untersucht, ob dieser regulatorische Mechanismus auch in der postnatalen Myogenese involviert ist.

Der adulte skelettale Muskel hat eine bemerkenswerte regenerative Kapazität, die eine Stammzellpopulation, sogenannte Satellitenzellen (SCs, engl. für "satellite cells") involviert. Diese befinden sich zwischen der Basallamina und der Muskelfaser. Beim Übergang zwischen juvenilen und adultem Stadium (zwischen 3-4 Wochen während des postnatalen Wachstums in Mäusen), gehen die Satellitenzellen in einen nicht-zyklischen Ruhezustand über. Satellitenzellen werden nach einer Verletzung schnell aktiviert, expandieren und stellen differenzierte Abkömmlinge für die Muskelreparatur zur Verfügung. Eine Subpopulation der Satellitenzellen erneuern sich selbst und gehen zurück in den Ruhezustand, so dass bei erneuter Muskelverletzung Unterstützung gewährleistet ist. Um die Mechanismen verstehen zu können, die den Übergang in den Ruhezustand regulieren, haben wir die Rolle der zuvor genannten CDKIs im adulten Muskel untersucht. Obwohl CDKIs in ruhenden Satellitenzellen abwesend sind, werden sie nach Aktivierung der Satellitenzellen schnell exprimiert, und ihr Expressionslevel bleibt in differenzierenden Muskelzellen aufrecht erhalten. Erstaunlicherweise transloziert p57 während des Differenzierungsprozesses vom Zytosol in den Zellkern. Da p57-defiziente Mäuse bei Geburt sterben, haben wir eine konditionelle Mausmutante generiert, um funktionale Studien im postnatalen Stadium durchführen zu können. Dieses neue Mausmodell hat ein modifiziertes p57-Allel, in dem die codierende Region von p57 durch loxP/Cre-Rekombination entfernt werden kann. Des Weiteren beinhaltet es auch das Reportgen β -Galactosidase, um p57-exprimierende Zellen identifizieren zu können. Wir haben durch eine ubiquitär exprimierte Cre-Rekombinase vollständige Knockout-Mäuse generiert und dabei entwicklungsorientierte und perinatale Phänotypen, ähnlich wie bei bereits beschriebenen Knockout-Mäusen, beobachtet. Des Weiteren konnten wir zeigen, dass der in das konditionelle p57-Allel eingefügte Reporter dem Expressionsprofil von p57 im embryonalen und adulten Gewebe entspricht. Konditionelle Ablation von p57 in Satellitenzellen resultierte in einer reduzierten myogenen

Differenzierung von primären Myoblastenkulturen. p21-defiziente-Myoblasten wiesen ähnliche Proliferation- und Differenzierungsdefekte in einzelnen Muskelfaser-Kulturen auf. In-vivo-Regenerationsstudien mit p21-Mutanten haben eine initiale Reduktion der Satellitenzellenanzahl gezeigt. Zum Ende des Regenerationsprozesses waren die Anzahl der Satellitenzellen sowie die Muskelstruktur jedoch re-etabliert. Meine Doktorarbeit lässt darauf schließen, dass p21 und p57 während der adulten Myogenese und des Zellzyklusaustritts eine Rolle spielen, jedoch unterscheidet sich der Mechanismus im Vergleich zum pränatalem Stadium. Beide fungieren bei frühzeitigen Schritten der Satellitenzellenaktivierung, aber kompensieren sich nicht für den gegenseitigen Verlust. Künftige Studien werden zeigen, ob p21 und p57 von MRFs und Notch während der postnatalen Myogenese reguliert werden.

INTRODUCTION

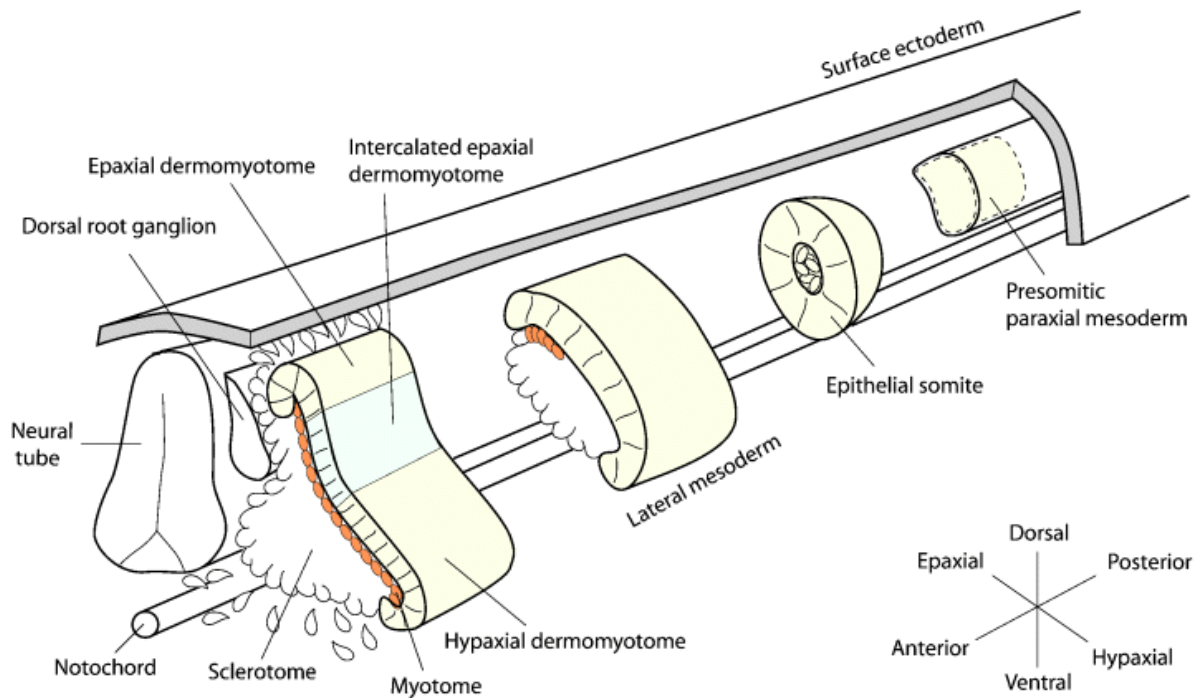


Figure 1.1. From somites to myotome.

Trunk and limb muscle derive from the paraxial mesoderm. The presomitic mesoderm will get progressively segmented into distinct, epithelially enclosed units, termed somites, which undergo several steps of specification and differentiation in a posterior-to-anterior orientation. As maturation proceeds they are compartmentalized into sclerotome and dermomyotome in response to signals from the adjacent structures. Cells from the dermomyotome migrate later to form the underlying myotome, the first differentiated skeletal muscle, as well as to distal myogenic sites, such as the developing limbs, diaphragm, and tongue.

Adapted from: Buckingham et al., 2003

Chapter 1. Skeletal muscle development, growth, and regeneration

1.1 Embryonic myogenesis: from somites to the first muscle masses

1.1.1 Somitogenesis: formation of multipotent mesodermal structures

Starting as a unicellular totipotent zygote the nascent organism develops into a multitude of differentiated, functionally interacting tissues and organs. During gastrulation, cells produced by consecutive divisions of the zygote and its descendants ingress through the blastopore/primitive streak to generate the three germ layers of the embryo, namely ectoderm, mesoderm and endoderm that will contribute all embryonic tissues. Skeletal muscle is of mesodermal origin and its emergence and organization is a complex process starting soon after gastrulation (E8 at mouse) [1].

Among the mesodermal compartments (axial, paraxial, intermediate, and lateral plate), the paraxial mesoderm gives rise to all the muscles of the limbs and ventral body [Christ & Ordahl, 1995], while branchiomic and ocular muscles derive from cranial/pharyngeal and prechordal cranial mesoderm [Scaal & Christ, 2004; Buckingham & Mayeuf, 2012]. The paraxial mesoderm constitutes two longitudinal columns of mesoderm on each side of the neural tube/notochord [Pourquié, 2001]. It will undergo somitogenesis, meaning generation of somites, which are segmented epithelial units that develop stepwise to give rise to the ventral mesenchymal sclerotome and the dorsal epithelial dermomyotome (**Fig. 1.1**). Apart from the differences in origin of body (trunk, limb) and head musculature, distinct genetic networks operate in each of them [Tajbakhsh, 2009; Bismuth & Relaix, 2010] and diverse turnover has been reported [McLoon et al., 2004; Keefe et al., 2015; Pawlikowski et al., 2015]. Limb and trunk musculature are, in general, more extensively studied and of more relevance to my PhD work and, thus, will be on focus in this section.

Somitogenesis occurs in coordination with embryo extension along the antero-posterior axis. Within the presomitic mesoderm, cellular arrangements of prospective somites point to a segmental pattern prior to somitogenesis and in response to intrinsic signals [Pourquié, 2001]. During gastrulation, new

Table 1.1. Somitogenesis across species [Hubaud & Pourquié, 2014].

Organism	Frequency of new pair addition	Final number of somite pairs
zebrafish	25 minutes	33
mouse	2 hours	65
human	4-5 hours	38-44

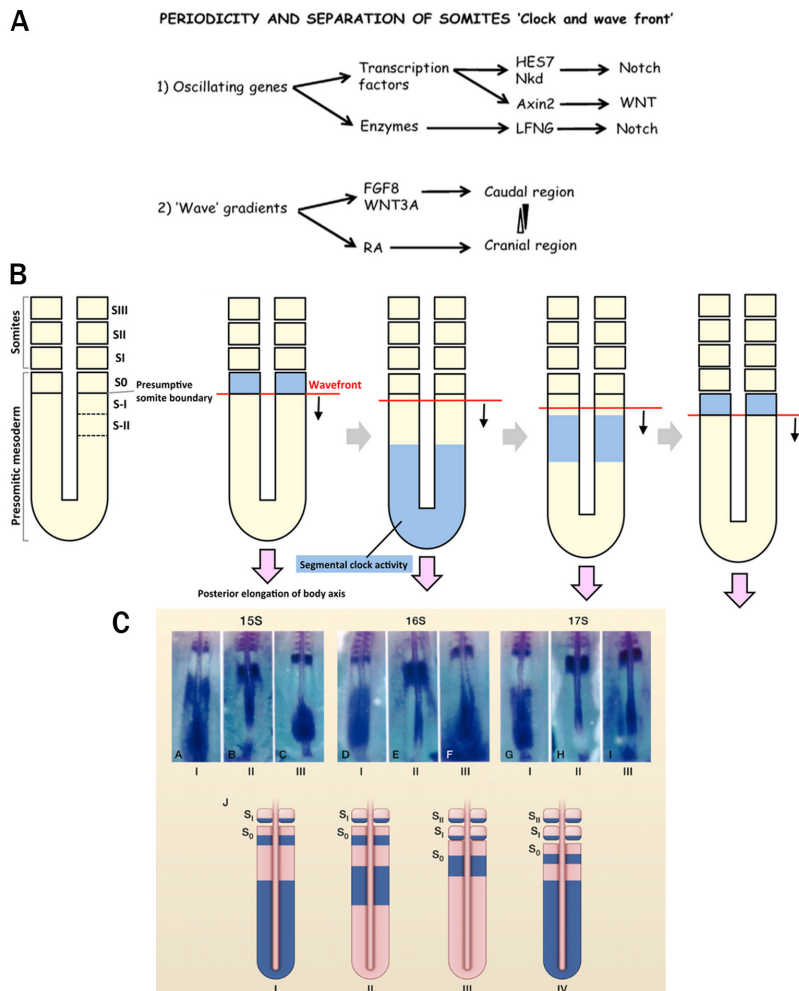


Figure 1.2. The “clock and wavefront” model for periodic somite generation

A. Major molecular contributors to the oscillation (clock) and gradient (wavefront) components of the model. B. Schematic representation of the location of activity of the clock and wavefront. C. Oscillatory expression of *c-hairy1* (*in situ* hybridization during formation of somites 15 to 17).

Adapted from: Pourquié, 2011; Musumeci et al., 2015; Yabe & Tanaka, 2016

mesenchyme cells enter the paraxial mesoderm leading to the addition of bilaterally symmetrical somite pairs at the anterior end of the presomitic mesoderm [Christ & Ordahl, 1995]. Convergence-extension movements of this phase produce the future anterior somitic mesoderm, while later on the tail bud contributes caudal somites [Pourquié, 2001]. Somitogenesis is a stepwise procedure involving periodic formation, separation, epithelialization, specification, and differentiation (with additional Epithelial–Mesenchymal Transition (EMT) transitions to form the sclerotome and myotome) [Musumeci et al., 2015].

Somites emerge periodically in a species-specific frequency (**Table 1.1**). To explain this feature, a theoretical model, the “clock and wavefront” model, was proposed in the 1970s (**Fig. 1.2**) [Cooke & Zeeman, 1976]. The clock refers to intrinsic oscillator(s) that make(s) presomitic mesoderm fluctuate between permissive and non-permissive states of somite formation (**Fig. 1.2B**) [Kalcheim & Ben-Yair, 2005]. The wavefront of competence to generate somites lays at a defined position with regard to the tail bud (**Fig. 1.2B**) [Saga, 2012]. On a molecular level, the idea of the clock was supported by the discovery of cyclic waves of *c-hairy1* linking it with somitogenesis (**Fig. 1.2C**) [Palmeirim et al., 1997]. This was followed by the identification of several other transcription factors (e.g. HES7, AXIN2) or enzymes (e.g. LFNG), mainly belonging to the Notch, FGF and Wnt pathways, which are periodically expressed at defined time intervals imitating the segmentation rounds (**Fig. 1.2A**) [Pourquié, 2011; Musumeci et al., 2015]. Notch, a key player of intercellular communication of neighboring cells, is further suggested to act by synchronizing the oscillations of individual cells [Horikawa et al., 2006]. The clock is mainly considered to act through the activation of the MESP transcription factors [Saga, 2012]. Different segment periodicities in the anterior and posterior presomitic mesoderm of zebrafish were revealed by recent live imaging experiments [Shih et al., 2015]. Another recent study challenged the role of the clock, counter-suggesting that cell-cell interactions drive somitogenesis [Dias et al., 2014]. However, they described somite-like structures [Dias et al., 2014] which were later criticized as being expected self-organizing differentiating derivatives, missing several somite characteristics [Hubaud & Pourquié, 2014]. The wavefront corresponds at the molecular level to a threshold of different signaling gradients, with Wnt and FGF signaling being highest at the posterior unsegmented paraxial mesoderm and retinoic acid following a counter-gradient (**Fig. 1.2A**) [Hubaud & Pourquié,

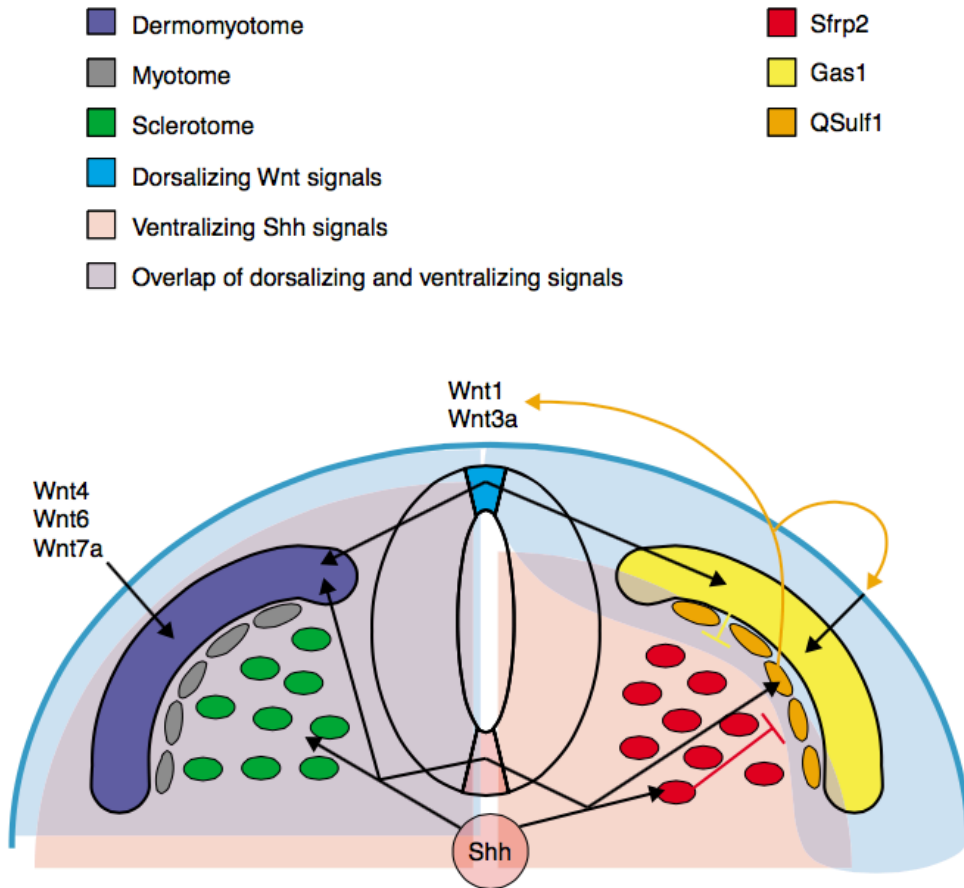


Figure 1.3. Dorsalizing and ventralizing signals involved in somite differentiation.

Left half simplistically focuses on the major dorsalizing (Wnt) and ventralizing (Shh) signals that act on the developing somites. Right half underlines the necessary activity of further factors for the final fine gradient driving the formation of dermomyotome in the dorsal somite and sclerotome in the ventral moiety.

Source: Brent & Tabin, 2002

2014; Mallo, 2016]. These gradients depend on mRNA decay or gradients of synthesizing and degrading enzymes, respectively [Aulehla & Pourquié, 2009]. Retinoic acid is also suggested to participate in synchronically generating left and right somites [Sirbu & Duester, 2006].

Once the new somite pair is signaled to be formed, it needs to undergo detachment and epithelialization. Separation of consecutive somites and formation of an acellular intersomitic border depend on the families of cadherins and ephrins and their receptors [Kalcheim & Ben-Yair, 2005]. Before complete detachment, mesenchymal-epithelial transition takes place, driven by the bHLH transcription factor PARAXIS and the GTPases CDC42 and RAC1 [Burgess et al., 1996; Nakaya et al., 2004]. Adhesion molecules (e.g. N-cadherin) and Fibronectin, an extracellular matrix protein were also found to participate in epithelialization [Duband et al., 1987; Linask et al., 1998]. Interestingly, in *Paraxis*-null mice epithelial somites are substituted by mesenchymal blocks, uncoupling segmentation and epithelialization [Burgess et al., 1996].

The next steps of somitogenesis involve specification and differentiation. Somite specification appears to depend on HOX genes, with maintenance of the HOX profile even after heterotopic transplantation [Nowicki & Burke, 2000]. Differentiation is largely attributed to interactions and molecular signals originating from the surrounding tissues. The ectoderm and notochord mediate dorsalization with signals such as Wnt proteins and Bone Morphogenetic Proteins (BMP), while the neural tube provides ventralizing signals, such as Sonic hedgehog homolog (Shh) or the BMP antagonist Noggin [Christ & Brand-Saberi, 2002; Scaal & Christ, 2004]. Further factors, such as SFRP2 or GAS1 antagonize Wnt and Shh, respectively, to prevent their long-range signaling and induce gradients of dorsalizing and ventralizing signals (**Fig. 1.3**) [Brent & Tabin, 2002].

Somites bud off the paraxial mesoderm at its rostral extremity, so that the caudalmost somite is the youngest or somite number I, according to the applied dynamic staging system with consecutive Roman numbers (**Fig. 1.2B**) [Scaal & Christ, 2004]. At early stages somites show developmental plasticity, as evidenced by single cell transplantation experiments [Kato & Gurdon, 1993]. Moreover, transplantations of groups of cells demonstrate a community effect [Gurdon et al., 1993]. Both cases

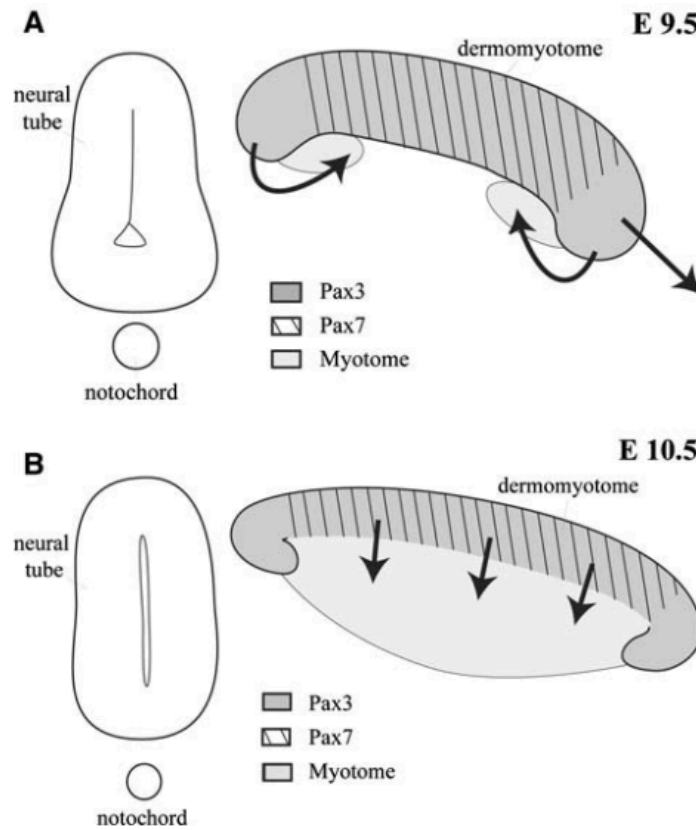


Figure 1.4. Myotome formation.

Myotome formation depends on cells migrating from the dermomyotome, initially from the lips (A) and subsequently from the central region (B).

Source: Buckingham et al., 2006

underline the influence of the surrounding microenvironment. Up to somites III-IV, they are shaped as epithelial spheres enclosing mesenchymal cells. Later on, the adjacent structure provide the aforementioned ventralizing and dorsalizing signals to compartmentalize somites into sclerotome and dermomyotome, respectively [Christ & Brand-Saberi, 2002; Scaal & Christ, 2004]. The ventral moiety undergoes EMT to form the sclerotome, which is mainly the source of the cartilage and bone of vertebral column and ribs (axial skeleton), but also contributes tendons, and joints [Musumeci et al., 2015]. The dorsal dermomyotome remains epithelial and stretches to form a sheet that roofs sclerotome. At later steps and while it adapts its rectangular-like form, dermomyotome develops a central mesenchymal sheet adjacent to the ectoderm and four inwardly curved epithelial lips. Dermomyotome gives rise to the dermis of the back and skeletal muscle, with their precursors originating in asymmetric divisions at the dorsal-ventral axis during EMT of the central part (see section 1.1.2) [Ben-Yair & Kalcheim, 2005]. Somites are divided in epaxial and hypaxial domains, lying dorsally and ventrally, respectively, to the horizontal septum of the vertebrae. The epaxial compartment generates muscles of the back, while the hypaxial part is the source of the muscles of the ventral body and limbs [Birchmeier & Brohmann, 2000; Musumeci et al., 2015].

1.1.2 Myotome: the first skeletal muscle

As the somite matures, cells delaminate and migrate underneath to form a third compartment, called myotome (**Fig. 1.4**), which corresponds to the first differentiated skeletal muscle. The groups of Ordahl and Kalcheim came to contradictory results when trying to elucidate the starting points and movements of myotomal precursors [Brent & Tabin, 2002], but a more recent stepwise model described by Gros et al. [2004] helped resolve the controversy. According to these findings, at a first phase cells translocate from the dorso-medial lip to the myotome and once there, they elongate along the rostral-caudal axis. At a second phase, cells invade myotome starting from all four lips and elongate along the anterior-posterior axis. Elongation can be unidirectional (cells from rostral or caudal lip) or bidirectional (cells from dorso-medial or ventro-lateral lip).

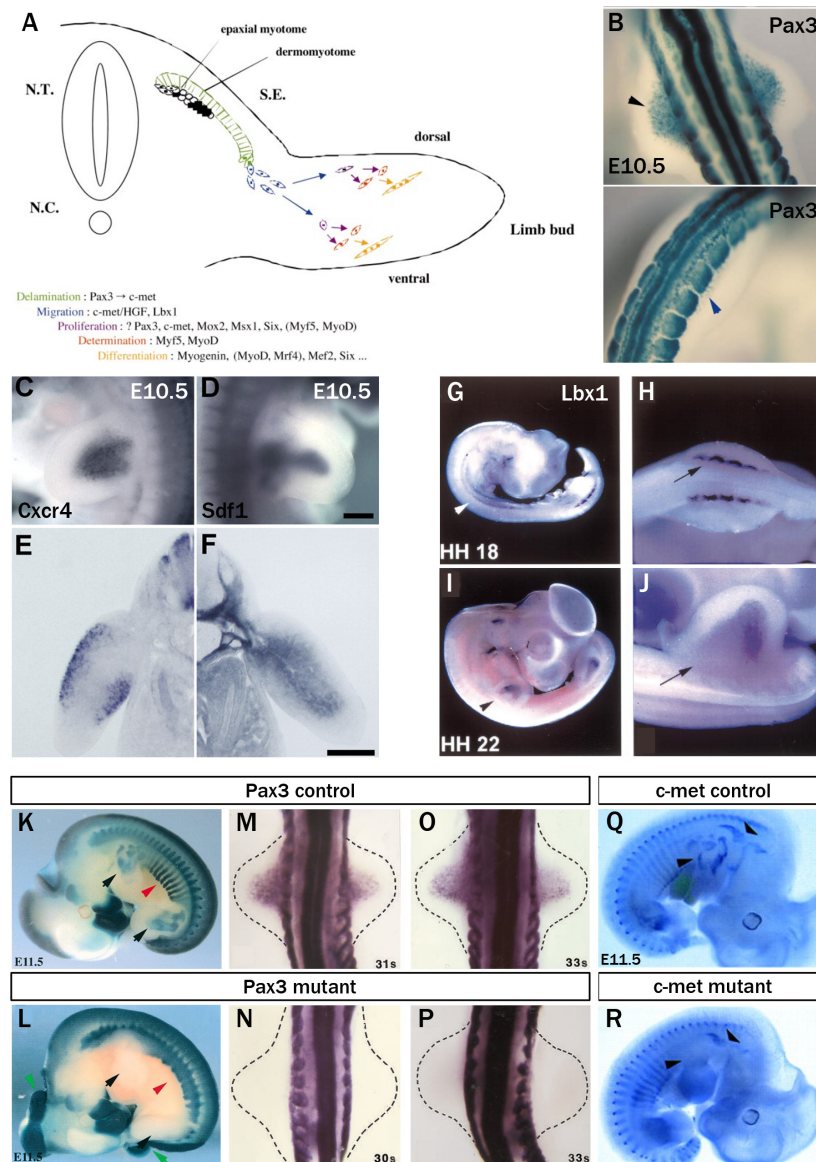


Figure 1.5. Progenitor migration for limb myogenesis.

A) Major factors controlling the consecutive steps of limb myogenesis. B) At E10.5 in the mouse embryo, Pax3-LacZ progenitors have migrated to the forelimb (up) and begin to migrate to the hindlimb (down). C,E) *Cxcr4* expression in limb at E10.5 mouse embryo. D,F) *Sdf1* expression in limb at E10.5 mouse embryo. G-J) *Lbx1* expression in somites and limbs of chick embryos at HH18 and HH22 stages. K-L) Limb muscles (black arrows) are absent from and somites are fused and truncated (red arrowheads) in *Pax3*^{IRESnLacZ/Sp} E11.5 mouse embryos as opposed to *Pax3*^{IRESnLacZ/+} control embryos at the same stage. M-P) PAX3-expressing cells do not colonize the forelimbs of *Spotch* mutants at 30-33 somite stages. Q-R) Lack of muscle (identified with *MyoD*) in the forelimbs of *c-Met* mutant E11.5 mouse embryos, in contrast to controls. NC: notochord, NT: neural tube, SE: surface ectoderm.

Adapted from: Bober et al., 1994; Maina et al., 1996; Mennerich et al., 1998; Buckingham et al., 2003; Relaix et al., 2003; Vasyutina et al., 2005.

Once the primary myotome is formed, a second population of myogenic progenitor cells originating from the central dermomyotome is colonizing the underlying myotome, rendering its initial name “dermatome” erroneous (**Fig. 1.4**). At later stages of embryonic and fetal life, muscle growth was found to depend on progenitors originating from the central dermomyotome, rather than the lips (see session 1.2.1) [Ben-Yair & Kalcheim, 2005; Gros et al., 2005; Kassam-Duchossoy et al., 2005; Relaix et al., 2005]. Of note, embryonic and fetal progenitors are mitotically active and have not engaged to the myogenic program. They maintain their proliferative status in embryonic and fetal muscles of trunk and limbs throughout development. They depend on transcription factors of the PAX family and they contribute to the forming muscles as well as their associated stem cells, as discussed in the following sections.

1.1.3 Migration of muscle progenitors to support limb myogenesis

Distant sites of myogenesis, such as the developing limb, depend on long-range migration of progenitors from the hypaxial dermomyotome to the limb buds, where they proliferate and subsequently commit to the myogenic lineage and undergo differentiation into skeletal muscle (**Fig. 1.5 A**) [Buckingham et al., 2003].

The transcription factor LBX1 is considered a bona fide marker of long-range migrating muscle precursors [Dietrich et al., 1998]. *Lbx1* starts to be expressed in the dispersing dermomyotomal lips, meaning prior to delamination. It then follows the migrating population (**Fig. 1.5 G-J**) and declines only when these progenitors arrive at the target sites and start to differentiate [Jagla et al., 1995; Dietrich et al., 1998; Mennerich et al., 1998]. In its absence, migratory precursors manage to form and delaminate but they display defective routing, demonstrating that LBX1 is critical for migration [Schäfer & Braun, 1999; Brohmann et al., 2000; Gross et al., 2000].

The migratory behavior of muscle progenitors is also controlled by PAX3, which is essential for the initiation of their migration. *Spotch* and other PAX3 mutant embryos show a number of developmental phenotypes in dorsal neural regions, neural crest cells and derivatives and muscle

tissues [Auerbach, 1954; Relaix et al., 2004]. Strikingly, PAX3-deficient embryos are devoid of myogenic migrating cells, leading to complete absence of muscular diaphragm, tongue and limb muscles (**Fig. 1.5 K-P**) [Mennerich et al., 1998; Relaix et al., 2004]. PAX3-expressing migrating progenitor cells also express LBX1 [Vasyutina et al., 2005].

Central in the genetic hierarchy controlling delamination and migration are the c-MET tyrosine kinase receptor - expressed by hypaxial muscle precursors - and its ligand scatter factor/ hepatocyte growth factor (SF/HGF) - lining the migratory route in the limb mesenchyme and other sites of migratory myogenesis [Birchmeier & Brohmann, 2000]. In their absence, migrating myogenic progenitors and, subsequently, muscle masses are missing from the limbs, tongue and diaphragm (**Fig. 1.5 Q-R**) [Bladt et al., 1995; Maina et al., 1996; Dietrich et al., 1999]. *c-Met* transcription depends on PAX3 [Epstein et al., 1996; Relaix et al., 2003]; *c-Met* expression as well as migratory progenitors and limb muscles are absent from *Spotch* embryos [Bober et al., 1994; Epstein et al., 1996; Yang et al., 1996; Tajbakhsh et al., 1997]. The *c-Met* promoter contains a PAX3 binding site [Epstein et al., 1996] and *c-Met* has been established as PAX3 target *in vitro* and *in vivo* [Epstein et al., 1996; Relaix et al., 2003]. *Cxcr4* and *Sdf1* constitute a further receptor-ligand pair affecting progenitor migration to the limbs [Vasyutina et al., 2005]. Similarly to c-MET and SF/HGF, CXCR4 receptor-expressing muscle progenitors are guided by a SDF1-paved route to the limb (**Fig. 1.5 C-F**). However, CXCR4/SDF1 seems to be required only for a subset of cells and to have a transient expression [Vasyutina et al., 2005].

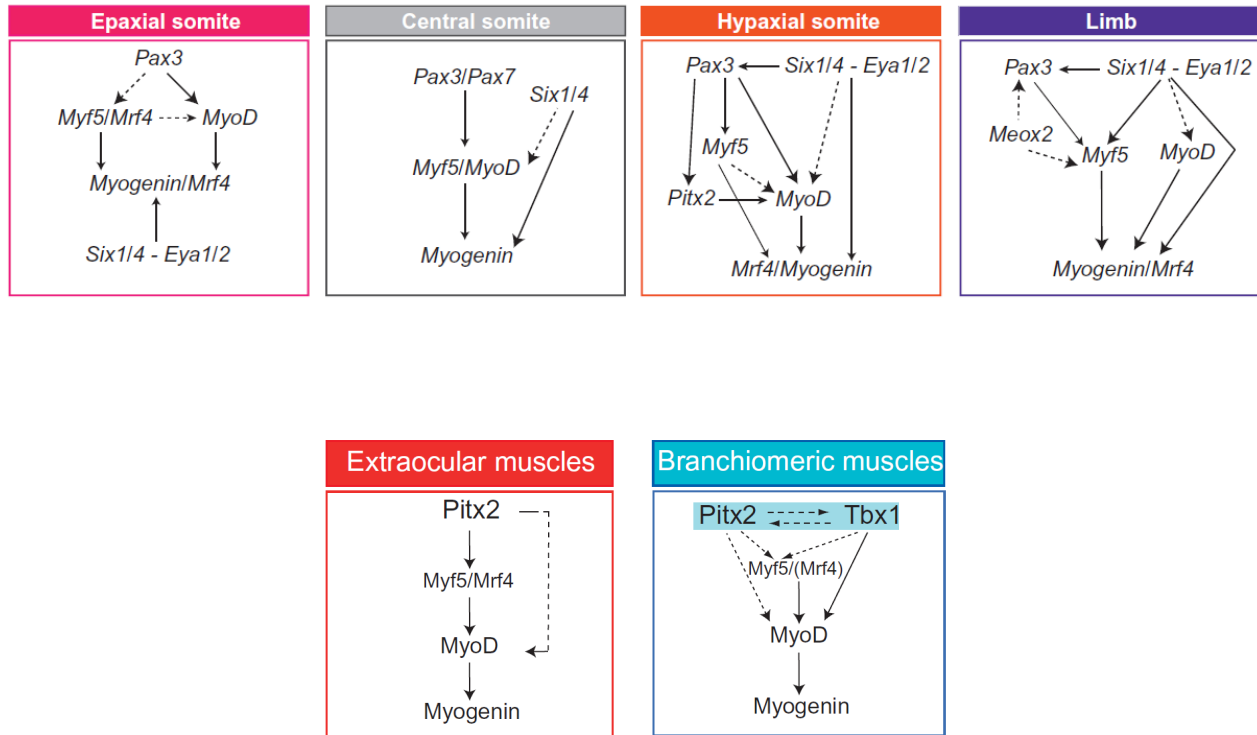


Fig. 1.6. Distinct genetic hierarchies control trunk, limb, and head myogenesis.

Adapted from: Buckingham & Mayeuf, 2012

1.2 Genetic hierarchies in head and body musculature establishment

A complex repertoire of transcription factor is crucial for the acquisition of the myogenic fate and skeletal muscle differentiation. Myogenic determination and differentiation rely on the Myogenic Regulatory Factors (MRFs), a family of basic helix-loop-helix (bHLH) transcription factors, including MYF5, MYOD, MRF4, and MYOGENIN. Upstream transcription factors act in the activation of MRFs as well as by regulating the proliferation and survival of progenitor cells. The upstream regulators differ between head/neck and trunk/limb muscles (**Fig. 1.6**) [Bismuth & Relaix, 2010; Braun & Gautel, 2011; Buckingham & Mayeuf, 2012].

In the body musculature PAX3/7 play a central role (see section 1.3), while a similar upstream role, linked to that of PAX3, was shown for the SIX homeodomain transcription factors and EYA cofactors [Buckingham & Rigby, 2014]. SIX1/4 or EYA1/2 deficient mice show a pronounced downregulation of MRFs and lacked limb and many trunk muscles [Grifone et al., 2005; Grifone et al., 2007]. Accordingly, SIX proteins were found to control *Myf5* [Giordani et al., 2007], *MyoD* [Relaix et al., 2013] and *Myogenin* [Spitz et al., 1998] expression, the first two in synergy with PAX3.

Head muscle development follows a distinct program, not requiring PAX3/7 but depending on four transcription factors -MYOR, Capsulin, PITX2, and TBX1- acting on different head muscle groups. PAX3 is not expressed in mesodermal derivatives in the head, while PAX7 is expressed in some head muscles, but its absence does not cause any head muscle phenotype [Bismuth & Relaix, 2010]. MYOR and Capsulin are bHLH transcription factors that redundantly function in specifying masticatory muscles [Bismuth & Relaix, 2010]. PITX2 is central in the regulation of non-somitic myogenic progenitors, controlling the survival and differentiation of muscle progenitors from the first branchial arches as well as progenitors that will form extraocular muscles [Buckingham & Mayeuf, 2012]. Finally, TBX1 has been described as “genetically equivalent to PAX3 during branchial arch development”, as it is expressed in the mesodermal cores of branchial arches and it is involved in bilateral branchiomic myogenesis [Bismuth & Relaix, 2010].

Table 1.2. Pax transcription factors structure and control of development [Buckingham & Relaix, 2007].

<i>Pax</i> genes	Structural characteristics	Expression in developing tissues/organs
<i>Pax3</i>	PD OP HD1/HD2/3	CNS, craniofacial tissue, trunk neural crest, somites/skeletal muscle
<i>Pax7</i>		CNS, craniofacial tissue, somites/skeletal muscle
<i>Pax4</i>		Pancreas, gut
<i>Pax6</i>		CNS, pancreas, gut, nose, eye
<i>Pax2</i>		CNS, kidney, ear
<i>Pax8</i>		CNS, kidney, thyroid
<i>Pax5</i>		CNS, B-lymphocytes
<i>Pax1</i>		Skeleton, thymus, parathyroid
<i>Pax9</i>		Skeleton, thymus, craniofacial tissue, teeth

CNS: central nervous system, HD: homeodomain, OP: octapeptide, PD: paired domain.

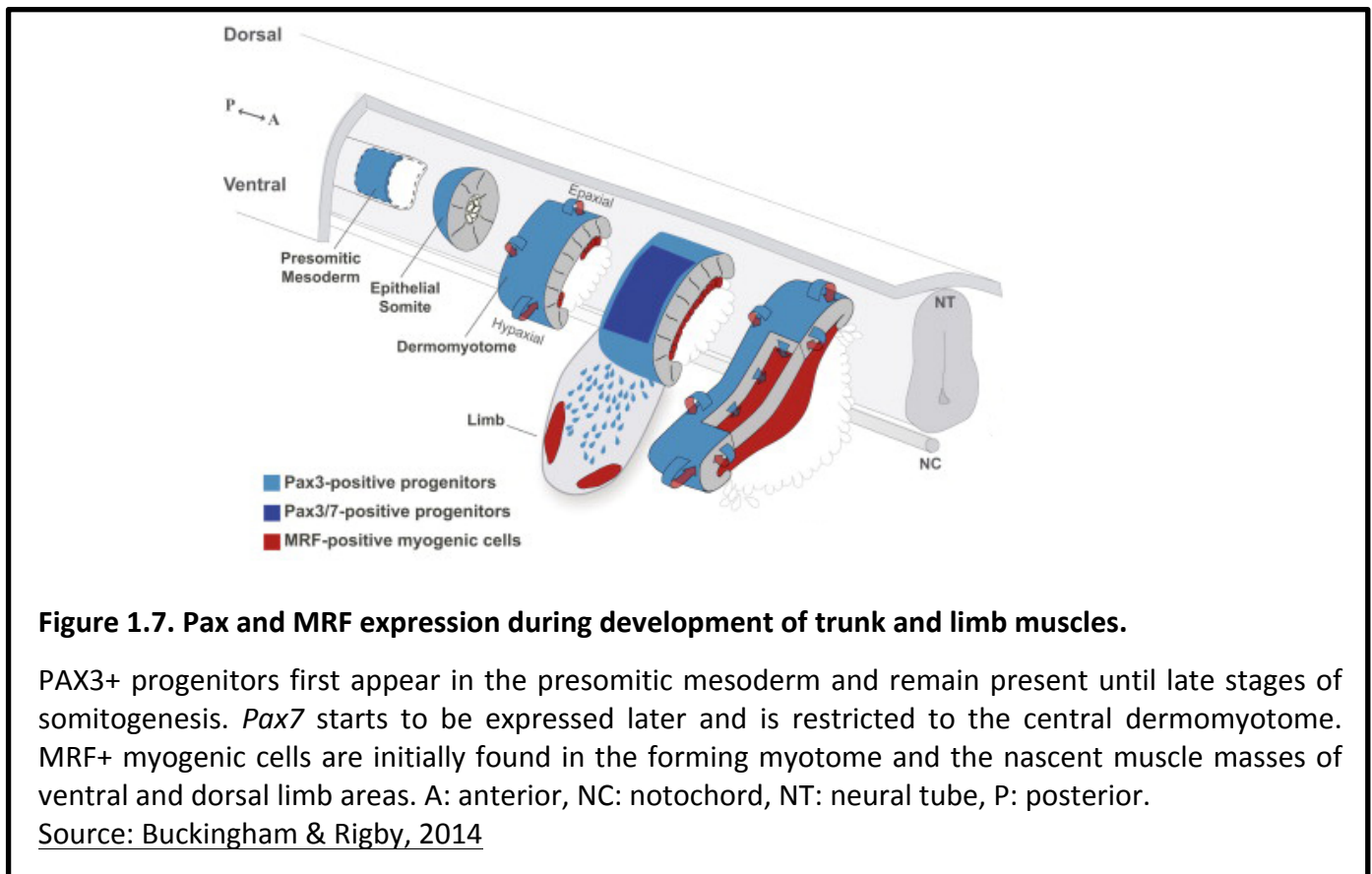


Figure 1.7. Pax and MRF expression during development of trunk and limb muscles.

PAX3+ progenitors first appear in the presomitic mesoderm and remain present until late stages of somitogenesis. *Pax7* starts to be expressed later and is restricted to the central dermomyotome. MRF+ myogenic cells are initially found in the forming myotome and the nascent muscle masses of ventral and dorsal limb areas. A: anterior, NC: notochord, NT: neural tube, P: posterior.

Source: Buckingham & Rigby, 2014

1.3 PAX proteins and bHLH MRFs play a central role in the myogenic program

My PhD work focuses on limb and trunk musculature and, thus, this section will cover general aspects of the function of PAX and MRFs in body musculature (**Fig. 1.7**) as well as their essential participation in embryonic myogenesis. Their role in postnatal growth and adult regeneration will be included in the session presenting satellite cells, which are the stem cells providing muscle precursors after birth.

1.3.1 PAX3 and PAX7 as upstream myogenic regulators

PAX proteins control the development of many lineages during embryogenesis (**Table 1.2**), with PAX3 and PAX7 acting as key regulators in the muscle lineage [Buckingham & Relaix, 2007]. In mammals, nine PAX proteins have been described, structurally characterized by a common paired box domain offering sequence-specific DNA binding. Some of them (including PAX3/7) also possess an octapeptide motif and an entire or truncated homeodomain (**Table 1.2**) [Buckingham & Relaix, 2007; Olguín & Pisconti, 2012]. *Pax* genes encode transcription factors and both PAX3 and PAX7 were shown to act as transcriptional activators *in vivo* [Relaix et al., 2003; Relaix et al., 2004] and orchestrate various biological aspects of myogenic progenitors and stem cells, including survival, proliferation, migration, self-renewal and triggering the myogenic program [Buckingham & Relaix, 2015]. Apart from their essential role in the muscle tissue, they are also important for neural crest derivatives and the central nervous system [Buckingham & Relaix, 2007].

As early as in the somite, compartmentalization and lineage specification are accompanied by alterations in the expression patterns of *Pax* genes [Christ & Ordahl, 1995]. PAX3 is mainly functioning during early embryonic myogenesis and gets downregulated in most muscles after birth, while PAX7 prevails in the post-natal growth phase as well as during adult muscle regeneration [Buckingham & Relaix, 2015]. Genetic replacement of PAX3 by PAX7 rescues most of the phenotypes of PAX3 mutants, but also shows that PAX7 cannot fully substitute PAX3 function in delamination, migration and proliferation of limb muscle progenitors (**Fig. 1.8**) [Relaix et al., 2004]. Furthermore, despite some overlapping functions of PAX3 and PAX7 in triggering the adult myogenic program, PAX7 has a distinct

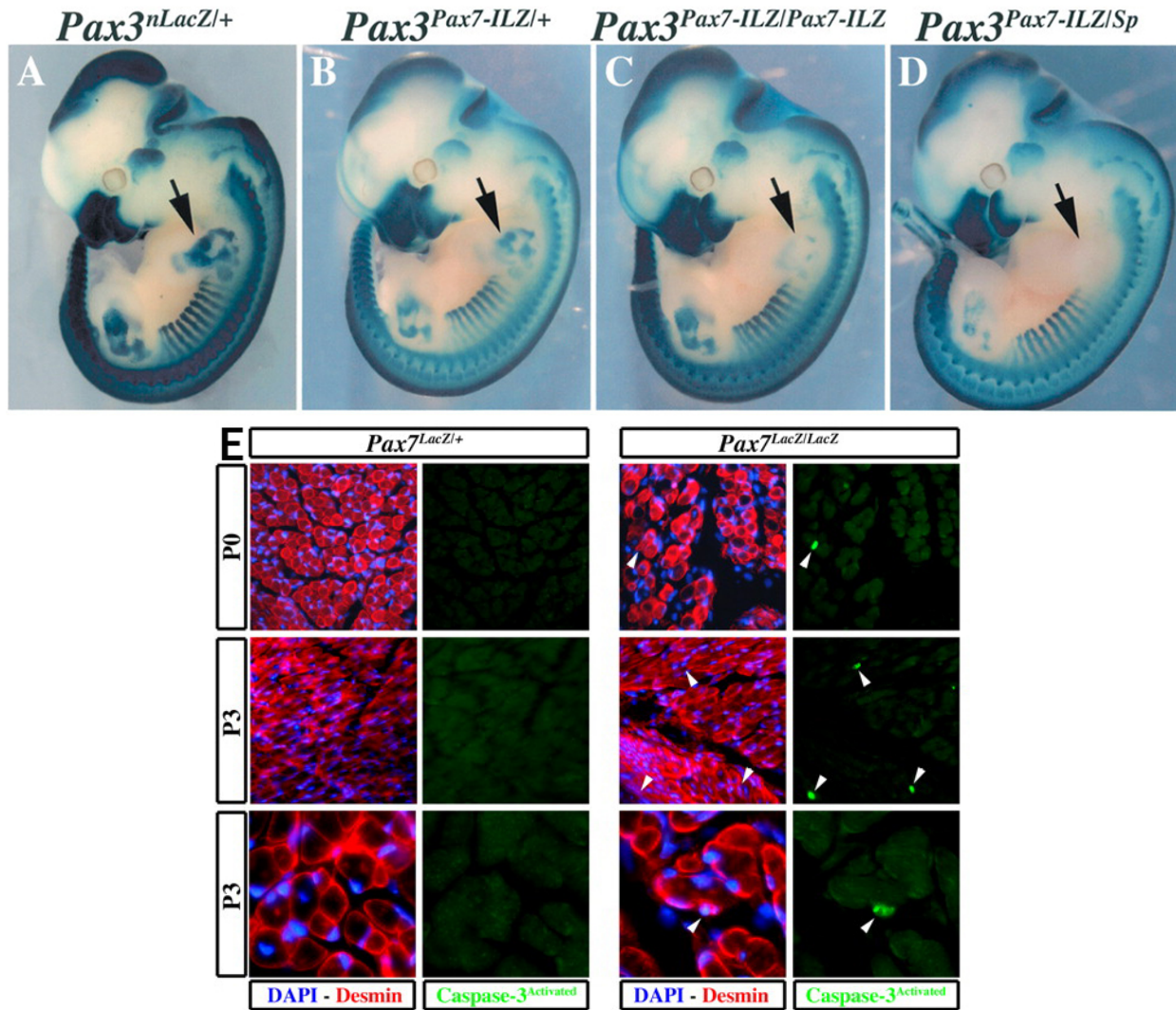


Figure 1.8. PAX3 and PAX7 have only partially overlapping functions.

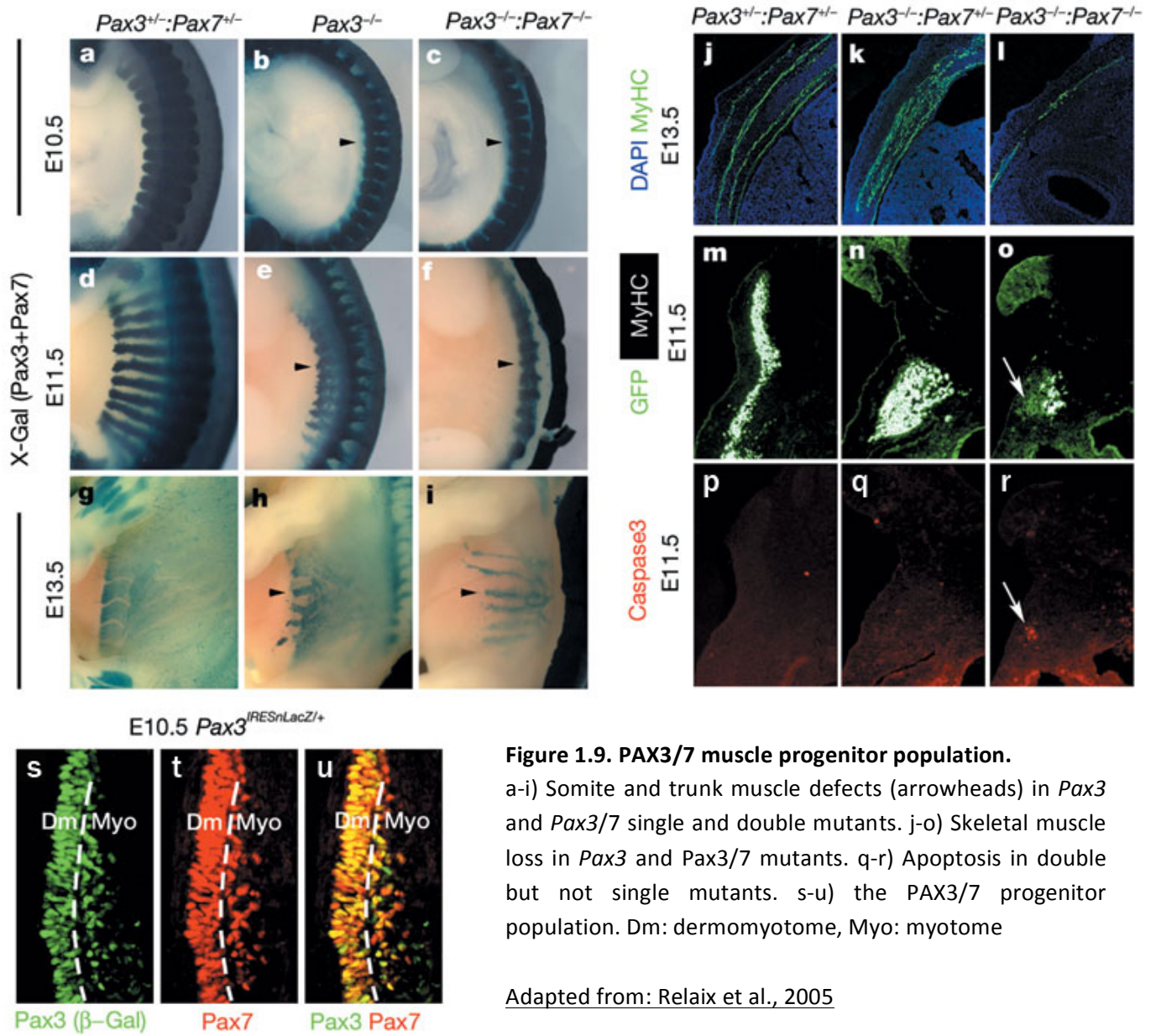
A-D) PAX7 cannot rescue limb defects of *Pax3*-mutant embryos, when knock-in in the *Pax3* locus. E) In the absence of PAX7, PAX3 cannot replace its antiapoptotic function.

Adapted from: Relaix et al., 2004; Relaix et al., 2006

role in survival (**Fig. 1.8**) and cell cycle progression [Relaix et al., 2006]. Large scale analysis of PAX3 and PAX7 binding profiles revealed several factors that could account for these differences, such as a) differential binding affinities for paired (PAX3) versus homeobox (PAX7) motifs, b) PAX3 binding only a subset of PAX7 targets (~5K sites for PAX3 vs ~53K sites for PAX7, with ~3.5K common sites), c) PAX7 occupying in the adult sites bound by PAX3 in the embryo, d) unique PAX3 targets involved in embryonic myogenesis (i.e. enrichment in ontology terms of skeletal muscle morphogenesis and neural and epithelial tube formation) [Soleimani et al., 2012].

PAX3 and PAX7 expression begins early in the nascent myogenic lineage and their absence leads to a complete arrest of skeletal muscle development. Transcript and reporter analyses revealed that PAX3 expression initiates in the presomitic mesoderm prior to segmentation (around E8 in the mouse) and its expression is progressively confined to the dermomyotome covering the epaxial and hypaxial extremities, while PAX7 appears later (around E9 in the mouse) and is concentrated in the central dermomyotome [Murphy & Kardon, 2011]. In the limb, PAX3+ progenitors migrate to the limb buds, where they are transiently present from E10 to E12.5 [Bober et al., 1994], while PAX7 appears later (E11.5), in PAX3-expressing myogenic progenitors and persists until fetal/neonatal stages [Relaix et al., 2004]. Later on, PAX3 gets downregulated but PAX7 persists [Kassar-Duchossoy et al., 2005]. In adult muscles, PAX7 is a universal marker of satellite cells, the progenitor/stem cell population responsible for postnatal growth/regeneration (see section 1.3), while PAX3 is restricted to a subset of trunk and limb muscle satellite cells [Seale et al., 2000; Relaix et al., 2006; Calhabeu et al., 2013].

Importantly, in 2005 a PAX3/PAX7+ progenitor population was identified in both chick and mouse embryos as the major source of myogenic cells for the forming muscle fibers in the trunk and limb [Ben-Yair & Kalcheim 2005; Gros et al., 2005; Kassar-Duchossoy et al., 2005; Relaix et al., 2005]. They are maintained in a proliferative state and lack MRFs or other muscle-specific markers. Reporter- or quail-to-chick-based genetic tracing place these dermomyotome-derived progenitors in lineage continuum with the forming MRF+, post-mitotic cells that enter the myogenic program. Strikingly, satellite cells also derive from the same dermomyotome population once cells become embedded under the basal lamina at E18.5 acquiring the characteristic “satellite” position [Gros et al., 2005;



Kassar-Duchossoy et al., 2005; Relaix et al., 2005]. PAX3/7 compound deficiency demonstrated their unequivocal role in the survival and specification of embryonic myogenic progenitors. Specifically, in the absence of PAX3/7, the progenitor cells either undergo cell death or fail to enter the myogenic program and get integrated to other tissues (**Fig. 1.9**) [Relaix et al., 2005]. PAX3 mutants, also known as splotch-mice (Auerbach, 1954), show defects in somitogenesis and segmentation, later affecting hypaxial trunk musculature. Furthermore, their migratory muscle progenitors are absent and they lack limb musculature (**Fig. 1.9**) [Bober et al., 1994; Tajbakhsh et al., 1997; Relaix et al., 2003; Relaix et al., 2004]. PAX7-null mice do not manifest any overt embryonic muscle phenotype and their trunk, limb, and facial muscles seem to develop normally [Mansouri et al., 1996; Relaix et al., 2004]. Embryonic lethality of splotch mice leaves PAX3 role uncharacterized in later stages; conversely, PAX7 exerts principal and indispensable functions at postnatal stages (see section 1.3).

1.3.2 MRFs play a central role in myogenic determination and differentiation

Skeletal muscle identity is conferred by the MRF family of transcription factors, which are expressed solely in skeletal muscle. In order to activate muscle-specific genes via direct binding to an E-box -a specific DNA sequence (CANNTG)-, MRFs heterodimerize with the ubiquitously expressed E proteins [Singh & Dilworth, 2013]. The MRF family consists of four members, MYOD [Davis et al., 1987], MYF5 [Braun et al., 1989], MRF4 [Rhodes & Konieczny, 1989], and MYOGENIN [Wright et al., 1989], which were originally identified by their ability to trigger conversion of non-muscle cell types into myogenic fate when ectopically expressed [Olson & Klein, 1994]. All four MRFs share a bHLH domain, mediating DNA binding as well as dimerization to form transcriptional complexes [Maroto et al., 2008]. The bHLH domain is characterized by ~80% amino acid identity among the four members, while limited sequence similarity is observed in the transcriptional activation domains, residing in the amino- and carboxyl-termini [Olson & Klein, 1994]. Target binding and expression profiling revealed shared targets between some members of the family and, in the case of MYOD and MYOGENIN, suggested a model whereby MYOD establishes an open chromatin structure at muscle-specific genes and MYOGENIN enhances transcription once chromatin is rendered accessible [Blais et al., 2005; Cao et al., 2006]. A further study implicated MYOD in chromatin loop dynamics regulation [Battistelli et al.,

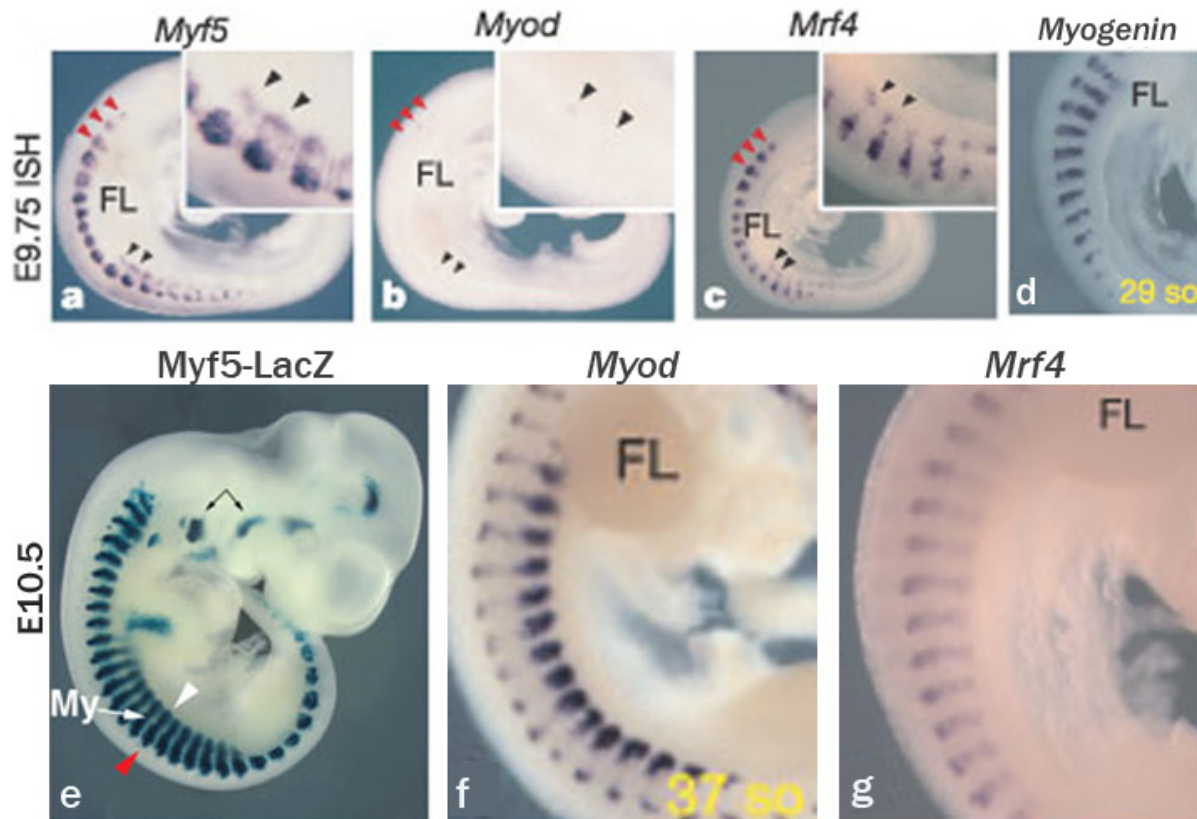


Figure 1.10. MRF expression profile at early embryonic myogenesis.

a-d) MRF transcripts at somites of E9.75 mouse embryos. e-g) MYF5 (reporter-based) and *MyoD/Mrf4* (transcript-based) expression at E10.5 mouse embryos. FL: forelimb, My: myotome.

Adapted from: Kassar-Duchossoy et al., 2004

2014], while MYOD and MYOGENIN targets include chromatin remodeling factors [Cao et al., 2006].

The specific expression of MRFs transcripts is initiated early during muscle development and follow distinct spatio-temporal patterns (**Fig. 1.10**) [summarized in Murphy & Kardon, 2011; Singh & Dilworth, 2013]. *Myf5* is the first MRF expressed, with its transcripts being observed from E8 in the epaxial dermomyotome and showing declining levels from recently formed (caudal) to mature (rostral) somites [Ott et al., 1991]. *Myf5* expression decreases from E14 onwards [Ott et al., 1991]. Of note, some PAX7+ cells do not express MYF5 and represent progenitors with slower proliferation and earlier exit from the cell cycle [Picard & Marcelle, 2013]. *Myogenin* is found from E8.5, accumulating in the most rostral somites and coinciding with differentiating muscle cells [Sassoon et al., 1989; Ott et al., 1991]. MYOD appears at E10.25-E10.5 [Sassoon et al., 1989; Kablar et al., 1997; Zabludoff et al., 1998]. In the limb, *Myf5* is detected in forelimb and hindlimb from E10.5 and E11, respectively, and gets downregulated by E11.5 when MYOD and MYOGENIN accumulate [Sassoon et al., 1989; Ott et al., 1991; Kablar et al., 1997]. *Mrf4* shows a biphasic pattern, with its transcripts appearing from E9 until E12 and then again from E16 onwards [Bober et al., 1991]. Adult myonuclei will maintain the expression of MRF4, which becomes the predominant MRF in adult muscle [Hinterberger et al., 1991; Gayraud-Morel et al., 2007].

Genetic ablation during embryonic and fetal myogenesis established MYF5, MYOD, and MRF4 as myogenic determination factors and MYOD, MYOGENIN, and MRF4 as myogenic differentiation factor [Murphy & Kardon, 2011]. MYF5-null mice form myotome with a 2-day delay [Braun et al., 1992; Tajbakhsh et al., 1997; Kassar-Duchossoy et al., 2004] and a MYF5-driven LacZ reporter revealed the presence of progenitors which activated *Myf5* in MYF5-null mice, but remained multipotent, failed to localize correctly and eventually differentiated into non-muscle derivatives according to their local environment [Tajbakhsh et al., 1996]. Despite the delayed myotome initiation in MYF5-deficient embryos, the myogenic program gets rescued around E11.5 by the delayed activation of MYOD [Braun et al., 1994] and muscles of MYF5 mutants become structurally and functionally normal until birth (**Figs. 1.11-1.12**) [Braun et al., 1992; Tajbakhsh et al., 1997]. It has been proposed that MYF5-independent MYOD-expressing myoblasts sustain myogenesis in the absence of a distinct, MYF5-

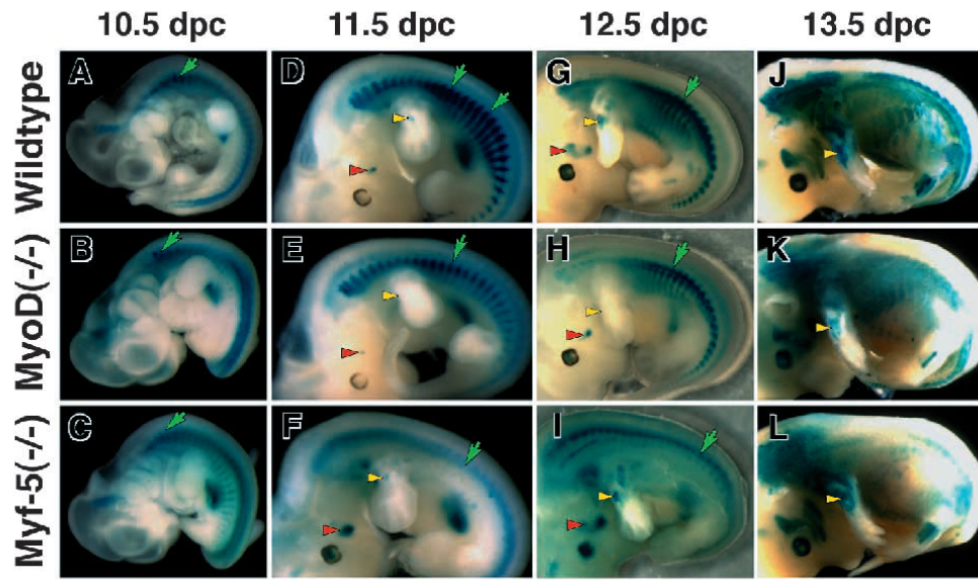


Figure 1.11. Embryonic myogenesis in the absence of *MyoD* or *Myf5*.

Developing muscles marked by MD6.0-LacZ reveal subtle differences in the absence of either *MyoD* (B, E, H, K) or *Myf5* (C, F, I, L) compared to age-matched control embryos (A, D, G, J).

Source: Kablar et al., 1997

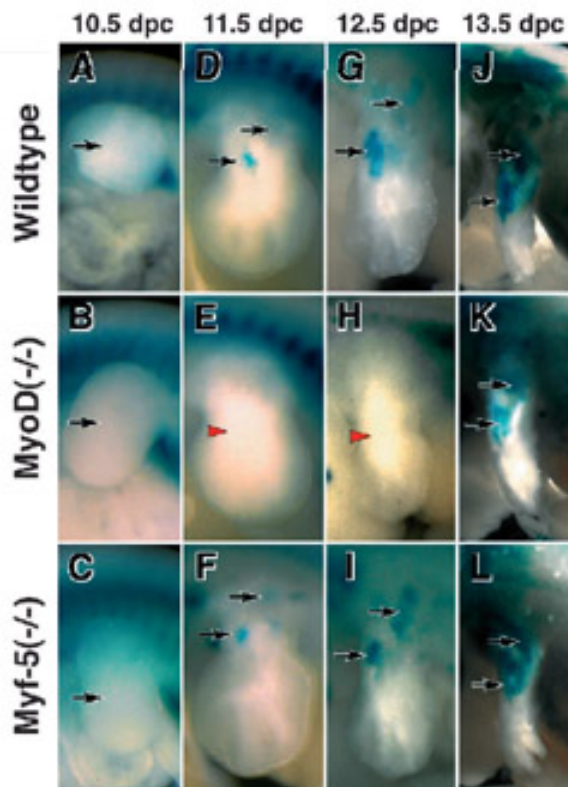


Figure 1.12. Limb myogenesis in the absence of *MyoD* or *Myf5*.

Delayed muscle development in the limbs of *MyoD* mutant (B, E, H, K), but not *Myf5* mutant embryos (C, F, I, L) compared to age-matched controls (A, D, G, J) as evidenced by MD6.0-LacZ expression. dpc: days post coitum

Source: Kablar et al., 1997

dependent lineage [Haldar et al., 2008]. Conversely, mice lacking MYOD have morphologically normal muscles (albeit showing 2-day delayed differentiation in the limb [Kablar et al., 1997]) (Figs. 1.11-1.12) and maintain high levels of MYF5 [Rudnicki et al., 1992; Kablar et al., 1998]. In the absence of both factors, newborns are completely devoid of skeletal muscles in both trunk and limbs [Rudnicki et al., 1993]. However, this effect appears to depend on compromised *Mrf4* expression, since skeletal muscle manages to differentiate in MYF5/MYOD double knockouts with functional MRF4 [Kassar-Duchossoy et al., 2004]. It has been proposed that the proximity of *Mrf4* and *Myf5* (the former residing 8kb 5' of the latter) likely account for cis-regulatory interactions [Olson et al., 1996], that are diversely affected in different MYF5 nulls. Thus, *Mrf4* was also identified as a determination gene, while genetic manipulation of the three factors placed both MYF5 and MRF4 upstream of *MyoD* [Kassar-Duchossoy et al., 2004]. Specific ablation of *Myf5*-expressing cells using *Myf5^{Cre}; R26R^{DTA/+}*, suggested the presence of distinct *Myf5*-dependent and *Myf5*-independent *MyoD*-expressing myoblasts [Gensch et al., 2008; Haldar et al., 2008]. However, when *Myf5*-expressing cells were eliminated in a *MyoD* null background, no muscles were formed, indicating that the previously observed *Myf5*-independent myoblasts were in fact MYF5+ escaper cells [Comai et al., 2014].

Myogenic differentiation was found to depend on MYOGENIN, MYOD, and MRF4 [Murphy & Kardon, 2011]. MYOGENIN did not overlap with MYOD or MYF5 in specification of the myogenic lineage [Rawls et al., 1995]. However, its deficiency led to compromised muscle-specific gene expression and differentiation, including a generalized fusion defect so that mutant mice presented with severely reduced muscle masses associated with lethality at birth [Hasty et al., 1993; Nabeshima et al., 1993; Rawls et al., 1995; Rawls et al., 1998]. These defects are phenocopied in MYOD/MRF4 double knockout mice [Rawls et al., 1998], implying that either MYOGENIN or MYOD and MRF4 need to be present to drive differentiation. Finally, single MRF4 loss-of-function overall did not jeopardize muscle development [reviewed in Olson et al., 1996], although different strategies for *Mrf4* disruption resulted in phenotypes as different as ranging from perinatal lethality to normal survival [Braun & Arnold, 1995; Patapoutian et al., 1995; Zhang et al., 1995], again likely due to interrelated cis-regulatory interactions with *Myf5* [Olson et al., 1996].

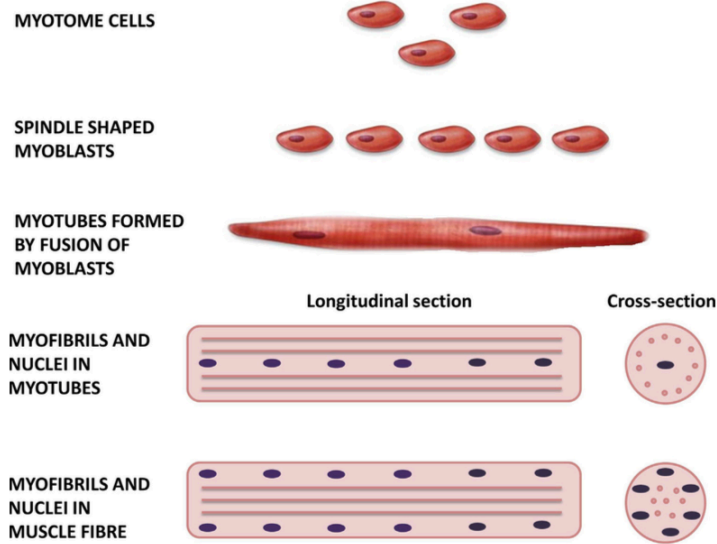


Figure 1.13. Myofiber formation.

Round myotomal cells undergo a step-wise differentiation and fusion procedure into long myotubes that further mature to myofibers with peripherally-positioned myonuclei.

Source: Musumeci et al., 2015

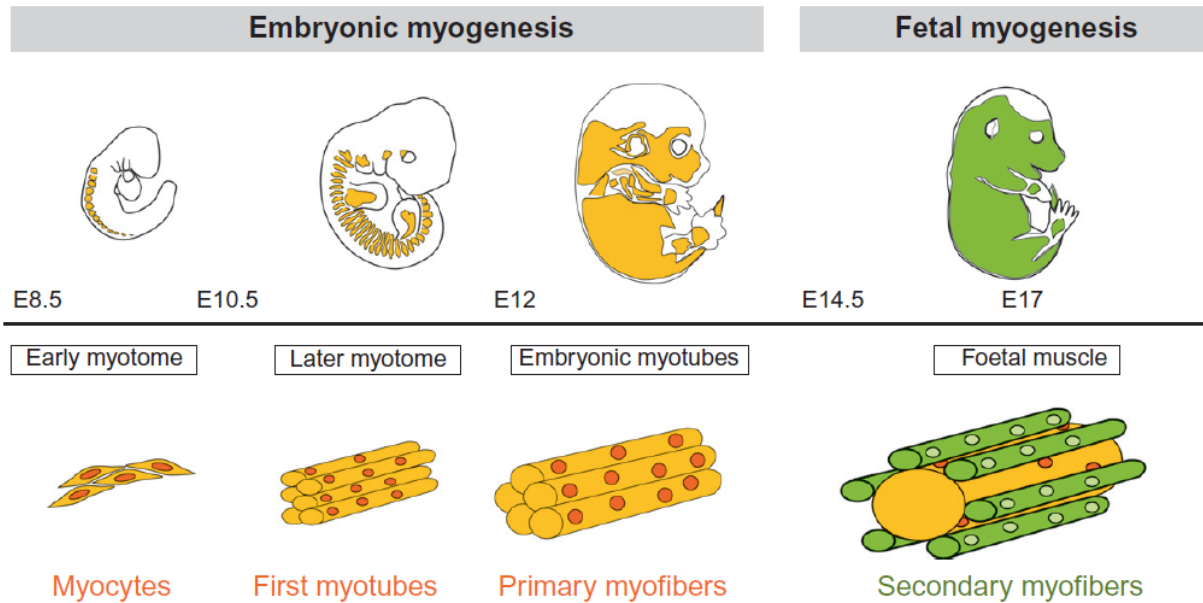


Fig. 1.14. Embryonic and fetal waves of myogenesis.

Source: Buckingham & Mayeuf, 2012

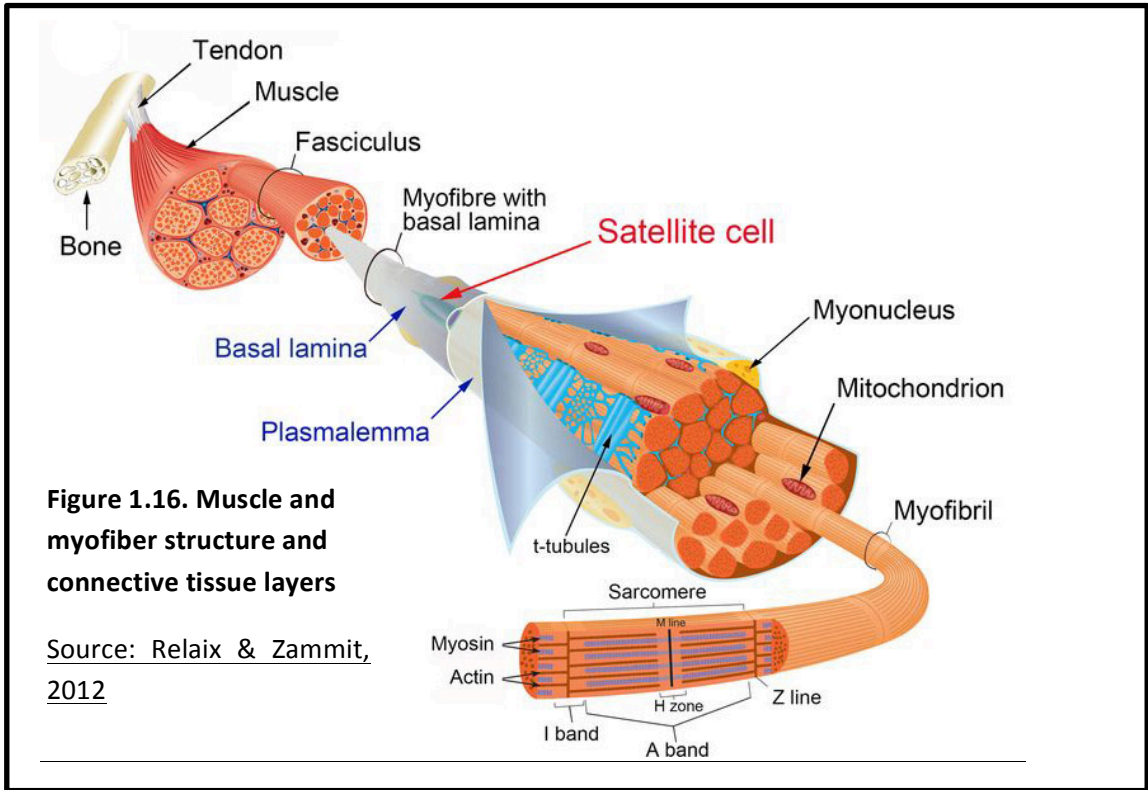
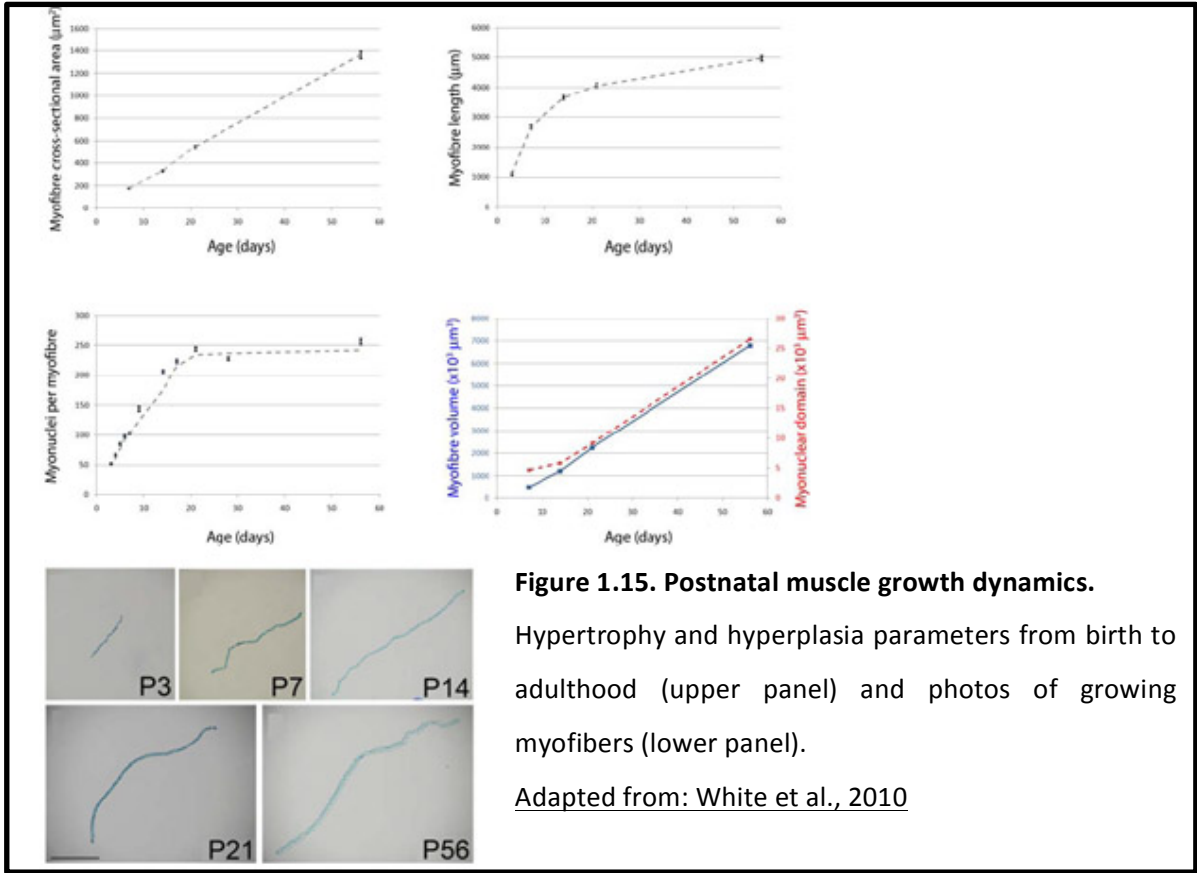
In vivo genetic manipulations abrogating PAX3, MYF5 or both, complemented by *in vitro* cultures in the presence of dominant negative forms, established that *Pax3* and *Myf5* define two distinct pathways, with *MyoD* lying genetically downstream [Tajbakhsh et al., 1997; Relaix et al., 2003]. *Pax3* and *Pax7* seem to act similarly in MYOD activation and skeletal muscle differentiation [Relaix et al., 2006].

1.4 From embryonic myogenic development to postnatal muscle

1.4.1 Embryonic and fetal waves of myogenesis

During development, myoblasts progressively differentiate and fuse to form the myofibers present in adult skeletal muscles [reviewed in Musumeci et al., 2015]. Differentiation includes increased synthesis of myofibrillar proteins (e.g. myosin heavy and light chains, skeletal muscle actin, tropomyosin, troponin) and increased activity of enzymes, such as creatine phosphokinase. Prior to fusion, myoblasts' shape and metabolic status undergo changes, including elongation, mitochondria amount increase, and sarcoplasmic reticulum development. The next step is a cadherin- and cell-adhesion-molecule-mediated adherence to fibronectin and alignment in chains. Eventually they fuse to give rise to myotubes, and further mature in myofibers, which are long syncytia consisting of postmitotic nuclei that will ultimately be located to the myofiber periphery (**Fig. 1.13**) [Musumeci et al., 2015]. Using the Fucci system (Fluorescent ubiquitynation-based cell cycle indicator), it was shown that myonuclei in terminally differentiated muscle fibers in fetal limb muscles as well as in adult myofibers have exited the cell cycle in G1 phase and are blocked in a G1-like state [Esteves de Lima et al., 2014].

The first muscle fibers start appearing at E11 in the mouse, and correspond to the so-called embryonic or primary myogenesis (**Fig. 1.14**) [Buckingham et al., 2003; Buckingham & Mayeuf, 2012]. This is followed by a second wave of myofiber formation after E14, around the previously formed primary myofibers [Buckingham et al., 2003; Buckingham & Mayeuf, 2012]. Primary and secondary fibers differ in their transcriptome and contractile proteins [Buckingham & Mayeuf, 2012].



Furthermore, Secondary/fetal myogenesis is associated with the establishment of definitive neuromuscular junctions and excitation-contraction coupling [Buckingham & Mayeuf, 2012]. Growth and patterning events prefigure the complex organization that persists from fetal to adult muscles [Relaix, 2006].

1.4.2 Postnatal muscle growth

Muscle growth is achieved by an increase in the number (hyperplasia) and/or size (hypertrophy) of myofibers. In the mouse, the number of myofibers is settled by birth, while subsequent muscle growth occurs by hypertrophy of existing myofibers, including increase in length, volume, and, importantly, myonuclei number (**Fig. 1.15**) [Moss & Leblond, 1971; White et al., 2010]. Myonuclei per myofiber increase by 5-fold between P3 and P21 [White et al., 2010]. Satellite cells support this hypertrophic phase (see section 1.5).

1.4.3 Adult muscle: structure & function

The entire adult muscle is ensheathed in a layer of connective tissue, called epimysium, while sets of neighboring myofibers are arranged in fasciculi covered by a further layer of connective tissue, called perimysium. Individual fibers are coated by the basal lamina (also called basement membrane or endomysium) [Tajbakhsh, 2009; Yin et al., 2013]. The interstitial stromal cell population residing between the epimysium and the basal lamina is mostly comprised by fibroblasts. Skeletal muscle also involves a microvascular network and neurons establishing neuromuscular junctions. Furthermore, a few immune cells reside within intact muscles [Yin et al., 2013].

Myofibers consist of myofibrils (bundles of myosin and actin filaments) subdivided in repetitive units (sarcomeres) giving a striated appearance. Within the myofiber, sarcoplasmic reticulum connects transverse tubules (t-tubules) and provides the calcium needed for muscle contraction and force generation (**Fig. 1.16**) [Cooper, 2000; Tajbakhsh, 2009; Yin et al., 2013]. Slow-contracting and fast-contracting muscle fibers (**Fig. 1.17**) differ in their myosin heavy chain expression profile and in

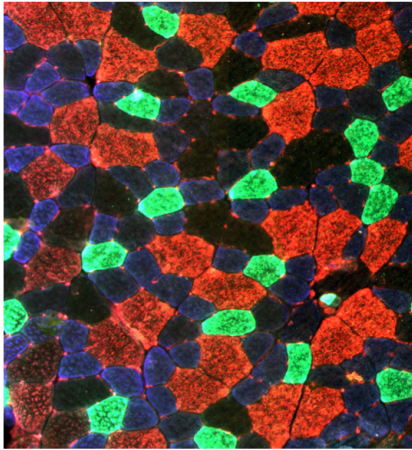


Figure 1.17. Skeletal muscle fiber types.

Muscle fiber typing based on different myosin heavy isoforms. Fiber types are as follows: MHC1 (green)-slow type I fibers, MHC2a (blue)-fast type IIA fibers, MHC2b (red)-fast type IIB fibers, MHC2x (unstained)-fast type IIX fibers.

Source: Liu et al., 2015

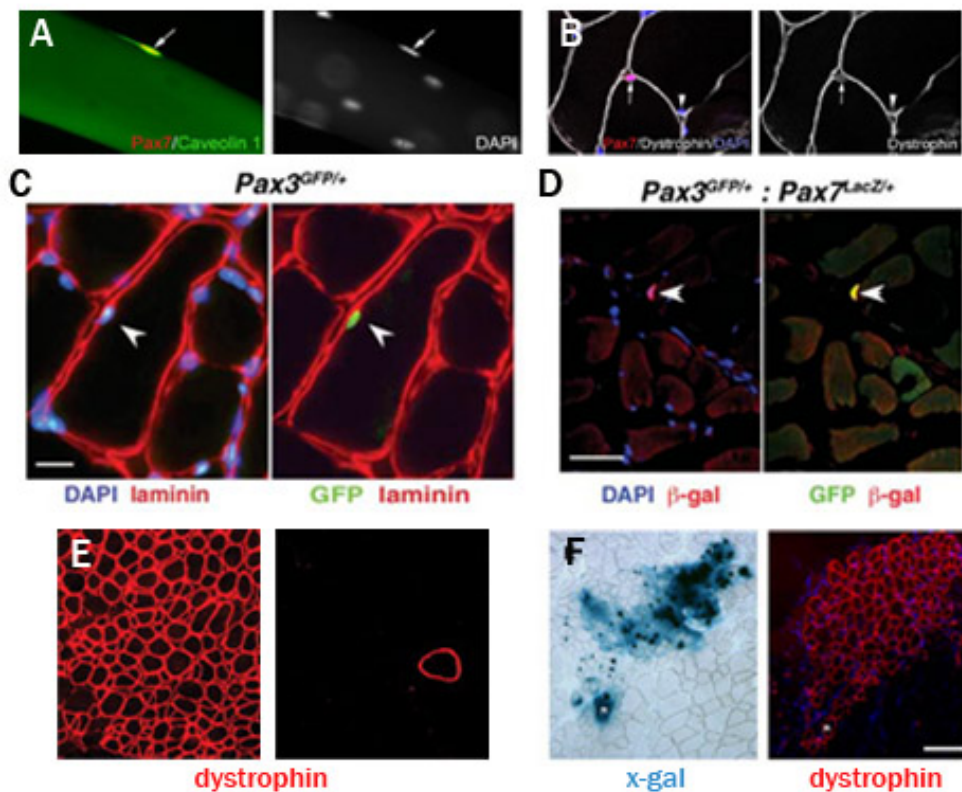


Figure 1.18. Satellite cells are skeletal muscle stem cells.

A) Satellite cell on isolated myofiber marked by PAX7/Calveolin-1. B) PAX7-marked satellite cell on muscle section, residing outside the dystrophin-outlined myofiber. C-D) Sublamellar PAX3⁺ or PAX3/7⁺ satellite cells in diaphragm. E-F) DYSTROPHIN restoration in *dystrophin*-deficient *mdx* mice three weeks following grafting of satellite cells (GFP⁺ cells of Pax3^{GFP/+} mice; E) or myofibers of 3F-nLacZ-2E mice (areas with donor-derived cells are detected by b-gal activity; F).

Adapted from: Collins et al., 2005; Montarras et al., 2005; Relaix & Zammit, 2012

physiological properties (e.g. oxidative metabolism and fatigue resistance in slow fibers) and they appear in varying proportions at different muscles [Haizlip et al., 2015; Luna et al., 2015]. Apart from contraction to support voluntary movement, skeletal muscle is also essential for breathing, posture maintenance, and metabolic aspects (e.g. heat production, storage of carbohydrates and amino acids) [Kharraz et al., 2013]. Loss of functionality – caused by disorders (i.e. myopathies) or naturally occurring during aging – has detrimental effects in strength, locomotion, and metabolic status, and can even result in lethality [Kharraz et al., 2013; Chang & Rudnicki, 2014].

Postnatal muscle growth, homeostasis, and repair depend on a population of resident muscle stem cells, termed satellite cells. They reside under the basal lamina of their adjacent myofiber, accounting for 2.5-6% of the muscle's nuclear content [Tedesco et al., 2010].

1.5 Satellite cells: the skeletal muscle stem cells

Originally described as “wedged between the plasma membrane of the muscle fiber and the basement membrane” by A. Mauro in 1961, satellite cells were named after their anatomical position in the periphery of the myofibers (**Fig. 1.18A-B**), under the basal lamina (**Fig. 1.18C**). They were predicted to represent dormant myoblasts ready to recapitulate the embryonic developmental myogenic program for myofiber repair [Katz, 1961; Mauro, 1961]. However, it was not until 2005 that their stem cell potential was proven, by virtue of a) providing differentiated progeny for muscle regeneration (**Fig. 1.18E-F**) and b) self-renewing their pool, upon engraftment of FACS-sorted satellite cells [Montarras et al., 2005] or single myofibers [Collins et al., 2005] into the muscles of *mdx* mice that lack dystrophin and undergo continuous regeneration. Recently, human satellite cells were also shown to support regeneration and repopulation of the satellite cell compartment after transplantation [Marg et al., 2014]. Importantly, muscle fails to regenerate in the absence of satellite cells. DTA (diphtheria toxin fragment A)-mediated ablation of satellite cells demonstrated the absolute requirement of satellite cells for muscle regeneration [Lepper et al., 2011; McCarthy et al., 2011; Murphy et al., 2011; Sambasivan et al., 2011]. The term stem cell often entails multipotency, although this is not an obligate criterion for stem cells [Tajbakhsh, 2003]. Indeed, satellite cells can be

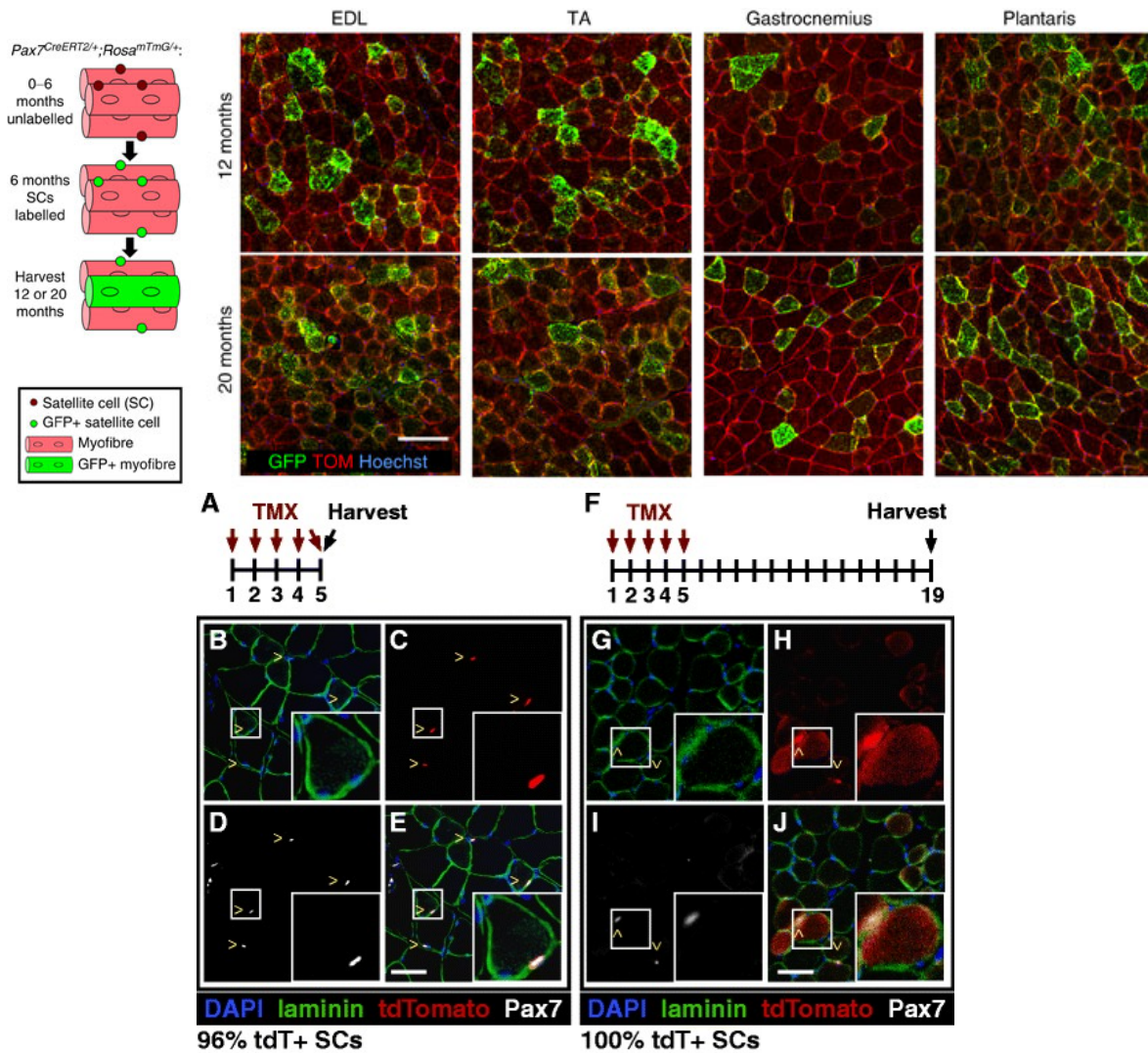


Figure 1.19. Adult satellite cell contribute to resting muscle.

Genetic tracing of PAX7+ satellite cells fusing with adult uninjured myofibers by TMX-inducible membrane-GFP (upper panel) or tdTomato (lower panel). EDL: *Extensor digitorum longus* muscle, TA: *Tibialis anterior* muscle, TMX: tamoxifen

Adapted from: Keefe et al., 2015; Pawlikowski et al., 2015

driven to adipogenic and osteogenic fates in culture [Asakura et al., 2001], arguing for multipotent nature. However, culture contamination by other lineages cannot be excluded and satellite cells are generally considered monopotent cells in physiological conditions [Relaix & Zammit, 2012].

1.5.1 Establishment during development

At late fetal stages (E16.5-E18.5) satellite cells become embedded under the basal lamina that forms and surrounds the muscle fibers, while remaining outside of the myofibers [Kassar-Duchossoy et al., 2005; Relaix et al., 2005]. Reporter-based tracing as well as quail-to-chick grafts place the origin of satellite cells back to the dermomyotome, similarly to the muscles in which they reside [Gros et al., 2005; Kassar-Duchossoy et al., 2005; Relaix et al., 2005]. These findings on shared developmental origin between muscles masses and their associated satellite cells were later extended to limb [Schienda et al., 2006] and head musculature [Harel et al., 2009]. PAX3/7 proteins are marking the progenitor cells that will give rise the satellite cells. While PAX7 is maintained and provides a reliable marker for the emerging and adult satellite cells, *Pax3* is downregulated in some muscles [Kassar-Duchossoy et al., 2005; Relaix et al., 2005; Relaix et al., 2006; Calhabeu et al., 2013].

1.5.2 Satellite cells in the control of postnatal growth and homeostasis

Postnatal muscle growth depends on myofiber size increase, in the mouse (see section 1.4.2). Of note, the number of myonuclei per myofiber undergoes a 5-fold increase within the first three weeks of life [White et al., 2010]. Satellite cells are the main contributors to this hypertrophic phenomenon. They proliferate rapidly and extensively and are the source of new myoblasts that fuse with existing myofibers, while myonuclei have stopped dividing [Moss & Leblond, 1971; Lepper et al., 2009; White et al., 2010].

Satellite cell fusion reaches a plateau around three weeks postnatally [White et al., 2010]. Thereafter, satellite cells were hypothesized to contribute to adult skeletal muscle homeostasis even in the

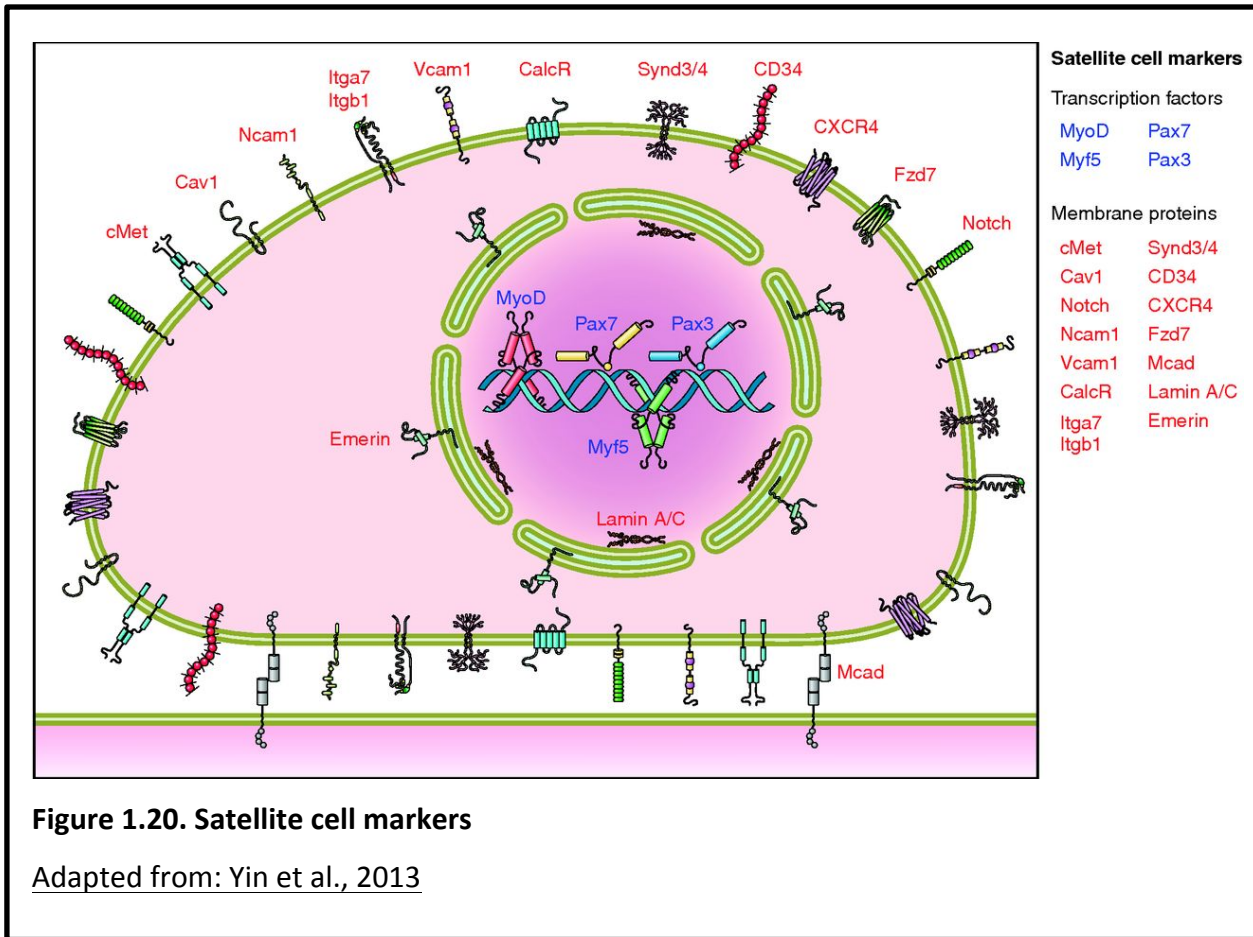
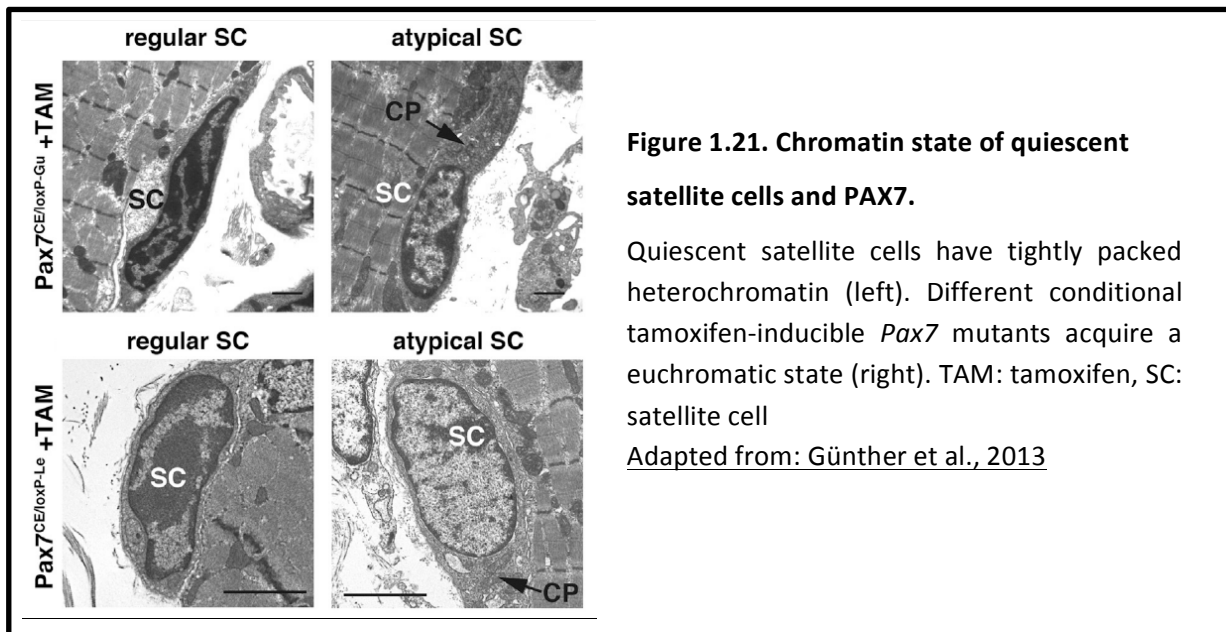


Figure 1.20. Satellite cell markers

Adapted from: Yin et al., 2013



absence of injuries. Indeed, genetic lineage studies provided experimental evidence for the predicted low rate fusion that constantly occurs (**Fig. 1.19**) [Keefe et al., 2015; Pawlikowski et al., 2015]. It should be stressed, however, that in general adult myofibers persist throughout the life in the absence of injury or myopathies that trigger a regeneration response resulting to new muscle formation [Grounds & Shavlakadze, 2011].

1.5.3 Acquisition of quiescence for function preservation

At three weeks of age (i.e. P21) there is a critical period of change from juvenile muscle/satellite cells to their form observed in the adult. Postnatally, satellite cells undergo a progressive number diminution and loss of proliferative capacity, which culminates in entering into a quiescent, non-cycling state around P21 [Lepper et al., 2009; White et al., 2010]. Long-standing efforts have described a series of markers (**Fig. 1.20**) to identify quiescent satellite cells, with PAX7 being central. Active Notch is fundamental to maintain quiescence (see Chapter 3), while Angiopoietin-1/TIE2 is a further signaling promoting this state [Abou-Khalil et al., 2009]. On the DNA level, the histone methyltransferase Suv4-20H1 was recently found to maintain satellite cell quiescence by promoting a heterochromatic state [Boonsanay et al., 2016]. Moreover, PAX7 has a rather unappreciated role in chromatin architecture modifications, with its loss leading to euchromatic morphology (**Fig. 1.21**) [Günther et al., 2013]. Quiescence preservation is also ensured by translation repression, via phosphorylation of the translation initiation factor eIF2a [Zismanov et al., 2016].

It is widely accepted, that the dormant quiescent state is adopted to preserve key functional features, since it is accompanied by low metabolism and higher resistance to DNA damage [Cheung & Rando, 2013; Wang et al., 2014]. As opposed to other forms of growth arrest (i.e. differentiation, senescence), quiescence is reversible (see section 2.1). This allows fast activation and reentry into the cell cycle upon specific needs, such as exercise or injury, in the case of muscle.

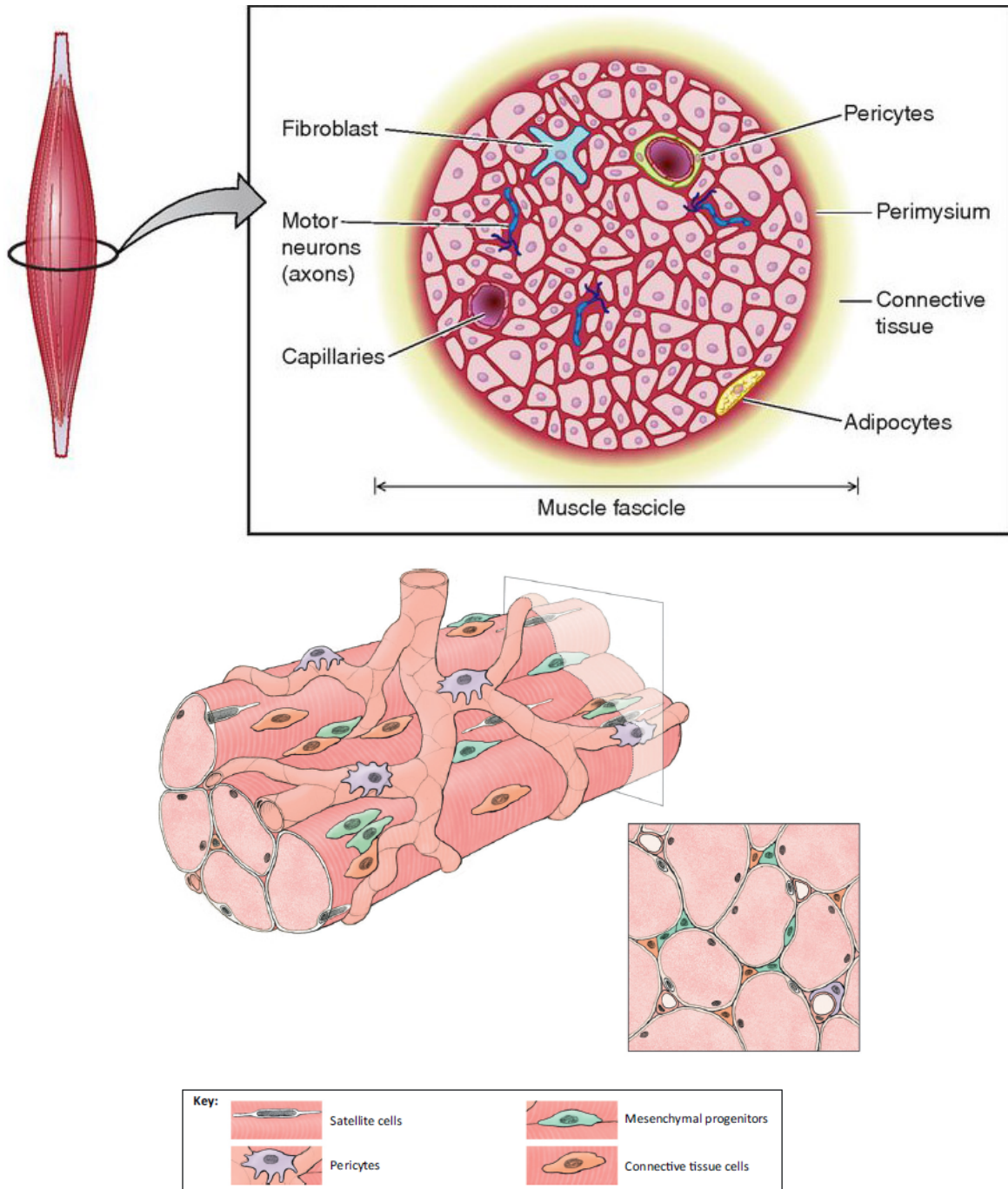


Figure 1.22. Satellite cell niche.

Adapted from: Pannérec et al., 2012; Yin et al., 2013

1.5.4 Satellite cell niche

Stem cells reside in a microenvironment that profoundly affects their properties and behavior, termed niche. It encompasses both anatomical and functional dimensions, meaning that it is in a defined anatomical location where a) the stem cells can proliferate yet remain in limited numbers and b) “stemness” maintenance is assured via differentiation inhibition [Scadden, 2006; Brack & Rando, 2012]. It is composed by heterologous cell types providing structural and biochemical cues that influence the stem cells. Satellite cell niche consists of extracellular matrix (ECM) and various types of surrounding cells, including interstitial cells, motor neurons, and blood vessels (**Fig. 1.22**) [reviewed in Pannérec et al., 2012; Yin et al., 2013]. ECM deposition defines its stiffness, which in turn impacts on satellite cell proliferation and differentiation as shown by *in vitro* studies with primary myoblast cultures and the C2C12 myogenic cell lines. Among the interstitial cells, there is a population of connective tissue fibroblasts, identified by TCF4, which interact with satellite cells during regeneration, while ablation of any of the two dysregulates the dynamics of the other. Platelet-derived growth factor receptor (PDGFR) α + mesenchymal progenitors and fibro/adipogenic progenitors (FAPs) also reside in the muscle interstitium, and may overlap with the TCF4 population. The balance between PDGFR α +cells/FAPs-mediated adipogenesis and satellite cell-dependent myogenesis drives normal muscle homeostasis and regeneration. Muscle contraction is directed by a neuronal network that signals to individual fibers. Finally, the muscle is nourished by a microvascular network, with satellite cells residing in close proximity to capillaries and their associated pericytes [session based on informative reviews by Pannérec et al., 2012; Yin et al., 2013].

1.5.5 Satellite cells in the control of regeneration

Skeletal muscle shows a remarkable regenerative capacity, with rapid functional and structural reestablishment (within three weeks) even after multiple rounds of severe injury that cause widespread necrosis [Relaix & Zammit, 2012]. Under both physiological (e.g. accidental injuries) and pathological (e.g. toxin-induced injury or diseases such as muscular dystrophies) conditions, satellite

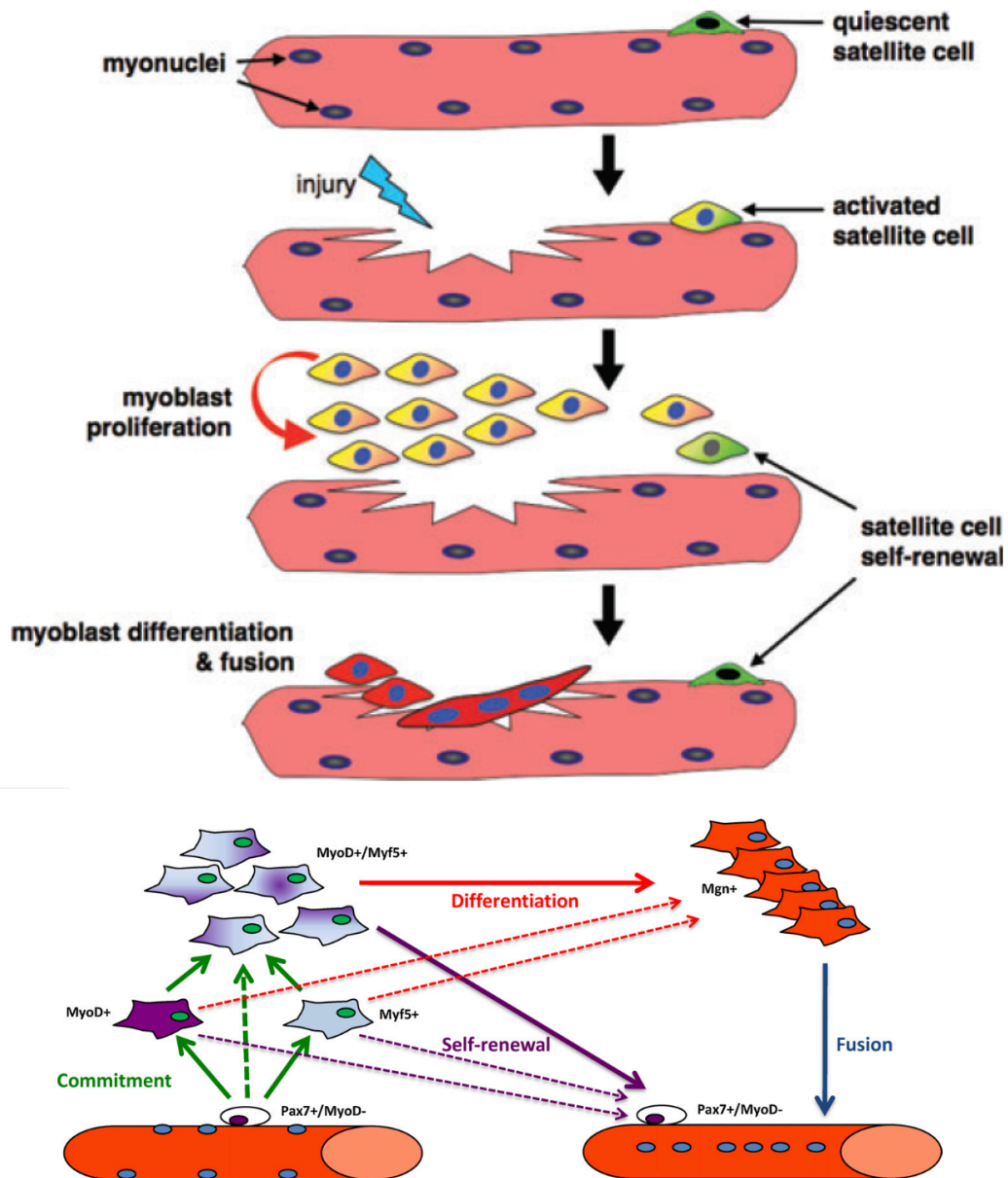


Figure 1.23. Activation of quiescence satellite cells supports muscle regeneration.

of PAX7/MYOD activated myoblasts proceeds to differentiation, downregulating *Pax7* and activating *Myogenin*, inducing fusion and new myofiber formation. A small subpopulation loses the MFRs but keeps PAX7 and self-renews the satellite cell pool.

Adapted from: Biressi & Rando, 2010; Olgúin & Pisconti, 2012

cells play a pivotal role during the remodeling and regeneration process. They are rapidly activated, reenter the cell cycle, expand, and express MRFs (MYF5 and MYOD, succeeded by MYOGENIN). The activated population initially co-expresses PAX7 and MYOD. Subsequently, most myoblasts downregulate *Pax7* but maintain MYOD, express MYOGENIN, and follow the myogenic program to supply differentiated progeny for muscle repair (**Fig. 1.23**). Importantly, a subpopulation of activated satellite cells does not engage into terminal differentiation, maintains PAX7 (while losing MYOD) and returns to the quiescent state. Hence, they replenish the satellite cell pool by self-renewal in order to support future needs (**Fig. 1.23**). Similarly, satellite cells activated *in vitro*, either by primary culture of satellite cell-derived myoblasts [Abou-Khalil et al., 2010] or by culture of isolated fibers with their associated satellite cells [Zammit et al., 2004], are induced to proliferate and (upon certain culture stimuli) differentiate, while a subpopulation bypasses differentiation and returns to quiescence. Signaling molecules and pathways crucial functioning during development are redeployed for regeneration, implying that this process recapitulates many, yet not all, aspects of embryonic myogenesis [discussed in Tajbakhsh, 2009].

While disturbed homeostasis and minor forms of muscle tissue injury can activate satellite cells (without myogenic fusion with myofibres), damage that results in myofibre necrosis can regularly result from accidents, surgical and orthopaedic situations and some degenerative neuromuscular disorders [Grounds, 2014]. The *in vivo* modelling and study of myofibre necrosis (and subsequent myogenesis and regeneration) employs many forms of experimental injury or transplantation, including intramuscular myotoxin administration or cryoinjury in rodents [compared in Lee et al., 2013; Hardy et al., 2016]. Muscle necrosis ignites the following four phases for reconstitution of structurally and functionally normal muscle tissue: degeneration/necrosis, inflammation, myogenesis and fusion as essential components of regeneration (that may sometimes also require revascularisation and re-innervation), and subsequent remodelling and maturation (**Fig. 1.24A**) [reviewed in Barberi et al., 2013; Yin et al., 2013]. Extracellular calcium influx induces myofiber proteolysis and necrosis, while muscle proteins and microRNAs are released from the cytosol. Necrotic material stimulates an inflammatory response to remove tissue debris from the lesion site and activate stem cell-mediated muscle repair and regeneration. Neutrophils, M1 macrophages, and

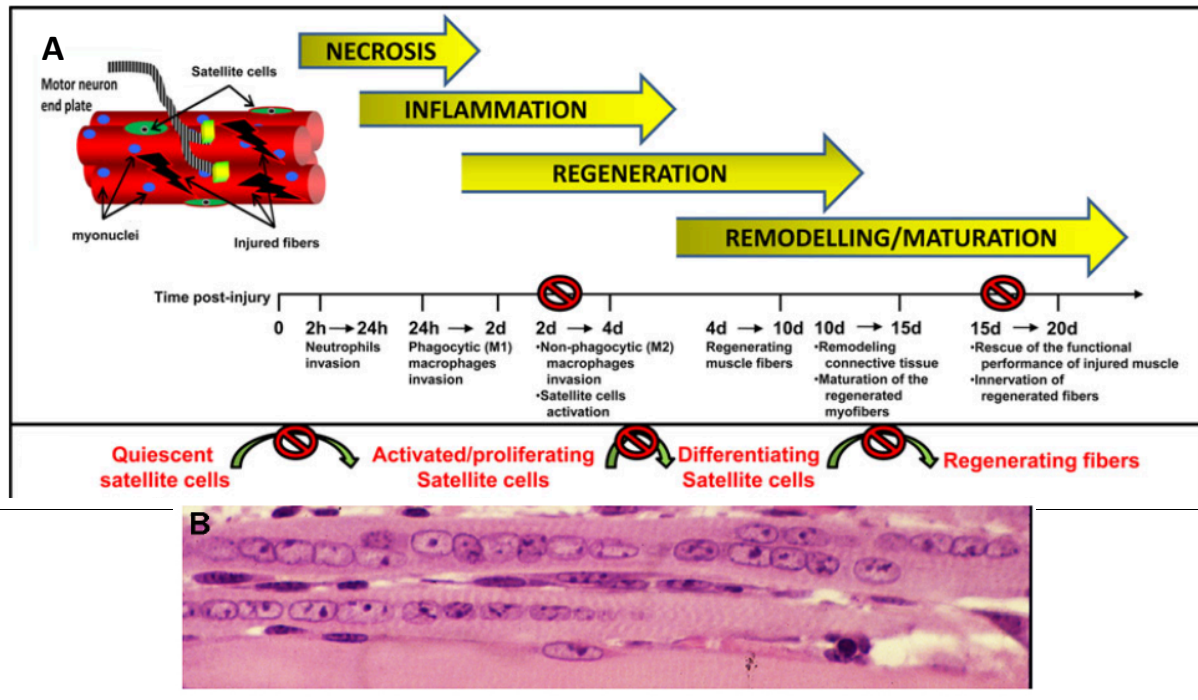


Figure 1.24. Muscle regeneration.

A) Timeline and satellite cells state of four major steps of the muscle regeneration procedure. Restrictive signs indicated stages susceptible to defects during aging. B) Characteristic central nuclei of regenerating muscle.

Adapted from: Barberi et al., 2013; Grounds, 2014

M2 macrophages sequentially invade the injured muscle, with macrophages being predominant. M2 macrophages progressively replace M1 and they are the ones that tune the inflammatory response, remove the debris, and promote remodeling. Inflammation is followed by a phase of active regeneration, characterized by satellite cell activation and expansion. Subsequently, a small proportion of satellite cells will replenish the quiescent pool, while the vast majority will provide committed myoblasts that fuse to form myotubes that later fuse with ends of remaining segments of damaged myofibres, or mature to form new myofibres as required, all with typical centrally located nuclei (**Fig. 1.24B**) [Grounds, 2014]. These steps of myogenesis with fusion to form myotubes can be reproduced in culture, helping thorough study of early myogenic events but often leading to misuse of the term regeneration [Grounds, 2014]. Recently, 3D time-lapse intravital imaging revealed that basal lamina remnants serve as “ghost fibers” to orient satellite cells and preserve muscle architecture [Webster et al., 2016]. Upon activation, satellite cells switch from an immobile to a migratory state *in vitro* [Siegel et al., 2009; Marg et al., 2014] and *in vivo* [Webster et al., 2016], which enables them to invade the lesion sites. The final steps of regeneration involve contractile apparatus maturation and reestablishment of innervation, vascular network, and extracellular matrix to ensure recovery of the functional performance. If any of these four phases is disrupted or misregulated, myofiber formation and muscle architecture are compromised, while excessive deposition of fibrotic and adipose tissue may also be observed, further hindering regeneration [Brzoska et al., 2011].

Satellite cells seem to be primed for myogenesis and quickly convert from a quiescent to an activated state. The observations of Crist et al. [2012] might explain this phenomenon, as they describe a model, whereby satellite cells transcribe *Myf5*, but post-translationally suppress it (through sequestration in messenger ribonucleoprotein granules by mir-31) to maintain quiescence. Accordingly, quiescent satellite cells in muscle sections and isolated myofibers of *Myf5^{nlacZ}* mice express β -galactosidase, further confirming the *Myf5* locus transcription [Gayraud-Morel et al., 2007; Kuang et al., 2007a; Tajbakhsh, 2009]. These results can also be interpreted as indicative of myogenic commitment by the majority of satellite cells before returning to quiescence. Satellite cells were shown *in vivo* to progress from G0 quiescence to the so-called G_{Alert} phase, in which they are primed for activation [Rodgers et al., 2014]. Upon activation, they upregulate *c-Met*, *Pax7*, and *M-cadherin* as

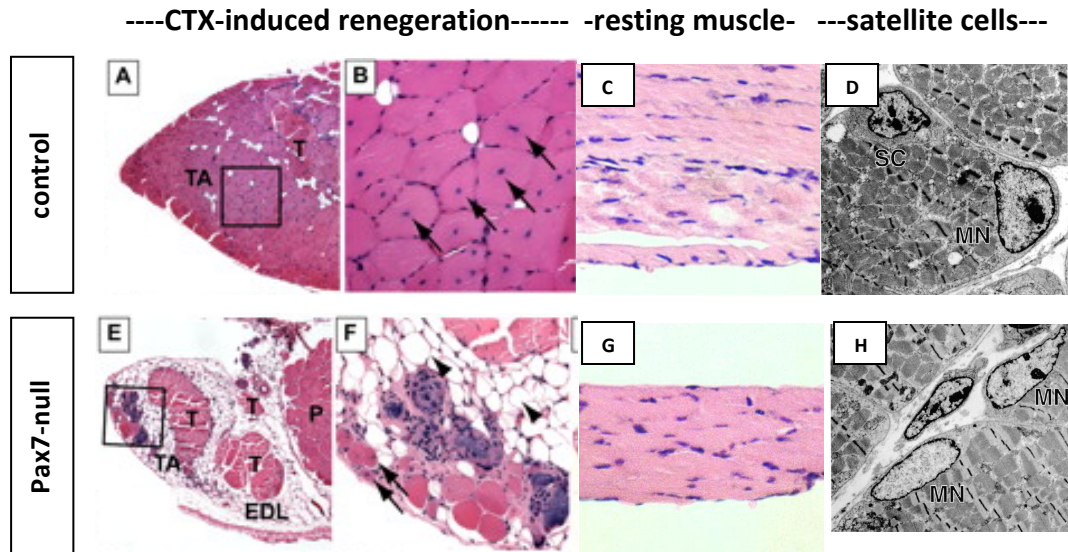


Figure 1.25. Regeneration deficit, muscle atrophy, and loss of satellite cells in the absence of *Pax7*.

Regenerative response, resting muscle and satellite cells of wild-type mice (A-D) or *Pax7*-deficient mice (E-H) reveal differences in muscle structure reestablishment post-injury (A, B, E, F), muscle mass (C, G) and satellite cell specification (D, H). CTX: cardiotoxin, EDL: *Extensor digitorum longus* muscle, MN: myonucleus, P: *Plantaris* muscle, SC: satellite cells, T: tendon, TA: *Tibialis Anterior* muscle.

Adapted from: Seale et al., 2000; Kuang et al., 2006

well as specification factors, such as *Myf5* and *MyoD* [summarized in Barberi et al., 2013]. *MyoD* upregulation correlates with post-injury [Grounds et al., 1992; Yan et al., 2003] or culture-triggered [Zammit et al., 2004] satellite cell activation. This possibly represents an early activation event, since *MyoD* is detectable as early as six hours post-trauma [Grounds et al., 1992]. Interestingly, when the methyltransferase Suv4-20h1, which is involved in heterochromatin formation and quiescence maintenance, is depleted, the *MyoD* locus is repositioned away from heterochromatic sites and transcriptionally activated [Boonsanay et al., 2016]. At the onset of differentiation, cells withdrawing from the cell cycle, express *Myogenin* and *Mrf4* and down-regulate *Pax7*, the latter possibly mediated by MYOGENIN [Seale et al., 2000; Zammit et al., 2004; Olguín & Olwin, 2004; Gayraud-Morel et al., 2007; Olguín et al., 2007]. Recent studies showed that the ubiquitin-ligase NEDD4 drives proteasome-dependent PAX7 degradation upon differentiation [Bustos et al., 2015], while casein kinase 2-dependent PAX7 phosphorylation ensures its maintenance in proliferating progenitors [González et al., 2016]. Either PAX3/7, acting through MYOD, or MYF5 is required for satellite cell-driven myogenesis [reviewed in Buckingham, 2007]. Intriguingly, PAX3/7+ satellite-like cells also support muscle regeneration in arthropods, extending the vertebrate findings and suggesting early evolution of this repair strategy [Konstantinides & Averof, 2014].

PAX7 is essential for satellite cells function during adult muscle growth and regeneration. In its absence, postnatal muscle growth and regeneration are severely affected, due to the progressive loss of satellite cells (**Fig. 1.25**), even in muscles expressing *Pax3*, the paralogue of *Pax7* [Seale et al., 2000; Oustanina et al., 2004; Kuang et al., 2006; Relaix et al., 2006]. Specifically, body weight and muscle mass are reduced, muscles consist of smaller myofibers with less myonuclei, and upon injury there is negligible fiber formation accompanied by extensive adipocyte, calcium, and fibrotic deposition. The Braun group reported that *Pax7* mutant phenotype is sensitive to the genetic background. In a sv129 background, *Pax7*-deficient juvenile muscles are nearly normal and do not present with a complete lack of satellite cells; nevertheless, adult muscles display severe deficiencies in homeostasis or regeneration associated with delayed loss of satellite cells [Oustanina et al., 2004]. Conditional ablation of PAX7+ satellite cells in adult mice using different tamoxifen-inducible Cre lines, demonstrated its critical role for adult satellite cell function, as in its absence skeletal muscle repair

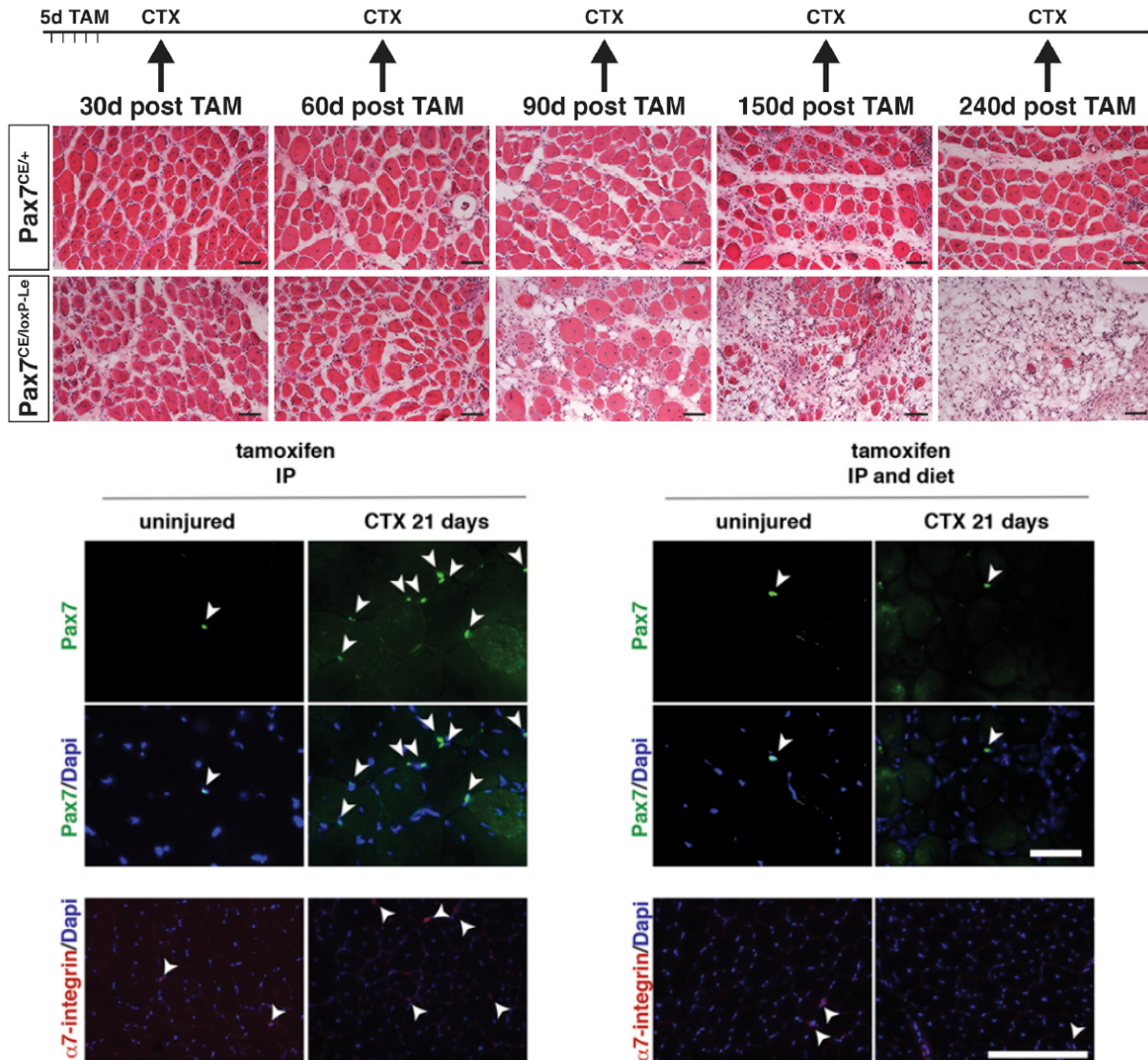


Figure 1.26. Applying tamoxifen scheme might mask the limitations of the loxP-Cre system.

Different chase periods (upper, lower panel) and tamoxifen administration during regeneration (lower panel) have diverse outcomes on PAX7+ satellite cell elimination (PAX7+ cells marked with arrowheads in lower panel) and subsequent regeneration. CTX: cardiotoxin, TAM: tamoxifen.

Adapted from: Günther et al., 2013; von Maltzahn et al., 2013

was strongly compromised [Günther et al., 2013; von Maltzahn et al., 2013]. These studies ignited a debate with the first work with conditionally Pax7-depleted muscles which reported functional satellite cells and efficient regeneration [Lepper et al., 2009]. The differences have been attributed to (a) the inefficient CreERT2-mediated recombination in quiescent satellite cells, (b) the limitations of the Cre-LoxP technology and the presence of “escaper” cells; suboptimal gene targeting occurred when the chase period was not long or tamoxifen was not administered during regeneration (**Fig. 1.26**), (c) the possible generation of truncated PAX7 by one of the used alleles [discussed in Brack, 2014; Buckingham & Relaix, 2015]. Even few wild-type satellite cells are capable of replenishing the satellite cell pool and out-compete the recombined majority [von Maltzahn et al., 2013].

A subset of MRFs is also important for muscle homeostasis and regeneration in the adult, as revealed by genetic manipulations in mice. Opposite to the embryo situation, MYF5 and MYOD do not seem to effectively compensate for each other in the adult. MYF5 deficiency results in delayed muscle regeneration after injury, possibly due to delayed transition from proliferation to differentiation [Gayraud-Morel et al., 2007; Ustanina et al., 2007]. Resting muscles of young MYF5-deficient adults are comparable to controls, while satellite cell amount and quiescent state (evaluated by electron microscopy) are preserved [Gayraud-Morel et al., 2007; Ustanina et al., 2007]. However, old MYF5 mutant muscles show increased fibrosis and centrally-located nuclei, indicative of a cumulative defect following chronic regeneration [Gayraud-Morel et al., 2007]. MYOD-null mice have more satellite cells, but they show a defective balance in proliferation/self-renewal and myogenic progression resulting in delayed regeneration [Megeney et al., 1996; Yablonka-Reuveni et al., 1999; White et al., 2000]. MYOD-deficient donor myoblasts show increased migrations, possibly as a side effect of the proliferation/differentiation dysregulation [Smythe & Grounds, 2001]. MYOGENIN deficiency at post-embryonic-myogenesis stages does not compromise muscle postnatal growth in terms of structure or function [Knapp et al., 2006]. Satellite cell-derived myoblasts lacking MYOGENIN grow and differentiate in culture as controls [Meadows et al., 2008]. However, muscles have not been challenged with injury to evaluate the full potential of their satellite cells in repair. Finally, MRF4 is absent from quiescent satellite cells or their activated progeny until differentiation initiation and there are no signs of upregulation upon MYF5 deficiency; thus, its participation at the early stages of

satellite cell function was excluded and no further analyses have been performed [Gayraud-Morel et al., 2007].

Apart from the critical contribution of satellite cells in muscle regeneration, several other populations have myogenic potential [reviewed in Pannérec et al., 2012; Yin et al., 2013]. *In vitro* or upon transplantation in compromised muscles, they have myogenic capacity, participate in the repair procedure by myofiber formation, and some even contribute to the satellite cell compartment. Initiated with experiments with bone-marrow-derived cells, these findings were later expanded in several more populations, including a hematopoietic population referred to as side population (SP), CD133+ cells, PW1+ interstitial cells (PICs), mesangioblasts, and the vessel-associated pericytes. However, none of these cell types is able to form new muscle or replenish the satellite cell pool in the absence of satellite cells, as demonstrated by regeneration analyses in muscles depleted of satellite cells. In fact, the defective regeneration was only rescued by wild-type satellite cell engraftment [Sambasivan et al., 2011].

Since satellite cell characterization as adult skeletal muscle stem cells, their studying progressed immensely, with implications in regenerative medicine. However, technical challenges hold their therapeutic promise back, since they are isolated in small amounts from muscle biopsies and expansion in culture is challenging due to rapid and irreversible differentiation *in vitro* [Bentzinger et al., 2012]. Similarly, *in vitro* genetic corrections and subsequent autologous transplantation cannot be accomplished [Bentzinger et al., 2012]. Amelioration of culture conditions resembling the characteristics of the niche might resolve these problems, raising hopes for implementation in humans. For instance, substrate elasticity mimicking the one of muscle favors self-renewal *in vitro* and efficient regeneration after transplantation [Gilbert et al., 2010]. Similarly, enhanced performance can be achieved by treatment with the bisperoxovanadium, a phosphotyrosine phosphatase inhibitor [Smeriglio et al., 2016].

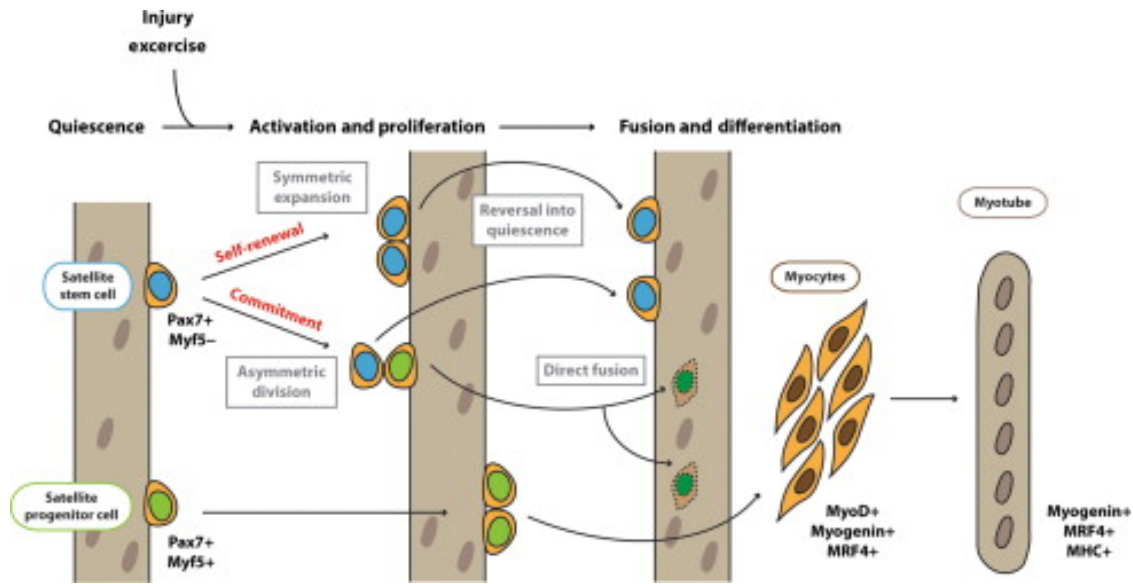


Figure 1.27. Symmetric and asymmetric divisions of Myf5⁻ or Myf5⁺ satellite cells.

PAX7+MYF5⁻ satellite cells represent a more stem population able to provide the more committed PAX7+MYF5⁺ satellite cells but also self-renew, undergoing apico-basal/asymmetric or planar/symmetric division, respectively. On the contrary, PAX7+MYF5⁺ satellite cells provide only committed progeny (by symmetric division) that will sustain differentiation.

Source: Chang & Rudnicki, 2014

1.5.6 Satellite cell heterogeneity

Satellite cells are considered a heterogeneous population, in terms of gene expression signatures, proliferation rate, differentiation propensity, and, importantly, stemness. Satellite cells originating from different muscles show variations (e.g. in amount, gene expression profile, proliferation/differentiation kinetics) [Ono et al., 2010]; however, the above properties are also subject to intra-muscle variations. Live imaging revealed differences in the rates of cell cycle reentry and subsequent divisions [Siegel et al., 2011; Webster et al., 2016]. Other groups have linked this variation to subpopulations defined by different markers. A subset of satellite cell markers are not expressed by the entire population [Beauchamp et al., 2000; Gnocchi et al., 2009], including *Pax3*, which is postnatally restricted to a subset of muscles [Relaix et al., 2006; Calhabeu et al., 2013]. Furthermore, markers that are considered universal, such as PAX7, do not have uniform levels [Rocheteau et al., 2012]. Higher *Pax7* expression characterizes a more stem/less committed subpopulation, with lower metabolic rates, delayed activation, and higher self-renewal ability and regeneration capacity. At cell division, PAX7^{Low} cells segregate their chromosomes randomly and provide two committed daughters, while PAX7^{High} cells perform asymmetric DNA segregation, with the template-DNA-retaining daughter exhibiting a more stem-like phenotype [Rocheteau et al., 2012]. This is in line with early reports of asymmetric divisions leading to inheritance of the older template strand to the daughter cell with more immature and self-renewing phenotype [Shinin et al., 2006; Conboy et al., 2007]. A further intrinsic factor with asymmetric distribution and preferential transmission to the self-renewing daughter cell is the cilium, a structure present in quiescent satellite cells but disassembled in activated ones [Jaafar Marican et al., 2016].

Variability in *Myf5* locus activity has also been noted among satellite cells, but their possible origin from the embryonic co-existing PAX7+MYF5+ and PAX7+MYF5- populations [Picard & Marcelle, 2013] has not been investigated. Following muscle damage and induction of regeneration, a subset of satellite cells undergoes activation, proliferation, and early differentiation phases, presumably without ever activating MYF5 [Cooper et al., 1999]. Similarly, only 87-90% of quiescent adult satellite cells are β -galactosidase+ and YFP+ in *Myf5*^{nLacZ} and *Myf5*^{Cre}; *Rosa26*^{YFP} mice, respectively [Kuang et

al., 2007a]. The reporter-negative cells may represent a more stem population [Kuang et al., 2007a], although after *Myf5*-Cre ablation of PAX7+ satellite cells, surviving satellite cells do not replenish the pool and cannot rescue the regeneration phenotype [Günther et al., 2013]. MYF5- satellite cells are able to give rise to both compartments, as opposed to their MYF5+ counterparts [Kuang et al., 2007a]. Asymmetric divisions seem to account for the generation of the latter (**Fig. 1.27**). Symmetric divisions give rise to two identical daughter cells, while asymmetric ones contribute daughter cells with distinct fates. Mitotic spindle orientation perpendicular to the myofiber drives an asymmetric apico-basal division, whereby a PAX7+MYF5- cell gives rise to an apical PAX7+MYF5+ and a basal PAX7+MYF5- daughter [Kuang et al., 2007a]. Non canonical Wnt pathway was shown to promote symmetric expansion of satellite cells [Le Grand et al., 2009], while dystrophin cooperates with Par proteins to drive asymmetric divisions [Dumont et al., 2015] in addition to its well-defined traditional role in maintaining sarcolemma integrity. Time lapse imaging revealed that cells resulting from vertical divisions remain associated longer than cells from planar divisions [Siegel et al., 2011]. Notably, recent intravital imaging studies render these observations unlikely in the *in vivo* situation, where divisions predominantly occur in parallel to the axis of the myofibers [Webster et al., 2016].

The histone H2B-GFP pulse/chase system allowed a further classification of satellite cells based on label retention [Chakkalakal et al., 2014]. Label-retaining cells (LRCs) present with a more stem signature, associated with ability to self-renew, earlier entry into quiescence, rare divisions, and lower propensity for expansion and differentiation. They can give rise to both LRCs and differentiation-competent non-LRCs. The two populations are already established at birth.

While different assays classify satellite cells into a more stem and/or a more committed compartment, the degree of overlap between the subpopulations defined by different methods still remains to be established.

1.5.7 Aging effect in muscle and satellite cells

Advanced age results in sarcopenic muscles, characterized by reduced mass and function, with implications on personal (e.g. independence, quality of life) and socio-economical levels (e.g. health costs) [Grounds, 2014]. Sarcopenia has a multifactorial etiology and both intrinsic (e.g. satellite-cell-related) and extrinsic (e.g. microenvironment, inflammatory status, innervation) factors contribute to the observed regenerative decline (**Fig. 1.22A**) with aging. Reduced numbers, proliferative capacity, and telomere length of satellite cells have been associated to the defective phenotype [reviewed in Barberi et al., 2013]. As opposed to young (2-6 months) or old (20-24 months) satellite cells, geriatric ones (28-32 months) undergo a so-called geroconversion, shown by p16-mediated switch into an irreversible pre-senescent state, which severely affects their intrinsic regenerative and self-renewal potential [Sousa-Victor et al., 2014]. The same group later linked age-related decline in autophagy with the quiescence-to-senescence shift upon aging [García-Prat et al., 2016]. Furthermore, compromised function of the neuromuscular function and myofiber denervation are associated to sarcopenia [Barns et al., 2014; Krishnan et al., 2016]. Heterochronic parabiosis experiments (linking the blood supply of young and old mice) rescue the regenerative defects of aged mice, underlining the importance of systemic factors [Conboy et al., 2005]. However, the intrinsic defects such as senescence marker expression and increased p38 α / β MAPK signaling cannot be counteracted by the young microenvironment and rather depend on pharmacological treatment [Bernet et al., 2014; Cosgrove et al., 2014]. Study design seems to be an unappreciated factor strongly influencing the reported conclusions. Several factors affect the final outcome, including short-term (5 days post-injury) or long-term (28 days post-injury, when regeneration procedure is complete) follow-up, applied methodologies to induce a regenerative response (preserving or not the neurovascular supply), age range (including or not geriatric individuals), and assessed endpoints (e.g. overt muscle architecture reestablishment, satellite cell status and self-renewal).

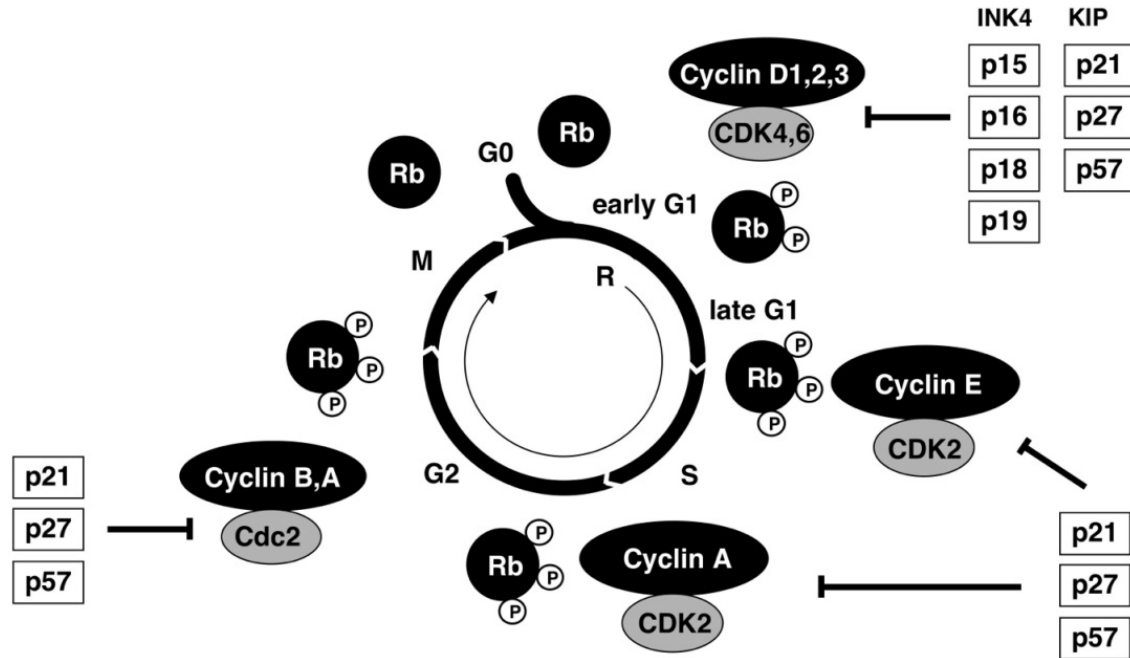


Figure 2.1. Cell cycle progression and arrest.

Non-cycling cells are maintained at the resting, non-proliferating phase called G0, while cycling cells undergo S-phase (Synthesis; DNA replication step) and M-phase (Mitosis; division step), which are separated by two "gap phases" (G1, G2). Cell cycle progression depends on the activity of various complexes of Cyclin-Dependent Kinases (CDKs) and cyclins, acting on different cell cycle transitions. When active, these complexes phosphorylate the retinoblastoma protein (Rb), blocking its inhibitory activity on proliferation. Two families of Cyclin-Dependent Kinase Inhibitors counteract the activity of CDK-cyclin complexes at G1 phase (family INK4, encompassing p15, p16, p18, p19) or throughout the cell cycle (family Cip/Kip, including p21, p27 and p57).

Source: Müller, 2010

Chapter 2. Cell cycle and growth arrest in skeletal muscle and beyond

Tightly controlling cell cycle is imperative throughout the life of complex organisms. Dysregulated cell cycle has been associated with developmental abnormalities [Yan et al., 1997; Zhang et al., 1997], severe organ malfunction and defective stem cell self-renewal [Matsumoto et al., 2011; Sherr, 2012] or stem cell loss [Kozar et al., 2004]. Moreover, perturbations of the cell cycle are linked to cancer [Sherr, 2012], aging [Matheu et al., 2009] and age-related pathologies [Sherr, 2012; Chandler & Peters 2013]. Cell cycle regulation and differentiation are intimately linked, although they can be uncoupled [Zalc et al., 2014]. Appropriate embryonic growth and patterning depend on a balance between proliferation and differentiation, to ensure sufficient propagation but not at the expense of forming differentiated structures [Ciemerych et al., 2011]. Once growth or tissue repair are completed, cell division is no longer required to guarantee a healthy homeostasis. Yet, in the absence of proper cell cycle control, cancer may develop. Notably, among the hallmarks of cancer are sustained proliferation, limitless replicative potential, and evasion of growth suppressors or apoptosis [Khabar, 2016]. Thus, interrupting cell cycle holds therapeutic promise to arrest cancer growth [Blachly et al., 2016].

Cell cycle progression is primarily achieved by an array of Cyclin-Dependent Kinases (CDKs) and their activating subunits named cyclins [Malumbres, 2014]. The regulation of these CDK-Cyclin complexes depends on the Cyclin-Dependent Kinase Inhibitors (CDKIs), which are classified in two families based on their structural homology and specificity of action (Borriello et al., 2011): INK4 [including p15^{Ink4b} (p15), p16^{Ink4a} (p16), p18^{Ink4c} (p18), p19^{Ink4d} (p19)] and Cip/Kip [including p21^{Cip1} (p21), p27^{Kip1} (p27), p57^{Kip2} (p57)]. In case of CDKI-mediated inhibition, the CDK-Cyclin complexes lose the capacity to phosphorylate the retinoblastoma protein (pRB) [Cobrinik, 2005]. In its hypophosphorylated form, pRB can target factors of the E2F family, negatively affecting E2F-responsive genes, which are maintaining the cell in cycle (**Fig. 2.1**) [reviewed in van den Heuvel & Dyson, 2008].

A more in-depth presentation of the molecules described above is provided in the sub-sessions to follow. Relevant discoveries in the muscle field are summarized in the end of each sub-session.

Substantial work has identified essential participants of *in vitro* cell cycle exit and differentiation of muscle cells. Nevertheless, less data are available to confirm this interplay *in vivo* and, most importantly, to unravel the mechanism of satellite cell entry into quiescence.

2.1 Cell cycle overview

During the cell cycle, cells consecutively pass through four phase (**Fig. 2.1**):

- G1 (Gap 1), during which the cells are metabolically being prepared for division. They grow in size, produce RNA and proteins and increase their supply in organelles. The G1 checkpoint safeguards that everything is ready for DNA synthesis.
- S (Synthesis), during which DNA replication occurs. The genetic material is duplicated guaranteeing that each daughter cell will inherit a complete copy of the mother cell genome.
- G2 (Gap 2), during which the cells continue to grow. The G2 checkpoint is the control mechanism ensuring that the cell is ready to divide.
- M, during which the cells undergo nuclear division (mitosis), followed by cell division (cytokinesis).

After division, cells return to G1 and will either proceed to a further division or enter G0 phase. The latter is a resting phase characterizing cells that withdraw from the cell cycle either permanently (differentiation, senescence) or reversibly (quiescence). It is a post-mitotic, non-proliferative state. Quiescence allows cells to start cycling upon specific needs. For instance, following tissue damage, quiescent myogenic stem cells get activated, proliferate to amplify their population, and are available for fusion and muscle regeneration where necrosis has occurred. In contrast, cells driven to terminal differentiation post-mitotically, such as myonuclei, do not re-enter the cell cycle in physiological conditions [Cheung & Rando, 2013]. Similarly, cells may cease to divide and enter into (irreversible) senescence, in response to developmental cues/regeneration (physiological context) [Muñoz-Espín et al., 2013; Storer et al., 2013; Le Roux et al., 2015] or damage/stress (pathological context) [Muñoz-Espín & Serrano, 2014].

Apart from the temporal orchestration of the molecules forming the core cell cycle machinery, chromatin changes are intimately linked to cell cycle regulation. Global and local chromatin architecture and packaging dictate transitions between chromatin compaction and decompaction and render the chromatin non-permissive or permissive, respectively, to DNA-templated processes, such as transcription, replication, and repair [Yu et al., 2016]. Thus, chromatin accessibility and condensation are coordinated with the cell cycle to tune specific genome activities.

2.2 CDK-Cyclin complexes: cell cycle progression

General features in cell cycle regulation

The cell cycle is orchestrated by a cascade of CDKs and their associated cyclin subunits. The importance of these complexes in promoting transitions through the cell cycle was established in the late 90s. Less than a decade later, the Nobel prize in Physiology or Medicine (LH Hartwell, T Hunt, PM Nurse; 2001) internationally acknowledged the characterization of these key cell cycle players of all eukaryotic organisms, identified across the eukaryote kingdom, from yeasts to plants and animals.

CDKs are serine/threonine kinases that acquire their catalytic activity via binding of a cyclin molecule and the resulting conformational modifications [reviewed in Malumbres, 2014]. There is considerable specificity in CDK-Cyclin interactions as well as in the cell cycle phase to be controlled (**Fig. 2.1**). The first, and most important, restriction point is the G1 checkpoint, that has to be overcome for cells to proliferate. At early G1, complexes of Cyclin D with CDK4 [Quelle, 1993] or CDK6 suppress the antiproliferative effects of pRb protein, while CDK2-Cyclin E complexes act at later G1 in a pRb-independent way [Resnitzky & Reed, 1995]. CDK2 further participates in S-phase interaction, but at that phase it binds with Cyclin A [Pagano et al., 1992]. Mitosis is then initiated by the complexes of CDC2 (CDK1) with Cyclin B or Cyclin A [Draetta et al., 1989; Pagano et al., 1992; Kishimoto & Okumura 1997].

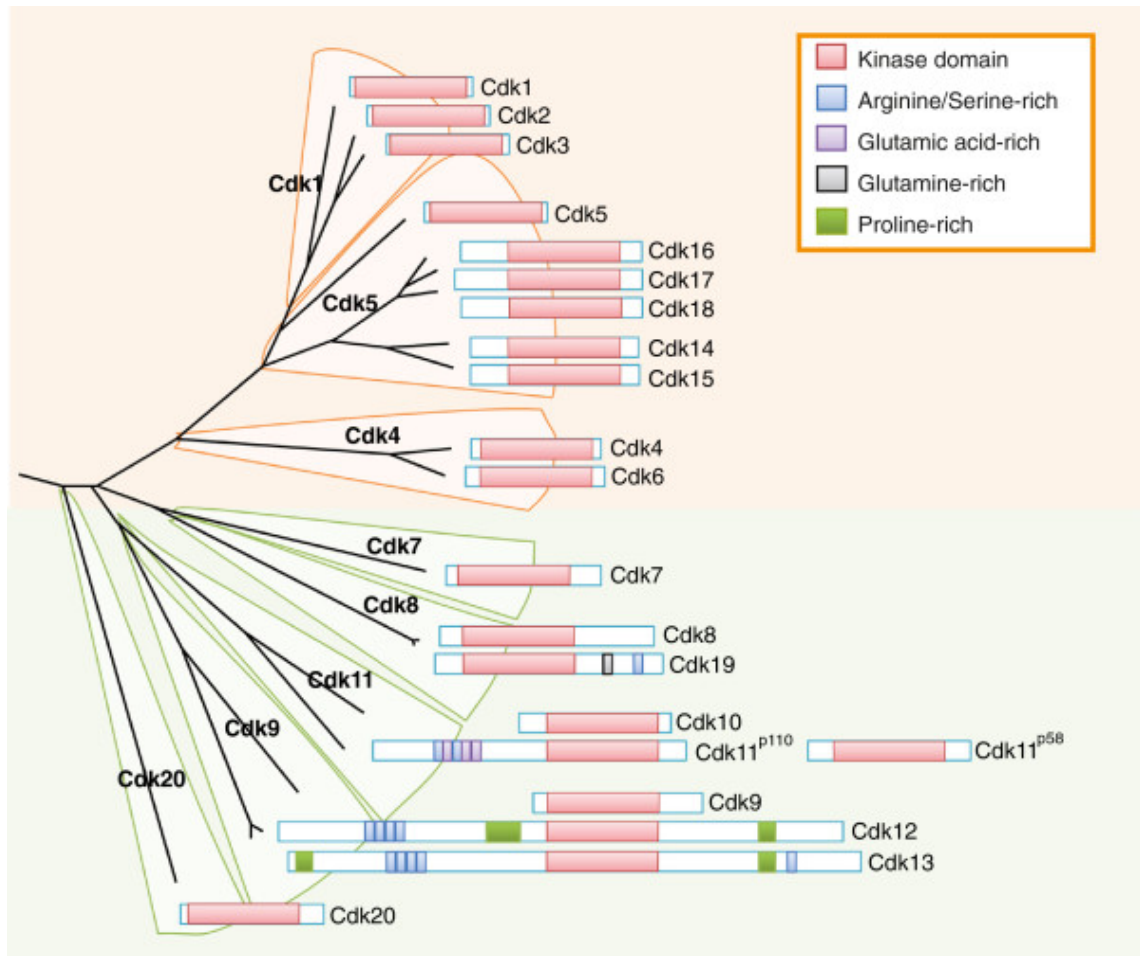


Figure 2.2. Phylogenetic tree of mammalian CDK families.

Eight CDK families (left part of the tree) encompass 20 CDKs (right part of the tree), functioning in cell cycle (orange; top) or transcription (green; bottom). All CDKs share a conserved kinase domain (red), while some possess further domains (as indicated in key).

Source: Malumbres, 2014

The mechanisms governing CDK activity involve i) activation by cyclin binding and by CDK phosphorylation at a conserved threonine and ii) repression by CDKI binding (subsession 2.3) or by phosphorylation at inhibitory sites near the N terminus. Like other kinases, CDKs have a two-lobed structure with the active site for ATP binding and catalysis being in a deep cleft at the junction of the lobes [de Bondt et al., 1993]. Although the biochemical features of different CDK-Cyclin complexes may vary, the model arising from CDK2 studies is well accepted to mirror how the CDK-Cyclin association is the key step for CDK activation. Structural analyses of monomeric and cyclin-bound CDK showed that the CDK catalytic cleft of the cyclin-free form is closed, preventing enzymatic activity. However, cyclin binding reorients key side chains, restoring active sites to their correct position [de Bondt et al., 1993]. To acquire full activation, the CDK-Cyclin complex is then phosphorylated by the Cdk-Activating Kinase (CAK), which is believed to be the complex CDK7-Cyclin H-Mat1 in higher eukaryotes [Fisher & Morgan, 1994; Devault et al., 1995]. Moreover, CAK-mediated phosphorylation promotes the formation of high-affinity complexes in some CDK-Cyclin cases [Desai et al., 1995]. Inhibitory phosphorylation is, by definition, non-CAK-induced. CDK4 and CDC2 are phosphorylated during G1 and G2 progression, respectively, and their dephosphorylation is required for S-phase and mitosis entry, respectively [Terada et al., 1995; Mueller et al., 1997]. Increased CDK4 phosphorylation is observed following DNA damage, causing G1 arrest [Terada et al., 1995].

Cyclins constitute a large, diverse family, ranging in size from 35 to 90 kDa [Malumbres, 2014]. Despite the substantial sequence differences, considerable structural similarities are suggested [Nugent et al., 1991]. Cyclins are structurally defined by a 100-residue domain called cyclin box, which is required to bind and activate CDKs [Nugent et al., 1991; Kobayashi et al., 1992; Lees & Harlow, 1993]. Different cyclins appear at different phases of the cell cycle, with D/E functioning at G1, E/A at S phase, and A/B at mitosis (**Fig. 2.1**). As implicated by their name, cyclins are subject to oscillations during the cell cycle. These fluctuations depend on gene transcription and protein degradation [Morgan, 1977]; remarkably, proteolysis of mitotic cyclins is essential for mitosis exit [King et al., 1996]. In the case of cyclin B there is additional regulation by subcellular translocation [Gallant et al., 1995].

Apart from the three CDK families regulating the cell cycle, there are a few more that are functioning in transcription. Those are more conserved (**Fig. 2.2**) and they are regulated by mechanisms different from cyclin level oscillations [Malumbres, 2014].

Implications in skeletal muscle cell cycle regulation

Dividing and differentiating myoblasts have been extensively analyzed for the presence and activity of various CDK-Cyclin complexes. Both early G1 CDKs (i.e. CDK4/6) levels were constant during myocyte differentiation [Wang & Walsh, 1996b]. Their cyclin partners showed divergent patterns: i) Cyclin D1 was high in the proliferating population and low in the differentiating one, ii) Cyclin D2 was not detectable in any population, and iii) Cyclin D3 unexpectedly increased upon differentiation [Rao et al., 1994; Kiess et al., 1995; Skapek et al., 1995; Wang & Walsh, 1996b]. Among the CDKs, CDK4 has been extensively studied. It was found to be expressed in both myoblasts and myotubes [Shapek et al., 1995]; however, in myotubes it appeared to not be associated with cyclins but interacted with the inhibitors p21 and p27 [Wang & Walsh, 1996b]. Moreover, Cyclin D1-dependent subcellular localization has been shown. CDK4 was nuclear in the dividing Cyclin D1-expressing myoblasts, but translocated to the cytoplasm of the forming myotubes, that lacked Cyclin D1 [Zhang et al., 1999a]. Chemical inhibition of CDK4/6 promoted G1 accumulation and enhanced muscle-specific expression [Saab et al., 2006]. Conversely, forced expression of CDK4 and cyclin D1 in C2C12 or primary myoblast cultures triggered DNA synthesis in myotubes [Latella et al., 2001]. As far as differentiation is concerned, ectopic CDK4-CyclinD impaired muscle-specific gene expression [Latella et al., 2001] and repressed MEF2 activity, without affecting all transcriptional activation [Lazaro et al., 2002], while ectopic Cyclin D1 inhibited MyoD-induced activation of muscle gene transcription [Rao et al., 1994]. Zhang and colleagues showed that CDK4 can bind MYOD and block its DNA binding capacity [Zhang et al., 1999a], while MYOD can bind CDK4 and inhibit its ability to phosphorylate pRb [Zhang et al., 1999c]. These seemingly contradictory observations might depend on differences in MYOD/CDK4 levels, on subcellular trafficking or on the involvement of further partners in the MYOD/CDK4 complexes.

Fewer studies focused on the other CDK/Cyclin complexes. The mitotic complex of CDC2/Cyclin B was found to phosphorylate MYOD, leading to repression of MYOD-dependent transactivation as well as to diminished MYOD amount during mitosis [Tintignac et al., 2004]. Muscle-specific expression of different CDKs-Cyclins in *C.elegans* activated the cell cycle transcriptional program, without loss of the differentiated state. Under those experimental conditions, cell cycle reentry and S-phase induction was not driven by CDK2/Cyclin E but rather CDK4/Cyclin D [Kotzelius et al., 2011]. In line with this observation, ectopic CDK2/Cyclin E did not reinforce cell cycle reentry in differentiated myotubes, as opposed to the other G1 CDK/Cyclin complex [Latella et al., 2001]. CDK2-associated kinase activity was observed in lysates from proliferating C2C12 myoblasts, but not differentiating ones, while inactive CDK2/Cyclin E seemed to persist in the latter [Mal et al., 2000]. CDK2 immunoprecipitated with Cyclin A in proliferating myoblasts [Chu & Lim, 2000], in agreement with this complex's role in S-phase (**Fig. 2.1**). However, upon differentiation there is evidence that CDK2 is sequestered from Cyclin A into inactive Cyclin D3-p27-CDK2 complexes [Chu & Lim, 2000]. As previously mentioned, Cyclin D3 was the only G1 cyclin surprisingly found to be upregulated upon myogenic differentiation [Rao et al., 1994; Skapek et al., 1995; Wang & Walsh, 1996b]. In fact, MYOD was found to activate its expression, assisted by the co-activator p300 [Cenciarelli et al., 1999]. Cyclin D3 was suggested to participate in the induction but not maintenance of muscle differentiation. In line with this, Cyclin D3 was highly expressed at late fetal stages, but disappeared in terminally differentiated skeletal muscle [Bartkova et al., 1998]. Furthermore, Cyclin D3 primes myoblasts for differentiation when overexpressed [Gurung & Parnaik, 2012]. *In vitro* and *in vivo* studies of Cyclin D3 knock-out muscles showed defects under homeostatic conditions as well as reduced satellite cell renewal following activation [de Luca et al., 2014].

Regarding the CDK families that are implicated in transcription rather than cell cycle progression, scarce data are available. In contrast to cell cycle-related CDK/Cyclin complexes, CDK9/Cyclin T2a seemed to enhance MYOD function and promote differentiation [reviewed in de Falco & de Luca, 2009], consistent with a role distant from supporting proliferation.

2.3 The Pocket Protein- E2F network: downstream effectors of Cdk/Cyclins

General features in cell cycle regulation

The pocket protein (PP) family is comprised by pRB and the pRB-related p107 and p130. They are known to inhibit G1-S transition by controlling E2F-induced genes. The classical model includes a) PP-mediated inhibition of activator E2F or b) formation of repressive E2F/PP complexes, both leading to cell cycle suppression. CDK-induced phosphorylation of PPs promotes its dissociation from E2F, allowing entry into S phase.

Among PPs, pRb has a leading role as implicated by single and compound knock-outs [reviewed in Lin et al., 1996]. Tissue-specific pRb loss contributes to various proliferation and differentiation defects [Cobrinik, 2005]. pRb-lacking mice die at mid-gestation (E13.5-E16.5), while *Mox2^{Cre}*-mediated ubiquitous pRb loss from E5.5 causes perinatal death [Lee et al., 1996; Berman et al., 2009]. In both cases mice display a series of phenotypes (e.g. apoptosis in liver and central nervous system) which are exacerbated by additional deletion of p107 [Lee et al., 1996; Berman et al., 2009]. Compound deficiency of pRb/p107 or pRb/p107/p130 results in earlier embryonic lethality (E11.5 and E9-E11, respectively) and in even more elevated proliferation and apoptosis compared to pRb abrogation [Lee et al., 1996; Berman et al., 2009; Wirt et al., 2010]. In contrast, p107 and p130 single knockouts are viable, healthy and fertile. Yet combined ablation of both factors results in perinatal lethality associated with breathing abnormalities and maternal rejection [Cobrinik et al., 1996; Lee et al., 1996]. One study reported strain-dependent phenotypes upon p130 ablation; p130 loss resulted in multiple defects (abnormal growth, increased apoptosis/proliferation and impaired myogenesis/neurogenesis) culminating to embryonic lethality, but this phenotype was completely suppressed in C57BL/6J background, in which the mice were viable, fertile and without detectable phenotypes [LeCouter et al., 1998].

Given the eminent role of pRb, data regarding this PP will mainly be presented hereafter. pRb is synthesized throughout the cell cycle, but its phase-specific Cdk-driven phosphorylation renders it

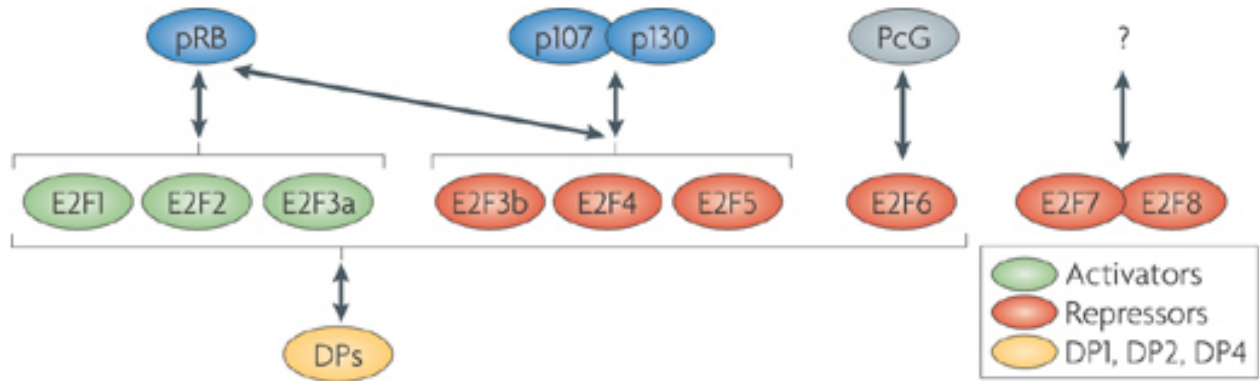


Figure 2.3. Interactions of Pocket Protein and E2F families.

The Pocket Protein (PP) family includes retinoblastoma protein [pRb; acting on several E2F factors (E2F1-5)], and p107 and p130 (limited to E2F3b-5). PP binding to activatory E2Fs (green) inhibits their positive effect on targets. On the other hand, PP or other protein binding to repressive E2Fs (orange) consists complexes that suppress target expression. Some E2F members need to associate with Differentiation-regulated transcription factor-1 Polypeptide (DP; yellow) to exert their action.

Source: van den Heuvel & Dyson, 2008

inactive at S/G2/M [Buchkovich et al., 1989; Zarkowska & Mittnacht, 1997]. Resting cells express the hypo-phosphorylated form of pRb, but it is increasingly phosphorylated during progression through G1 and is maintained hyperphosphorylated until late mitosis, while dephosphorylation precedes total growth arrest during differentiation [Buchkovich et al., 1989; Chen et al., 1989]. pRb is shown to restrict cell cycle progression at a specific, early G1 point [Goodrich et al., 1991]. Loss of pRb function is associated with loss of proliferative control, which is rescued by pRb restoration [Huang 1988; Qin et al., 1992]. Furthermore, reintroduction of pRb in carcinoma cell lines not only suppresses the neoplastic phenotype *in vitro*, but it also halts the tumor forming ability in nude mice [Huang et al., 1988; Bookstein et al., 1990].

The two growth-suppressing pRb domains are also identified as necessary for E2F binding [Qin et al., 1992]. Direct physical association of the two factors is revealed by a series of experiments including overexpression of pRb or mutant pRb as well as transactivation assays with E2F1 mutants incapable of pRb binding [Flemington et al., 1993; Helin et al., 1993]. Although PPs mostly restrain the cell cycle by affecting E2F-regulated transcription, there is also evidence that they directly (p107) or indirectly (pRB) inhibit Cdk activity [Zhu et al., 1995; Alexander and Hinds, 2001].

Eight mammalian E2F transcription factor genes have been identified. *In vivo* abrogation of one or multiple closely related E2F proteins suggested partial redundancy [Gaubatz et al., 2000; Li et al., 2003]. As illustrated in **Fig. 2.3**, the family contains activators (E2F1-E2F3a) and repressors (E2F3b-E2F8), some of which (E2F1-E2F6) form heterodimeric complexes with Differentiation-regulated transcription factor-1 Polypeptide (DP) before binding to DNA in a sequence-specific manner. E2F1-E2F3a are transcriptional activators, preferentially inhibited by pRb. They occupy promoters when the genes are being transcribed and they stimulate cells to pass the G1/S transition [reviewed in van den Heuvel & Dyson, 2008]. On the other hand, E2F3b-E2F8 form repressive complexes with unphosphorylated PPs (E2F1-5) or further factors (E2F6-8) at quiescence or at early G1 and they bind to promoters, while the targets are being repressed [reviewed in van den Heuvel & Dyson, 2008]. It is further known that E2F4 and E2F5 are expressed throughout the cell cycle, but -due to their lack of

nuclear localization signal- they remain cytoplasmic until p107 and p130 bind them and recruit them to the nucleus to repress targets [Allen et al., 1997].

E2F proteins regulate genes important for multiple processes during the cell cycle. Since *myc*, the first E2F cellular target to be described [Hiebert et al., 1989; Thalmeier et al., 1989], a multitude of cell cycle-associated factors has been demonstrated to be regulated by E2F, including cyclins, CDKs, CDKIs, Ki67, PCNA, and E2F themselves [summarized in Bracken et al., 2004]. Furthermore, E2F as well as PPs control cell fate decisions, impact stem cell maintenance and differentiation, and participate in procedures such as DNA damage response/apoptosis/development, often without simultaneous effect on cell cycle dynamics [Cobrinik, 2005; Julian & Blais, 2015].

Implications in skeletal muscle cell cycle regulation

The expression levels and phosphorylation status of pRb changed upon myogenic differentiation, with MYOD participating in both processes. More precisely, MYOD was found to enhance pRb expression during myogenic differentiation [Martelli et al., 1994] and to bind CDK4 to inhibit its ability to phosphorylate pRb [Zhang et al., 1999c]. Although pRb was present in both growing myoblasts and differentiating myocytes, the hypo-phosphorylated form prevailed in the latter populations [Kieck et al., 1995; Wang & Walsh, 1996b; Carnca et al., 2000].

The requirement for pRb in myogenesis is well-established. Unphosphorylated pRb was shown to be bound by MYOD, with the interaction requiring the bHLH domain of MYOD and the C-terminal half of pRb [Gu et al., 1993a]. In differentiating myoblasts pRb was found to cooperate with MYOD to promote MEF2 transcriptional activity [Novitch et al., 1999]. pRb overexpression blocked S-phase entry and induced myogenesis, as evidenced by MYOGENIN expression [Carnca et al., 2000]. Abrogation of pRb was studied *in vitro* in C2C12 muscle cell line, primary myoblasts, and MYOD-mediated myogenically converted fibroblasts. pRb-null mice die at early embryonic stages, before myogenesis is completed, by severe anemia attributed to placental abnormalities. Thus, *in vivo* pRb ablation was studied by (a) generation of mice carrying a hypomorphic pRb minigene expressed in

placenta and nervous system but not in skeletal muscle [Zacksenhaus et al., 1996; Ciavarra et al., 2011], (b) conditional knock-out mice, in which pRb excision was induced by the Mox-Cre (acting from E5.5 onwards) [de Bruin et al., 2003] or the muscle-specific Myf5-Cre and MCK-Cre [Huh et al., 2004]. A series of abnormalities was observed in all *in vitro* and *in vivo* systems. Firstly, no post-mitotic stage was reached and growth factor re-stimulation of Rb^{-/-} differentiated myotubes resulted in S-phase re-entry [Schneider et al., 1994; Novitch et al., 1996; Camarda et al., 2004; Huh et al., 2004; Ciavarra et al., 2011]. pRb deletion in differentiating C2C12 was sufficient to cause cell cycle reentry without growth factor re-stimulation [Pajcini et al., 2010]; however, in primary myoblast cultures, BrdU was incorporated in myotubes only when pRb and Arf (an alternative product encoded by the *Ink4a* locus) were concomitantly suppressed [Camarda et al., 2004; Pajcini et al., 2010]. The non-maintained cell cycle arrest was evidenced *in vivo* by accumulation of elongated nuclei that actively synthesized DNA within the myotubes of hypomorphic pRb mice [Zacksenhaus et al., 1996]. Reversibility of the terminally differentiated state is a unique phenotype, which was never noted upon the modulation of any other cell cycle factor. Secondly, pRb removal was associated with enhanced proliferative potential. It resulted in molecular events associated with transition into S phase, including reactivation of E2F, upregulation of Cyclins A and E, re-induction of PCNA, MCM2 and replication protein A, accumulation of Cyclin A, Cyclin B1, CDK2 and CDC2 in the nuclei [Peschiaroli et al., 2002; Camarda et al., 2004]. Thirdly, there was increased susceptibility to apoptosis [Zacksenhaus et al., 1996; Wang et al., 1997; Peschiaroli et al., 2002; Ciavarra et al., 2011]. Finally, the myogenic differentiation was severely affected *in vitro* and *in vivo* [Novitch et al., 1996; Zacksenhaus et al., 1996; de Bruin et al., 2003; Huh et al., 2004; Ciavarra et al., 2011]; even though differentiation could be initiated (e.g. *Myogenin* expression), there was a failure to progress and complete the myogenic program (e.g. reduction in late muscle-specific genes, including MyHC, MCK, MRF4, Troponin T). Mutant mice exhibited shorter myotubes with fewer myofibrils and enlarged nuclei (pRb minigene), hypoplastic and dysplastic myofibers within the intercostal muscles, diaphragm, limbs (Mox2-Cre;Rb^{f/-}) or dramatic reduction in muscle mass accompanied by complete absence of mature fibers (Myf5-Cre;Rb^{f/f}). Some of these phenotypes were even associated to the respiratory problems and early perinatal death of the mutants. Interestingly, pRb did not seem to be required for the maintenance or regeneration of differentiated skeletal muscle as implied by two independent systems. Firstly, MCK-

Cre;Rb^{f/f} mice are viable, healthy, with normal muscle tissue and without deficits in regenerative capability of damaged muscles [Huh et al., 2004]. Secondly, when pRb deletion was performed in myotubes rather than before differentiation initiation, the post-mitotic status was not challenged by growth factor re-stimulation [Camarda et al., 2004].

Other pocket proteins have also been shown to play some functions in muscle differentiation. p107 is high in proliferating myoblasts but sharply drops upon differentiation [Kiess et al., 1995; Carnca et al., 2000]. In the absence of pRb this downregulation did not occur, but neither did it inverse the defective phenotype, implying lack of compensation [Schneider et al., 1994]. p130 exhibited the opposite pattern of p107 [Kiess et al., 1995; Carnca et al., 2000]. Unlike pRb, the differentiation-induced upregulation of p130 mainly occurred in reserve cells rather than multinucleated myotubes [Carnca et al., 2000]. Indeed, when overexpressed it blocked S-phase entry but repressed myogenesis, through inhibition of MYOD and its transactivation ability [Carnca et al., 2000]. Initial studies of *in vivo* p130 loss did not report any muscle phenotype [Cobrinik et al., 1996], while later reduction in myotomal myocytes was demonstrated [LeCouter et al., 1998]; however, the latter effect was strongly dependent on the genetic background and was completely suppressed in certain strains.

The complexes of pocket proteins with E2F transcription factors were analyzed in proliferation and differentiation conditions *in vitro*. p107 was the most prominent component in the complexes appearing in undifferentiated cells, while it was reported to be replaced by p130 in E2F complexes upon differentiation [Corbeil et al., 1995; Kiess et al., 1995; Shin et al., 1995]. E2F4 appeared to be sequestered in a complex with pRb2/p130 and to accumulate in the nucleus when growth arrest occurred [Puri et al., 1997; Puri et al., 1998]. Undifferentiated C2C12 lacked pRb-E2F complexes [Corbeil et al., 1995], consistent with the presence of hyper-phosphorylated pRb in this proliferating group [Kiess et al., 1995; Wang & Walsh, 1996b; Carnca et al., 2000]. Similarly, pRb/E2F repressor complexes did not form in myotonic dystrophy patients, whose muscle differentiation is affected, but they are abundant in differentiated cells of control subjects [Timchenko et al., 2001]. However, only few such complexes appeared in fully differentiated C2C12 myotubes [Corbeil et al., 1995], in

agreement with the proposed unimportant role of pRb in maintenance of that population [Camarda et al., 2004; Huh et al., 2004]. Permanent silencing of E2F targets during muscle differentiation was driven by Suv39h1-dependent H3K9 tri-methylation and positioning close to heterochromatin nuclear compartment [Guasconi et al., 2010], but this tri-methylation and silencing was not directly specified by pRb binding [Vandromme et al., 2008], as is the case for senescent cells.

E2F levels do not show a uniform behavior upon differentiation. E2F1 and E2F3a were strongly down-regulated, dropping to undetectable levels in mature myotubes, while E2F2, E2F3b and E2F4 remained unchanged [Wang et al., 1995; Asp et al., 2009]. Furthermore, the subcellular distribution of these factors was subject to changes. When myogenic cells were induced to differentiate, E2F1 and E2F5 became exclusively cytoplasmic, E2F3 remained cytoplasmic, E2F2 was primarily nuclear and E2F4 appeared in both compartments [Gill & Hamel, 2000]. In contrast, even weak mitogenic signals could cause nuclear import of E2F1 and E2F4 [Gill & Hamel, 2000]. Ablation of each of these factors established E2F3b as a critical regulator of myogenic differentiation through the transcriptional control of developmental and differentiation genes in myotubes [Asp et al., 2009]. E2F4 ablation caused a milder differentiation impairment [Asp et al., 2009]. In contrast, E2F1 overexpression inhibited cell cycle exit and MYOD- or MYOGENIN-mediated transcription activation [Wang et al., 1995; Wang et al., 1996] or caused S-phase entry in myocytes [Chen & Lee, 1999]. Furthermore, ectopic nuclear E2F1 and E2F4 promoted cell cycle reentry and opposed the differentiation program [Gill & Hamel, 2000]. Whether the same effects occur *in vivo* remains to be confirmed. Upon *in vivo* E2F1 ablation and muscle injury, the regeneration procedure was severely compromised, as shown by rough histological analysis [Yan et al., 2003]. However, the reasons causing this defect as well as the potential role of E2F factors during embryonic and postnatal myogenesis are still uncharacterized.

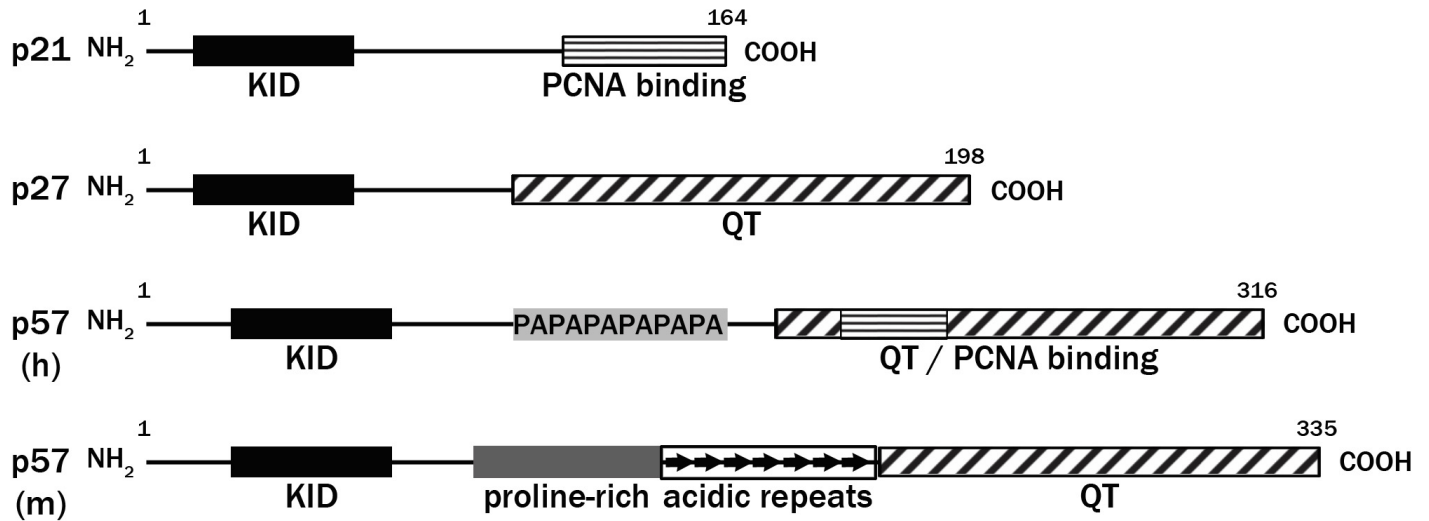


Figure 2.4. Domain structure of Cip/Kip family of CDKIs.

p21, p27 and p57 share conserved regions (indicated as boxes), including an N-terminal CDK inhibitory domain (KID-black box; all members), a C-terminal PCNA binding domain (horizontally striped box; p21 and p57) and a C-terminal QT box (diagonally striped box; p27 and p57). Human and mouse p57 further possess a unique central domain, implicated in functions not shared with p21 and p27. This region consists of PAPA repeats (human p57 - h) or a proline-rich and acidic domain (mouse p57 - m). Numbers indicated first and last amino acid of each protein [Scheme based on: Matsuoka et al., 1995; Galea et al., 2008; Pateras et al., 2009].

2.4 CDKIs: major negative regulators of Cdk-Cyclin activity

General features in cell cycle regulation

The control of formation and activity of CDK-Cyclin complexes largely depends on the CDKIs, which belong to two broad categories on the basis of their structural and functional profile. The INK4 family includes p15, p16, p18, and p19, which contain conserved ankyrin repeats [Serrano et al., 1993; Hannon & Beach, 1994; Hirai et al., 1995], a motif participating in protein recognition and interaction [Michaely & Bennett, 1992]. The Cip/Kip family involves p21, p27, and p57, which show partial structural homology (**Fig. 2.4**). They share a 60-residue kinase inhibitory domain in their N-terminal region and a nuclear localization signal (NLS) within their C-terminal region [Polyak et al., 1994; Lee et al., 1995; Matsuoka et al., 1995]. In addition, p21 and p57 contain a C-terminal PCNA-binding domain, allowing interaction with PCNA and prevention of DNA replication [Li et al., 1994; Waga et al., 1994; Watanabe et al., 1998]. Moreover, p27 and p57 include a C-terminal QT box with a consensus CDK phosphorylation site [Polyak et al., 1994; Lee et al., 1995; Matsuoka et al., 1995], involved in their SCF/Skp2-mediated degradation [Lu & Hanter, 2010]. Finally, p57 possesses a unique central region (**Fig. 2.4**), consisting of PAPA repeats (human p57) or proline-rich and acidic-rich domains (mouse p57) [Lee et al., 1995; Matsuoka et al., 1995].

These structural differences underlie the divergent specificity of action between the two CDKI families. This functional variation is two-fold. Firstly, INK4 members are selective inhibitors of CDK4 and CDK6, leaving other CDKs unaffected [Serrano et al., 1993; Hannon & Beach, 1994; Guan et al., 1994; Chan et al., 1995; Hirai et al., 1995]. In contrast, Cip/Kip proteins bind and inhibit all CDK-Cyclin complexes that are formed throughout the cell cycle [Gu et al., 1993b; Harper et al., 1993; Xiong et al., 1993; Polyak et al., 1994a, b; Lee et al., 1995; Matsuoka et al., 1995], in line with the finding that they show intrinsic flexibility and they are largely disordered prior to binding to other proteins [Galea et al., 2008]. In fact, they belong to a group of intrinsically unstructured proteins, meaning that they are devoid of secondary and/or tertiary structure under physiological conditions. Structural analysis of free and CDK/Cyclin-bound p21 and p27 demonstrates a disorder-to-order transition following

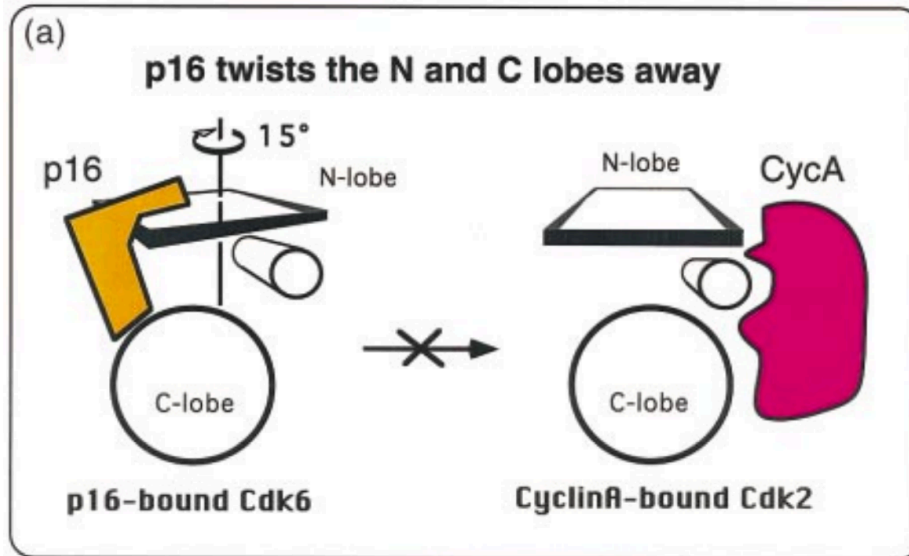


Figure 2.5. Differential CDKI and Cyclin binding to Cdk.

For CDK binding, p16 (left) and Cyclin A (right) interact with both Cdk lobes but depending on different relative orientation. p16-mediated allosteric changes inhibit ATP and cyclin binding, interfering with CDK kinase activity.

Source: Pavletich, 1999

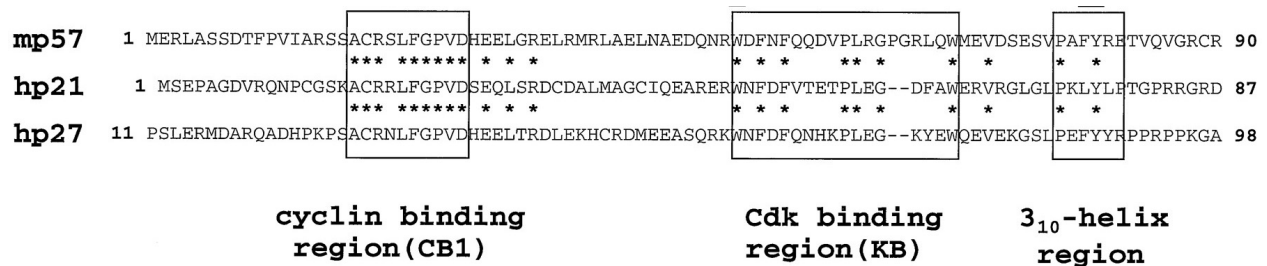


Figure 2.6. Regions of Cip/Kip Cdk-inhibitory domain.

Aligned N-terminal regions of p21, p27 and p57 proteins with highlighted subregions of the CDK-inhibitory domain, includes regions for CDK and cyclin binding and a 3₁₀-helix. (modified from Hashimoto et al., 1998)

Source: Hashimoto et al., 1998

binding [Kriwacki et al., 1996; Russo et al., 1996; Lacy et al., 2004] that is achieved in a stepwise manner [Lacy et al., 2004]. Even though most structural data come from p27 studies, acquisition of secondary structure is observed in several subdomains with homology among the three family members. Secondly, INK4 inhibitors can bind to untethered or cyclin-associated CDK subunits, causing a strong inhibition of CDK catalytic activity and inducing G1 arrest [Serrano et al., 1993; Hannon & Beach, 1994; Guan et al., 1994; Chan et al., 1995; Hirai et al., 1995]. They promote conformational changes that interfere with both ATP and cyclin binding, by changing the relative orientation of the two CDK lobes (**Fig. 2.5**) [Brotherton et al., 1998; Russo et al., 1998]. In contrast, p21, p27, and p57 bind CDK-Cyclin heterodimers, with the affinity for CDKs increasing by Cyclin association [Gu et al., 1993b; Harper et al., 1993; Xiong et al., 1993; Polyak et al., 1994a, b; Harper et al., 1995; Matsuoka et al., 1995]. In fact, their N-terminal CDK inhibitory domain involves a cyclin binding site, a CDK binding site and a 3_{10} helix (**Fig. 2.6**); however, the last is indispensable only in p57-dependent inhibition [Hashimoto et al., 1998]. Moreover, structural examination of p27-bound complexes reveals that p27 intrudes the CDK catalytic cleft and interferes with ATP binding [Russo et al., 1996]. Considering their ability to bridge CDKs and cyclins, members of the Cip/Kip group would be expected to promote CDK-cyclin assembly. Indeed, at low concentrations p21 stabilized this interaction, while addition of extra p21 molecules to the complex leads to inhibition [Michieli et al., 1994; Zhang et al., 1994; Harper et al., 1995; LaBaer et al., 1997]. p57 is found to stabilize early G1 complexes only in the absence of p21 and p27 [Cerqueira et al., 2014], while CDK4-Cyclin D1 inhibition premises p57 homodimerization [Reynaud et al., 2000a]. Furthermore, p27 binding to CDK-Cyclin complexes was compatible with kinase activity [Soos et al., 1997]. Thus, Cip/Kip might function as titratable buffers and repress CDKs in a concentration-dependent manner.

CDKI expression is mostly linked to cell cycle withdrawal that accompanies differentiation, but additional functions have been described. An expression pattern that correlates with terminal differentiation of multiple lineages during organogenesis (e.g. skeletal muscle, kidney, brain, gonads, cartilage, cranio-facial structures) has been described for all INK4 and Cip/Kip inhibitors except p15 and p16 [Parker et al., 1995; Zindy et al., 1997; Westbury et al., 2001; Susaki et al., 2009]. Furthermore, all CDKIs have been related to cellular senescence [Erickson et al., 1998; Zhu et al.,

1998; Tsugu et al., 2000; Alexander & Hinds, 2001; Gargica et al., 2012; Giovannini et al., 2012; García-Fernández et al., 2014], with the major CDKI-involving pathways being the p16/pRb, the p19^{ARF}/p53/p21, and the PTEN/p27 pathways [reviewed by Bringold & Serrano, 2000]. Although it is traditionally associated with aging, senescence has also been detected during embryonic development, with p21 and p15 playing a leading role [Muñoz-Espín et al., 2013; Storer et al., 2013]. Moreover, CDKIs, mainly from the Cip/Kip family, have been reported to modulate apoptosis [reviewed in Besson et al., 2008] and transcription [Pippa et al., 2012; Orlando et al., 2015].

Implications in skeletal muscle cell cycle regulation

The expression of CDKIs is, generally, upregulated during myogenesis, as evidenced by developing mouse embryos and myogenic cells. When the equilibrium is shifted to or against one of these factors, the differentiation and proliferation capacities are rapidly affected *in vitro*. More recently, *in vivo* data started to support the unique and redundant roles of CDKIs for muscle development, homeostasis and regeneration. Further studies are expected to elucidate the currently poorly understood role of CDKIs in the entry (early postnatally) or re-entry (post-regeneration) of satellite cells into quiescence.

A limited number of studies have evaluated the impact of the INK4 family on the muscle lineage. Terminally differentiated adult muscles or C2C12-derived myotubes lacked p15 [Franklin & Xiong, 1996]. However, p15 was found elevated in satellite cells from aged animals along with other CDKIs and this could contribute to the defective regeneration of old muscle [Li et al., 2015]. Same as p15, p16 was also absent from proliferating C2C12 cells, myotubes or adult muscle samples [Franklin & Xiong, 1996; Reynaud et al., 1999]. However, ectopic p16 was associated with increased muscle-specific gene expression [Skapek et al., 1995] as well as with protecting differentiating C2C12 from apoptosis [Wang & Walsh, 1996a], the latter only in the presence of pRb [Wang et al., 1997]. p16 transcripts and protein levels increased with aging, with the promoter exhibiting a more open chromatin pattern following injury-induced satellite cell activation [Li et al., 2015]. In addition, p16 derepression ignited a switch of resting geriatric satellite cells from reversible quiescence to

irreversible pre-senescence state, compromising muscle regenerative capacity [Sousa-Victor et al., 2014]. p18 showed a remarkable increase upon C2C12 differentiation, that was regulated transcriptionally (promoter switch to produce a different transcript) and translationally [Franklin & Xiong, 1996; Phelps et al., 1998]. Furthermore, there was a continuing increase of p18-associated CDK4 and CDK6 [Franklin & Xiong, 1996]. In adult muscle tissue, p18 was very abundant and it was associated with all CDK6 and half CDK4 (shared with p27) [Franklin & Xiong, 1996]. *In vivo* regeneration studies showed an early transient increase in p18 levels, although this effect was not further followed [Yan et al., 2003]. p19 was the only CDKI found at high levels in proliferating myoblasts and declining following differentiation [Franklin & Xiong, 1996]. The reciprocal p19 decrease and p18 increase raised the possibility of p19 replacement by p18 as CDKI during differentiation.

Members of the Cip/Kip family have been more extensively explored. Historically, most studies focused on p21, the first family member to be identified. When C2C12 myoblasts were maintained in proliferation conditions, p21 was generally low [Mal et al., 2000]. p21 reached detectable levels close to G1, but at mitosis entry it rapidly decreased [Tintignac et al., 2004]. After switching to differentiation conditions, p21 sharply increased [Halevy et al., 1995; Franklin & Xiong, 1996; Reynaud et al., 1999; Mal et al., 2000]. Consistent with a role in differentiation-associated growth arrest, myotubal p21 (a) immunoprecipitated with CDK2 and CDK4, (b) showed increased interaction with CDK2, CDK4, and CDK6 compared to myoblasts, and (c) exceeded the quantity of active CDK4/Cyclin [Franklin & Xiong, 1996; Wang & Walsh, 1996b; Wang et al., 1997; Figliola & Maione, 2004]. In fact, p21 overexpression in C2C12 was sufficient for cell cycle withdrawal under growth conditions [Guo et al., 1995], while p21-deficient primary myoblasts showed increased proliferation [Hawke et al., 2001]. Forced p21 promoted muscle-specific gene expression, rescued Cyclin D1-mediated *MyoD* repression and exerted protection against apoptosis [Skapek et al., 1995; Wang & Walsh, 1996a]. Conversely, primary myoblast culture of p21^{-/-} animals showed problematic differentiation and increased apoptosis [Hawke et al., 2001]. MYOD transcriptionally activated *p21* [Halevy et al., 1995; Tintignac et al., 2004] and p21 was able to inhibit MYOD phosphorylation, stabilizing it [Reynaud et al., 1999]. Non-phosphorylatable MYOD sustained *p21* expression, interfering

with M-phase entry [Tintignac et al., 2004], while in the absence of MYOD, p21 expression was delayed during muscle development [Parker et al., 1995]. MYOD-mediated p21 induction and p21 expression in the developing muscle are p53-independent [Halevy et al., 1995; Parker et al., 1995], although p21 is transcriptionally regulated by p53 [El-Deiry et al., 1993]. Overexpression and silencing approaches established Cyclin D3 as a further factor that enhanced p21 expression [Gurung et al., 2012; de Luca et al., 2014], in agreement with the notion that Cyclin D3 is the only G1 cyclin to promote myogenic differentiation rather than proliferation. A direct MYOGENIN-mediated induction of p21 would be anticipated because (a) MYOGENIN binding sites were detected in the *p21* promoter [Singh & Dilworth, 2013] and (b) on a protein level, p21 expression follows MYOGENIN upon differentiation [Andres & Walsh, 1996]. However, when myoblasts are induced to differentiate, *p21* transcripts appear before *Myogenin*, while MYOGENIN knock-out embryos provided unequivocal evidence that MYOGENIN is not required for p21 induction [Parker et al., 1995].

The *in vivo* profiling of p21 expression in muscle revealed age-related differences. During embryonic development and the establishment of the muscle lineage, *p21* mRNA first appears in the forming myotome of E8.5 somites [Parker et al., 1995]. From E10 it followed the *Myogenin* pattern [Parker et al., 1995]. In the adult, no protein was found in skeletal muscles [Franklin & Xiong, 1996; Mademtoglou et al., submitted-2], while the mRNA expression is controversial. Very high [Parker et al., 1995; Park & Chung, 2001] or undetectable [Macleod et al., 1995] transcripts were reported in adult skeletal muscles; differences in analyzed muscles, mouse background, age window, or probes could partially explain these differences. *p21* mRNA in skeletal muscle increased up to 5 months, followed by a continuing decline until 23 months [Park & Chung, 2001]. However, the protein seemed to be absent from adult skeletal muscle. Satellite cell-focused analysis revealed p21 rise with aging along with increased chromatin accessibility post-injury [Li et al., 2015]. Notably, p21 loss did not affect the development of muscle or the establishment of the satellite cell compartment [Deng et al., 1995; Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015; Mademtoglou et al., submitted-2], while AAV-mediated transient *p21* suppression did not impact the genetic status or differentiation capacity of myoblasts [Biferi et al., 2015]. During injury-induced activation of satellite cells and muscle regeneration, p21 seemed to exert its function during the early period of the

regeneration procedure [Yan et al., 2003; Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015; Mademtzoglou et al., submitted-2].

Searching for alternative cell cycle regulators in cells undergoing G1 arrest and myogenic differentiation, the groups of Maione, Lanfranchi and Leibovitch independently identified p57 [Reynaud et al., 1999; Figliola & Maione, 2004; Bean et al., 2005]. p57 increased remarkably as C2C12 cells were stimulated to differentiate [Reynaud et al., 1999] or as embryonic muscle progenitors proceeded to myogenic determination and differentiation [Zalc et al., 2014]. During development it was present in most nuclei of limb and abdominal wall muscles [Zhang et al., 1997; Zalc et al., 2014], but p57 knockout mice did not manifest any severe muscle phenotype [Yan et al., 1997; Zhang et al., 1997], implying compensatory pathways (see below). Postnatally, p57 progressively decreased as mice aged [Park & Chung, 2001] and it was barely detectable in satellite cells [Chakkalakal et al., 2014; Mademtzoglou et al., submitted-2]. However, p57 peaked soon after muscle regeneration was initiated and it was maintained at higher than baseline levels throughout regeneration [Yan et al., 2003; our unpublished observation].

p57 and MYOD have been implicated in a positive feedback loop *in vitro* and *in vivo*, whereby MYOD induced p57 which then enhanced MYOD activity and stabilization. Although three putative MYOD -boxes proximal and downstream from the transcription start site were not functional [Bean et al., 2005], MYOD -responsive regions upstream of the transcription start site were identified by us and others [Bean et al., 2005; Zalc et al., 2014]. *In vitro* studies, based mostly on fibroblasts converted to the myogenic lineage, suggested that MYOD plays a dual function in inducing p57. Firstly, it counteracted a cis-acting repression, by sequentially interacting with CTCF, disrupting the CTCF-mediated chromatin loop and releasing the p57 promoter, and secondly, it assisted up-regulation by the intermediate factors p73, SP1 and EGR1 [Vaccarello et al., 2006; Figliola et al., 2008; Busanello et al., 2012; Battistelli et al., 2014]. Our data provide evidence for E-box-dependent MYOD binding and transactivation of a muscle-specific p57 regulatory element [Zalc et al., 2014]. *MyoD* knockdown in zebrafish embryos reduced p57, while MYOD over-expression promoted ectopic p57 in somatic and head mesoderm [Osborn et al., 2011]. Notably, MYOD -induced myogenic conversion of fibroblasts

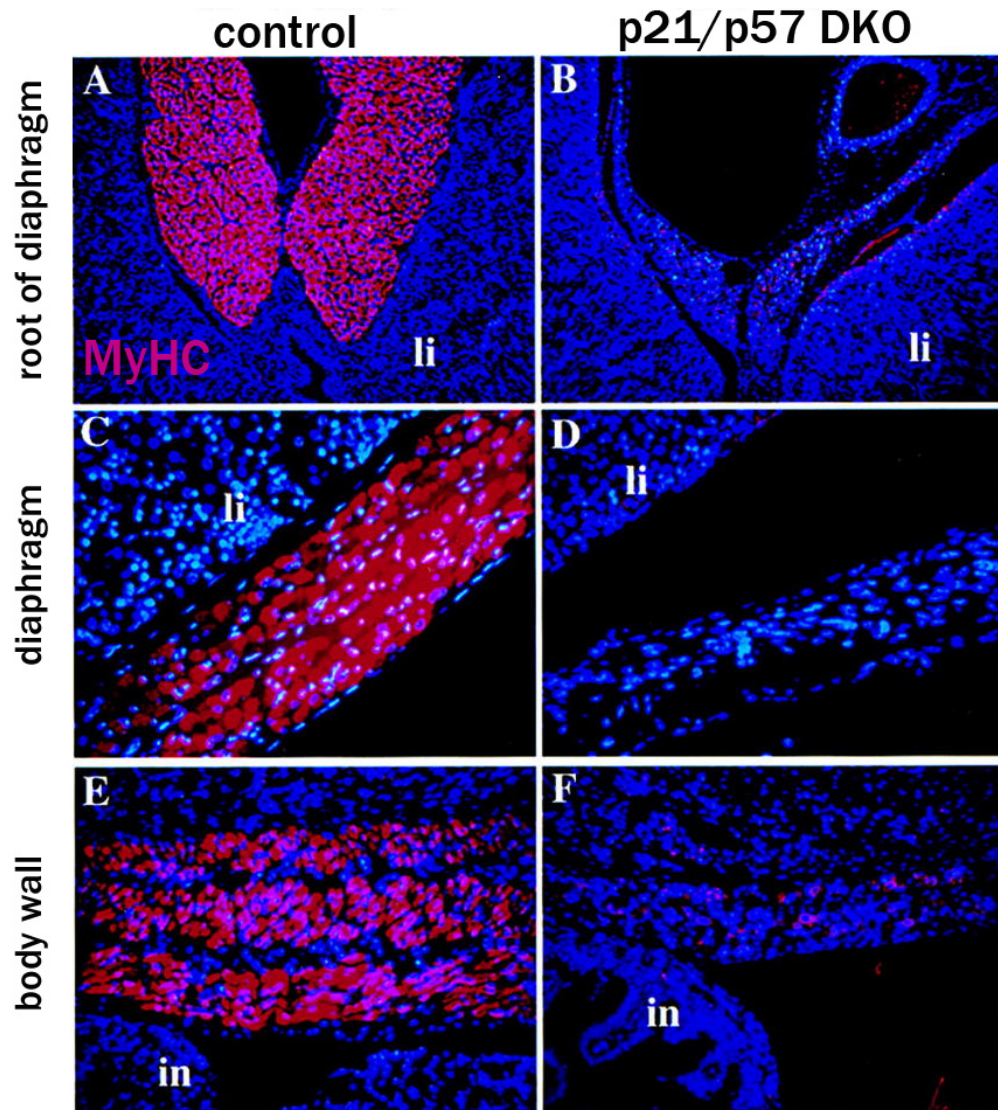


Figure 2.7. p21 and p57 loss severely affects skeletal muscle during embryogenesis.

p21/p57 double knock-out (DKO; right) fetuses had diminished muscle masses compared to controls (left), as evidenced by myosin heavy chain staining (MyHC). Muscle masses from the root of diaphragm (A, B), diaphragm (C, D), and body wall (E, F) are shown. in: intestine; li: liver.

Adapted from: Zhang et al., 1999b

upregulated p57 only in the absence of p21 [Figliola & Maione, 2004; Zhang et al., 1999c], implying intrinsic differences between that lineage and muscle. On the other way round, p57 was found to promote MyoD accumulation in adaxial cells of zebrafish embryos [Osborn et al., 2011]. Co-immunoprecipitation assays showed that p57 directly binds to MYOD, through the basic region of MYOD and the amino-terminal domain of p57 [Reynaud et al., 2000b]. Remarkably, the same p57 helix domain (**Fig. 2.6**) that was indispensable for CDK/Cyclin inhibition was also important for MYOD binding [Hashimoto et al., 1998; Reynaud et al., 2000b]. *In vitro* dissection of the underlying mechanism revealed that p57 increased MYOD stability by inhibiting its phosphorylation by CDK2/CyclinE complexes [Reynaud et al., 1999], while p57- MYOD binding was competed by CDK4/Cyclin D1 complexes [Reynaud et al., 2000b].

Based on the similarities of the profiles of p21 and p57 during myogenesis as well as the lack of muscle phenotypes in p21 or p57 knock-out embryos, their possible redundancy during development was evaluated by concomitantly ablating both factors [Zhang et al., 1999b]. Simultaneous abrogation of p21 and p57 boosted CDK2 activity toward pRb, increased myoblast proliferation and apoptosis, and induced endoreduplication of residual myotubes. Head, trunk and limb muscles as well as diaphragm of double knock-out embryos were severely diminished (**Fig. 2.7**). Fewer and smaller myotubes were present, indicative of fusion defects. Somitogenesis and primary myogenesis appeared normal, suggesting a secondary myogenesis defect. p21/p57 mutant muscles phenotypically resemble the ones from embryos lacking *Myogenin*. p21/p57 levels in MYOGENIN mutants were comparable to controls, while *Myogenin* expression was not impaired in p21/p57 double knock-outs. Furthermore, MYOGENIN failed to induce p21 or p57 when applied to fibroblasts for myogenic conversion. Thus, the two CDKIs are placed in parallel to MYOGENIN rather than upstream or downstream of this MRF. Whether the redundant functions of p21 and p57 in myogenic cell growth arrest are at play during adult myogenesis remains to be shown and is part of my PhD studies.

Studies on the last member of the Cip/Kip family in muscle development and maintenance revealed p27 participation both in the differentiation program and in maintaining a certain satellite cell

subpopulation. During *in vitro* myogenic differentiation, p27 protein levels increased, even though the mRNA amount remained constant, implying posttranslational regulation [Halevy et al., 1995; Franklin & Xiong, 1996; Reynaud et al., 1999; Chu et al., 2000]. Ectopic p27 inhibited proliferation and promoted differentiation, while p27 repression had the opposite effect [Chakravarthy et al., 2000; Messina et al., 2005]. An in-depth *in vivo* analysis of satellite cell subpopulations revealed that p27 was required to maintain the primitive fate of a subpopulation with label-retaining dormant behavior [Chakkalakal et al., 2014]. The authors also showed that p21 was acting in the non-label-retaining satellite cells to promote lineage commitment [Chakkalakal et al., 2014]. These *in vivo* findings are in agreement with early *in vitro* observations, whereby p21 was enriched in differentiated myotubes and p27 in renewing reserve cells [Cao et al., 2003]. Further supporting this model, p27 was only transiently expressed in the developing muscle masses between E10.5 and E11.5 and all p27+ cells co-expressed MYOD, while only a subset had MYOGENIN [Zabludoff et al., 1998].

2.5 p57 – “KI P”layer in cell physiology and pathology

p57 was identified in 1995 as the last member of the Cip/Kip family of CDKIs [Lee et al., 1995; Matsuoka et al., 1995]. Structural (e.g. N-terminally-located Cdk inhibitory domain, PCNA binding domain, C-terminal QT box) and functional (inhibition of a broad spectrum of CDK-Cyclin complexes) criteria place it in the Cip/Kip family, close to p21 and p27. However, it is distinguished from its siblings in terms of structure (non-shared central domain; **Fig. 2.4**), distribution pattern throughout development and adult life, and multifunction. Furthermore, it is the only CDKI that is subjected to imprinting, with preferential expression of the maternally transmitted allele [Hatada & Mukai, 1995; Matsuoka et al., 1996; Li et al., 2012].

Apart from its well-established role in cell cycle by G1 arrest (see section 2.4), it is emerging as a multifaceted protein participating in several cellular processes [reviewed in Pateras et al., 2009; Rossi & Antonangeli, 2015]. p57 function is critical during embryogenesis, evidenced by the gross developmental defects and perinatal lethality of mice lacking or having excess p57 [Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000; Andrews et al., 2007; Susaki et al., 2009]. In contrast to p21

Table 2.1. Defects of p57 knock-out mice

TISSUE	DEFECTS	REFERENCES*
survival	perinatal lethality	1-4
palate	apoptosis on the surface of palatal shelves (E14.5) cleft palate	1-4
skeleton	sternum closure defect limb bone shortening and thickening delayed differentiation/ossification	1-4
gastro-intestinal tract	absent, shortened and/or inflated stomach/intestine increased apoptosis in smooth muscle of intestinal wall omphalocele	1-4
abdominal wall	thinner body wall muscle dysplasia	2-4
umbilical region	umbilical hernia	1, 4
kidney	renal medullary dysplasia	2, 3
eye lens	vacuolization cataracts	2, 3
adrenal gland	hyperplasia cytomegaly	2, 3

*1, Yan et al., 1997; 2, Zhang et al., 1997; 3, Susaki et al., 2009; 4, Mademtzoglou et al., submitted-1

and p27, p57 is broadly expressed in developing mice, with levels peaking at key differentiating steps of specific organs [Matsuoka et al., 1995; Westbury et al., 2001]. Loss of p57 affects the proliferation, apoptosis, and differentiation status of a multitude of tissues and organs, including skeletal elements, cranio-facial structures, kidney, adrenal gland, reproductive system, gastro-intestinal tract, and sensory organs (**Table 2.1**) [Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000; Susaki et al., 2009]. p27 knock-in rescues many of these abnormalities, supporting the idea that their conserved regions are essential during organogenesis. Nevertheless, many defects, including perinatal lethality, persist, suggesting additional roles for p57 or cell type-dependent differences in p27 and p57 stability [Susaki et al., 2009]. Although in adult life p57 seems less abundant, it has an emerging importance in stem cell homeostasis, promoting acquisition of quiescence [Matsumoto et al., 2011; Zou et al., 2011; Furutachi et al., 2013].

The putative roles in senescence and apoptosis of p57 have been explored. p57 expression increases upon progressive cell passages and it induces senescence when over-expressed in a variety of cell types (e.g. human astrocytoma cells, human hepatocellular carcinoma cells, human prostate epithelial cells, and human uroepithelial cells) [Schwarze et al., 2001; Tsugu et al., 2000; Giovannini et al., 2012; Valcheva et al., 2014]. It seems to have a dual role in programmed cell death, since both pro-apoptotic and anti-apoptotic function have been described. On one hand, p57 is found to sensitize cells to apoptosis [Samuelsson et al., 2002; Gonzalez et al., 2005; Vlachos et al., 2007], by translocating to mitochondria and triggering the intrinsic apoptotic pathway [Vlachos et al., 2007]. Interestingly, the methylation status of its promoter could influence the impact on apoptosis [Kuang et al., 2007b]. On the other hand, increased apoptosis of palate, intestine, and lens upon p57 loss imply anti-apoptotic potential [Yan et al., 1997; Zhang et al., 1997; Zhang et al., 1998]. p57 anti-apoptotic role is mediated by inhibition of the JNK/SAPK pathway, with p57 C-terminal QT box being central to this function [Chang et al., 2003]. However, JNK/SAPK suppression is not observed with p27, despite possessing a QT box as p57 [Chang et al., 2003]. Notably, pro-apoptotic effects are mostly observed in a cancer context, while the anti-apoptotic function is associated with physiological conditions (i.e. development, JNK pathway regulation).

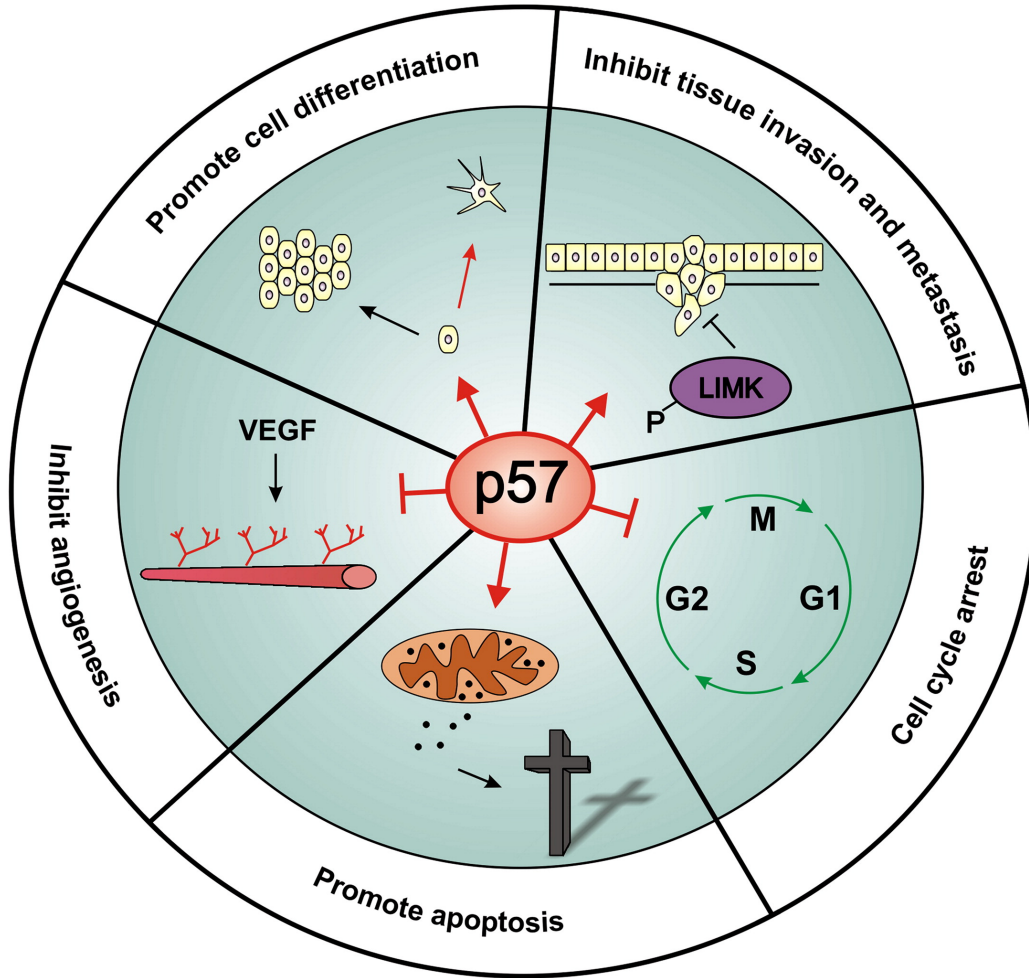


Figure 2.8. p57 implication in cancer hallmarks.

p57 functions that associate its dysregulation with carcinogenesis include growth arrest induction, apoptosis promotion via the mitochondrial pathway, angiogenesis inhibition by negatively regulating Vascular Endothelial Growth Factor (VEGF), cell differentiation advancement, invasion/metastasis repression by interaction with LIM domain Kinase (LIMK) [Kavanagh & Joseph, 2011].

Source: Kavanagh & Joseph, 2011

Cell cycle-independent p57 functions also include participation in actin cytoskeleton dynamics and cell migration. p57 negatively affects cell mobility by interacting with LIM-kinase 1 and regulating cofilin, an actin depolymerization factor [Vlachos & Joseph, 2009; Chow et al., 2011; Guo et al., 2015]. This activity appears independent of the CDK inhibitory ability but connected to the unique central region that p57 does not share with any other CDKI [Yokoo et al., 2003; Vlachos & Joseph, 2009]. The effect on mobility is associated with cytoplasmic p57 presence and is linked to p57 pro-apoptotic effect [Vlachos & Joseph, 2009; Kavanagh et al., 2012]. In contrast, cytosolic p21 and p27 are putative oncogenic factors, involved in tumor invasion and metastasis, and their retention in the cytoplasm is negative prognostic factor in certain tumors [reviewed in Besson et al., 2004; Besson et al., 2008].

p57 implication in carcinogenesis and growth disorders underlines its impact on human pathology [reviewed in Guo et al., 2010; Borriello et al., 2011; Kavanagh & Joseph, 2011; Soejima & Higashimoto, 2013]. Participating in cell cycle arrest, apoptosis, and cell mobility makes p57 a putative tumor suppressor (**Fig. 2.8**) [Kavanagh & Joseph, 2011]. Remarkably, p57 was downregulated in several malignancies and its deficiency or overexpression correlated with proliferative and invasive capacities of cancer cells or tumor formation [Jin et al., 2008; Guo et al., 2011; Xu et al., 2012; Guo et al., 2015]. p57 loss is associated with poor patient prognosis [Xu et al., 2012; Hu et al., 2013; Yang et al., 2015]. Furthermore, p57 is located at a site of frequent loss of heterozygosity in sporadic and familial cancers [Matsuoka et al., 1995]. In fact, p57 loss and gain of function were found to cause two clinically opposite growth disorders, namely BWS (Beckwith-Wiedemann syndrome) and IMAGE (intrauterine growth restriction, metaphyseal dysplasia, adrenal hypoplasia congenita, and genital anomalies), respectively [Soejima & Higashimoto, 2013].

p57 is the most recently discovered CDKI and remains the least studied, partially owing to the lack of genetic tools. Its emerging importance in several cell traits, in stem cell function and in human pathology, prompted us to generate a new mouse model, allowing conditional ablation and reporter-dependent tracking of p57-expressing cells. This mouse is further expected to facilitate *in vivo* studies of p57 impact in postnatal myogenesis, circumventing the early perinatal lethality of germline knock-outs.

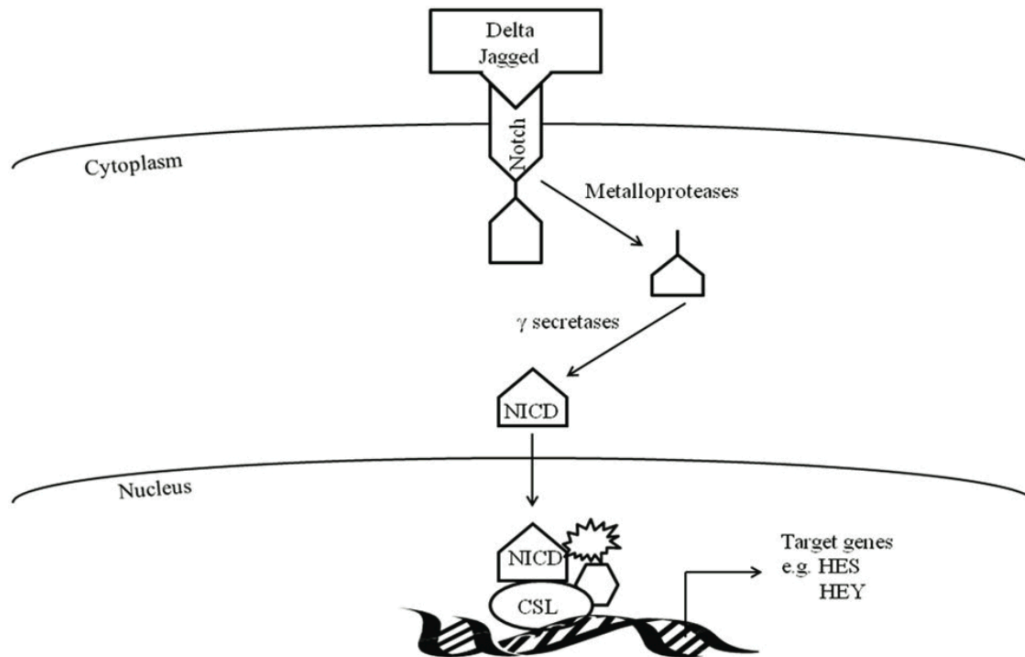


Figure 3.1. Core elements of Notch signaling pathway.

Transmembrane Notch receptors (Notch1-4 in mammals) and ligands (Delta-like, Jagged in mammals) interact to activate the pathway. As a result, a series of metalloprotease/ γ -secretase-driven cleavages releases part of the receptor (Notch IntraCellular Domain, NICD) and allows it to translocate to the nucleus and form activatory complexes with CSL DNA-binding proteins leading to transcription of Notch-responsive genes, such as members of the Hes or Hey families.

Source: Tsivitse, 2010

**Chapter 3. Notch signaling pathway:
pleiotropic role of a master cell fate regulator in myogenesis**

The Notch signaling pathway exerts fundamental functions in most cellular decision-making, including stem cell fate and maintenance in adult tissues, while its perturbation is associated with several genetic disorders and cancer [Guruharsha et al., 2012]. By regulating the balance of proliferation and differentiation, it profoundly affects cell fates in multiple metazoan tissues [Artavanis-Tsakonas & Muskavitch, 2010]. Despite its crucial activity in almost all developing tissues and organs, its effects are highly pleiotropic and the final outcome is context-dependent. The spatial and temporal developmental context and the dosage of Notch activity will dictate the resulting cell fates [Artavanis-Tsakonas & Muskavitch, 2010]. For instance, its activation has an oncogenic effect in T-cells and breast cancer, but tumor suppressive effect in acute myeloblastic leukemia [Guo et al., 2011a; Hernandez Tejada et al., 2014].

Notch signaling depends on the physical interaction of ligand-expressing and receptor-expressing cells, linking the fates of cellular neighbors [Hori et al., 2013]. Thus, Notch was cleverly described as segregating specific lineages from developmentally equivalent cells as well as specifying borders between cellular fields [Hori et al., 2013]. Trans-interactions of apposing cells are activatory, while cis-interactions of receptors and ligands of the same cell are inhibitory [Guruharsha et al., 2012]. The core elements of Notch signaling include the trans-membrane Notch receptors (NOTCH1-4 in mammals), the ligands [Delta-like (DLL1, DLL4) and Jagged (JAG1, JAG2) in mammals], and CSL [CBF1–Su(H)–LAG1] DNA-binding proteins (e.g. CBF1, also known as RBPJ) mediating the transcriptional output of Notch signaling. Receptor-ligand binding ignites proteolytic cleavage and liberation of the Notch IntraCellular Domain (NICD), leading to its nuclear translocation and in the formation of a transcriptional complex including the CSL transcription factors. Upon NICD interaction with CSL, co-repressors are released and co-activators are recruited, leading to transcription of Notch-responsive genes, such as members of the Hes and Herp families (**Fig. 3.1**) [Iso et al., 2003; Guruharsha et al., 2012; Hori et al., 2013].

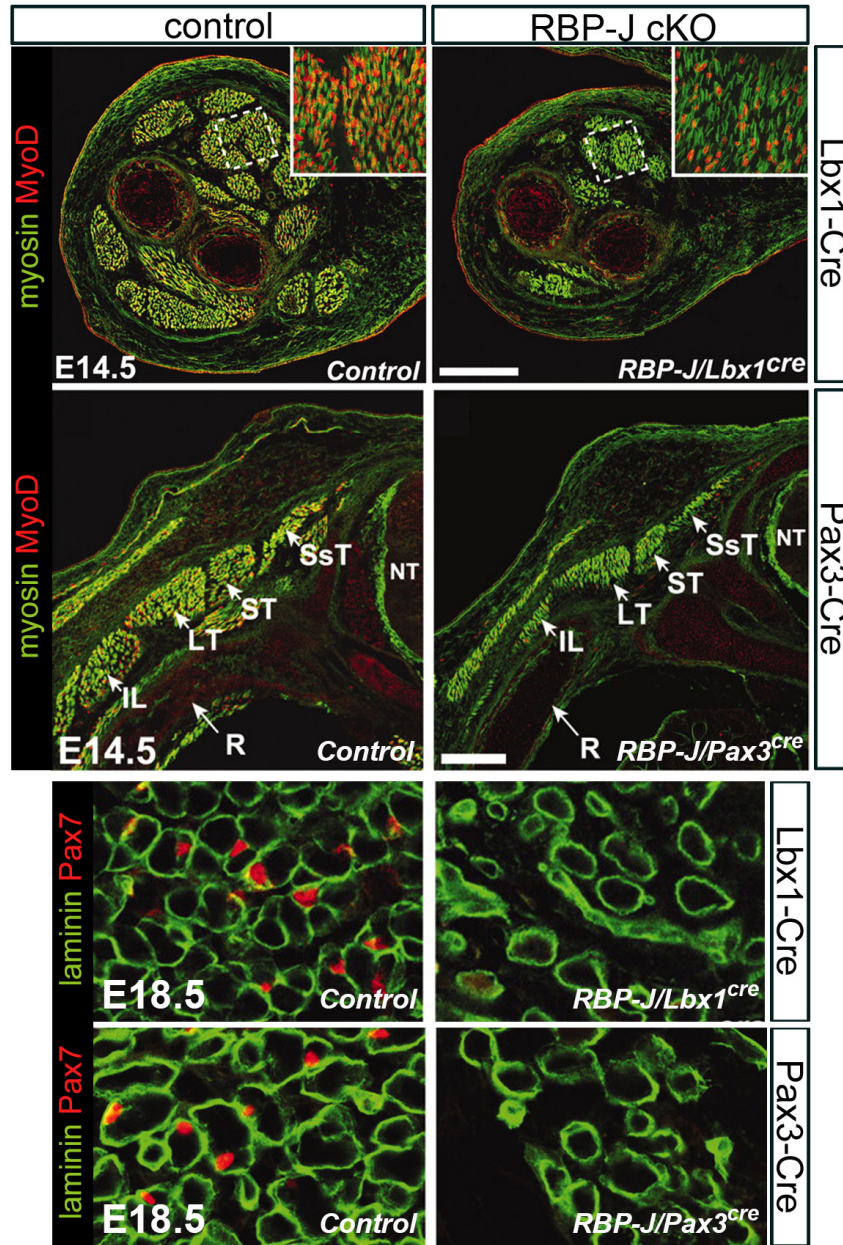


Figure 3.2. Defective myogenesis in Notch mutants.

Rbpj conditional ablation (cKO) in Lbx1+ migrating muscle progenitors or Pax3+ non-migrating hypaxial and epaxial muscle progenitors resulted in diminished muscle mass formation (upper panel) and lack of satellite cells (lower panel). NT: neural tube; R: rib; SsT: *semispinalis thoracis*; ST: *spinalis thoracis*; LT: *longissimus thoracis*; IL: *ilicostalis lumborum*.

Adapted from: Vasayutina et al., 2007

Early *ex vivo* studies suggested that Notch controls myogenesis, by suppressing differentiation of cultured myoblasts. Exposure to activated NOTCH1 or DLL1 or activated RBPJ severely affected myotube formation as well as MYOD, MYOGENIN, and MYOSIN levels [Shawber et al., 1996; Nofziger et al., 1999; Kuroda et al., 1999; Buas et al., 2009]. The genetic manipulation of Notch elements *in vivo* in mouse and chick models rendered this notion indisputable, strongly supporting that the fine tuning of this pathway is central in the development, homeostasis and regeneration of skeletal muscle [reviewed in Vasyutina et al., 2007a; Mourikis & Tajbakhsh, 2014].

In both chick and mouse development models it was shown that postmitotic cells and muscle fibers expressed various ligands (e.g. *Dll1*, *Delta1*, *Serrate2*), while mononucleated immature myoblasts expressed receptors (e.g. *Notch1*, *Notch3*) [Delfini et al., 2000; Hirsinger et al., 2001; Schuster-Gossler et al., 2007; Mourikis et al., 2012a]. Some dorsal dermomyotomal chick myogenic progenitors transiently activated Notch by *Dll1*-expressing passing neural crest cells, pointing to a different ligand source [Rios et al., 2011]. Furthermore, in that case differentiation was promoted by Notch [Rios et al., 2011], in contrast with all the other reports of developmental or adult myogenesis. Given that Notch is context-dependent, these results imply that the time window and progenitor subpopulation might affect the final outcome.

The Notch signaling pathway plays key roles during developmental myogenesis. In the chicken embryo, DLL1-triggered Notch reduced the *MyoD*-expressing region, ultimately leading to smaller and disorganized muscles [Delfini et al., 2000; Hirsinger et al., 2001]. Similar conclusions were drawn from loss-of-function (i.e. interfering with DLL1 or RBPJ) and gain-of-function (NICD overexpression) approaches in mouse embryos. *Dll1* hypomorphs [Schuster-Gossler et al., 2007] and conditional RBPJ mutants [Vasyutina et al., 2007b] were used to overcome early embryonic lethality of DLL1 and RBPJ knock-out mice. In both cases, muscle masses were diminished and the remaining muscles were deprived of satellite cells (**Fig. 3.2**). This was due to early depletion of the progenitor pool by precocious differentiation [Schuster-Gossler et al., 2007; Vasyutina et al., 2007b]. On the contrary, constitutive Notch was sufficient to maintain a self-renewing progenitor population, to promote transition into a state with quiescent signature, and to inhibit lineage progression and myogenic

differentiation [Mourikis et al., 2012a]. Interestingly, the PAX3+MYF5+ fraction was unaltered both in the gain-of-function chick experiments and in the loss-of-function mouse approach [Delfini et al., 2000; Vasyutina et al., 2007b].

Apart from the well-characterized Notch-mediated regulation of muscle progenitor maintenance by differentiation inhibition, Notch exerts additional functions in the emerging muscle masses. It has been implicated in the correct homing of emerging satellite cells; Notch elimination affected satellite cell adhesion to myofibers and basal lamina assembly around them, a phenotype rescued by concomitant ablation of *MyoD* [Bröhl et al., 2012]. Furthermore, Notch promoted the endothelial/vascular versus skeletal muscle fate in PAX3+ multipotent progenitors of the somites prior to migration to the limb [Mayeuf-Louchart et al., 2014], consistent with the role of this pathway in cell fate decisions of early developmental precursors [Buas & Kadesch, 2010]. However, Notch is dispensable for dermis versus muscle fate of precursors of the central dermomyotome [Mourikis et al., 2012a], underlining again the context-dependent outcome of Notch activity.

Similarly to the developmental scenario, adult satellite cell populations showed a prevalence of DLL1 ligand in committed PAX7+MYF5+ progenitors and NOTCH3 receptor in the upstream PAX7+MYF5- cells [Kuang et al., 2007a]. In fact, under those conditions, Notch inhibition resulted in loss of the PAX7+MYF5- population [Kuang et al., 2007a]. Furthermore, it is reported that precursor divisions lead to daughter cells with asymmetric distribution of the Notch inhibitor NUMB [Conboy & Rando, 2002]. The asymmetrical NUMB localization linked Notch cessation with myogenic progression and sustained Notch with the maintenance of undifferentiated state [Conboy & Rando, 2002]. Furthermore, NUMB was shown to prevent p53-dependent senescence in injured skeletal muscle, while *Numb* elimination impaired muscle regeneration [Le Roux et al., 2015].

In the adult, Notch is essential for the satellite cell compartment. Conditional ablation of RBPJ leads to progressive depletion of the satellite cell pool of resting or regenerating muscles and severely impairs regeneration of injured muscles. Spontaneous activation, precocious differentiation, and failure to self-renew accounted for the satellite cell loss [Bjornson et al., 2012; Mourikis et al., 2012b].

Interestingly, it was shown there is a tendency to differentiate bypassing division [Bjornson et al., 2012; Mourikis et al., 2012b]. In a complementary approach, satellite cell-specific NICD overexpression promoted self-renewal and reduced proliferation and differentiation *ex vivo*. Moreover, satellite cell-specific NICD overexpression impaired regeneration and increased PAX7+ mononuclear cells post-injury [Wen et al., 2012]. Notch was found active in quiescent satellite cells, while it declined upon *in vitro* activation or right after muscle injury [Bjornson et al., 2012; Mourikis et al., 2012b]. Notch was restored around 20-30 days post-injury [Mourikis et al., 2012b], a time point when differentiated myofibers have been formed and the satellite cell compartment is being self-renewed. However, one report associates Notch activation with transition from quiescent to activated satellite cells *ex vivo* [Conboy & Rando, 2002]. *Notch3* receptor was found upregulated in quiescent and self-renewing versus activated satellite cells [Fukada et al., 2007; Kitamoto & Hanaoka, 2010; Mourikis et al., 2012b], suggesting that it might mediate Notch effects in satellite cells. However, knock-out mice had an expanded satellite cell compartment of resting or regenerating muscles, showed increased myoblast proliferation *ex vivo* and exhibited hypertrophic muscles following regeneration [Kitamoto & Hanaoka, 2010]. During aging, Notch activity declines, impairing muscle regeneration. This was rescued by forced Notch activation in old mice; on the contrary, Notch inhibition in young muscle diminished the regenerative capacity [Conboy et al., 2003]. Conditional deletion of the different Notch receptor, as well as combined and conditional ablation should help clarify the temporal and functional requirement of this pathway during myogenesis.

In conclusion, it is argued that Notch signaling allows maintenance and expansion of muscle progenitors by preventing precocious differentiation. It is a pathway with crucial role on myogenesis and satellite cell homeostasis, while cross-talks with other pathways is not excluded. Specifically, Notch and TGF- β /pSmad3 were shown to antagonize each other by favoring satellite cell proliferation and growth arrest, respectively [Carlson et al., 2008]. The declining competence of old satellite cells was partially attributed to Notch and TGF- β /pSmad3 imbalance, occurring with aging [Carlson et al., 2008]. The balance between Notch and Wnt signaling is also decisive and a transition to Wnt was associated with differentiation steps during postnatal myogenesis [Brack et al., 2008].

RESULTS

Aims and Hypotheses

Balanced cell proliferation and differentiation are crucial for embryonic tissue formation (during development) as well as for stem cell-mediated tissue regeneration (in adulthood). Muscle differentiation relies on tissue-specific gene expression and irreversible cell cycle exit, while their dysregulation might cause apoptosis or cancer. Although several key regulators of myogenic differentiation have been identified, less information is available on the molecular circuits controlling cell cycle arrest. Thus, the present project was planned to elucidate the molecular and cellular mechanisms of growth arrest in muscle progenitor and stem cells, with specific focus upon factors and signals involved in the cell cycle exit, such as the Cyclin-Dependent Kinase Inhibitors (CDKIs) p21 and p57.

The specific aims and hypotheses outlined in the following three papers are:

Aim 1: Identification of networks controlling growth arrest at embryonic and fetal stages of muscle development.

Hypothesis: The CDKIs p21 and p57 are suppressed in the proliferating muscle progenitors to allow sufficient propagation, while they are induced in the differentiating myoblasts to ensure cell cycle exit and subsequently differentiation.

Aim 2: Development of a genetic tool allowing postnatal functional studies of p57, given that p57 mutants show perinatal lethality.

Aim 3: Evaluation of cell cycle control following activation of adult quiescent satellite cells.

Hypothesis: When quiescent satellite cells are activated upon specific homeostatic or regenerative needs, the CDKIs p21 and p57 are implicated i) in the return into quiescence (to replenish the quiescent pool) and/or ii) in irreversible cell cycle exit (to provide differentiated progeny for muscle repair).

Preface

During organogenesis, normal tissue development relies on the equilibrium between cell proliferation and cell differentiation, while overt proliferation or precocious differentiation can jeopardize correct embryogenesis. In the developing skeletal muscle, we addressed how the balance of these two processes is coordinated by growth arrest signals (i.e. the CDKIs p21 and p57), the Notch signaling pathway, and Myogenic Regulatory Factors (MRFs). Using mouse molecular genetics we showed that although cell cycle exit and myogenic differentiation occur synchronously, they can be uncoupled. Furthermore, we demonstrated that progenitors and differentiating myoblasts of the nascent muscle masses interact via the Notch signaling pathway, which is essential to repress p57 and maintain the cycling status of muscle progenitors. Finally, we identified a muscle-specific regulatory element of p57, directly regulated by an interplay between Notch downstream targets and MRFs to influence the decision between progenitor propagation and myogenic differentiation.

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RESEARCH ARTICLE

STEM CELLS AND REGENERATION

Antagonistic regulation of p57^{kip2} by Hes/Hey downstream of Notch signaling and muscle regulatory factors regulates skeletal muscle growth arrest

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ABSTRACT

A central question in development is to define how the equilibrium between cell proliferation and differentiation is temporally and spatially regulated during tissue formation. Here, we address how interactions between cyclin-dependent kinase inhibitors essential for myogenic growth arrest (p21^{cip1} and p57^{kip2}), the Notch pathway and myogenic regulatory factors (MRFs) orchestrate the proliferation, specification and differentiation of muscle progenitor cells. We first show that cell cycle exit and myogenic differentiation can be uncoupled. In addition, we establish that skeletal muscle progenitor cells require Notch signaling to maintain their cycling status. Using several mouse models combined with *ex vivo* studies, we demonstrate that Notch signaling is required to repress p21^{cip1} and p57^{kip2} expression in muscle progenitor cells. Finally, we identify a muscle-specific regulatory element of p57^{kip2} directly activated by MRFs in myoblasts but repressed by the Notch targets Hes1/Hey1 in progenitor cells. We propose a molecular mechanism whereby information provided by Hes/Hey downstream of Notch as well as MRF activities are integrated at the level of the p57^{kip2} enhancer to regulate the decision between progenitor cell maintenance and muscle differentiation.

KEY WORDS: Myogenesis, Cell cycle regulation, p57^{kip2}, Cdkn1, Notch signaling, MRF

INTRODUCTION

The formation of functional organs of an appropriate size is highly controlled during development. Organ transplantation and regeneration studies have revealed that organ size relies on both intrinsic and extrinsic mechanisms (reviewed by Cook and Tyers, 2007). Systemic factors, such as growth hormones and nutritional status, have been known for many years to regulate organ size, while more recently the role of the Hippo and insulin/TOR pathways has emerged (Tumaneng et al., 2012). Of note, increasing evidence links these pathways with stem cell self-renewal and differentiation (Cherrett et al., 2012). Nevertheless, how cell fate decisions and differentiation programs are coordinated with cell cycle progression and arrest remains poorly understood.

Skeletal muscle provides a suitable model for such studies because the molecular pathways regulating differentiation and growth arrest have been identified. Muscle formation relies on a proliferating population of progenitor cells that express and require the Paired homeobox transcription factors Pax3 and Pax7 (Buckingham and Relaix, 2007). These resident progenitors are maintained in the developing muscles, where they provide a source of cells for muscle growth during development and eventually generate the adult stem cells population, termed satellite cells (Gros et al., 2005; Kassam-Duchossoy et al., 2005; Lepper and Fan, 2010; Relaix et al., 2006). Initially, muscle progenitor cells are located in the somite where they give rise to the trunk musculature of the myotome (Ben-Yair and Kalchauer, 2005; Kassam-Duchossoy et al., 2005; Relaix et al., 2005) or migrate out of the somitic dermomyotome to form limb skeletal muscles (Birchmeier and Brohmann, 2000; Schienda et al., 2006). During limb embryonic myogenesis, Pax3/7⁺ progenitor cells undergo consecutive steps of differentiation via sequential expression of bHLH myogenic regulatory factors [MRFs; Myf5, MyoD1 and myogenin (Myog)], and first form committed progenitor cells that express Pax3/7 and Myf5, which correspond to a transit amplifying population (Picard and Marcelle, 2013), followed by the generation of myoblasts that express Myf5 and MyoD1, culminating in the appearance of differentiating myoblasts marked by Myog (Fig. 1) (Murphy and Kardon, 2011). The Myog⁺ cells then fuse to form multinucleated muscle fibers. In the absence of MyoD1, despite upregulated Myf5 expression, myogenic differentiation is delayed during early limb development, resulting in a transient absence of differentiating (Myog⁺) myoblasts and fibers prior to E14.5 (Kablar et al., 1998). When both Myf5 and MyoD1 are impaired, Pax3/7⁺ cells do not enter the myogenic program and skeletal muscle formation is abolished at all sites of myogenesis (Rudnicki et al., 1993).

Building a tissue requires the coordination of cell cycle exit with differentiation. Despite the identification of key molecular regulators of myogenic specification and differentiation (Buckingham and Relaix, 2007), how cell cycle exit is synchronized with skeletal muscle differentiation is not well understood. Cell cycle exit in muscle cells is orchestrated by cyclin-dependent kinase inhibitors (CDKIs) belonging to the CIP/Kip family: p21^{cip1} (Cdkn1a, p21^{waf1}), p27^{kip1} (Cdkn1b) and p57^{kip2} (Cdkn1c), abbreviated here as p21, p27 and p57, respectively. These CDKIs can bind and inhibit all combinations of cyclin-CDK complexes (reviewed by Besson et al., 2008). Most notably, in the absence of both p21 and p57, skeletal muscle development is severely affected and fiber formation is impaired, with myogenic cells undergoing apoptosis. This points to an essential function of p21 and p57 in cell cycle arrest during myogenesis (Zhang et al., 1999). *In vitro*, MyoD1 has been suggested to be a direct regulator of p21, thus controlling cell cycle exit during

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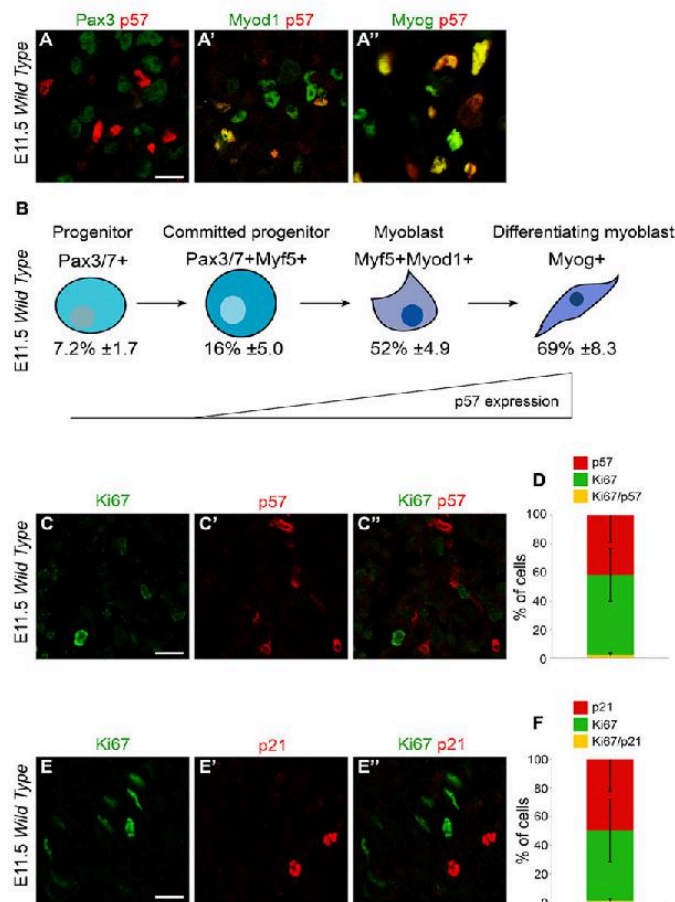


Fig. 1. Cell cycle exit occurs at the determination stage.

(A-A'') Co-immunostaining for Pax3 (A), Myod1 (A') and myogenin (Myog, A'') in green, and p57 (A', A'') in red in E11.5 embryonic limb muscles. (B) Percentage of p57-expressing cells during forelimb myogenesis is given for each population. Progenitors and committed progenitors are mostly proliferating, whereas myoblasts and differentiating myoblasts are exiting the cell cycle. (C-C'') Co-immunostaining for Ki67 (C, C'', green) and p57 (C', C'', red) in E11.5 embryonic limb muscles. (D) Quantification of C-C''. (E-E'') Co-immunostaining for Ki67 (E, E'', green) and p21 (E', E'', red) in E11.5 embryonic limb muscles. (F) Quantification of E-E''. Ki67 is not expressed in cells expressing p21 or p57. For all experiments $n=3$ embryos; error bars indicate s.d. Scale bars: 10 μ m.

adult muscle differentiation (Halevy et al., 1995). It has also been shown, both in mammalian cells (Reynaud et al., 2000) and in zebrafish (Osborn et al., 2010), that p57 interacts and stabilizes Myod1 to promote muscle differentiation, demonstrating a role for CDKIs beyond that in growth arrest. Analysis of *p21*; *p57* double-mutant mouse embryos suggested that cell cycle exit occurs in parallel to, but independently of, Myog-dependent terminal differentiation, while the lack of Mef2c expression in these mice suggested that late differentiation is defective (Zhang et al., 1999).

Previous studies have implicated the Notch signaling pathway as a key regulator of proliferation and differentiation of muscle progenitor cells (Buas and Kadesch, 2010; Mourikis and Tajbakhsh, 2014). This pathway is highly conserved during evolution and plays key roles during development, including the regulation of cell fate decisions, differentiation and homeostasis of progenitor cells in a wide variety of tissues (reviewed by Artavanis-Tsakonas and Muskavitch, 2010). Notch signaling requires direct interaction between a cell expressing at least one of the ligands [δ -like 1 (Dll1) and 4 and jagged 1 and 2 in mammals] with a cell expressing one of the receptors (notch 1-4 in mammals). This interaction leads to a proteolytic cleavage of the receptor that releases the Notch intracellular domain, which translocates into the nucleus and interacts with the Rbpj transcription factor to induce downstream effectors, such as the Hes/Hey family of bHLH transcriptional repressors (reviewed by Borggreffe and Liefke, 2012).

The role of Notch signaling in skeletal muscle development has been assessed in two mouse models: in a hypomorphic *Dll1* mutant (Schuster-Gossler et al., 2007) or in mice in which *Rbpj* expression was conditionally abrogated specifically in the myogenic lineage (Vasyutina et al., 2007). These *in vivo* models, along with studies performed in chick embryos, have demonstrated that Dll1-triggered canonical Notch signaling is required for the maintenance of muscle progenitor cells (Delfini et al., 2000; Hirsinger et al., 2001; Mourikis et al., 2012a; Schuster-Gossler et al., 2007; Vasyutina et al., 2007). Dll1 absence leads to early onset differentiation (Schuster-Gossler et al., 2007; Vasyutina et al., 2007), resulting in rapid exhaustion of the muscle progenitor cell pool and near complete absence of skeletal muscles at the fetal stage (Schuster-Gossler et al., 2007; Vasyutina et al., 2007). This is in part mediated by the repression of Myod1 target genes through direct binding of Hey1 to their promoters (Bröhl et al., 2012; Buas et al., 2010). Interestingly, the role of Notch can be context dependent, since in the young somite of the chick embryo, Dll1⁺ neural crest cells provide a transient stimulation of Notch activity that is important for the initiation of early myogenesis (Rios et al., 2011).

Here, we evaluated the *in vivo* expression of p57 and its link with muscle cell differentiation. Although cell cycle exit is normally synchronous with cell differentiation, we show that these events can be uncoupled. In fact, we found that during embryonic myogenesis p57-mediated cell cycle arrest occurs earlier than

previously recognized, namely in determined muscle cells. Moreover, we demonstrate that in the absence of terminal differentiation muscle progenitor cells aberrantly induce p57 expression, leading to growth arrest. We further show that this growth arrest is associated with a loss of Notch signaling. This is confirmed by conditional genetic ablation of *Rbpj* that leads to upregulation of p21 and p57 in muscle progenitors associated with increased growth arrest. We finally identify a muscle-specific p57 regulatory element and show that this enhancer is the target of both positive regulation by MRFs in myoblasts and negative regulation by Hes/Hey repressors downstream of Notch in progenitor cells. Our data therefore demonstrate that the regulation of cell cycle exit integrates both negative (via Hes/Hey downstream of Notch signaling) and positive (by MRFs) regulation at the same p57 regulatory element during muscle differentiation, and that Notch signaling acts upstream, but independently, of both differentiation and cell growth arrest.

RESULTS

Cell cycle exit and differentiation can be uncoupled during skeletal muscle development

We first assessed whether myogenic progenitors leave the cell cycle at specific steps of the MRF-mediated differentiation program, by comparing p57 expression with that of MRFs in E11.5 mouse limbs by immunofluorescence (Fig. 1A-A''). As expected, p57 expression was very low in Pax3/7⁺ progenitors (7.2 ± 1.7%). By contrast, a proportion of the Pax3/7⁺/Myf5⁺ committed progenitor cells did express p57 (16 ± 5%), and this proportion

increased significantly in Myf5⁺/Myod1⁺ (52 ± 4.9%) and Myog⁺ (69 ± 8.3%) populations (Fig. 1B). Similar results were obtained with p21 (data not shown). We verified that p21 and p57 are accurate markers of cell cycle exit of myogenic progenitors as their expression almost never co-localized with that of Ki67, a marker of cycling cells (Fig. 1C-F). Our data are consistent with the results of previous *in vivo* studies analyzing the proliferation of myogenic cells during development (Gros et al., 2005; Lagha et al., 2008; Relaix et al., 2005).

In order to test the existence of a link coupling cell cycle arrest with muscle differentiation, we first investigated whether muscle differentiation is affected when cell cycle exit is impaired. We examined whether the differentiation program proceeds normally in *p21*:*p57* double-null embryos, in which growth arrest is abolished (Zhang et al., 1999). In limb muscles of control mice, 4.7 ± 1.4% of Myog-positive cells underwent proliferation as assessed by phosphohistone H3 (P-H3) (Fig. 2A-A'',C). By contrast, *p21*^{-/-}; *p57*^{+/-m} double-mutant embryos displayed a marked increase in Myog⁺/P-H3⁺ cells (25.2 ± 3.2%; Fig. 2B-C). Taken together, we conclude that p21- and p57-mediated cell cycle exit and MRF-mediated myogenic differentiation can occur independently of each other.

We then examined whether the uncoupling of proliferation and differentiation that we observed in the *p21*^{-/-}; *p57*^{+/-m} double-mutant embryos holds true in a complementary condition. Delayed myogenesis in *Myod1* mutant embryos provides a useful model for such analysis (Kablar et al., 1997). As expected, Myog and p57 co-localized in the forelimbs of control *Myod1*^{+/-} mice at E12.5 (Fig. 2D-D''). By contrast, in the E12.5 *Myod1*^{-/-} forelimbs, even

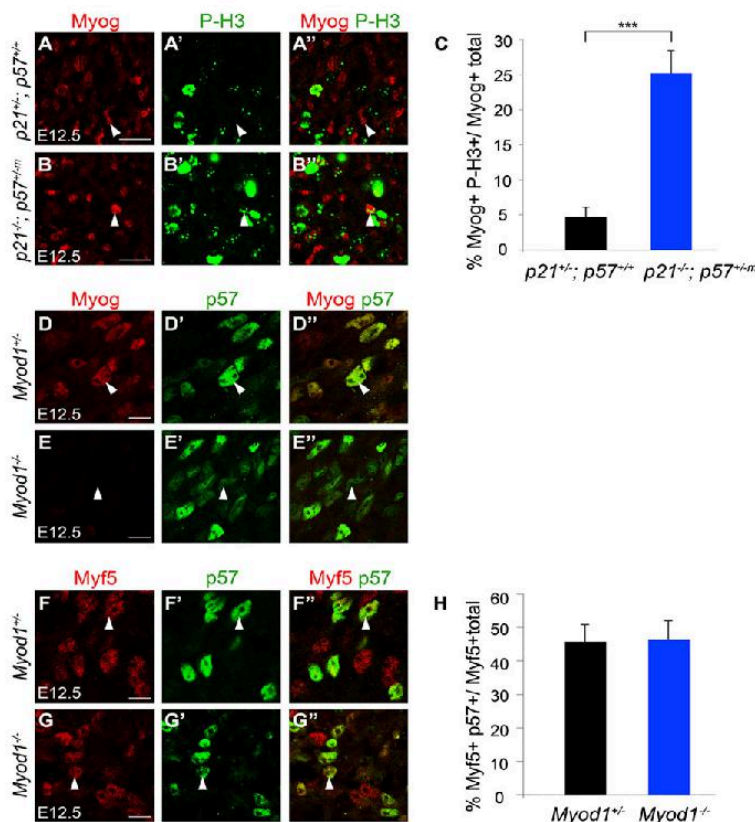


Fig. 2. Cell cycle exit can be uncoupled from cell differentiation. (A-B'') Co-immunostaining for Myog (A, A'', B, B'', red) and P-H3 (A', A'', B', B'', green) in *p21*^{+/-}; *p57*^{+/-} (A-A'') or *p21*^{-/-}; *p57*^{+/-m} (B-B'') forelimbs at E12.5. Myog⁺ cells (A) do not normally express P-H3 (A', A''), whereas in *p21*^{-/-}; *p57*^{+/-m} embryos Myog⁺ cells aberrantly proliferate (B-B''). (C) Quantification of A, B''. (D-G'') Co-immunostaining for Myog (D, D'', E, E'', red), p57 (D', D'', E', E'', F', F'', G', G'', green) and Myf5 (F, F'', G, G'', red) in *Myod1*^{+/-} (D-D'', F-F'') or *Myod1*^{-/-} (E-E'', G-G'') embryonic limb muscles at E12.5. Myog⁺ cells express p57 in *Myod1*^{-/-} embryos (D-D'', arrowheads). p57 is expressed in *Myod1*^{-/-} embryos (E) despite the absence of Myog (E). Myf5 is co-expressed with p57 in both *Myod1*^{+/-} (F-F'', arrowheads) and *Myod1*^{-/-} (G-G'') embryos. (H) Quantification of F', G''. For all experiments *n*=3 embryos for each genotype; error bars indicate s.d.; ****P*<0.001. Scale bars: 10 μm.

though Myog is not expressed, p57 is detected in the forming muscle masses (Fig. 2E-E''), where it labels nearly half of the Myf5⁺ cells in both *Myod1*^{+/+} (Fig. 2F-F'',H) and *Myod1*^{-/-} (Fig. 2G-H) forelimb (45.6±5.1% versus 46.3±5.5%). These data suggest that cell cycle exit coincides with Myf5 expression in myoblasts and is unaffected when Myod1/Myog-mediated differentiation is impaired.

In the absence of differentiated myoblasts, muscle progenitors precociously express p57 and exit the cell cycle

It has been previously shown that differentiating myoblasts are required for the survival of muscle progenitor cells throughout development (Kassar-Duchossoy et al., 2005). We examined in more detail the impact of differentiating myoblasts on the proliferation state of Pax3⁺ cells by analyzing different allelic combinations of *Myod1*:*Myf5* double-null embryos to allow key steps during myogenic commitment to be separated. In the absence of Myod1⁺ myoblasts but in the presence of Myf5⁺ myoblasts in *Myod1*^{-/-}; *Myf5*^{+nlacZ} mice (Rudnicki et al., 1993; Tajbakhsh et al., 1997) (supplementary material Fig. S1), the proliferation rate of Pax3⁺ cells was comparable to that observed in control mice at E12.5 (23.6±3.9% versus 25.6±4.6%; Fig. 3A-B'',D). By contrast, in the double-mutant *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} forelimbs, which lack both committed progenitors and myoblasts (supplementary material Fig. S1), we observed a significant decrease in the proliferation of Pax3⁺ cells (12.8±3.6% versus 25.6±4.6%; Fig. 3C-D). These data suggest that committed progenitors are required to maintain the proliferation of muscle progenitor cells, whereas differentiated myoblasts are dispensable.

Consistent with the proliferation profile, the cell cycle inhibitor p57 was aberrantly expressed in Pax3⁺/MRF⁻ progenitor cells of *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} embryos compared with control embryos (28.4±2.7% versus 2.3±2.7%; Fig. 3E-G). These data suggested that myoblasts are required to maintain cycling muscle progenitor cells by preventing p57 expression and cell cycle arrest.

Impaired Notch signaling in *Myod1*; *Myf5* mutant embryos

Our analysis of *Myod1*; *Myf5* mutant embryos reinforced the notion that functional interactions are taking place between myoblasts and muscle progenitor cells. A strong candidate pathway to mediate these interactions is Notch signaling. It has been previously shown that differentiating myogenic cells express Dll1 and possibly signal to the upstream population that expresses higher levels of Notch receptors (mainly notch 1, 2 and 3) (Delfini et al., 2000; Hirsinger et al., 2001; Mourikis et al., 2012b; Schuster-Gossler et al., 2007). This feedback mechanism of receptor/ligand regulation is supported by many independent *in vivo* studies. However, it has not been formally shown that such cell-cell interactions occur during development, a prerequisite for Notch signaling.

To demonstrate an interaction between myoblasts and muscle progenitor cells, we analyzed the cellular organization on sections of embryonic forelimb muscle masses by co-immunostaining, and found that the majority of Pax7⁺ progenitor cells are in close proximity to Myod1⁺ myoblasts (Fig. 4A-B). Our analysis therefore suggests that direct cell-cell signaling via Notch can occur between progenitors and myoblasts.

To further assess the significance of differentiating muscle cells in Notch activation, we measured endogenous pathway activity in E12.5 *Myod1*; *Myf5* double-mutant embryos that lack differentiated muscle due to the MRF deficiency. It was previously shown that Pax7 expression is lost when Notch signaling is abrogated in myogenic progenitor cells (Vasyutina et al., 2007). Consistent with impaired Notch activity, Pax7 protein was undetectable by immunofluorescence at E12.5 in *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} forelimbs (Fig. 4C-E), whereas it was expressed in *Myod1*^{-/-}; *Myf5*^{+nlacZ} embryos (Fig. 4C-E). In addition, we found downregulation of the Notch target genes *Hes1* and *Hey1* in the forelimbs of *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} compared with *Myod1*^{-/-}; *Myf5*^{+nlacZ} or with *Myod1*^{-/-}; *Myf5*^{+nlacZ} at E12.5 (Fig. 4F,G).

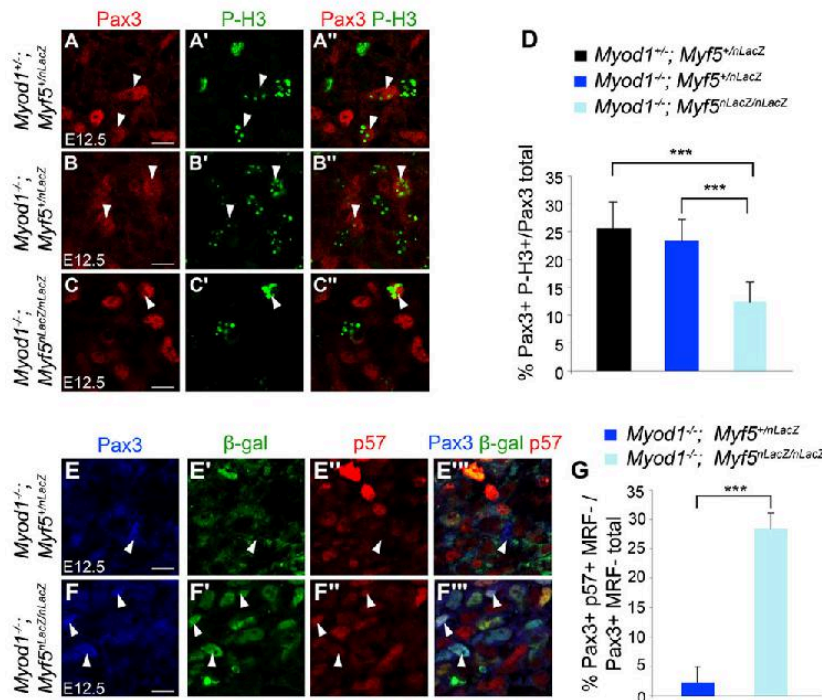


Fig. 3. Myoblasts control muscle progenitor cell proliferation by preventing cell cycle exit. (A-C'') Co-immunostaining for Pax3 (red) and P-H3 (green) in *Myod1*^{+/+}; *Myf5*^{+nlacZ} (A-A''), *Myod1*^{-/-}; *Myf5*^{+nlacZ} (B-B'') and *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} (C-C'') embryos at E12.5. Arrowheads indicate Pax3⁺ cells undergoing mitosis. (D) Quantification of A'', B'', C''. (E-F'') Co-immunostaining for Pax3 (blue), β -gal (green) and p57 (red) in *Myod1*^{+/+}; *Myf5*^{+nlacZ} (E-E'') or *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} (F-F'') embryos at E12.5. *Myf5*⁻ β -gal⁻ cells do not express p57 (arrowheads in E-E'') in *Myod1*^{+/+}; *Myf5*^{+nlacZ} embryos, whereas in *Myod1*^{-/-}; *Myf5*^{nlacZ/nlacZ} embryos Pax3⁺ cells are p57⁺ (arrowheads in F-F''). (G) Quantification of E'', F''. For all experiments $n=3$ embryos for each genotype; error bars indicate s.d.; *** $P<0.001$. Scale bars: 10 μ m.

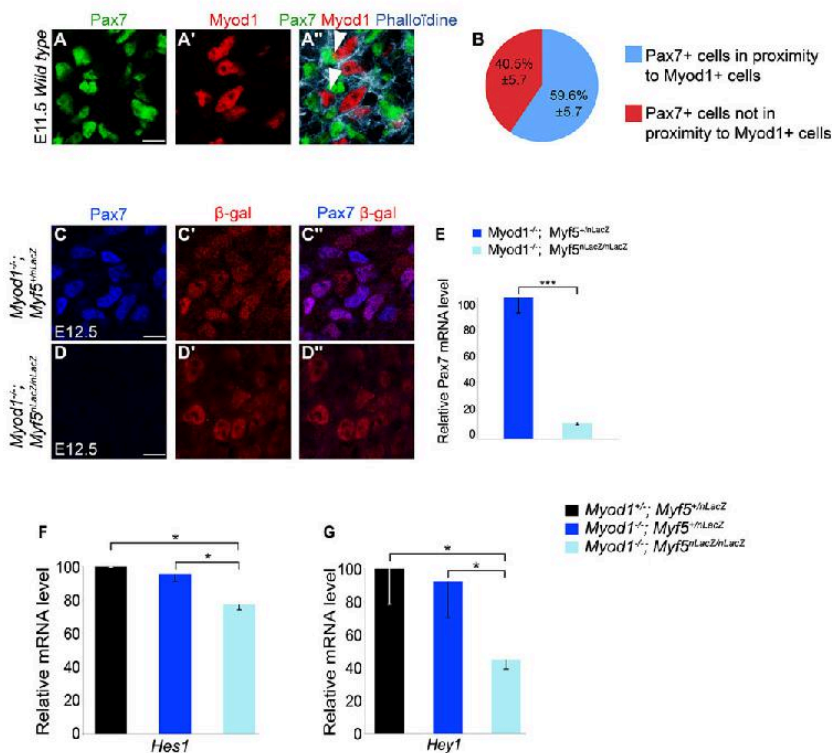


Fig. 4. Close proximity of Pax7⁺ and Myod1⁺ cells, with decreased Pax7 and Hes1/Hey1 expression in muscle progenitor cells of the Myod1; Myf5 double mutant. (A-A'') Co-immunostaining for Pax7 (green) and Myod1 (red), with phalloidin (cyan) to label actin to visualize cell membranes, in wild-type limb muscles at E11.5. (B) Percentage of Pax7⁺ cells in proximity to Myod1⁺ cells in limb muscle masses. (C-D'') Co-immunostaining for Pax7 (blue) and β -gal (red) in *Myod1*^{-/-}; *Myf5*^{+/rlacZ} (C-C'') and *Myod1*^{-/-}; *Myf5*^{rlacZ/rlacZ} (D-D'') embryos at E12.5. (E-G) qRT-PCR for *Pax7* (E), *Hes1* (F) and *Hey1* (G) on E12.5 forelimbs of the genotypes indicated. For all experiments $n=3$ embryos for each genotype; error bars indicate s.e.m.; * $P<0.05$, ** $P<0.01$, *** $P<0.001$. Scale bars: 10 μ m.

Notch signaling prevents activation of p57 in muscle progenitor cells

Based on our results (Fig. 4) and previous reports (Georgia et al., 2006), we hypothesized that myoblasts control progenitor cell proliferation by activating the Notch/Hes1/Hey1 pathway, which would then repress p57 expression.

First, to establish whether Notch signaling participates directly in the coordinated control of cell cycle exit and differentiation, we used an *ex vivo* whole limb culture system (Zúñiga et al., 1999). We cultured E11.5 mouse forelimbs for 28 h, with or without 20 μ M γ -secretase inhibitor DAPT, an inhibitor of Notch signaling. As expected, we saw decreased expression of the Notch target genes *Hes1* and *Hey1* after DAPT treatment (Fig. 5A). In addition, inhibition of Notch signaling led to reduced numbers of Pax7⁺ cells (56.8 \pm 5.6% in control versus 27.7 \pm 7.0% in DAPT-treated limb explants; Fig. 5B',C',D), whereas the Myod1⁺ cell population was increased (62.7 \pm 9.0% compared with 32.6 \pm 5.3% in control DMSO-treated explants; Fig. 5B'',C'',D), confirming previous reports (Schuster-Gossler et al., 2007; Vasyutina et al., 2007) and the robustness of our *ex vivo* model. Accordingly, we found decreased levels of *Pax7* mRNA and increased levels of *Myod1* mRNA in DAPT-treated samples (Fig. 5A). We next examined whether pharmacological inhibition of Notch signaling induces cell cycle arrest in cultured muscle progenitor cells. We found a 5-fold increase in p57 expression in Pax3⁺/MRF⁻ cells in DAPT-treated limb explants compared with controls (Fig. 5E-G).

To confirm these results *in vivo*, we genetically abrogated Notch signaling in progenitor cells by conditionally deleting *Rbpj*. RbpJ is a DNA-binding transcription factor and the major effector of all four Notch receptors (Fortini and Artavanis-Tsakonas, 1994; Jarriault et al., 1995; Kopan and Ilagan, 2009; Schweisguth and Posakony, 1992).

We performed a conditional deletion of *Rbpj* in the Pax3 lineage by crossing *Rbpj*^{flax/flax} mice (Han et al., 2002) with a *Pax3*^{Cre/+} allele (Engleka et al., 2005). Ablation of *Rbpj* led to increased myogenic differentiation as previously reported (Vasyutina et al., 2007), with a severe loss of progenitor cells leading to tiny limb muscles at a fetal stage. Strikingly, both p57 and p21 were upregulated in the Pax3⁺/Myf5⁻ muscle progenitor cells in the forelimbs of *Rbpj*^{flax/flax}; *Pax3*^{Cre/+} mice at E11.5, whereas Pax3 and these CDKIs were rarely co-expressed in such cells in control mice (Fig. 6A-D, see also Fig. 1). To demonstrate that expression of p21 and p57 is associated with growth arrest in these mutants, we analyzed the co-expression of Ki67 with either p57 or p21 in Pax3⁺ muscle progenitors (Fig. 6E,F) in the forelimbs of *Rbpj*^{flax/flax}; *Pax3*^{Cre/+} mice at E11.5. We found a small but significant increase of Pax3⁺ cells co-expressing p21 or p57 with Ki67 in the mutant embryos; nevertheless, the large majority of the Pax3⁺/p57⁺ cells did not express Ki67, as predicted.

Altogether, these results demonstrate that in embryonic muscle progenitor cells Notch signaling antagonizes cell cycle exit by repressing p57 expression.

A p57 muscle-specific enhancer is directly regulated by Notch signaling and MRFs

To gain insight into the molecular mechanisms of p57 regulation, we used data generated by a Myod1 ChIP sequencing experiment (Cao et al., 2010) to identify Myod1 binding sites in the vicinity of the p57 locus. A previous study had predicted that p57 muscle-specific regulatory elements are located between +35 and +225 kb from the p57 transcription start site (John et al., 2001). In keeping with this, a high density of Myod1 binding sites was found in a conserved region located +59 kb from p57. We isolated an evolutionarily conserved

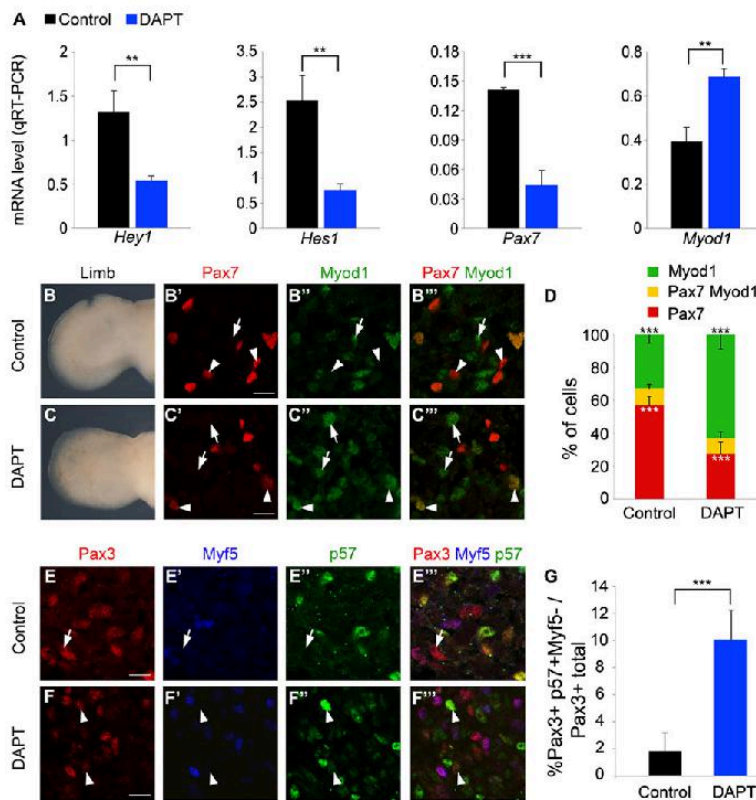


Fig. 5. The Notch pathway prevents activation of p57 in progenitor cells. (A) qRT-PCR for *Hey1*, *Hes1*, *Pax7* and *Myod1* mRNA in control (DMSO-treated) and DAPT-treated *ex vivo* whole limb culture. (B,C) An E11.5 forelimb kept in culture for 28 h treated with DMSO (B) or 20 μ M DAPT (C). (B'-B'', C'-C'') Co-immunostaining for Pax7 (red) and Myod1 (green) in DMSO-treated (B'-B'') or 20 μ M DAPT-treated (C'-C'') explants from E11.5 limb muscles. Arrowheads indicate Pax7⁺/Myod1⁻ cells in B'-B'' and Pax7⁺/Myod1⁺ cells in C'-C''; arrows indicate Myod1⁺/Pax7⁻ cells. (D) Quantification of B'', C''. (E-F'') Co-immunostaining for Pax3 (red), Myf5 (blue) and p57 (green) in DMSO-treated (E-E'') or 20 μ M DAPT-treated (F-F'') explants from E11.5 limb muscles. Arrow in E-E'' indicates a Pax3⁺/Myf5⁻/p57⁻ cell. Arrowheads in F-F'' indicate Pax3⁺/Myf5⁻/p57⁺ cells. (G) Quantification of E'', F''. For all experiments $n=3$; error bars indicate s.d.; ** $P<0.01$, *** $P<0.001$. Scale bars: 10 μ m.

686 bp fragment that contains 15 E-boxes, which are binding sites for MRFs, Hey1 and Hes1 (supplementary material Fig. S2).

We first validated this *p57* muscle regulatory element (*p57MRE*) as a functional enhancer *in vivo* by generating transgenic embryos carrying a *p57MRE-tk-nlacZ* construct. Following analysis of *lacZ* expression at E12, we detected robust reporter expression in all myogenic domains (Fig. 7A,A'), with an expression profile that matched that of Myod1. Interestingly, this element is skeletal muscle specific, since no other sites of p57 expression, such as parenchymal organs and intestine (Westbury et al., 2001), were observed. In order to characterize the myogenic cell type that expresses the p57 reporter, we performed immunohistochemical analyses on limb buds from these transgenic embryos. β -Gal⁺ cells co-expressed p57 (Fig. 7B-B'') and Myod1 (Fig. 7C-C'') but not Pax7 (Fig. 7D-D''), defining the cellular specificity of the *p57MRE*.

We next hypothesized that this regulatory element integrates negative regulation by Hes/Hey proteins and positive regulation via direct activation by the MRFs. We performed ChIP experiments on E12.5 wild-type forelimbs and found that both Myod1 and Hes1 were bound *in vivo* to the *p57MRE* fragment (Fig. 8A). To ensure that our assay was specific, and given the lack of known positive controls for Hes1 in the myogenic lineage, we performed ChIP experiments in HEK293 cells transfected with either Hes1 or Myod1 and either wild-type *p57MRE* or containing mutations in the MRF and Hes binding sites (*p57MREAE-Boxes*). Robust binding was observed for Hes1 (Fig. 8B) and Myod1 (Fig. 8C) on the *p57MRE* and this binding was abrogated on *p57MREAE-Boxes* (Fig. 8B,C).

Finally, to further establish this interplay between positive and negative regulation, we tested the transcriptional activity of Myod1,

Hes1 and Hey1 on *p57MRE-tk-nlacZ* in transient transfection experiments in C2C12 muscle cells. Myod1 enhanced the activation of the *p57MRE* (Fig. 8D), but was not able to activate the *p57MREAE-Boxes* element. Furthermore, Myod1 transcriptional activation was abolished when exposed to increasing concentrations of Hes1 or Hey1 (Fig. 8D), suggesting that both are able to repress the Myod1-dependent activation of *p57MRE*.

We propose a model in which the integration of Notch and MRF activities at the level of a muscle-specific enhancer of the key cell cycle arrest gene *p57* provides a means to control the equilibrium between progenitor pool amplification and the establishment of definitive functions of skeletal muscle (Fig. 8E).

DISCUSSION

The generation of organs of a defined size requires a balance between proliferation and differentiation. This balance is ensured by regulated cell growth, which prevents prolonged proliferation or premature differentiation, both of which are deleterious for normal development.

During skeletal muscle development and postnatal regeneration, Notch signaling activity is crucial for sustaining stem/progenitor cell self-renewal and its downregulation is required to allow myogenic differentiation. Cell cycle exit was previously thought to be controlled by the differentiation program (Halevy et al., 1995). In this report we show that growth arrest is also negatively regulated by Notch signaling and demonstrate that these two events, despite appearing synchronous, can be uncoupled. In *Myod1*^{-/-} forelimbs, myogenesis is paused between E11.5 and E14.5 (Kablar et al., 1998). Although Myf5 is unable to drive myogenesis and activate *Myog* at these stages,

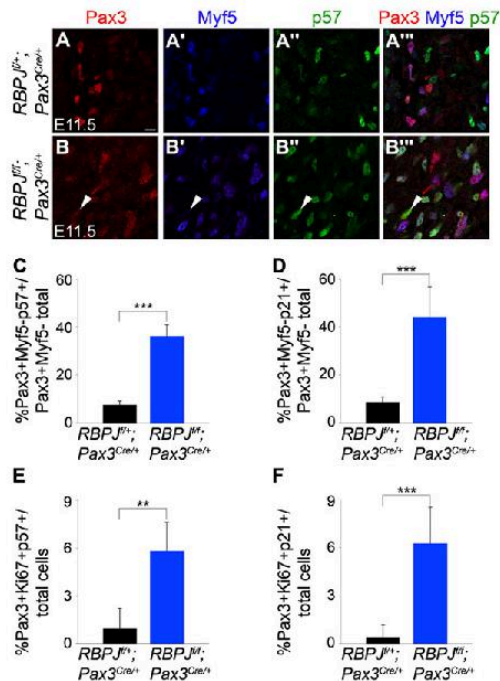


Fig. 6. Conditional ablation of *Rbpj* leads to upregulation of p57 and p21 and to cell cycle arrest in muscle progenitor cells. (A-B) Co-immunostaining for Pax3 (red), Myf5 (blue) or p57 (green) in *Rbpj*^{fl/+}; Pax3^{Cre/+} (A-A') or *Rbpj*^{fl/-}; Pax3^{Cre/+} (B-B') forelimbs at E11.5. Arrowhead indicates a Pax3⁺/Myf5⁺/p57⁺ cell. Scale bars: 10 μ m. (C) Quantification of A', B'. (D) Quantification of co-immunostaining for Pax3, Myf5 or p21 in *Rbpj*^{fl/+}; Pax3^{Cre/+} or *Rbpj*^{fl/-}; Pax3^{Cre/+} forelimbs at E11.5. (E) Quantification of co-immunostaining for Pax3, Ki67 or p57 in *Rbpj*^{fl/+}; Pax3^{Cre/+} or *Rbpj*^{fl/-}; Pax3^{Cre/+} forelimbs at E11.5. (F) Quantification of co-immunostaining for Pax3, Ki67 or p21 in *Rbpj*^{fl/+}; Pax3^{Cre/+} or *Rbpj*^{fl/-}; Pax3^{Cre/+} forelimbs at E11.5. For all experiments n=3 embryos for each genotype; error bars indicate s.d.; **P<0.01, ***P<0.001.

we found that Myf5⁺/Pax3/7⁺ cells expressed p57 at E12.5 and this did not prevent them from resuming differentiation at E14.5 (presumably when Mrf4 is activated). Given our finding that MyoD1 directly binds and activates *p57* via the *p57MRE* sequence, we believe that Myf5 operates in the same way, thereby providing a functional uncoupling between MRF myogenic activity and growth arrest. Moreover, our study and those of others indicate that cell cycle exit occurs at the transition from committed progenitors to determined myoblasts (Fig. 1A). Consistently, we found that committed progenitor cells express Pax3/7 and Myf5, but neither p21 nor p57. This finding is consistent with the robust repressive activity exerted by Hes/Hey on MRF-mediated transactivation (Fig. 8D). The cycling status of committed progenitor cells is therefore of interest. A recent study showed that whereas the undifferentiated resident progenitor cells that express Pax7 represent a slow-cycling pool, the Pax3/7⁺/Myf5⁺ committed progenitors correspond to a fast-cycling population (Picard and Marcelle, 2013). Our study did not address the subtle cell cycle regulation of these progenitor cell populations and future studies will be required to determine whether these changes in cell proliferation are linked to Myf5 or to other, as yet unidentified, factors.

The model of coordinated regulation that we propose, with a single *p57* element integrating positive (from the MRFs) and negative (from Hes/Hey) regulatory information suggests that the interplay

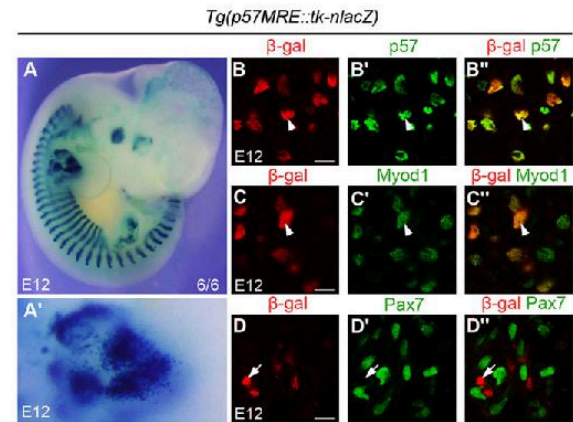


Fig. 7. Expression of a *p57* muscle regulatory enhancer (MRE) in transgenic mice. (A,A') X-Gal staining on a transgenic *p57MRE-tk-nlacZ* embryo. A' is a higher magnification of the forelimb region from A. (B-D') Co-immunostaining for β -gal (B,C,D,B',C',D', red), p57 (B',B'', green), MyoD1 (C',C'', green) and Pax7 (D',D'', green). Arrowheads indicate β -gal⁺/p57⁺ (B-B') and β -gal⁺/MyoD1⁺ cells (C-C'); arrows indicate β -gal⁺/Pax7⁻ cells. Scale bars: 10 μ m.

between Notch repression of *p57MRE* in Pax3/7 progenitors and its activation by MRFs in myoblasts is crucial for growth arrest. The molecular mechanisms regulating Notch signaling components during myogenesis are not fully characterized. It was reported that during *Xenopus* development Dll1 expression is regulated by MyoD1 (Wittenberger et al., 1999) and that MyoD1 expression is repressed by Hairy-1 (Umbhauer et al., 2001). It is unclear if these regulatory mechanisms also exist in amniotes, but our data are compatible with such a sequence of events. Resolving the precise molecular interplay between Pax gene expression, cell growth arrest, MRF regulation and the switch in Notch signaling will require additional investigations.

Notch signaling plays a key role in maintaining the homeostasis of muscle stem cells in the adult (Bjornson et al., 2012; Carlson et al., 2008; Fukada et al., 2011; Kitamoto and Hanaoka, 2010; Mourikis et al., 2012b) and in colonization of the satellite cell niche (Bröhl et al., 2012). In particular, Notch controls quiescence of muscle satellite cells (Bjornson et al., 2012; Mourikis et al., 2012b). This activity might be mediated by Hey1 and HeyL, which are required in the adult lineage for satellite cell homeostasis and skeletal muscle regeneration (Fukada et al., 2011). Conditional deletion of *Rbpj* in Pax7⁺ satellite cells led to spontaneous differentiation without activation or division of the cells (Bjornson et al., 2012; Mourikis et al., 2012b). Strikingly, *Rbpj* ablation does not lead to an immediate and complete differentiation or growth arrest in the Pax3⁺ population during embryonic development, leaving open the possibility that other pathways are involved. For instance, Notch activity on adult muscle stem cells is counteracted by TGF β signaling (Carlson and Conboy, 2007). This is mediated through the activation of phosphorylated Smad3, which can directly bind and activate the *p15* (*Cdkn2b*), *p16* (*Cdkn2a*), *p21* and *p27* promoters (Carlson and Conboy, 2007) to favor muscle stem cell differentiation. Interestingly, during chicken myogenesis myostatin, which is a member of the TGF β family, has also been implicated in the control of terminal differentiation through indirect activation of p21 (Manceau et al., 2008).

In addition to driving cell cycle exit during adult myogenesis, p57 has also been implicated in stabilization of MyoD1 through direct association in C2C12 cells, resulting in enhanced myogenesis

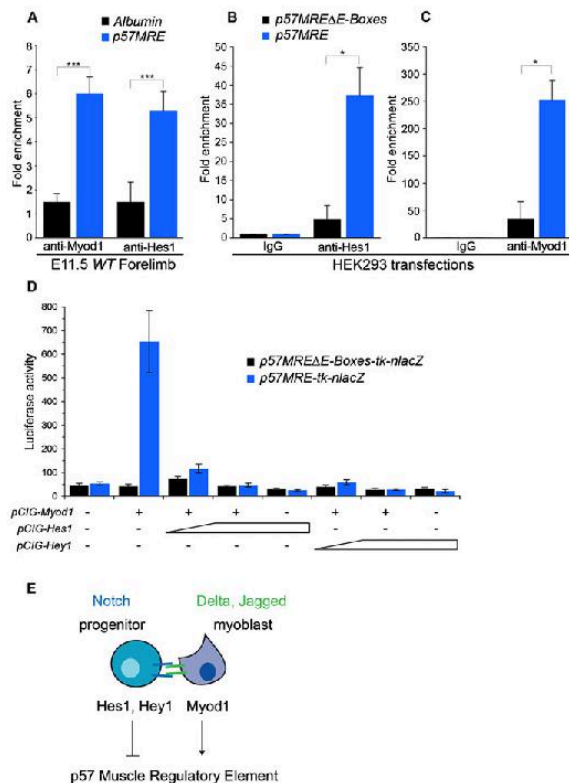


Fig. 8. Direct regulation of the *p57MRE* by Myod1 and Hes1/Hey1. (A) Chromatin immunoprecipitation followed by qPCR on wild-type forelimbs at E11.5. *p57MRE* is enriched when precipitated with anti-Myod1 or anti-Hes1 antibodies compared with an albumin gene control. (B,C) Validation of antibody ChIP capacities on transfected HEK293 cells: enrichment with anti-Hes1 (B) or anti-Myod1 (C) is obtained with the *p57MRE* compared with the construct in which all putative E-boxes have been mutated (*p57MREΔE-Boxes*). $n=3$; error bars indicate s.e.m.; * $P<0.05$, *** $P<0.001$. (D) Transactivation assay on C2C12 cells with the expression plasmids and reporters indicated ($n\geq 3$). (E) Schematic representation of the regulation of cell cycle exit during myogenesis. In muscle progenitors, Notch downstream effectors Hes1 and Hey1 repress the activation of p57 to allow the amplification of the pool, while in the neighboring myoblasts that express the Notch ligands, Myod1 directly activates p57 expression.

(Reynaud et al., 2000). A similar mechanism has also been identified in zebrafish, in which p57 cooperates with Myod1 to drive the differentiation of several early zebrafish muscle fiber types (Osborn et al., 2010). It is not known if this positive-feedback loop also operates during early murine skeletal muscle formation. One could propose that, although the initiation of myogenic differentiation and growth arrest are independent, these events may synergize subsequently, for instance to enhance Myod1 activity and reinforce terminal differentiation. In zebrafish, p57 cooperates with Myod1 to drive *myog* expression (Osborn et al., 2010); nevertheless, proliferating Myod1⁺ and Myog⁺ cells are detected in *p21^{-/-}*; *p57^{+/-}* mice (see Fig. 2A-C; our unpublished observations). Interestingly, expression of *Mef2c* is impaired in these mutant mice (Zhang et al., 1999), raising the possibility that p57 may also be involved in terminal differentiation in murine myogenesis during development.

In our study, the expression of p57 is firmly linked to an absence of cell cycle progression, since we observe no overlap between p57

(or p21) expression and Ki67 (Fig. 1C-F) under normal conditions. Strikingly, a small but significant proportion of the Pax3⁺/p21⁺ or Pax3⁺/p57⁺ cells are Ki67⁺ in the Pax3^{Cre/+}; *Rbpj^{lox/flox}* mutant context. Although this might correspond to a transitory state due to the differentiation phenotype of these mutant embryos, one cannot exclude the possibility that Notch might also be involved in both cell cycle progression and cell cycle arrest via a complex regulatory loop.

p57 expression has been reported previously in adult satellite cells (Fukada et al., 2007), but the precise timing of expression has yet to be characterized. The identification of *p57MRE* through a Myod1 ChIP-seq screen performed in C2C12 cells raises the possibility that this element is reused in adult muscle cells *in vivo*. Owing to the perinatal death of *p57* mutant mice, the role of p57 in postnatal myogenesis cannot be studied *in vivo*. *p21*-deficient mice display normal muscle development but impaired skeletal muscle regeneration (Hawke et al., 2003). Given the functional overlap between p21 and p57 during development, it would be interesting to evaluate the combined role of these two proteins in postnatal satellite cell homeostasis and skeletal muscle regeneration.

The recent identification of the role of p57 in the maintenance of quiescent hematopoietic (Matsumoto et al., 2011), neural (Furutachi et al., 2013) and lung (Zacharek et al., 2011) stem cells indicates that p57, along with other CDKIs, is important for stem cell function. Whether such a regulatory mechanism for CDKI expression is redeployed in other systems remains to be investigated. For example, Notch has been implicated in maintaining progenitor cell proliferation in intestinal stem cells (Riccio et al., 2008), in adult neural stem cells (Imayoshi et al., 2010) and in Rathke's pouch progenitors of the pituitary (Monahan et al., 2009) and, indeed, one proposed mechanism is the repression of CDKIs by the product of the Notch target gene *Hes1* (Monahan et al., 2009; Riccio et al., 2008). Unfortunately, these studies did not define which cells provide the ligands. Nevertheless, our data and the role of Notch and Hes1 in intestinal stem cells, neural stem cells and pituitary progenitor cells might suggest a general mechanism whereby the expansion of the progenitor cell population is regulated via modulation of CDKI genes. Such a regulatory mechanism could be used as a safeguard to prevent tumor formation by progenitor/stem cells, for instance when differentiation is impaired. It is also tempting to speculate that fine-tuning of this system could also be used for intrinsically regulating organ size.

MATERIALS AND METHODS

Mouse lines and harvest of embryos

Myf5^{+nlacZ}, *Myod1^{+/-}*, *p21^{+/-}*, *p57^{+/-m}* (*p57* is an imprinted gene; we indicate maternal origin of the allele by a superscript *m*), *Pax3^{Cre/+}* and *Rbpj^{lox/+}* lines have been described previously (Deng et al., 1995; Engleka et al., 2005; Han et al., 2002; Rudnicki et al., 1992; Yan et al., 1997). For explant and ChIP experiments, C57BL/6J embryos were used (Janvier). For timed pregnancies, the morning when a vaginal plug was found was defined as embryonic day (E) 0.5. All experiments were performed on three independent embryos for each genotype.

Immunohistochemistry and X-Gal staining

Embryos and forelimbs were harvested and fixed for 2 h and for 20 min, respectively, in PBS/4% paraformaldehyde at 4°C. Cryoprotection was performed by equilibration in PBS/15% sucrose overnight at 4°C. Frozen sections were permeabilized in PBS/0.1% Triton X-100, blocked in PBS/2% bovine serum albumin for 1 h at room temperature, then immunolabeled with primary antibodies overnight at 4°C. For X-Gal staining, embryos were collected in PBS, fixed 20 min in PBS/4% paraformaldehyde at room temperature and incubated in X-Gal solution (Life Technologies) overnight at 37°C on a rotary shaker.

Antibodies

The following antibodies were used: mouse anti- β -galactosidase 1/500 (Promega, Z378), mouse anti-MyoD1 5.8A 1/200 (DAKO, M3512), mouse anti-Myog F5D 1/200 (DSHB, F5D), mouse anti-p21 1/100 (BD Pharmingen, 556431), mouse anti-p57 1/100 (Santa Cruz, sc-56341), mouse anti-Pax7-c 1/100 (DSHB, Pax7-c), mouse anti-Pax3-c 1/100 (DSHB, Pax3-c), rabbit anti- β -galactosidase 1/1000 (Life Technologies, A-11132), rabbit anti-MyoD1 M318 1/100 (Santa Cruz, sc-760), rabbit anti-Myf5 C20 1/500 (Santa Cruz, sc-302), rabbit anti-p57 H91 1/100 (Santa Cruz, sc-8298), rabbit anti-phospho-histone 3 Ser10 1/1000 (Cell Signaling, 9701), goat anti-p57 M20 1/50 (Santa Cruz, sc-1039) and goat anti-Pax3 1/100 (Santa Cruz, sc-34916). Phalloidin (649 nm) 1/500 was from Life Technologies. Secondary antibodies were coupled to Alexa Fluor 488 1/250, 594 1/1000 (Life Technologies) or 649 1/250 (Jackson ImmunoResearch).

Explant and cell culture

Forelimbs from E11.5 wild-type embryos were cultured in 12-well plates in BGJb medium (Life Technologies), without serum, with 200 μ g/ml ascorbic acid (Sigma) and 100 μ g/ml penicillin/streptomycin (Life Technologies). For Notch inhibition, forelimbs were immediately treated with 20 μ M N-[N-(3,5-difluorophenacetyl)-L-alanyl]-S-phenylglycine t-butyl ester (DAPT; Sigma) or DMSO carrier (Sigma) for 28 h. Treated and control forelimbs originating from the same embryo were compared in each experiment. C2C12 and HEK293 cells were cultured in proliferating medium comprising DMEM with 10% fetal bovine serum and 100 μ g/ml penicillin/streptomycin (Life Technologies).

Plasmid construct for transgenesis

The *p57* muscle regulatory element (*p57MRE*) (chr7: 150,587,238-150,587,924) was isolated by PCR. For cloning convenience, *EagI* restriction sites were added to the forward and reverse primers used for amplification: forward, 5'-AAGCGGCCGACCCAGTTTGGCCAGTGTAG-3'; reverse, 5'-AACGGCCGCCAGGTAAGACACCCAGAG-3'. After *EagI* digestion, the 686 bp fragment was cloned, respecting its genomic orientation, into the *NotI* site of *ptkmlacZ(-)* plasmid (Hadchouel et al., 2000) (*tk*, thymidine kinase). The *p57MRE-tk-nlacZ* fragment was released by *SacII/XhoI* digestion and gel purified using the Nucleobond plasmid purification kit (Macherey-Nagel) before injection into pronuclei.

β -galactosidase assay

Hey1, *Hes1* cDNAs [gifts from S. Tajbakhsh (Pasteur Institute, Paris, France) and R. Kageyama (Institute for Virus Research, Kyoto University, Japan), respectively] and *MyoD1* cDNA were cloned into the pCIG plasmid (Megason and McMahon, 2002). C2C12 cells were transfected with a total of 1.2 μ g DNA using Lipofectamine LTX plus reagent (Life Technologies). Fixed concentrations of *p57MRE-tk-nlacZ* or *p57MREAE-Boxes-tk-nlacZ* (0.6 μ g), or pCIG-MyoD1 (0.15 μ g) were used. For pCIG-Hes1 and pCIG-*Hey1*, 0.15 or 0.3 μ g was used. Each sample was co-transfected with 0.1 μ g *tk*-Luciferase reporter for sample-to-sample normalization. Forty-eight hours after transfection, the cells were collected and the proteins were extracted and assayed for β -galactosidase activity (β -Gal assay Kit K1455-01, Life Technologies) and for luciferase activity (Luciferase assay system E1500, Promega) to normalize transfection variation. Measurements were made at least in triplicate and expressed as the mean (with s.e.m.) of the amount of β -galactosidase substrate (ONPG) hydrolyzed.

Reverse transcription and quantitative PCR (qPCR)

Total RNA from embryo forelimbs was extracted using the RNeasy mini kit (Qiagen). 1 μ g RNA was used to generate cDNA using the Superscript II reverse transcriptase kit (Life Technologies). qPCR was performed using the Lightcycler 480 SYBR Green mix (Roche) and Lightcycler 480 II (Roche). RT-qPCR on FACS-isolated cells was performed using the Superscript III cell direct cDNA kit (Life Technologies). qPCR results are expressed as relative ratios of target cDNA to *Hprt*. The following oligonucleotides were used (5'-3'; forward and reverse): *Hes1*, ACACCGACAAACCAAGAC and AATGCCGGGAGCTATCTTTC;

Hey1, CACCTGAAAATGCTGCACAC and ATGCTCAGATAACGGG-CAAC; *MyoD1*, GGCTACGACACCGCTACTA and GAGATGCGCT-CCACTATGCT; *Pax7*, AGGCCTCGAGAGGACCCAC and CTGA-ACCAGACCTGGACGCG.

Chromatin immunoprecipitation (ChIP)

MyoD1 ChIP-seq has been described in detail (Cao et al., 2010). For qPCR ChIP experiments, forelimbs from E11.5 embryos were frozen in liquid nitrogen and processed for ChIP according to the manufacturer's protocol (Active motif). 150 μ g of chromatin was used for each experiment. 2 g of a rabbit anti-MyoD1 M318 (Santa Cruz, sc-760) and 2 g of a goat anti-Hes1 (Santa Cruz, sc-13844) were used; 2 g of a rabbit anti- β -galactosidase (Life Technologies, A-11132) or 2 g of a goat anti- β -galactosidase (Santa Cruz, sc-19119) were used as the corresponding IgG negative control. The precipitated and input chromatin were analyzed by qPCR using *p57MRE* primers (forward, 5'-ATGTGCACACAG-CTCAGAGG-3'; reverse, 5'-GGAAGGATGGAGGGCTTTAC-3') with albumin primers as negative control (forward, 5'-GGGACGAGATGGT-ACTTTGTG-3'; reverse, 5'-GATCAGTCCAACTTCTTTCTG-3').

For ChIP on transfected cells, HEK293 cells were transfected with a total of 7.5 μ g DNA using FuGENE6 (Promega). A mutant *p57MRE* sequence, *p57MRE Δ E-Boxes*, was synthesized (GeneART) in which all putative E-boxes were mutated according to Iso et al. (2003). Fixed concentrations of *p57MRE-tk-nlacZ* or *p57MRE Δ E-Boxes-tk-nlacZ* (4 μ g) were used together with either pCig-MyoD1 or pCig-Hes1 (2 μ g). After 48 h, chromatin was extracted and processed as above; 100 μ g chromatin was used for each experiment. For ChIP, 2 μ g normal mouse (Santa Cruz) and goat (Santa Cruz) IgG were used for negative controls for the MyoD1 and Hes1 antibodies mentioned above. Results are expressed as fold change compared with IgG control.

Statistical test

Immunostainings were performed on at least three embryos of each genotype. Quantifications were performed using images of all muscle masses present in an embryo section (6-8 sections per slide, 2-3 frames per masse). All qPCR experiments were performed at least three times independently. Cell counting and qPCR results were analyzed by Mann-Whitney or Student's *t*-test. In Fig. 3D and Fig. 4F,G, quantifications were analyzed by ANOVA. In Fig. 5D, quantifications were analyzed by a chi-square test.

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Competing interests

The authors declare no competing financial interests.

Author contributions

A.Z. designed and performed experiments, analyzed data and wrote the paper. S.H. and F.A. designed and performed experiments, analyzed data and edited the manuscript. T.C., D.M. and P.M. designed and performed experiments, analyzed data. Z.Y. and Y.C. provided data on MyoD1 ChIP-seq. D.B. provided *Rbpj* mutant embryos. C.B. analyzed data and edited the manuscript. F.R. oversaw the entire project, designed experiments, analyzed data and wrote the paper.

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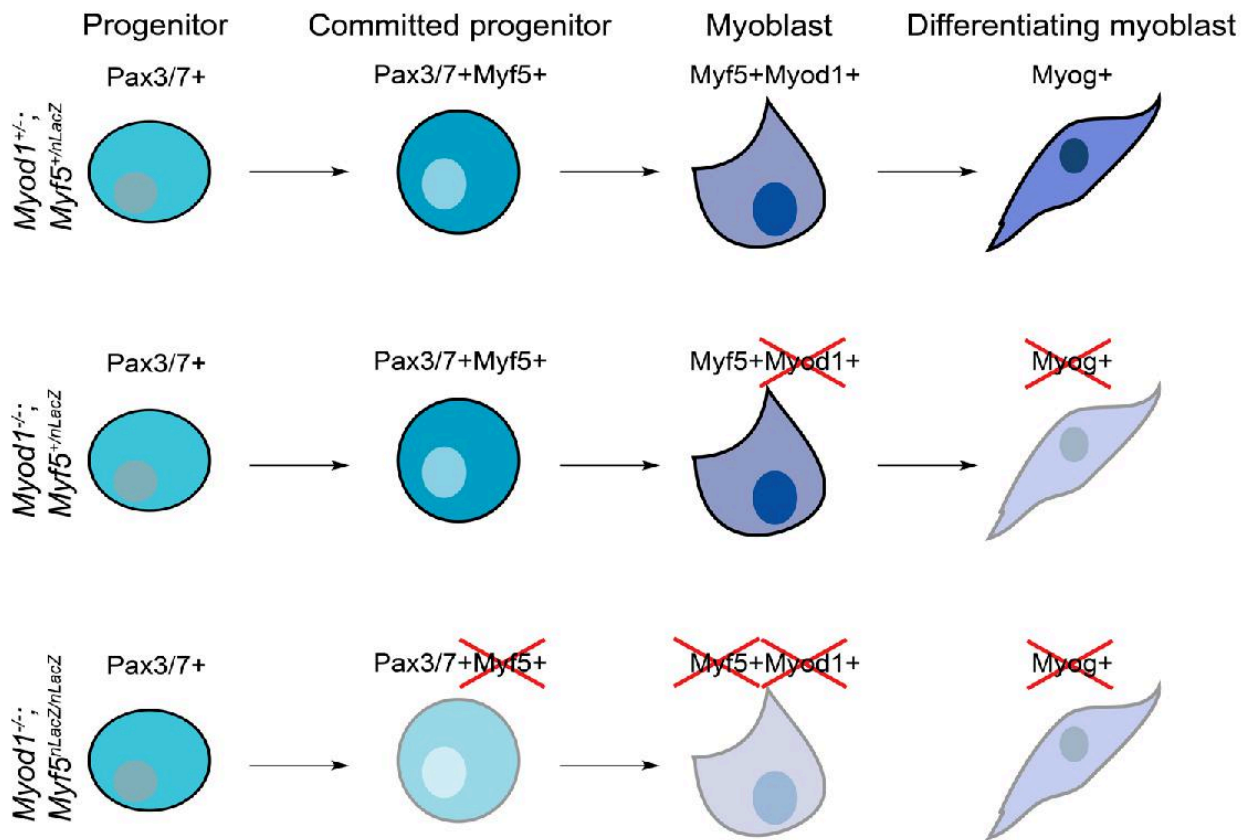
Supplementary material

Supplementary material available online at
<http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.110155/-DC1>

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Supplementary Figure 1. Stages of myogenic differentiation impairment using the different *MyoD1: Myf5* mutant compounds.

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      .           .           .           .           .
1-  ACCCAGTTTGCCCAAGTGTAGAGTGCCCAAAGCCCTCAGAAAAGACAAAGAAAGAGACCT
      .           .           .           .           .
61-  TTCAGACTAGAAAACAGAAGCAGGATACTTTGACCATATGTCACCATGTGCCTGGGCCACT
      .           .           .           .           .
121- TGGCTTGGCTACCCAAGCAAGGGCCCAAGAGAGTGCCCCCTTTGTCCCTTCTTCACAAAT
      .           .           .           .           .
181- GCTGTGTATCTACTCCACCACATGTGCACACAGCTCAGAGGCTCTTGCCTTTGGAATGCAG
      .           .           .           .           .
241- TCCACCCACCTCCAGTTTTCCCCTAAGCAGTTCAGCTGCTAAGCCCTGAACTCTCCCAC
      .           .           .           .           .
301- CTGCCGGGTCCTTAGCACCAGCTGCCCAGAGCTCCTGGACCCAGGCACCAGCTGGATGACC
      .           .           .           .           .
361- ACCCCTCCGAGCTGGCTAGGCCTGCTGTCAGCTGCCAGTAAAGCCCTCCATCCTTCCCAA
      .           .           .           .           .
421- CAGGTGTCCCCACAACTGCTGGGGGGGTTGCTTAGCTAGGCAGCTATTCCACTGACCCAG
      .           .           .           .           .
481- GCCTGCAGAATGATGTCACACTTACCCTGTGGAAGGAGGTAGCCCGAGGGATCTGACAG
      .           .           .           .           .
541- ACTCCACGGAAGTTATTGGTTCTAGAGAAACGGCACACCAGTCATACTCAGCTCCAGCTG
      .           .           .           .           .
601- GGTGGGTCAGACCTATCCTTTTAAAGACCAACTGGACATTGGCCAGAGAAGTTTGGCCC
      .           .           .           .           .
661- TGAAGGGTCTGGGGTGTCTTTACCTGG -687

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Supplementary Figure 2. Genomic sequence of p57MRE enhancer cloned in p57MRE-tk-nlacZ reporter.

Putative E-boxes are underlined with grey background. To generate *p57MRE Δ E-Boxes* mutant construct, every CANNTG motif has been replaced by CGNNAG.

Preface

Recent studies from us and others showed that the CDKI p57 participates in cell cycle control during muscle development. The perinatal lethality of p57 mutants precludes postnatal functional analyses to elucidate its role in adult muscle and, thus, we generated a conditional knock-out allele. We inserted loxP sites encompassing the coding region of p57 to allow tissue-specific and adult-specific excision of p57 using the loxP/Cre recombination system. By using the ubiquitous PGK-Cre, we uniformly ablated p57 and showed that the resulting mice exhibited the same defects as previously reported for the null allele. Specifically, we observed perinatal lethality and growth defects in several tissues, including the palate, the gastro-intestinal tract, the abdominal wall, and the skeletal system. Furthermore, in the new p57 allele we inserted an IRES-linked β -galactosidase reporter, to allow tracking of p57-expressing cells. We confirmed that the reporter faithfully recapitulates the expression profile of p57 in embryonic and adult tissues. Hence, we here describe a new genetic tool for expression and functional analyses of p57.

A *p57* conditional mutant allele that allows tracking of *p57*-expressing cells

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ABSTRACT

p57^{Kip2} (*p57*) is a maternally expressed imprinted gene regulating growth arrest which belongs to the CIP/KIP family of Cyclin-Dependent Kinase Inhibitors. While initially identified as a cell cycle arrest protein through inhibition of cyclin and cyclin-dependent kinase complexes, *p57* activity has also been linked to differentiation, apoptosis, and senescence. In addition, *p57* has recently been shown to be involved in tumorigenesis and cell fate decisions in stem cells. Yet, *p57* function in adult tissues remains poorly characterized due to the perinatal lethality of *p57* knock-out mice. In order to analyze *p57* tissue-specific activity, we generated a conditional mouse line (*p57^{FL-ILZ/+}*) by flanking the coding exons 2-3 by LoxP sites. In order to track *p57*-expressing or mutant cells, the *p57^{FL-ILZ}* allele also contains an IRES-linked β -galactosidase reporter inserted in the 3' UTR of the gene. Here, we show that the β -galactosidase reporter expression pattern recapitulates *p57* tissue-specificity during development and in postnatal mice. Furthermore, we crossed the *p57^{FL-ILZ/+}* mice with *PGK-Cre* mice to generate *p57^{CKO-ILZ/+}* animals with ubiquitous loss of *p57*. *p57^{CKO-ILZ/+}* mice display developmental phenotypes analogous to previously described *p57* knock-outs. Thus, *p57^{FL-ILZ/+}* is a new genetic tool allowing expression and functional conditional analyses of *p57*.

INTRODUCTION

Tightly controlling cell cycle is imperative for development and tissue homeostasis of complex organisms. Cell cycle progression is primarily achieved via an array of Cyclin-Dependent Kinases (CDKs) and their activating subunits named cyclins (Malumbres, 2014). Growth arrest is mediated by the interaction of these CDK-cyclin complexes with Cyclin-Dependent Kinase Inhibitors (CDKIs). CDKIs are subdivided into two families based on their structural homology and specificity of action (Borriello et al., 2011): INK4 [including p15^{Ink4b} (p15), p16^{Ink4a} (p16), p18^{Ink4c} (p18), p19^{Ink4d} (p19)] and Cip/Kip [including p21^{Cip1} (p21), p27^{Kip1} (p27), p57^{Kip2} (p57)]. The latter counteracts cell cycle progression more broadly by inhibiting all CDK-cyclin complexes, in addition to activities outside of cell cycle regulation (Borriello et al., 2011).

p57 is the most recently discovered and least studied member of the Cip/Kip family of CDKIs. It is distinguished from its siblings, p21 and p27, by its unique structure (including a divergent central domain), distinct distribution pattern, and imprinting status. It is emerging as a multifaceted protein involved in several cellular processes, such as proliferation, differentiation, senescence, apoptosis, and motility (reviewed in Pateras et al., 2009; Rossi & Antonangeli, 2015). The gross developmental defects and perinatal lethality of mice that lack or overexpress p57 highlight its critical role during organogenesis (Yan et al., 1997; Zhang et al., 1997; Andrews et al., 2007). In addition to its role in growth arrest during differentiation, p57 plays distinct functions in several organs, including quiescence acquisition in hematopoietic and neural stem cells (Matsumoto et al., 2011; Zou et al., 2011; Furutachi et al., 2013). Its impact on human pathology is illustrated by p57 implication in carcinogenesis, which was initially hypothesized due to its capacity to prevent proliferation. Remarkably, p57 is downregulated in several malignancies and loss of p57 is associated with poor patient prognosis. Noticeably, p57 is mutated or inactivated in the cancer-predisposing disorder Beckwith-Wiedemann syndrome (reviewed in Guo et al., 2010; Borriello et al., 2011).

In order to perform tissue-specific functional studies coupled with analysis of p57 expression, we generated a *p57* allele (*p57*^{FL-ILZ/+}), allowing LacZ-dependent visualization and Cre-inducible ablation of p57. Moreover, in contrast to a previously reported conditional *p57* mutant mice (Matsumoto et al., 2011), *p57*^{FL-ILZ} allele allows complete ablation of the coding region, while preserving potential regulatory elements of the surrounding sequences. Here, we describe the generation of the *p57*^{FL-ILZ/+} mouse line and demonstrate that the LacZ reporter recapitulates the p57 expression profile during development and in the adult. We also show the

efficiency of Cre-dependent ubiquitous removal of p57 function and the subsequent recapitulation of knock-out phenotypes.

RESULTS AND DISCUSSION

We generated a mouse line ($p57^{FL-N-ILZ/+}$), in which the entire p57 coding region (exons 2 and 3) is flanked by LoxP sites, to allow spatial and temporal conditional ablation of p57 by Cre recombinase (Fig.1A-D). A *FRT-neo* cassette was introduced for selection and subsequently excised using mice containing FLP recombinase activity, generating the $p57^{FL-ILZ}$ allele (Fig. 1C). The $p57^{FL-ILZ}$ allele also contains a downstream *IRES-nLacZ* gene inserted in exon 4, encoding a nuclear-localized β -galactosidase (Fig. 1C). To evaluate the LacZ expression profile, we used $p57^{FL-ILZ/+}$ heterozygotes with maternal inheritance of the targeted allele, since p57 is subjected to genomic imprinting and the paternal allele is silenced (Hatada & Mukai, 1995). Co-immunostaining experiments demonstrated that p57 and β -galactosidase co-localized in forming bone tissues during development and in postnatal kidney (Fig. 1E), two sites where p57 expression was previously documented (Yan et al., 1997; Zhang et al., 1997; Westbury et al., 2001).

Next, we performed an analysis of the *nLacZ* reporter expression in embryonic (Fig. 2) and adult (Fig. 3) tissues. During development, *p57* is widely expressed during organogenesis with high levels peaking at key differentiation steps of specific organs (Matsuoka et al., 1995; Westbury et al., 2001). Specifically, we detected high reporter expression in the developing musculoskeletal system, neural tissues and parenchymal organs of X-Gal-stained whole mount embryos (Fig. 2). At early stages (E10.5-E11.5), we observed a strong expression in brain, heart, and limbs as well as in the somites (Fig. 2A, B). Sensory organs, including the developing ears, lens and nasal processes, are also stained (Fig. 2A, B). Remarkably, the reporter pattern thoroughly mirrors p57 expression visualized by *in situ* hybridization (ISH) (Fig. 2A', B'). In the developing limbs the reporter and p57 transcripts are initially concentrated in central masses, which contain differentiating muscle and skeletal cells (Fig. 2A-B'). Strong reporter expression extends to the forming digits at later stages (Fig. 2C, D). From E13.5 the staining becomes more widespread and less intense, covering almost the entire exterior of the embryos (Fig. 2D-H). Sections of E13.5 $p57^{FL-ILZ/+}$ embryos revealed numerous X-Gal-positive cells in the heart, lung, body wall, musculoskeletal parts of the limb, and neural tissues, including the neural tube, dorsal root ganglia and thoracic vertebral body (Fig. 2I), consistent with previous reports of p57 expression (Matsuoka et al., 1995; Yan et al., 1997; Zhang et al., 1997; Westbury et al., 2001). β -galactosidase-positive cells likely correspond to cell cycle exit linked with terminal differentiation.

In contrast to embryonic tissues, p57-LacZ expression was more restricted in adult tissues. Strikingly, LacZ expression was limited to small regions or cell populations of specific organs, including the kidney, testis, skeletal muscle, heart, brain, intestine, and lung (Fig. 3), in agreement with previous studies (Matsuoka et al., 1995; Zhang et al., 1997). Specifically, we observed robust expression in the glomeruli of the kidney (Fig. 3A), where p57 is expressed at essential stages of glomerulogenesis and during the establishment of the mature phenotype (Nagata et al., 1998; Hiromura et al., 2001). β -galactosidase is also detected in the seminiferous tubules of the testis, where p57 was previously suggested to promote the meiotic progression of early spermatocytes along with cell cycle arrest and differentiation of spermatids (Kim et al., 2006). In general, β -galactosidase signal is highly localized and is, for instance, not detected in the spleen or the liver (data not shown), in line with previous reports (Matsuoka et al., 1995; Yanagida et al., 2015). Strikingly, β -galactosidase expression was also observed in organs where p57 expression was not previously described, such as the salivary gland (Fig. 3D).

We further wanted to evaluate the ability to recombine the $p57^{FL-ILZ}$ allele (via the LoxP sites) to generate p57 loss of function animals ($p57^{cko-ILZ}$). To this end, we used *PGK-Cre* mice, which induce ubiquitous recombination (Lallemand et al., 1998). We focused our study on heterozygotes with a maternally transmitted p57-null allele (*PGK-Cre;p57^{CKO-ILZ} (m)/+*), since the paternal allele is transcriptionally repressed (Hatada & Mukai, 1995), and previous studies revealed indistinguishable phenotypes between heterozygote and homozygote knock-outs (Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000). As expected, p57 protein was no longer detected in all tissues examined from recombined *PGK-Cre;p57^{CKO-ILZ} (m)/+* fetuses (Fig. 4). As we crossed the $p57^{CKO-ILZ}$ allele with a *PGK-Cre* allele inducing early and ubiquitous recombination, we expected to phenocopy previously reported p57 knock-out mice. Indeed, as previously reported (Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000), we observed a prominent perinatal lethality of mutants (Fig. 5A), associated with a marked size reduction of the fetus (Fig. 5B, C), similarly to the knock-out mice reported by Nakayama and colleagues (Takahashi et al., 2000). Perinatal death of p57 mutants has been attributed to cleft palate that leaves the oral and nasal cavities connected along with difficulties in suckling/breathing and an abnormal gastrointestinal tract (Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000). Likewise, in $p57^{CKO-ILZ} (m)/+$ mutants the two palatal shelves failed to grow and meet in the midline (Fig. 5D), associated with inflated stomach and intestine in a number of cases (Fig. 5E). We also observed body wall dysplasia associated with abdominal slit at birth or with thin body wall during development (Fig. 5F), in agreement with previous studies on mice lacking p57 (Zhang et al., 1997). Finally, we noticed embryonic and perinatal skeletal deformities evidenced by sternum

fusion defects, limb shortening and thickening, and delayed ossification (Fig. 5G), consistent with the skeletal defects of previous reports (Yan et al., 1997; Zhang et al., 1997; Susaki et al., 2009). Thus, we were able to successfully recombine the $p57^{FL-ILZ/+}$ allele, ablating p57 expression and reproducing the spectrum of p57 knock-out phenotypes. However, β -galactosidase activity is hardly detected in recombined $p57^{cKO-ILZ (m)/+}$ mutants embryos, likely due to destabilized mRNA in the deleted allele (Fig. S1).

In conclusion, the $p57^{FL-ILZ/+}$ allele provides a genetic tool that allows conditional ablation of p57 by removal of the complete coding region, encoded by exons 2-3, in a spatiotemporal manner using tissue-specific and/or inducible Cre lines. Moreover, the $p57^{FL-ILZ/+}$ allele provides a faithful reporter to track the expression pattern of p57, including the possibility to follow epigenetic effects and imprinting changes, and identify new sites of expression that may require p57 activity.

MATERIALS AND METHODS

Generation of cKO mice

A 61.8kb BAC clone from a mouse 129Sv (129S7AB2.2; Adams et al., 2005) genomic library centered around the p57 locus (bMQ-38L12; SourceBioscience #WTSIB741L1238Q) has been chosen to carry out locus modifications using λ -red recombination (Lee et al., 2001). A LoxP site has first been inserted into p57 5'UTR in exon 1 and then a *LoxP-FRT-PGK-NEO-BGHpA-FRT-IRESnlLacZ* cassette into p57 3'UTR in exon 4. Finally, the 19.5kb cKO targeting cassette containing recombined p57 locus, 5.8kb of genomic sequence upstream of exon 1, and 5kb downstream of exon 4, has been extracted from the BAC clone by gap repair.

BAC targeting vectors construction

The first BAC targeting construct was generated by PCR amplification of two homology arms, using bMQ-38L12 as template DNA. The first arm upstream of ATG contained 98bp of 5' exon 1; the second 21bp of 3' exon 1, intron 1, and 173bp of exon 2 including beginning of CDS (pair 1: 5'-CATTATGCTAATCGTGAGGAGGC-3' and 5'-CTTAAGTCTGGATCGCTTGTCTCTG-3', 203bp; pair 2: 5'-CTTAAGAGCCGTCCATCACCAATCAG-3' and 5'-AAGTTGAAGTCCCAGCGGTTGTAGAC-3', 380bp). Amplicons were subcloned in *pGEMTeasy* (Promega, # A3610) with a newly introduced AflIII site in-between. Then a *LoxP-kanamycin-LoxP* cassette was inserted blunt in AflIII. Similarly, homology arms for the second BAC targeting construct were amplified from the BAC DNA and

subcloned in *pGEMTeasy* with a HindIII site in-between. The first arm contains 119bp at the end of exon 3, including STOP codon, intron 3 and 216bp of 5' exon 4. The second homology arm contains the following 361bp of exon 4 (pair3: 5'-GAGAACTGCGCAGGAGAACAAG-3' and 5'-AAGCTTTACACCTTGGGACCAGC-3', 424bp; pair 4: 5'-AAGCTTTAAATCATTATGTAAAATGTTTAATCTCTACTCG-3' and 5'-GCAATCTAATGAAGTGGGGGAC-3', 361bp). In parallel, a 1.9kb EcoRV-SacII *FRT-PGK-Neo-BGHpA-FRT-LoxP* fragment from *pL451* (Liu et al., 2003) was cloned blunt into Sall of *pIRES-nLacZ* (Relaix et al., 2003). Then, the 5.9kb NotI fragment-containing *LoxP-FRT-PGK-Neo-BGHpA-FRT-IRESnLacZ* was inserted blunt into the homology arms carrying vector in HindIII. The complete 1.8kb and 6.8kb BAC targeting linear cassettes directed against exon 1 and exon 4, respectively, were released by NotI digestion and gel purified (Macherey-Nagel, NucleoSpin® Gel and PCR Clean-up, #740609).

The gap repair plasmid was built from PCR generated homology arms (5.9kb upstream of exon 1 and 5kb downstream of exon 4) using bMQ-38L12 as template, assembled in *pGEMTeasy* with EcoRV created in central position (pair 5: 5'-CTACTTACCAGCCTCTGAGGA-3' and 5'-GATATCCAAGCCAGACCTCCCTGC-3', 529bp; pair 6: 5'-GATATCCTGGTTTCCCACCTGTGATG-3' and 5'-GTTGTGATCCACACCTGACTCC-3', 520bp). The full insert was then excised with EcoRI and cloned blunt in *pBluescriptSK-* (*pBSK-*, Stratagene) to give *p57-gap-repair* plasmid. An hygromycin resistance cassette was PCR amplified from *pSKT.HygroBACe3.6Lox511* (kindly given by Dr. J. Hadchouel) with SacI introduced at each extremities and cloned into *p57-gap-repair* SacI site to give the gap repair plasmid *p57gaprepairHygro(-)*.

All cloning PCR were carried out using Advantage-2 (Clontech, #639201) or PHUSION DNA polymerases (Thermoscientific, #F530L). All DNA digestions were performed using NEB enzymes.

BAC recombinations

BAC recombinations were performed by first electroporation of bMQ-38L12 BAC DNA (BTX ECM399 electroporator; 0.1cm cuvettes, 1.3kV, 100ng DNA) into the *E. coli* Cre-inducible SW106 strain and selection on chloramphenicol (cam^R ; 10 $\mu\text{g}/\text{ml}$) LB plates. DNA from five transformants was analyzed by restriction profiling using NotI + SpeI on 0.6% large agarose gels ran overnight. One clone, L12A, with the expected restriction profile, satisfactorily completed a deeper restriction profiling with EagI, XhoI+NruI, Aat2, AgeI+NotI, HpaI+NotI, RsrII+NotI. To perform DNA recombination, 500 μl of this clone's overnight culture were used to inoculate 25ml of LB+ cam^R , further grown for 135 minutes at 32°C. Then, the λ -red recombinase was induced by 15 minute-incubation of the culture at 42°C. Bacteria were made electro-competent by two washes

in 10ml cold dH₂O and gentle resuspension on ice. The 1.8kb purified exon 1 targeting cassette was introduced by electroporation (0.2cm cuvettes, 1.75kV, 20ng DNA), and transformants were selected on kanamycin (kan^R; 25µg/ml) plates. Extracted DNA from four kan^R clones, L12AK1 to 4, and analyzed with NotI+SpeI was effectively recombined. To pop-out the kanamycin cassette located between the two LoxP sites and leave one LoxP site at the wanted location, Cre recombinase expression was induced in L12AK1 clone using 0.1% L(+) arabinose (Sigma-Aldrich, #A3256) in liquid culture for 2 hours. Bacteria were then plated on cam^R LB plates. Three clones were positively tested for loss of kanamycin resistance, BglII restriction profiling and PCR across deletion on BAC extracted DNA, and one of them was then chosen for further modifications. L12AK#NE1 DNA was electroporated in *E. coli* Flp-inducible SW105 strain as described above, and DNA from two clones growing on cam^R, L12AK#OF1 and OF2, were subjected to BglII restriction profiling to assess integrity following the transformation process. A L12AK#OF1 culture was induced for λ-red recombinase expression and made electro-competent and, then, it was transformed with the 6.8kb purified exon 4 targeting cassette (0.2cm cuvettes, 1.75kV, 25ng). DNA from three kan^R transformants, L12A#FC1-3, was extracted and tested for NotI+RsrII restriction profiling, showing successful recombination for clones L12A#FC1 and 2, further confirmed by PCR at 5' and 3' extremities of the insertion, and deep restriction profiling of L12A#FC1 DNA with NheI, NdeI, SpeI, NruI, AgeI, EcoRI, and AseI+NotI. To assess FRT sites functionality (*FRT-PGK-Neo-pA-FRT* cassette), Flp recombinase was induced in L12A#FC1 bacterial clone with arabinose as described above. DNA from five clones, which have lost kanamycin resistance, was extracted and subjected to AatII profiling and showed that *PGK-Neo-pA* sequence was missing and FRT sites were functional. Similarly, LoxP sites functionality was tested. DNA from L12A#FC1 was electroporated in SW106, then Cre recombinase was induced with arabinose, and DNA from five cam^R-kanamycin sensitive clones was extracted and profiled with AatII. Again, the analysis showed that the sequence from the 3' end of exon 1 to 5' of exon 4 was properly floxed-out, thus proving that LoxP sites are functional.

Gap repair and final cKO construct

Bacterial clone L12A#FC1 was made electro-competent, λ-Red recombinase induced as described above, and transformed with 10ng of circular *p57gaprepairHygro(-)*, and then selected on hygromycin (hyg^R; 125µg/ml) LB plates. 22 out of the 24 DNA extracted (Macherey-Nagel, NucleoSpin[®] plasmid #740588) from hyg^R clones showed that the expected 19.5kb insert was present following gap repair. To provide a negative selectable marker in ES cells, a 1.6kb cassette containing the PGK promoter driving the expression of the Diphtheria toxin A subunit (*Pgk-DTApA*) was PCR amplified from a *pSK-pDTA* plasmid (Relaix et al., 2003) with Sall and XhoI

introduced in 5' and 3', respectively, and subcloned in one of the gap repair clones into Sall site. Resulting transformants were screened for *PGK-DTA-pA* in reverse orientation.

Generation of the p57^{FL-N-ILZ/+} mouse model

The targeting vector was electroporated in CK35 embryonic stem (ES) cells of 129S2 genetic background (Kress et al. 1998). 30 µg of a Pme1-linearized construct were electroporated (240 V, 500 µF - Gene Pulser II, Biorad) into 5x10⁷ cells in 0.8ml PBS. Cells were then seeded on a layer of embryonic fibroblast feeders in DMEM (Life Technologies, #31966047) supplemented with 15% ES-culture-tested FBS (Life Technologies #10270-106), non-essential amino acids (Life Technologies, #11140-035), Penicillin-Streptomycin (Life Technologies, #15140-114), 10⁻⁴ M β-mercaptoethanol (Sigma-Aldrich, #M6250); 1000 UI LIF (Merck-Millipore, #ESG1107). Geneticin (300 µg/ml) selection started on day 2 (Life Technologies, #10131-019) and picking was done on day 9.

ES cells were preselected by two 3' external PCRs (pair 7: 5'-CACTGCATTCTAGTTGTGGTTTGTCCAAAC-3' and 5'-GGGACAGTTGCTAGCTGTGA -3', 5.6kb product; pair 8: 5'-CCCCCTGAACCTGAAACATA-3' and 5'-GGTTCGCTACCATTACCAGTTGGTCTGG-3', 6kb). The first PCR product was digested with BglII into two bands of 3.3kb and 2.3kb. Clones were then screened for recombination events by Southern blot analysis using HindII digestion and a 5'-flanking probe (wt allele: 10.8kb, KO: 8.6kb). The positive clones were further verified by SacI digestion and LacZ internal probes. Targeted ES cells were recovered with 9.3% frequency. Positive clones were karyotyped and euploid clones were used to produce chimeras and germline transmission.

The resulting mice are kept in a mixed 129S2 X C57Bl/6N background and will be available to the research community upon manuscript acceptance.

Mouse lines

FLP and *PGK-Cre* transgenic mice have been previously described (Lallemand et al., 1998; Rodriguez et al., 2000). For *in situ* hybridization experiments C57Bl/6N (Janvier) embryos were used.

Embryo harvest

Female and male mice were crossed overnight and separated the following morning after macroscopical vaginal examination for plug presence. In case of plug, that morning was defined as embryonic day (E) 0.5. At least three embryos per genotype and per time point were analyzed.

Immunohistochemistry and X-Gal staining

Whole mount samples: Embryos and adult organs were harvested and fixed for 30 minutes to 2 hours (depending on the stage) in 4% paraformaldehyde/PBS at 4°C. Incubation with X-Gal solution (0.4 mg/ml X-Gal in 2 mM MgCl₂, 0.1% Tween20, 1X PBS, 4 mM K₄Fe(CN)₆, and 4 mM K₃Fe(CN)₆; Sigma-Aldrich) was performed at 37°C on a rotary shaker. Incubation duration varied from 2 hours to overnight depending on the age. Specimens were placed in 4% paraformaldehyde/PBS at 4°C for long-term storage.

Sections: Embryos and newborns were harvested and fixed for 25 minutes to 2 hours (depending on the age) in 4% paraformaldehyde/PBS at 4°C, followed by overnight incubation in 15% sucrose/PBS for cryoprotection before freezing. Organs of adult mice were frozen fresh and fixation was performed directly on sections with 4% paraformaldehyde/PBS for 20 minutes at room temperature. For X-Gal staining, frozen sections were incubated with X-Gal solution as previously mentioned overnight at 37°C. Adult tissues were counterstained with eosin (ThermoFisher Scientific, #6766007) for 30 seconds to highlight their structure. For hematoxylin-eosin staining, frozen sections were incubated with hematoxylin (Sigma-Aldrich, # MHS32) for 11 minutes to stain nuclei, counterstained with eosin for 30 seconds to highlight cytoplasm, and dehydrated with passages through increasing concentrations of ethanol (30%, 50%, 70%, 85%, 95%, 100%) and two washes in xylene. For immunofluorescence, frozen sections were permeabilized in 0.1% Triton X-100/PBS and blocked in 1% bovine serum albumin/PBS at room temperature. Immunolabeling was performed overnight at 4°C for primary antibodies and 1 hour at room temperature for secondary antibodies. Nuclei were counterstained with DAPI. The following antibodies were used: mouse anti-β-galactosidase 1:200 (Promega, #Z378), rabbit anti-p57 1:150 (Santa Cruz, #sc-8298), Fluorescein (FITC)-AffiniPure goat anti-mouse IgG2a secondary antibody (Jackson ImmunoResearch, #115095206), and AlexaFluor594-conjugated F(ab')₂-goat anti-rabbit IgG (H+L) secondary antibody (ThermoFisher Scientific, #A11072).

At least three animals were analyzed in each experiment.

Skeleton staining

E14.5 and newborn mice were eviscerated and fixed in 100% ethanol for 48 hours. The carcasses were stained with alcian blue (0.15% in 4:1 ethanol / acetic acid; “alcian blue 8GX”, Sigma-Aldrich, # A3157) for 48 hours, postfixed in ethanol for 24 hours, cleared in 2% KOH, stained with alizarin red (0.15% in 0.5% KOH; “alizarin red S monohydrate”, Riedel-de Haën, #33010), and decolorized in increasing concentrations of glycerol (20%, 50%, 70%) for one week. Specimens were stored in 70% glycerol. At least three animals per genotype and per time point were analyzed.

Whole-Mount *in situ* hybridization

p57 (Genbank accession number: BC005412.1) riboprobe was synthesized using PstII/T7. Primers for probe amplification were designed with the NCBI-Primer BLAST tool and ordered from Eurogentec. The probes were amplified using as template one of the BAC repair clones described above. The PCR product was purified with the NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel, #740609) and inserted into the *pGEM-T Easy* vector (Promega, #A1360) according to the manufacturer’s instructions. Following overnight transformation of competent cells, colonies were analyzed by restriction digestion and verified by sequencing. 2µg of the plasmid was linearized with appropriate restrictions enzymes at 37°C for 30 minutes and purified with the NucleoSpin Gel and PCR Clean-up kit. Transcription and Digoxigenin-labeling were performed with the DIG RNA labeling Kit (SP6/T7) (Roche, #11175025910) and the product was purified with illustra MicroSpin G-50 Columns (GE Healthcare, #27533001) following the manufacturer’s protocol.

Harvested embryos were fixed in 4% paraformaldehyde/PBS overnight, dehydrated through 25%, 50%, 75% methanol/0.1% Tween20/PBS and stored in 100% methanol at -20°C. Before use, embryos were rehydrated with descending concentrations of methanol, treated with 10µg/ml proteinase-K in 0.1% Tween20/PBS (15 minutes at E10.5, 25 minutes at E11.5), post-fixed with 4% paraformaldehyde/0.1% glutaraldehyde/PBS for 20 minutes and incubated with 1µg/ml DIG-labeled RNA probe overnight at 68°C. After several steps of washes, the embryos were incubated with 1:2000 anti-DIG-AP Fab fragment (Roche, #11175025910) overnight at 4°C. Following three days of intense washes, the embryos were incubated with BM purple AP substrate (Roche, #11442074001) until color was developed to the desired extent. For long-term storage, specimens were kept in 4% paraformaldehyde/PBS at 4°C. Three animals per time point were analyzed.

Graphic editing

Graphs and representative photos were arranged in Figure format with the graphics editor Photoshop CS5. Uniform background was added in whole mount sample photos to cover shadows or background differences.

Statistical test

Body weight differences were compared with the Mann-Whitney U-test. Differences at $p < 0.05$ were considered significant.

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FIGURE LEGENDS

Figure 1. Generation of $p57^{FL-ILZ/+}$ mice

(A-D) Schematic representation of wild-type (wt) mouse $p57^{Kip2}$ (p57) allele (A), floxed p57 allele with *Neo*-selection cassette ($p57^{FL-N-ILZ}$; B), floxed p57 allele after removal of the *Neo*-selection cassette by FLP recombinase ($p57^{FL-ILZ}$; C), and recombined p57 allele after removal of the floxed fragment by Cre recombinase ($p57^{KO-ILZ}$; D). The four exons are represented by empty boxes, marked as E1-E4. Exons 2 and 3 contain the

coding regions (black box), with start codons (arrowheads) and stop codon (asterisk) indicated below the exons. LoxP and FRT sites are also shown as indicated. (E) Immunofluorescence analysis of β -galactosidase (green) and p57 (red) showing forming skeletal elements of the vertebral column (upper panel), and perinatal kidney section (lower panel). Heterozygote animals with maternally transmitted transgene were used, due to imprinting silencing of the paternally transmitted *p57* allele. Maternal inheritance is indicated with superscript (m). Nuclei were counterstained blue with DAPI. Scale bars: 50 μ m.

Figure 2. Time-course of *LacZ* expression in *p57^{FL-ILZ/+}* embryos

(A-H) Whole mount X-Gal staining on *p57^{FL-ILZ (m)/+}* embryos at E10.5 (A), E11.5 (B), E12.5 (C), E13.5 (D), E14.5 (E), E15.5 (F), E16.5 (G), E17.5 (H). Heterozygous animals with maternal inheritance of the transgene were used, due to imprinted silencing of the paternally transmitted *p57* allele. Maternal transmission is indicated by superscript (m). The skin was removed from E15.5 onwards, while small remaining pieces in the limb and tail tip resulted in lack of staining. (I-J) Whole mount *in situ* hybridization for *p57* at E10.5 (I) and E11.5 (J), wild type embryos are provided for comparison with X-Gal profiles. Scale bars: 2mm. (K) Representative X-Gal stained sections at trunk level of E13.5 *p57^{FL-ILZ (m)/+}* embryo. Scale bar: 400 μ m. bw: body wall; di: digits; E: embryonic day; ea: ear; fb: forebrain, FL: forelimb; h: heart; HL: hindlimb; le: lens; mb: midbrain, nt: neural tube; som: somites; sv: sensory vibrissae; tvb-nc: thoracic vertebral body and notochord.

Figure 3. *p57^{FL-ILZ/+}* adult mice display tissue-specific *LacZ* reporter expression.

(A-H) Heterozygote *p57^{FL-ILZ/+}* animals with maternal inheritance of the transgene were used to evaluate postnatal *LacZ* reporter in kidney (A-A'), testis (B-B'), skeletal muscle (C-C'), salivary gland (D-D'), heart (E), brain (F), intestine (G) and lung (H) of adult mice (8-12 weeks of age). (A-H) Whole mount X-Gal staining. Asterisks indicate the magnified part appearing within the dashed box on the lower right corner. Scale bars: 2mm in non-magnified whole mount organs; 100 μ m in magnified panels of whole mount organs. (A'-D') X-Gal and eosin-stained sections of the corresponding organ. Scale bars: 100 μ m.

Figure 4. *p57* protein loss in *p57^{CKO-ILZ/+}* mice

(A-D) X-Gal staining of *p57^{FL-ILZ(m)/+}* mice for thoracic vertebral body (A), forelimb cartilage (B), heart (C) and body wall (D). Asterisks indicate the magnified panel shown on the right of each photo. (E-L') Immunofluorescence for p57 (red) in control (E-H') and *p57^{CKO-ILZ(m)/+}* (I-L') E13.5 tissues, corresponding to (A-D). Nuclei were counterstained blue with DAPI in (E-H) and (I-L). Scale bars: 100 μ m.

Figure 5. Phenotypes of $p57^{cKO-ILZ/+}$ mice

(A-F) Comparison of $p57^{cKO-ILZ (m)/+}$ mutants to control littermates. Heterozygote $p57^{cKO-ILZ (m)/+}$ animals with maternal inheritance of the transgene were used due to epigenetic silencing of the paternally transmitted p57 allele. Maternal transmission is indicated by superscript (m). (A) *In utero* and perinatal frequencies (fraction of control per total animals) of the indicated genotypes are shown. (B-C) Average body weight (B) and representative animals (C) for the indicated genotypes at birth. Error bars represent SD. Asterisk indicates significance; $p < 0.05$. (D) Closed and cleft palate in control (upper panel) and $p57^{cKO-ILZ (m)/+}$ (lower panel) mice, respectively, at birth. Sections were stained with hematoxylin (nuclei) and eosin (cytoplasm). Scale bars: 2mm *in toto* samples; 100 μ m in sections. (E) Representative images of control gastrointestinal tract with milk in stomach and intestine (control, upper panel) and $p57^{cKO-ILZ (m)/+}$ inflated gastrointestinal tract (lower panel) at birth. Scale bar: 5mm. (F) Umbilical region of control (upper panel) and $p57^{cKO-ILZ (m)/+}$ mice (lower panel) at embryonic stages (E13.5; left panel) and at birth (P0; right panel). Sections were stained with hematoxylin and eosin. Scale bars: 2mm *in toto* samples; 100 μ m in sections. (G) Sternum and limb (forelimb, foot) of control (upper panel) and $p57^{cKO-ILZ (m)/+}$ mice (lower panel) at embryonic stages (E14.5; left panel) and at birth (P0; central and right panels). Skeletons were stained with alcian blue (cartilage) and alizarin red (bone). Black (E14.5) or white (P0) asterisk indicates sternum fusion defects. Arrows indicate limb thickening and shortening. Arrowheads indicate delayed ossification. Scale bars: 2mm at E14.5 sternum and P0 foot; 5mm at P0 sternum and forelimb. E13.5: embryonic day 13.5; Int: intestine; P: palate; PS: palatal shelve; P0: post-natal day 0; P1: post-natal day 1; S: stomach.

Supplementary figure 1. LacZ expression in $p57^{FL-ILZ/+}$ and $p57^{cKO-ILZ/+}$ mice

Reporter expression in the indicated tissues of $p57^{FL-ILZ/+}$ (left panel) and $p57^{cKO-ILZ (m)/+}$ (right panel) animals. Arrowheads indicate X-Gal-stained glomeruli in perinatal kidney cortex. Scale bars: 200 μ m. E13.5: embryonic day 13.5; P0: post-natal day 0.

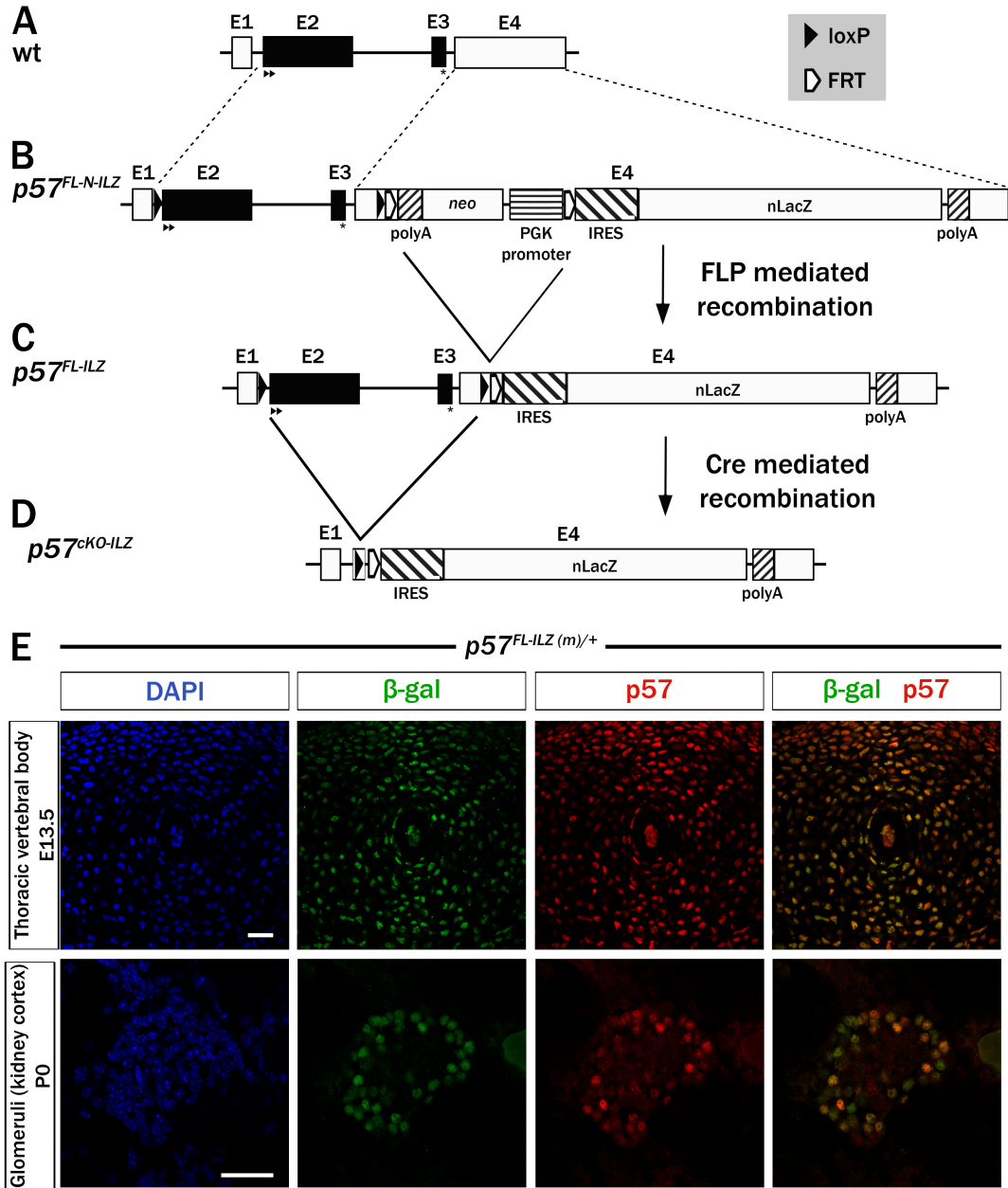


Figure 1

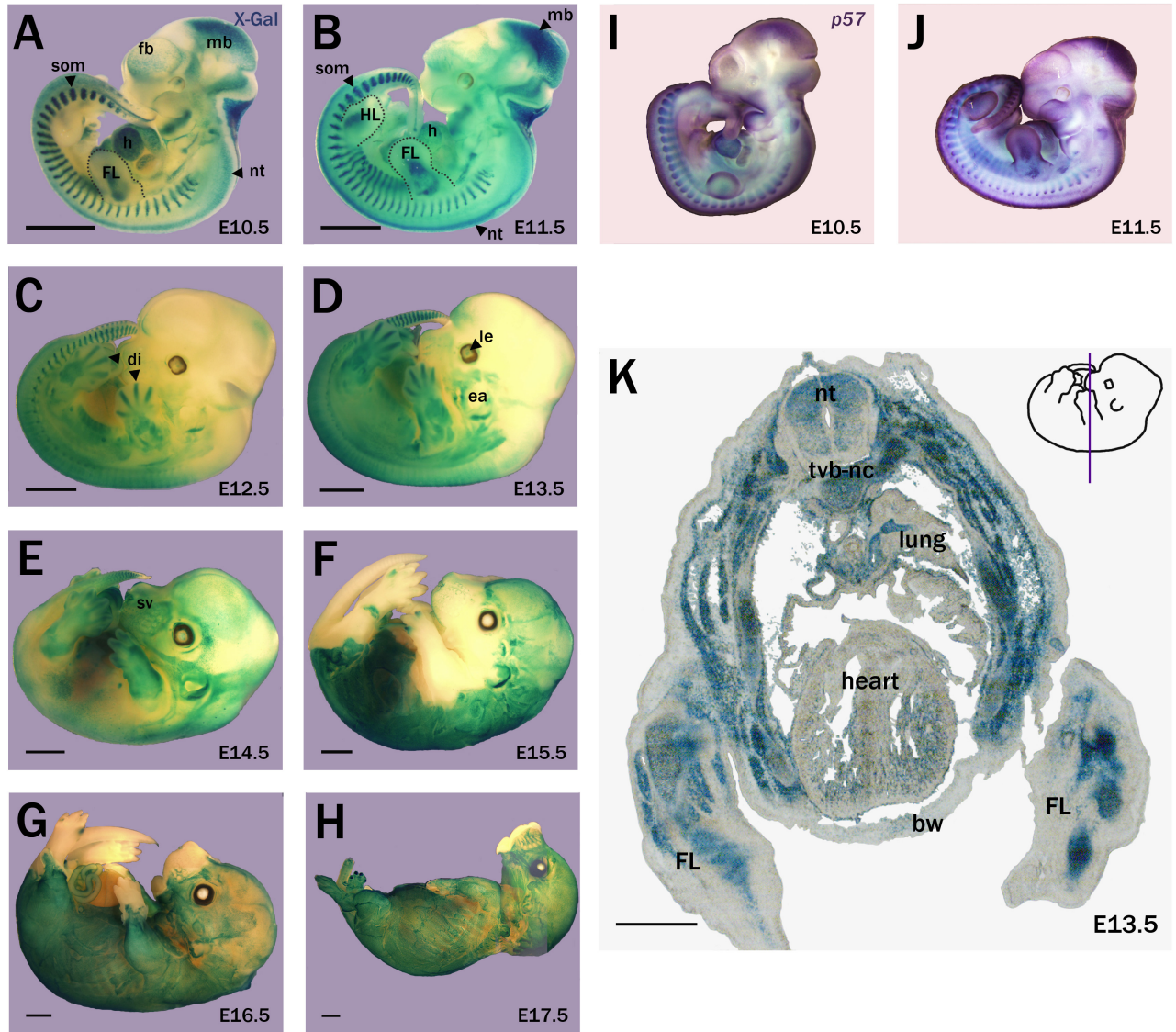


Figure 2

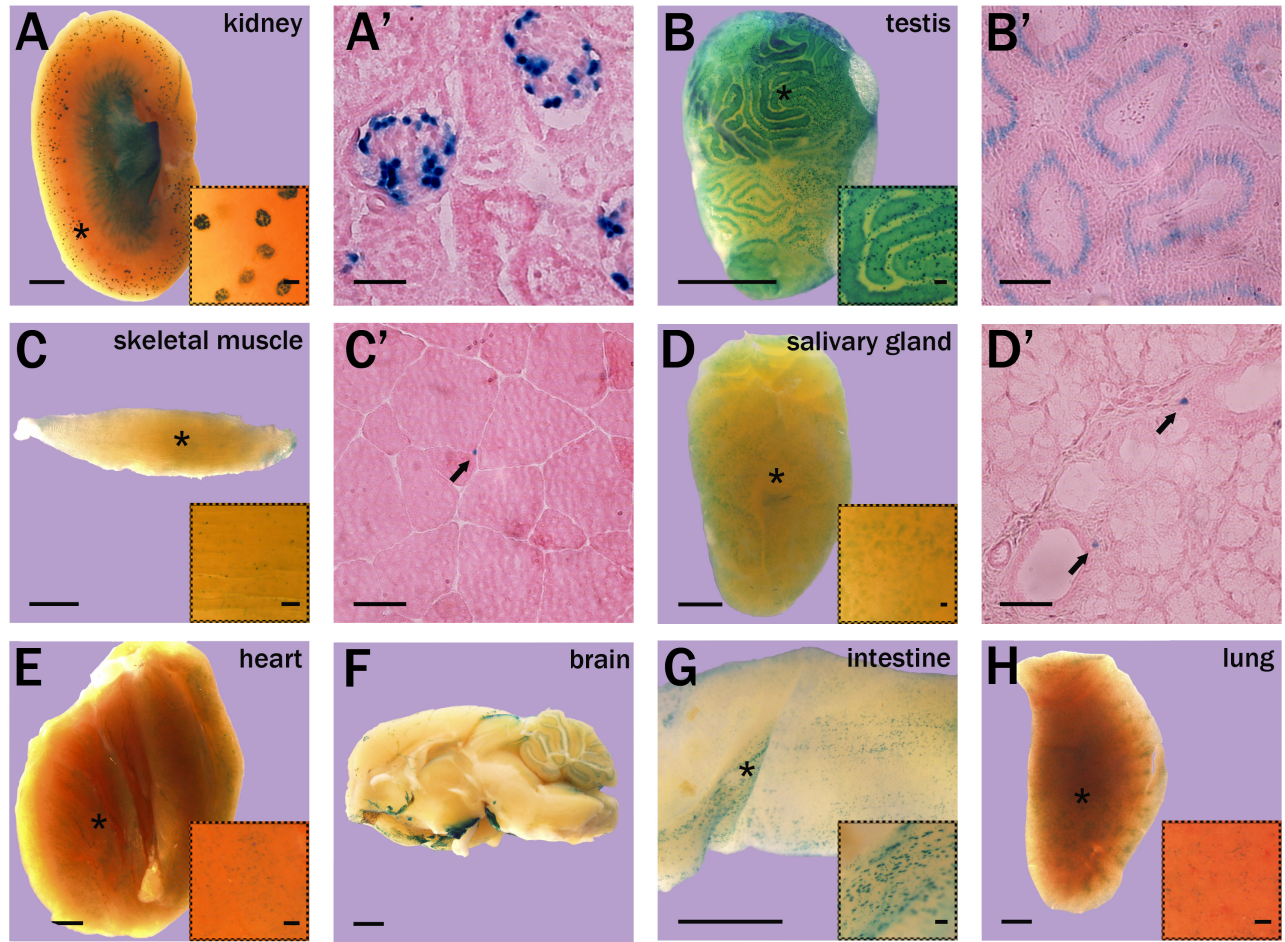


Figure 3

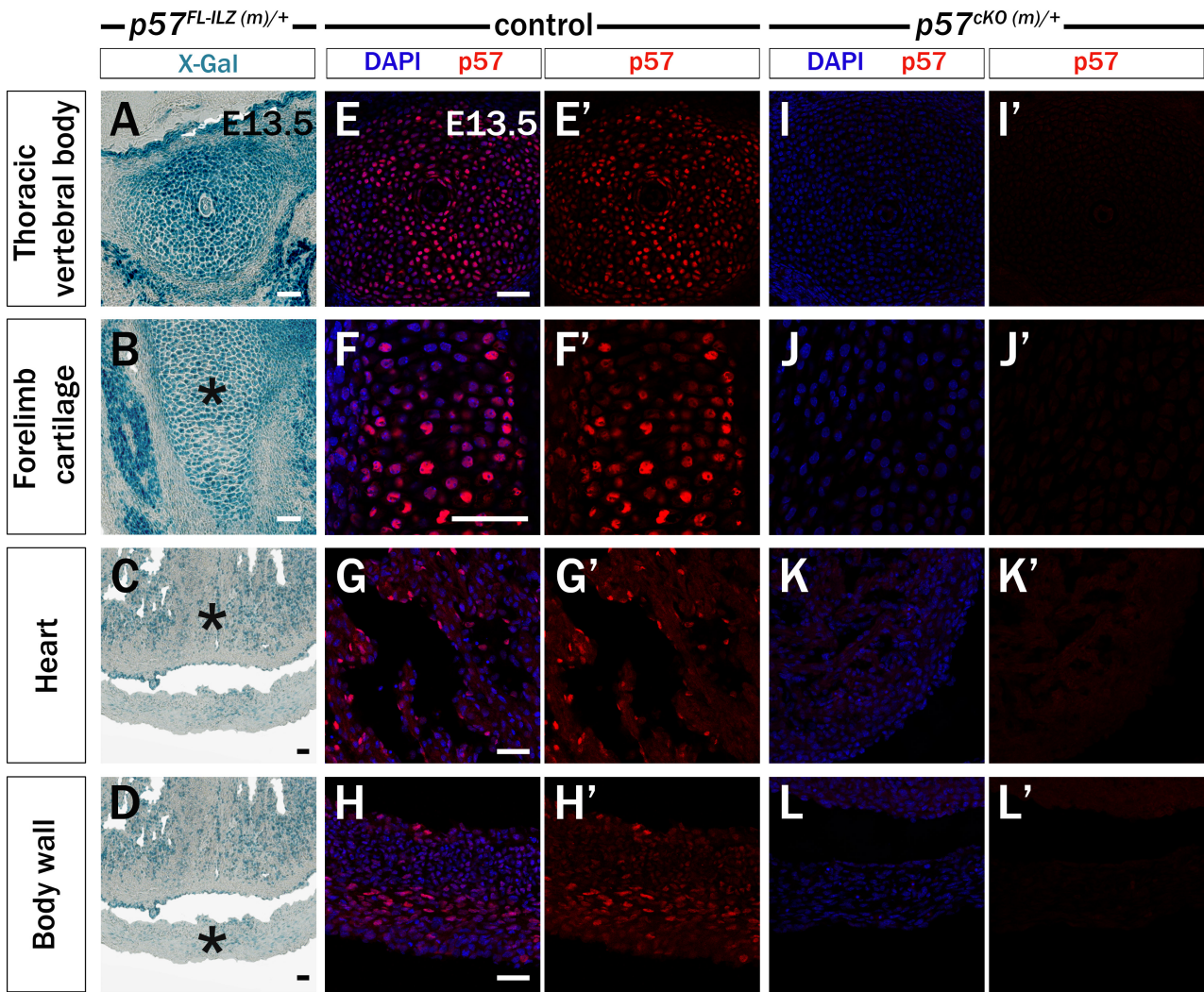


Figure 4

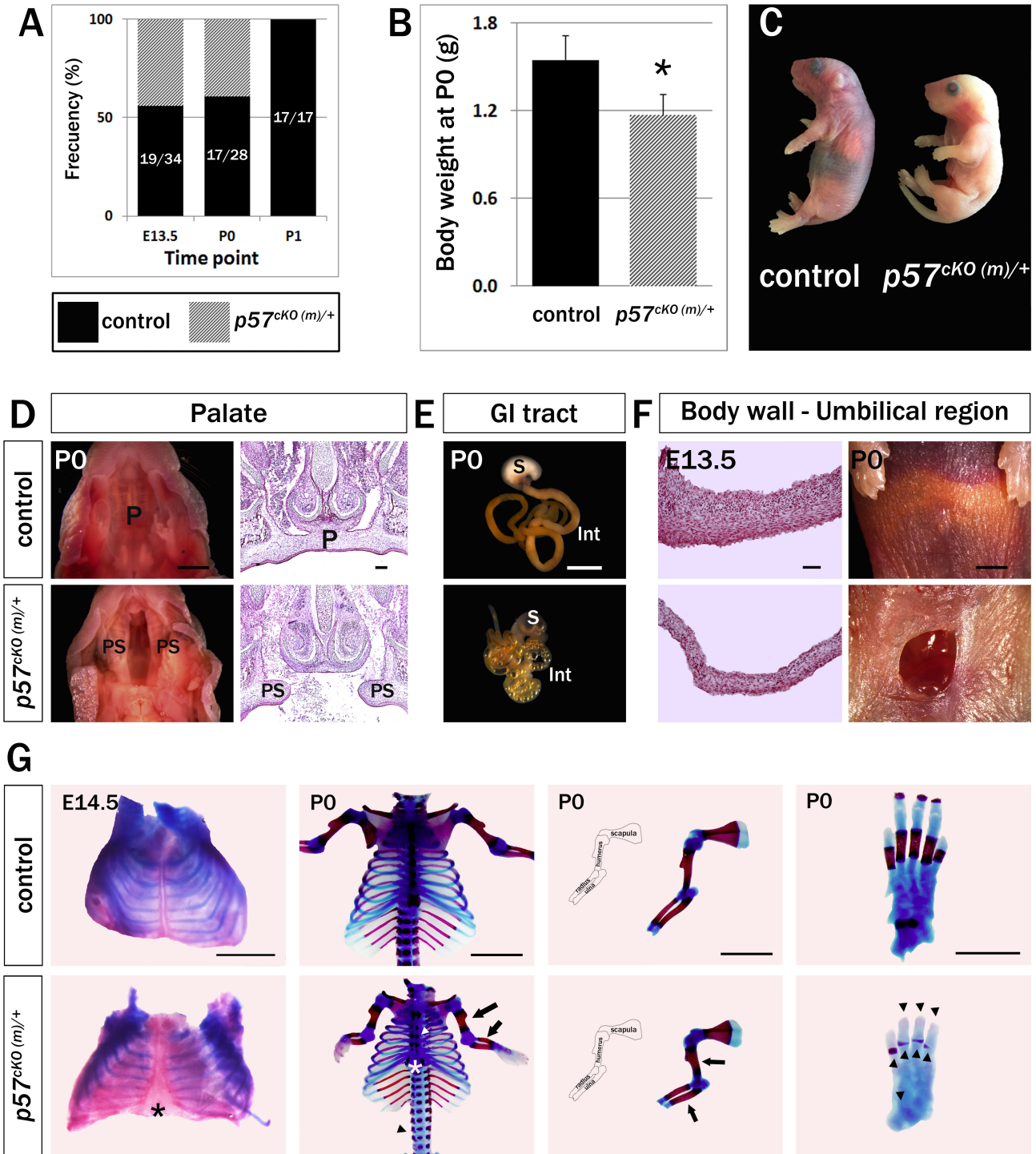


Figure 5

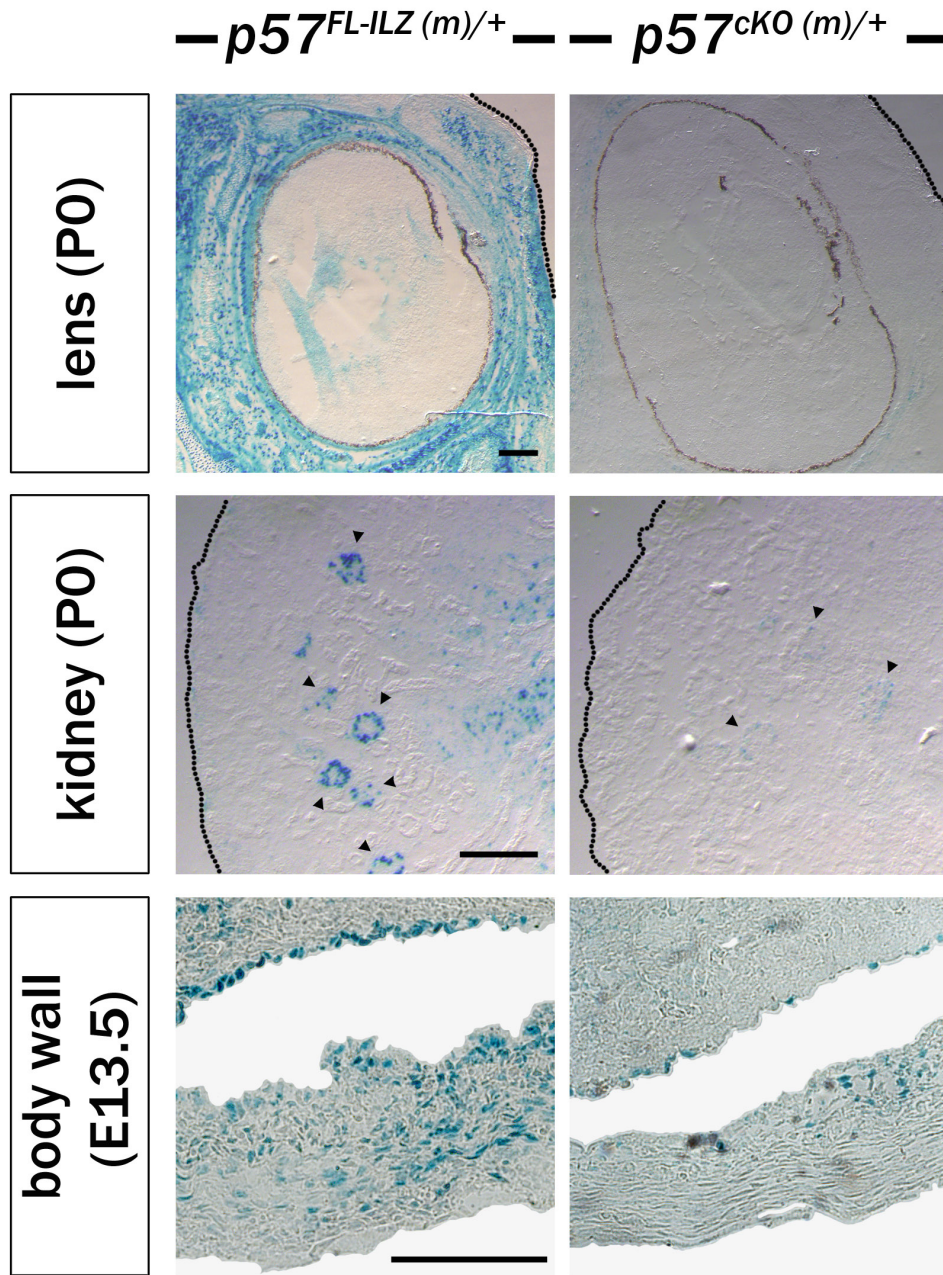


Figure S1

Preface

The CDKIs p57 and p21 have been shown to play crucial roles in cell cycle regulation of the nascent musculature. To identify their role in postnatal myogenesis we investigated their expression, function, and regulation in adult muscle stem cells, called satellite cells. The quiescent satellite cells that reside in resting adult muscles were devoid of p21, while this factor was upregulated following satellite cell activation (even in proliferating myoblasts) and differentiation. In single myofiber cultures and at early post-injury muscle regeneration time points, p21 deficiency affected the dynamics of satellite cells, indicating p21 importance in the early post-activation events. Specifically, *ex vivo*, proliferation/activation were increased at the expense of differentiation, while *in vivo*, the satellite cell compartment was significantly reduced. However, both the satellite cell numbers and the muscle architecture were restored at the end of injury-induced regeneration. Given p21 and p57 overlapping function in embryonic myogenesis, we have investigated p57 in the adult context. Same as for p21, p57 was absent from quiescent satellite cells. In contrast, we detected it in the cytoplasm of activated cells with increasing nuclear translocation upon differentiation. Using the new conditional p57 knock-out, we induced satellite cell-specific p57 ablation and observed a marked differentiation deficit of satellite cell-derived myoblasts *in vitro*. Thus, our data implicate p21 and p57 in muscle cell cycle dynamics postnatally.

Distinct regulation and function of p21 and p57 during muscle stem cell activation and differentiation

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KEYWORDS

Satellite cells, cyclin-dependent kinase inhibitors, p21, p57, cell cycle, muscle regeneration.

ABSTRACT

Adult tissue maintenance and regeneration depends on efficient stem cell self-renewal and differentiation. In skeletal muscle the contributors to postnatal muscle growth, maintenance, and repair are the muscle satellite cells (mSCs), residing on the periphery of the syncytial post-mitotic myofibers. The mechanisms coordinating the regulation of cell cycle exit with activation, renewal and differentiation of the mSCs remain poorly understood in adult muscle. Here, we investigated the role of specific cyclin-dependent kinase inhibitors in adult mSCs that up to date have been linked to embryonic myogenesis. We show that p21 is not detected in quiescent adult mSCs, but it is induced in activated and differentiating myoblasts. Using an *ex vivo* single myofiber explant system, we examined the kinetics of myoblast production/differentiation of p21-null mice, and we observed an increase in activation and proliferation accompanied by a diminution of differentiation. *In vivo* muscle regeneration studies in the absence of p21 revealed a decrease in mSCs at day 5 post-injury, confirming a role of p21 in the early activation phase of mSCs. On the contrary, muscle architecture and mSC numbers were restored by day 28, indicative of complete regeneration. Since p21 and p57 have been shown to redundantly control embryonic myogenesis, we evaluated if p57 functions in adult myogenesis. Although p57 was absent from quiescent mSCs, it was expressed in their activated and differentiating progeny. We further demonstrate that p57 expression and subcellular localization are dynamic; exclusively cytoplasmic p57 is first observed in activated/proliferating myoblasts, while progressive nuclear translocation is associated with differentiation and growth arrest. p57-deficient myoblasts displayed a marked differentiation deficit, manifested by reduction in MYOGENIN+ cells and, subsequently, formation of smaller myotubes. Our data suggest that p21 and p57 play distinct functions both at the early steps of mSC activation and during differentiation.

INTRODUCTION

Adult regeneration is of vital importance for restoring tissue structure and function following damage. Skeletal muscle has a remarkable capacity to self-repair after severe injuries, a process dependent on muscle stem cells, the so-called muscle satellite cells (mSCs) [Relaix & Zammit, 2012]. mSCs originate from a PAX3/7+ progenitor population that in late fetal life acquires their characteristic anatomical position between the basal lamina and the plasma membrane of muscle fibers [Mauro, 1961; Relaix et al., 2005]. mSCs are involved in postnatal muscle growth [White et al., 2010; Pawlikowski et al., 2015], maintenance [Keefe et al., 2015; Pawlikowski et al., 2015], and regeneration upon injury or in many neuromuscular disorders [Lepper et al., 2011; McCarthy et al., 2011; Murphy et al., 2011; Sambasivan et al., 2011]. Juvenile mSCs acquire a non-proliferative, quiescent state around three weeks of post-natal life to preserve key functions [White et al., 2010; Cheung & Rando, 2013]. However, once stimulated by homeostatic demand or damage, mSCs become activated, re-enter cell cycle and provide differentiated progeny for muscle repair, while a subpopulation self-renews the quiescent pool [Relaix & Zammit, 2012]. The myogenic regulatory factors MYOD and MYOGENIN orchestrate mSC commitment and progression through the myogenic lineage [Wang et al., 2014], while the signals that trigger cell cycle exit and (re-)entry into quiescence remain more elusive.

Cell cycle is a synchronized process responding to positive and negative signals, while inappropriate growth arrest can result in cancer, in malformations during development, and in defective stem cell renewal [Zhang et al., 1997; Matsumoto et al., 2011; Sherr, 2012]. The common negative cell cycle regulators are Cyclin-Dependent Kinase Inhibitors (CDKIs). CDKIs are classified in two structurally and functionally defined families [Borriello et al., 2011]: the INK4 family [including p15^{Ink4b} (p15), p16^{Ink4a} (p16), p18^{Ink4c} (p18), p19^{Ink4d} (p19)] and the Cip/Kip family [including p21^{Cip1} (p21), p27^{Kip1} (p27), p57^{Kip2} (p57)]. Although members of the Cip/Kip family have been shown to control proliferation and differentiation in embryonic muscle or in myoblasts *in vitro*, their potential involvement in mSC quiescence and cell fate decisions has not been documented [Halevy et al., 1995; Reynaud et al., 1999; Zhang et al., 1999; Messina et al., 2005; Chakkalakal et al., 2014; Zalc et al., 2014].

Given that p21 and p57 redundantly control embryonic skeletal muscle differentiation (Zhang et al., 1999), we explored their role in adult myogenesis and regeneration. We show here that p21 and p57 are not detected in quiescent mSCs, but are induced upon activation and maintained in differentiating myogenic cells. We further show that p57 subcellular localization is specifically regulated during adult myogenesis, with a progressive cytoplasmic to nuclear translocation as activated myoblasts proceed to differentiation. Using mouse mutants

and *ex vivo* analysis, we provide evidence that both p21 and p57 are distinctly involved in the early activation events, while their genetic ablation led to decreased differentiation, increased proliferation, and mSC pool diminution. Our results imply that growth arrest is dynamically controlled, with p21 and p57 playing distinct functions during adult myogenesis.

RESULTS

p21 is expressed in mSCs following their activation

To analyze the expression of p21 in muscle satellite cells (mSCs) in different states, including self-renewal and differentiation, while they are still exposed to their physical niche, we used the well-established myofiber culture system [Zammit et al., 2004; Moyle & Zammit, 2014]. We isolated single myofibers with their associated PAX7⁺ mSCs from *Extensor digitorum longus* (EDL) muscles of adult wildtype mice (Figs. 1A). Quiescent mSCs of freshly isolated fibers (time point 0; T0) were not labeled using antibodies against p21 (Fig. 1A-B, I). In line with this, we could not detect expression of p21 in the PAX7⁺ mSCs in cross-sections of resting adult *Tibialis anterior* (TA) muscles (Fig. 1C-D).

To evaluate the expression of p21 during activation, self-renewal and differentiation, we performed myofiber culture for 24-72 hours. The culture conditions allow the mSCs to get activated at T24-48 and proceed to myogenic differentiation or self-renewal of the quiescent pool (T72) [Zammit et al., 2004]. During the initial proliferation state (T24-T48) the activated mSCs co-express PAX7 and MYOD [Zammit et al., 2004]. In contrast to the quiescent population, we detected increasing levels of p21 in activated mSCs (Fig. 1I-J). By 24 hours nearly half the activated PAX7⁺/MYOD⁺ cells express p21, while almost all the PAX7⁺ or MYOD⁺ cells are p21⁺ at 48 hours. At T72 divergent mSC fates can be monitored including differentiation (downregulation of *Pax7* associated with expression of *MyoD* and *Myogenin*) and self-renewal (maintenance of PAX7, loss of MYOD, and lack of MYOGENIN) [Zammit et al., 2004]. p21 remained high at that time point of the culture (Fig. 1E-L). PAX7⁺MYOD⁺ cells at T72 represent activated cells [Zammit et al., 2004], suggesting that some p21⁺PAX7⁺ and p21⁺MYOD⁺ cells are proliferating myoblasts which express a cell growth arrest factor as in T48 (Fig. 1I-J). To confirm this unexpected observation, we labelled cycling cells with an antibody against KI67 and observed that indeed a high percentage of KI67⁺ myoblasts were co-expressing p21 (Fig. 1L). Finally, as expected, p21 presented a solid expression in differentiating MYOD⁺ and MYOGENIN⁺ cells at T72 (Fig. 1G-H, J-K).

Loss of p21 affects early phase of mSC activation

To evaluate the role of p21 on muscle homeostasis, we quantified the ratio of muscle weight per body weight and the relative number of mSCs in *p21* mutant mice compared to controls. We did not observe significant variations in the size and weight of limb muscles or diaphragm in mutant mice compared to controls (Fig. 2A). Trunk muscles did not exhibit any difference either (data not shown), although dissection limitations render this comparison less reliable. The number PAX7+ cells per 100 fibers was similar in *p21* mutant mice compared to controls, suggesting that the mSC compartment has been correctly established in the absence of p21 (Fig. 2B-D). We did not observe centrally located nuclei, indicative of ongoing regeneration, in muscle cross-sections (data not shown). Furthermore, freshly isolated myofibers of mutant and control EDLs had comparable amounts of mSCs and no PAX7+ cells were expressing KI67 at T0 (data not shown), implying that mSCs correctly exited cell cycle postnatally and entered into quiescence.

We then examined the kinetics of myoblast production and differentiation upon p21 depletion using single myofiber cultures. At T72, mSC-derived myoblasts represent a mixed population (see above) and PAX7/MYOD immunostaining can reveal self-renewing (PAX7+MYOD-), activated/proliferating (PAX7+MYOD+), and differentiating (PAX7-MYOD+) myoblasts [Zammit et al., 2004]. In the absence of p21, we observed an increased proliferation, as evidenced by the fraction of activated PAX7+MYOD+ myoblasts (Fig. 2E) and confirmed by quantification of the cycling, KI67+ myoblasts per fiber (Fig. 2F). Self-renewal (PAX7+MYOD-) did not seem to be affected (Fig. 2E), but we observed prolonged activation (PAX7+MYOD+) at the expense of differentiation (PAX7-MYOD+) (Fig. 2E). Moreover, we found less MYOGENIN+ differentiating myoblasts per fiber in *p21* mutants (Fig. 2G). These data suggest that p21 functions at the early phase of satellite cell activation to limit proliferation during the proliferative phase, and p21 deficiency affects their proliferation and differentiation.

Next, we evaluated the impact of p21 loss of function for skeletal muscle regeneration. We performed intramuscular cardiotoxin injections into the TA and sacrificed the mice at 5, 10 and 28 days post-injury, to evaluate early, intermediate and late time points of the regeneration procedure. Once muscle degeneration is induced, mSCs undergo a sequence of events similar to the myofiber culture: (1) get activated, (2) proliferate to expand their population, and (3) self-renew their quiescent pool for future needs and provide differentiated progeny for new fiber formation and muscle repair [Relaix & Zammit, 2012]. Our histological analysis of injured muscle did not show abnormalities in *p21* mutant mice (Fig. 3A-F). During the first post-injury days, damaged muscle show extensive inflammatory cell infiltration and muscle progenitor proliferation [Paylor et al., 2011]. Accordingly, we observed increased cellular content of control and mutant muscles at D5 of regeneration (Fig.

3A-B). Normal tissue structure, with well-formed myofibers and diminished interstitial space, was restored at similar time in controls and mutants (Fig. 4C-F). Deposition of fibrotic and adipose tissue was similar between controls and mutants at all time points analyzed (data not shown). Since our *ex vivo* data suggest an early activation defect upon p21 ablation (Fig. 3), we evaluated the mSC population throughout regeneration. Injury-triggered mSC pool amplification led to a remarkable increase of the PAX7+ population at D5 post-cardiotoxin in controls and mutants (Fig. 3M). However, significantly less mSCs were present in the *p21*-null animals at D5 (Fig. 3G-H, M). At later time points no significant difference was observed (Fig. 3I-M). In both controls and mutants, mSC numbers decreased back to baseline levels once their contribution to muscle regeneration was finished at D28 (Fig. 2D, 3M).

Combining the observed *ex vivo* defective differentiation/proliferation with the *in vivo* normal muscle regeneration and mSC pool restoration, we conclude that p21 regulates the early phase of mSC activation, leading to a delay but not a complete impairment of their performance. Our results suggest that p21 is involved in the fine tuning of activated satellite cells proliferation.

p57 is expressed in activated but not quiescent mSCs

Since embryonic muscle growth arrest was redundantly controlled by p21 and p57, we next evaluated if p57 also operates during adult myogenesis. We first examined p57 expression in mSCs in myofiber cultures and *in vivo* resting muscle (Fig. 4). On freshly isolated myofibers (T0), quiescent PAX7+ mSCs did not express p57 (Fig. 4A), while culture-mediated mSC activation induced p57 expression in PAX7/MYOD+ myoblasts at T24-T48 (Fig. 4A-B). In line with this, mSCs of resting adult TA muscles were p57-negative (Fig. 4C). However, p57 was detected in non-myogenic interstitial cells, consistently with previous reports of p57 presence in adult muscle extracts [Matsuoka et al., 1995; Park & Chung, 2001].

We next evaluated the kinetics of p57 expression in mSCs upon *ex vivo* activation in myofiber cultures. Activated PAX7+ (Fig. 5A) and MYOD+ (Fig. 5B) myoblasts at T24-T48 presented with increasing amounts of p57. Remarkably, during the early activation/proliferation phase, p57 was restricted to the cytoplasm (Figs. 4A-B, 5A-B). At T72, there is a mixed population of self-renewing (corresponding to part of Pax7+ cells), activated/proliferating (part of PAX7+ and MYOD+ cells), and differentiating (MYOGENIN+ and part of MYOD+ cells) myoblasts [Zammit et al., 2004]. We observed high percentages of p57 in each of these populations (Fig. 5A-C). Of note, as differentiation proceeded (MYOD+ followed by MYOGENIN+ at T72), p57 expression was becoming increasingly nuclear (Fig. 5B-C). Although p57 was mostly cytoplasmic in PAX7+p57+ T72 myoblasts,

p57 exhibited nuclear presence in around 25% of MYOD+p57+ and 55% of MYOGENIN+p57+ T72 myoblasts (Fig. 5A-C). Finally, similarly to the T24-T48 activated/cycling populations, p57 continued to be present in KI67+ proliferating cells at T72, yet limited to their cytoplasm (Fig. 5D). These results suggest that in contrast to p21, nuclear expression of p57 is not compatible with the proliferative status of activated satellite cells.

Loss of p57 affects myogenic differentiation

To circumvent the early perinatal lethality of *p57* knock-out mice, we have generated a floxed *p57* allele (*p57^{Flox-ILZ}*) to enable conditional abrogation of p57 using the Cre/loxP system [Mademtzoglou et al., submitted]. To selectively ablate p57 from the mSCs and their progeny, we intercrossed *p57^{Flox-ILZ}* mice with the *Pax7^{CreERT2}* line [Lepper et al., 2009]. In the resulting mice, Cre expression and p57 excision are dependent on tamoxifen (TMX) induction. Administration of TMX (intraperitoneal injections) or 4-hydroxytamoxifen (in culture) failed to efficiently recombine *p57* locus (Fig. S1A-C), hindering subsequent *in vivo* analyses of *p57*-deficient muscles in *Pax7^{CreERT2}; p57^{Flox-ILZ}* animals. To improve the efficiency of recombination, we crossed floxed animals with *Pax3^{Cre}* mice [Engleka et al., 2005], targeting the muscle lineage from early embryonic stages. However, high levels of p57 protein in the mSC progeny were still observed in myofiber cultures (Fig. S1D). We then inserted *p57^{Flox-ILZ}* and *Pax7^{CreERT2}* in the background of the Cre-responsive two-color fluorescent reporter *ROSA^{MT/mG}*, to allow the selection of recombined GFP+ cells. FACS-sorted cells grew in high serum conditions and then were serum-deprived to differentiate for 1 or 3 days and stained with early (e.g. Myogenin) or late (i.e. Myosin Heavy Chain-MyHC) differentiation markers, respectively (Fig 6A). p57 transcript and protein were induced by differentiation in the control cells (Fig. 6B-C, E). On the contrary, they were almost undetectable in cells from mutant animals (Fig. 6B, D-E), while p21 showed same levels in controls and mutants (Fig. 6B). In primary cultures of FACS-sorted satellite cells from *p57*-deficient mice, myogenic differentiation was impaired. One-day post-differentiation, myogenin expression was significantly decreased (Fig. 6F-H). Furthermore, myotube formation three days-post-differentiation was severely compromised (Fig. 3G-H). In conclusion, muscle regeneration in the absence of p57 could not be evaluated, despite promising *in vitro* results (Fig. 6) which strongly implicated p57 in the regulation of postnatal myogenesis.

DISCUSSION

Regenerative adult myogenesis is crucial for recovery from injuries, but can be compromised by degenerative or disease states that affect the functional capacity of skeletal muscle stem cells, the satellite cells (mSCs). The maintenance of mSCs functions largely depends on the entry and maintenance of a non-cycling, reversible

quiescent state. The molecular mechanisms that control mSC cell cycle transitions and adult myogenesis have gained significant interest in recent years, as a way to understand post-trauma tissue restoration and, subsequently, to design efficient innovative therapies when it is defective.

Given the essential, yet redundant, role of p21 and p57 in embryonic and fetal myogenesis [Zhang et al., 1999], we hypothesized that they may also control cell cycle and differentiation of mSCs in adult muscle. We found that p21 protein is not expressed in resting muscle cells, including quiescent mSCs (Fig. 1). This observation is in agreement with previous reports for negligible p21 mRNA [Macleod et al., 1995] or protein [Franklin & Xiong, 1996] in adult muscle, but contradicts reports for high *p21* transcripts [Parker et al., 1995; Park & Chung, 2001]; variance in analyzed muscles, mouse background, age window, and probes, could account for these differences. *p21* null mice do not exhibit any muscle phenotypes during development [Zhang et al., 1999], and as we did not observe p21 expression in resting adult muscles (Fig. 1C-D), we expected no differences in resting muscles of mutant and control animals. Indeed, muscle weight, architecture, and mSC numbers were not affected in the absence of p21 (Fig. 2A-D), consistent with previous reports [Deng et al., 1995; Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015].

Similarly to p21, we did not detect p57 in quiescent mSCs on sections or isolated myofibers of resting adult (8-12 weeks) muscle (Fig. 4A, C), while it was abundant in interstitial cells (Fig. 4C). Our finding is consistent with previous reports of high p57 levels in adult muscle [Matsuoka et al., 1995; Park & Chung, 2001] and lack of p57 in FACS-sorted postnatal mSC populations at different ages and dormancy states [Chakkalakal et al., 2014]. In contrast, an early study detected p57 in quiescent mSCs [Fukada et al., 2007] which were isolated with a FACS protocol using an antibody previously described by the same group [Fukada et al., 2004]. However, this antibody immuno-reacts with bone marrow cells [Fukada et al., 2004], where p57 has a well-established role and presence in the hematopoietic lineage [Matsumoto et al., 2011; Zou et al., 2011]. Furthermore, p57 immunostaining in Fukada et al. [2007] were performed with an antibody against the p57 carboxy-terminus, which might cross-react with the respective domain of p27 [Matsuoka et al., 1995; Galea et al., 2008; Pateras et al., 2009]. Combining our observation with previous reports [Fukada et al., 2004; 2007; Chakkalakal et al., 2014; present study], we conclude that quiescent mSCs do not express p57. Instead, it is established that they express p27, the other CDKI of the family including p21 and p57 [Chakkalakal et al., 2014; our unpublished data]. However, this observation does not preclude p21 and p57 participation in mSC cell cycle dynamics; in the pituitary p57 was found to promote cell cycle exit, while p27 prevented cell cycle re-entry [Bilodeau et al., 2009].

Previous reports demonstrated that p21 and p57 are required to drive embryonic muscle progenitors out of the cell cycle during terminal differentiation [Zhang et al., 1999]. Consistently, we found that p21 and p57 are not expressed in quiescent mSCs (Figs. 1, 4, 5). Yet we show that they are unexpectedly upregulated upon *in vitro* activation in proliferating mSCs (Figs. 1, 4, 5). Although T24-T48 cells are still cycling [Zammit et al., 2004], they were expressing cell cycle inhibitors, such as p57 (Fig. 5A-B) and p21 (Fig. 1I-J). Furthermore, many KI67+ cells at T72 expressed p57 (Fig. 1L) or p21 (Fig. 5D). Accordingly, post-regeneration activation of mSCs has been shown to induce p21 and p57 with their levels peaking at D3-4 [Yan et al., 2003; our unpublished observations]. The appearance of p21 and p57 might be associated with MYOD expression. MYOD is expressed soon after mSC activation [Zammit et al., 2004; Zhang et al., 2010] and sustains the transition from quiescence to cell cycle via the replication-related factor CDC6 [Zhang et al., 2010]. MYOD induces growth arrest in non-myogenic cell lines [Crescenzi et al., 1990; Sorrentino et al., 1990] and has a well-established role for the entry into the myogenic lineage. Remarkably, despite MYOD robust expression, myoblasts continue to proliferate and do not proceed to differentiation for several days [Tajbakhsh, 2009]. In fact, in dividing myoblasts MYOD activity is inhibited by Id proteins and CDK/Cyclin complexes [Wei & Paterson, 2001]. Furthermore, additional factors, including MYOGENIN, were suggested to initiate or enhance transcription in part of MYOD targets [Blais et al., 2005; Cao et al., 2006].

Both p21 and p57 have been implicated in positive feedback loops with MYOD, whereby MYOD induces them and they in turn enhance MYOD activity and stabilization in cultured cells or *in vivo*. Specifically, MYOD promoted *p21* transcription [Halevy et al., 1995; Tintignac et al., 2004], while in its absence p21 expression was delayed during muscle development [Parker et al., 1995]. p21 stabilized MYOD by inhibiting its phosphorylation [Reynaud et al., 1999], while non-phosphorylatable MYOD sustained p21 expression [Tintignac et al., 2004]. MYOD induced *p57* both by disrupting a chromatin loop to release the *p57* promoter and by upregulating intermediate factors [Vaccarello et al., 2006; Figliola et al., 2008; Busanello et al., 2012; Battistelli et al., 2014]. Furthermore, we previously identified a muscle-specific *p57* regulatory element that MYOD binds and transactivates [Zalc et al., 2014]. Co-immunoprecipitation assays revealed direct p57-MYOD binding [Reynaud et al., 2000]. Loss- and gain-of-function experiments in zebrafish embryos established a positive feedback loop between p57 and MYOD [Osborn et al., 2011].

Upon mSC activation, p21 and p57 were expressed in proliferating myoblasts in single myofiber cultures (Figs. 1, 4, 5), including T24-T48 cycling populations as well as KI67+ cells at T72. These observations might contradict

their traditional role as cell cycle exit factors [Gu et al., 1993; Harper et al., 1993; Xiong et al., 1993; Lee et al., 1995; Matsuoka et al., 1995]. However, at low concentrations, p21 promotes assembly and nuclear localization of CDK/Cyclin complexes, whereas at high amounts it exhibits its inhibitory activity towards them [Michieli et al., 1994; Harper et al., 1995; LaBaer et al., 1997]. Similarly, p21 is involved in nuclear accumulation and activity of CDK/Cyclin complexes in fibroblasts converted to myogenic fate [Peschiaroli et al., 2002], while differentiation of the C2C12 myogenic cell line leads to increased inhibition of CDK4 by p21 [Wang & Walsh, 1996]. It remains to be verified whether these interactions are at play in our experimental system. The fact that p21 ablation caused increased proliferation in the myofiber cultures (Fig. 4) suggests that p21 is involved in restraining the cell cycle.

The role of p57 in cycling myoblasts of isolated myofiber cultures seems to be more complex. p57 is cytosolically restricted in activated myoblasts but translocates to the nucleus as differentiation progresses (Fig. 5). We have not identified the molecular events underlying this shuttling, although some observations might explain this subcellular localization pattern. Firstly, p57 might be regulated on the cellular level, by cytoplasmic restriction during proliferation. A similar mechanism was shown for ERK signaling in muscle progenitors, whereby switching from proliferation to differentiation was associated with subcellular ERK localization [Michailovici et al., 2014]. Secondly, p57 could be implicated in cell cycle progression through CDK/Cyclin assembly, similarly to its Cip/Kip siblings (see above). In the absence of p21 and p27, p57 resumes their role in CDK/Cyclin complex stabilization in mouse embryonic fibroblasts [Cerqueira et al., 2014]. However, this might be less likely in myoblasts, where p57-MYOD binding engages the p57 helix domain [Reynaud et al., 2000] that was found indispensable for CDK/Cyclin binding and inhibition [Hashimoto et al., 1998; Reynaud et al., 2000]. Thirdly, p57 could be involved in nucleo-cytoplasmic distribution of cyclins or CDKs, as previously observed in other cell types. p57 has been shown to interfere with the nuclear translocation of cyclin D1 [Zou et al., 2011] and to relocalize fraction of CDK2 in the cytoplasm [Figliola & Maione, 2004]. Fourthly, while in the cytoplasm, p57 might participate in mSC mobilization, one of the earliest manifestations of their activation [Siegel et al., 2009]. Cytoplasmic p57 was described to regulate cell motility together with LIM-kinase1 [Vlachos & Joseph, 2009; Chow et al., 2011; Guo et al., 2015]. Although we did not detect LIM-kinase1 in our myofiber cultures, we cannot exclude association with other, yet uncharacterized, partners. Future studies are expected to elucidate the roles of p57 in different sub-cellular compartments of myoblasts.

p21 did not undergo nucleo-cytoplasmic translocation in myoblasts of myofiber cultures (Fig. 1). Similarly, p21 is strictly nuclear in forming myotubes [Figliola & Maione, 2004] and p21, but not p57 or p27, appears in the

nucleus of KI67+ proliferating b-cells [Fiaschi-Taesch et al., 2013]. Moreover, cytoplasmic presence of p21 is linked to degradation [Hwang et al., 2009] or oncogenesis [Besson et al., 2004; Besson et al., 2008]. Finally, p21 might not be subjected to regulation on the cellular level (e.g. initial cytoplasmic restriction), because of its role in the early events following mSC activation phase (Figs. 2-3), and at a moderate level, p21 expression is compatible with proliferation, which might not be the case for p57.

p21 and p57 were shown to function in a redundant manner during differentiation of embryonic myogenic cells. On the contrary, in the adult we observed early differentiation and proliferation defects in single mutants, implying lack of such compensation in the initial post-activation phase. Monitoring of myoblast kinetics at the early post-activation period showed that in the absence of p21 there was a differentiation deficit and proliferation increase *in vitro* as well as diminution of the mSC pool *in vivo* (Figs. 2-3). However, muscle regeneration was completed by 28 days post-injury, leading to indistinguishable muscle structure and mSC numbers between controls and mutants (Fig. 3). Similarly, mSCs and muscle architecture were restored by 28-30 days post-injury in previous studies [Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015], despite occasional defects at earlier time points [Hawke et al., 2003; Chinzei et al., 2015]. This implies a crucial role during the early activation events, while at later stages p21 either becomes dispensable or its role is masked by compensatory upregulation of other factors. Future studies with *p21/p57* double knock-out mice are expected to be more enlightening. In contrast to the myogenic differentiation defects observed with *p21*-null mice, siRNA-mediated p21 knock-down in primary myoblast cultures left myogenic differentiation unaffected [Biferi et al., 2015]. The difference was attributed to the effect of acute versus chronic p21 loss, while conditional genetic ablation of p21 with the loxP-Cre system would allow test this hypothesis.

p57 deficiency hindered myoblast differentiation and myotube formation (Fig. 6). p57 correlates with differentiation in many tissues [Westbury et al., 2001], while in myogenic cultures is sometimes considered as a differentiation marker [Reynaud et al., 1999; Mounier et al., 2011]. Nevertheless, the consequences of its ablation on adult muscle have not been examined, notably because of the perinatal lethality of p57 mutant mice, which our new conditional knock-out allele should allow to bypass [Mademtzoglou et al., submitted]. Flanking the coding exons 2-3 by LoxP sites ensured postnatal survival while leaving the possibility of Cre-induced excision [Mademtzoglou et al., submitted]. Our data implicate p57 in adult myogenesis, with the caveat that such observations are limited to an *in vitro* system lacking the structural, neurogenic, and metabolic fidelity of the muscle tissue [Grounds, 2014]. *In vivo* regeneration studies would decipher if and how p57 influences the re-establishment of mSC pool post-injury, given its emerging importance in the quiescence

and renewal of other stem cells [Matsumoto et al., 2011; Zacharek et al., 2011; Furutachi et al., 2013]. However, *in vivo* analysis was hindered by inefficient recombination (Fig. S1). Previous studies suggest that recombination resistance/success in mSCs is related to the cell cycle state (quiescence versus activation/proliferation) [Lepper et al., 2009; Günther et al., 2013; von Maltzahn et al., 2013]. In our set, various schemes TMX or 4-hydroxytamoxifen administration failed to recombine p57 locus (Fig. S1A-C). p57 is an imprinted gene with preferential expression of the maternal allele [Hatada & Mukai, 1995]. Thus, to exclude the possibility of paternal allele reactivation, we produced animals in which both p57 alleles were floxed, but we again could not observe successful recombination following tamoxifen treatment (Fig. S1B-C). Inefficient recombination persisted when we used *Pax3^{Cre};p57^{Flox-ILZ}* mice (Fig. S1D), while no homozygous *p57^{CKO/CKO}* animals were obtained in the crosses with *Pax3^{Cre}* mice. In addition, mice from the *Pax3^{Cre}* crosses did not follow mendelian frequencies at genotyping age and part of the mutants died at birth due to cleft palate, in agreement with Pax3 and p57 involvement in the development of craniofacial structures [Yan et al., 1997; Zhang et al., 1997; Zalc et al., 2015; Mademtoglou et al., submitted].

In conclusion, our data indicate that p21 and p57 play essential roles at the early phase following mSC activation. The presence of p21 and p57 is compatible with activation/ proliferation and possibly represents an early activation event. Their loss profoundly affects *ex vivo* myogenic differentiation. Our data so far indicate that p21 and p57 function in distinct ways during adult and embryonic myogenesis, in terms of early versus continuous myogenesis support and (possibly) compensation for each other's loss. It remains to be established whether at later stages of differentiation/regeneration p21 and p57 take a leading role in *p57* and *p21* mutants, respectively, to rescue the observed preliminary defects. mSC-specific double *p21/p57* knock-out will elucidate their relative contributions and putative redundancies in the adult.

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MATERIALS AND METHODS

Mouse lines

The following mouse lines have been previously described: *Pax3^{Cre/+}*, *Pax7^{CreERT2/+}*, *Rosa^{mTmG}* (The Jackson Laboratory, stock 007576), *p21^{+/-}*, *p57^{CKO (m)/+}* (p57 is imprinted with preferential expression of the maternal allele; superscript (m) indicates maternal inheritance) [Brugarolas et al., 1995; Engleka et al., 2005; Lepper et al., 2009; Mademtoglou et al., submitted]. Non-mutant littermates were used as controls. For recombination induction with the *Pax7^{CreERT2}* allele, mice were injected with tamoxifen (T5648, Sigma-Aldrich) as recommended [Lepper et al., 2009]. C57BL/6J (Janvier) mice were used as wildtype animals for Figs. 1, 4, 5. Adult mice of 8 to 12 weeks of age were used. At least three mice per genotype were assessed.

All animals were maintained inside a barrier facility, and all *in vivo* experiments were performed in accordance with the French and European Community guidelines for the care and use of laboratory animals.

Single myofiber isolation and culture

Single muscle fibers were isolated by enzymatic digestion and mechanical disruption of *EDL* muscles [Moyle & Zammit, 2014]. For enzymatic digestion muscles were incubated for 90 minutes at 37°C with 0.2% collagenase type I (C0130, Sigma-Aldrich) in Penicillin/Streptomycin(P/S)-supplemented DMEM (41966, ThermoFisher Scientific). For mechanical disruption, muscles were transferred to 5%-horse-serum-coated deep petri dishes (Z692301, Sigma-Aldrich) with P/S-supplemented DMEM and medium was flushed against the muscle. Detached fibers were transferred into new dishes with P/S-supplemented DMEM. For timed culture, all fibers were transferred after finishing isolation into 5%-horse-serum-coated 6-well plates and cultured in the presence of 10% horse serum and 1% chicken embryo extract (#092850145, MP Biomedicals). If indicated, 5μM 4-hydroxytamoxifen (H6278, Sigma-Aldrich) was added at the beginning of the culture and renewed 48 hours later.

Cell sorting and culture

Using the tamoxifen-inducible Cre line *Pax7^{CreERT2}*, membrane-GFP is expressed in muscle satellite cells (mSCs) of *Rosa^{mTmG}* mice. Hindlimb muscles were dissociated by 0.2% w/v collagenase A (11088793001, Roche) and 2.4U/ml dispase II (04942078001, Roche) in digestion buffer [HBSS (14025, Thermo Scientific), 1%

Penicillin/Streptomycin, 0.1µg/ml DNase I (11284932001, Sigma), 0.4mM CaCl₂, 5mM MgCl₂, 0.2% bovine serum albumin (BSA; 0010001620, Jackson ImmunoResearch)] with 90 minute incubation at 37°C. Dissociated muscles were filtered through 100µm and 40µm cell strainers. GFP+ cells were collected with FACS Aria II based on gating of GFP signal.

Sorted cells were plated on matrigel (354230, Corning Life Sciences)-coated chamber slides (177445, Nalge Nunc International). They were initially cultured in high-serum conditions (referred to as “growth phase”) with 20% fetal bovine serum, 10% horse serum and 2.5pg/ml bFGF (450-33B, PeproTech) in DMEM+Glutamax (61925, ThermoFisher Scientific) supplemented with 1% P/S, 20mM L-glutamine (25030, Thermo Scientific), 10mM pyruvate (11360, Thermo Scientific), and 0.1M HEPES (15630, Thermo Scientific). Upon reaching 70% confluence, they were switched to low-serum conditions (5% horse serum in P/S-supplemented DMEM+Glutamax) to differentiate (referred to as “differentiation phase”).

Gene expression analysis

RNA was extracted with the RNeasy Micro kit (74004, Qiagen), according to the manufacturer’s instructions. cDNA was synthesized with the Transcriptor First Strand cDNA Synthesis kit (04379012001, Roche). RT-qPCR reactions were carried out in triplicate using the LightCycler 480 Sy Green Master (04887352001, Roche). Hypoxanthine Phosphoribosyltransferase 1 (HPRT) transcripts were used for normalization. Oligonucleotide sequences are available upon request.

Muscle regeneration

Adult (8-12 week old) mice were intramuscularly injected with 45 µl of cardiotoxin solution (10 µM; L8102, Latoxan) into the *Tibialis Anterior* (TA) after being anesthetized. Muscles were recovered 5, 10 or 28 days post-injury.

Immunohistochemistry

Sections: Muscles were frozen fresh in liquid nitrogen-cooled isopentane and sectioned at 8µm. Frozen sections were fixed with 4% paraformaldehyde/PBS for 20 minutes at room temperature. For hematoxylin-eosin staining, nuclei were stained with hematoxylin (MHS32, Sigma-Aldrich) for 11 minutes and cytoplasmes were counter-stained with eosin (6766007, Thermo Scientific) for 30 seconds. The sections were then dehydrated with brief passages through increasing concentrations of ethanol (30%, 50%, 70%, 85%, 95%, 100%). For two-color immunofluorescence, frozen sections were treated with methanol for 6 minutes at -20°C,

immersed to 0.01M citric acid pH 6.0 at 90°C for antigen retrieval and blocked with 5% BSA/PBS at room temperature. For three-color immunofluorescence, sections were permeabilized and blocked with 3% BSA, 10% lamb serum, 0.25% TritonX-100/PBS for 30 minutes at room temperature. In both cases, immunolabeling was performed at 4°C overnight for primary antibodies and at room temperature for 1h for secondary antibodies. When fibers were outlined with Alexa-conjugated anti-laminin, incubation was performed for 3 hours at room temperature, after washing out the secondary antibody. Nuclei were counterstained blue with DAPI. When mouse-raised antibodies were applied, endogenous mouse IgG was blocked by incubation with goat anti-mouse fab fragment affinity-purified antibody (115-007-003, Jackson ImmunoResearch) for 30 minutes at room temperature.

Single myofibers: After isolation (T0) or following culture (T24, T48, T72), myofibers were fixed with 37°C-preheated 4% paraformaldehyde/PBS for 10 minutes at room temperature. Fixed fibers were permeabilized with 0.5% TritonX-100/PBS for 8 minutes, blocked with 10% goat serum, 10% swine serum in 0.025% Tween20/PBS for 45 minutes and incubated with primary antibody (overnight at 4°C) and secondary antibody (1h at room temperature). Nuclei were counterstained blue with DAPI.

Primary myoblast culture: Cell cultures were fixed with 4% paraformaldehyde/PBS for 15 minutes at room temperature, permeabilized with 0.5% TritonX-100/PBS for 5 minutes, blocked with 5% BSA, 10% goat serum and immunolabeled with primary antibody (overnight at 4°C) and secondary antibody (1h at room temperature). Nuclei were counterstained blue with DAPI.

Antibodies

The following antibodies were used: mouse anti-Ki67 1:80 (#556003, BD Pharmingen), mouse anti-PAX7 1:100 (Pax7-c, DSHB), mouse anti-MYOD 1:80 (M3512, DAKO), rabbit anti-MYOD 1:100 (sc769, Santa Cruz), mouse anti-MYOGENIN 1:100 (F5D-c, DSHB), rabbit anti-p57 1:100 (sc8298, Santa Cruz), goat anti-p57 1:50 (sc1039, Santa Cruz), rabbit anti-p21 1:100 (ab 2961, Abcam), mouse anti-MyHC 1:100 (mf20-c, DSHB), rabbit anti-laminin 1:400 (L9393, Sigma-Aldrich), rabbit AlexaFluor647-conjugated anti-laminin 1:200 (NB300-144AF647, Novus Biological), AlexaFluor-coupled secondary antibodies (Life Technologies, Jackson ImmunoResearch).

Graphic editing

Graphs and representative photos were arranged in Figure format with the graphics editor Photoshop CS5. Color intensities of hematoxylin-eosin photos were adjusted to acquire uniform result among different sections.

Statistical test

Data of control and mutant mice in Figs. 3-4 were compared with the Mann-Whitney U-test. Differences at $p < 0.05$ were considered significant.

FIGURE LEGENDS

Figure 1. p21 expression during muscle satellite cell (mSC) activation and differentiation.

(A-B) Quiescent PAX7⁺ (green) mSCs satellite cells (A) lack p21 (red; B) in single myofibers of *Extensor digitorum longus* (EDL) muscle. (C-D) Lack of p21 (red; D) in sections of *Tibialis anterior* muscle. Arrowhead indicates mSC, marked with PAX7 (green; C). Myofibers are outlined with laminin (gray). (E-H) p21 expression (red; F, H) 72 hours post-culture of single myofibers of *Extensor digitorum longus* muscle. Myoblasts were stained with PAX7 (E) or MYOD (G) in green. (I-L) Quantification of p21 expression in myoblasts stained for PAX7⁺ (I), MYOD⁺ (J), MYOGENIN⁺ (K) or KI67⁺ (L) cells. Single myofibers of EDL and their associated mSCs were cultured and fixed at 24-hour intervals. Data show mean+SD, n=3 animals (20-33 fibers/animal). Nuclei were counter-stained with DAPI (blue). Scale bars 40 μ m.

Figure 2. p21 deficiency affects early activation phase of muscle satellite cells (mSCs).

(A) Muscle weight per body weight ratios for *p21* mutants and control littermates. Analyzed muscles were from hindlimbs (*Extensor digitorum longus*, EDL; *Tibialis anterior*, TA; gastrocnemius, GCN), forelimbs (biceps, BC; triceps, TC), and diaphragm (DIA). (B-C) TA sections of control (B) and *p21*-null (C) animals stained with PAX7 (red) and laminin (green) to mark mSCs (arrowheads) and fiber outline, respectively. Nuclei were counter-stained with DAPI (blue). (D) Quantification of (B-C). (E-G) Myoblast kinetics in single myofiber cultures of EDLs of *p21* mutant and control animals at 72 hours. Proportions of PAX7⁺ and/or MYOD⁺ myoblasts (E), proliferating KI67⁺ myoblasts (F), and differentiating Myogenin⁺ cells (G) were quantified. Data show mean+SD (A-D: n \geq 4 animals, E-G: n \geq 3 animals, 18-28 fibers/animal). Asterisks indicate significance; $p < 0.01$ (**) or $p < 0.001$ (***). Scale bars 50 μ m.

Figure 3. Muscle regeneration time-course in the absence of p21.

(A-F) Hematoxylin (nuclei)-eosin (cytoplasm) stained sections of regenerating TA muscles of controls (A, C, E) and *p21* mutants (B, D, F) at 5 (A, B), 10 (C, D), and 28 (E, F) days post-cardiotoxin injury. (G-L) Regenerating muscles of controls (G, I, K) and *p21* mutants (H, J, L) were stained with PAX7 (red) and laminin (green) to mark muscle satellite cells (mSCs) and myofibers, respectively, at 5 (G, H), 10 (I, J), and 28 (K, L) days post-cardiotoxin. Arrowheads indicate mSCs at D28 (K, L). (M) Quantification of (G-L). Data show mean+SD, $n \geq 3$ animals. Asterisks indicate significance; $p < 0.001$ (***)). Scale bars 50 μ m.

Figure 4. p57 expression in single myofiber culture and muscle sections.

(A) Muscle satellite cells (mSCs; T0) and myoblasts (T24-T48) stained with PAX7 (green) and p57 (red) in single myofiber cultures of *Extensor digitorum longus* (EDL) muscles. Arrowheads indicate Pax7+ cells. (B) Myoblasts of single EDL myofibers stained with MYOD (green) and p57 (red). Arrowheads indicate MYOD+ cells. (C) p57 (red) presence in *Tibialis anterior* muscle section. mSCs were marked with PAX7 (green) and fibers were outlined with laminin (gray). Arrowheads indicate p57+ cells. Asterisks indicate satellite cells. Nuclei were counter-stained with DAPI. Scale bars 40 μ m.

Figure 5. p57 expression and subcellular localization during satellite cell activation and differentiation.

(A) Immunofluorescence for PAX7 (green) and p57 (red) at T72 in single myofiber cultures of *Extensor digitorum longus* muscles and quantification of PAX7+ cells that co-expressed over the time-course of the culture. (B) Immunofluorescence for MYOD (green) and p57 (red) at T72 in single myofiber cultures and quantification of MYOD+ cells that co-expressed p57 over the time course of the culture. (C) Immunofluorescence for MYOGENIN (green) and p57 (red) at T72 in single myofiber cultures and quantification of MYOGENIN+ cells that co-expressed p57 at T72, when MYOGENIN+ cells are abundant. (D) Immunofluorescence for KI67 (green) and p57 (red) at T72 in single myofiber cultures and quantification of KI67+ cells that co-express p57 at T72. Nuclei were counter-stained with DAPI (blue). Cytoplasmic (light gray) or nuclear (dark gray) localization of p57 is indicated in the graphs. Data show mean+SD, $n \geq 3$ animals, 20-32 fibers/animal. Scale bars 40 μ m.

Figure 6. p57 deficiency impairs myogenic differentiation.

(A) Time-course of intraperitoneal tamoxifen injections (TMX arrows), chase, and muscle satellite cell harvest (FACS arrow) and culture (light gray bar for growth culture conditions, dark gray bar for differentiation culture conditions). Analyzed animals were *Pax7^{CreERT2/+};p57^{flox/flox};Rosa^{mTmG}* (*p57* cKO) and *Pax7^{CreERT2/+};p57^{+/+};Rosa^{mTmG}* (control). (B) *p57* and *p21* transcript levels of control and *p57* cKO myoblast cultures three days post-

differentiation. (C-K) Control (C, F, I) and *p57* cKO (D, G, J) myoblast cultures were examined for p57 protein (red) three days post-differentiation (C, D), MYOGENIN+ cells (green) one day post-differentiation. (F, G), and myotube formation three days post-differentiation (I, J). Nascent myotubes were marked with myosin heavy chain (MyHC; green; I, J). Nuclei were counter-stained with DAPI (blue). Graphs show quantification of p57 expression (E), Myogenin expression (H), and nuclei/myotube (K). Data show mean+SD, n=3 animals. Asterisks indicate significance; $p < 0.001$. Scale bar 40 μ m.

Supplementary Figure 1. Cre-mediated recombination strategies of floxed *p57* allele.

(A-C) Animals with *p57* floxed allele(s) (*p57^{CKO/+}* or *p57^{CKO/CKO}*) were crossed with tamoxifen-inducible *Pax7^{CreERT2}* mice. Schemes show the tamoxifen (TMX) or 4-hydroxytamoxifen (4-OHT) regimens and chase periods before (TMX) or during (4-OHT) single myofiber culture of *Extensor digitorum longus* (EDL) muscle. Photos show p57 (red) expression in single myofibers at 72 hours, implying insufficient recombination. (D) Single myofiber from *Pax3^{Cre/+};p57^{CKO/+}* mouse, showing p57 expression (red) at 72 hours post-culture of EDL myofibers. (E-F) Heads of *Pax3^{+/+};p57^{CKO/+}* (control) and *Pax3^{Cre/+};p57^{CKO/+}* (cKO) animals at birth showing closed (control; E) or cleft (cKO; F) palate. When heterozygous *p57* animals (*p57^{CKO/+}*) were used, they had maternally transmitted transgene, due to imprinting silencing of the paternally transmitted *p57* allele. Homozygous animals were used to exclude possible paternal allele activation in the crosses with *Pax7^{CreERT2}* mice. Nuclei in (A-D) were counter-stained with DAPI (blue). P: palate; PS: palatal shelf. Scale bars 40 μ m (myofibers) or 2mm (head).

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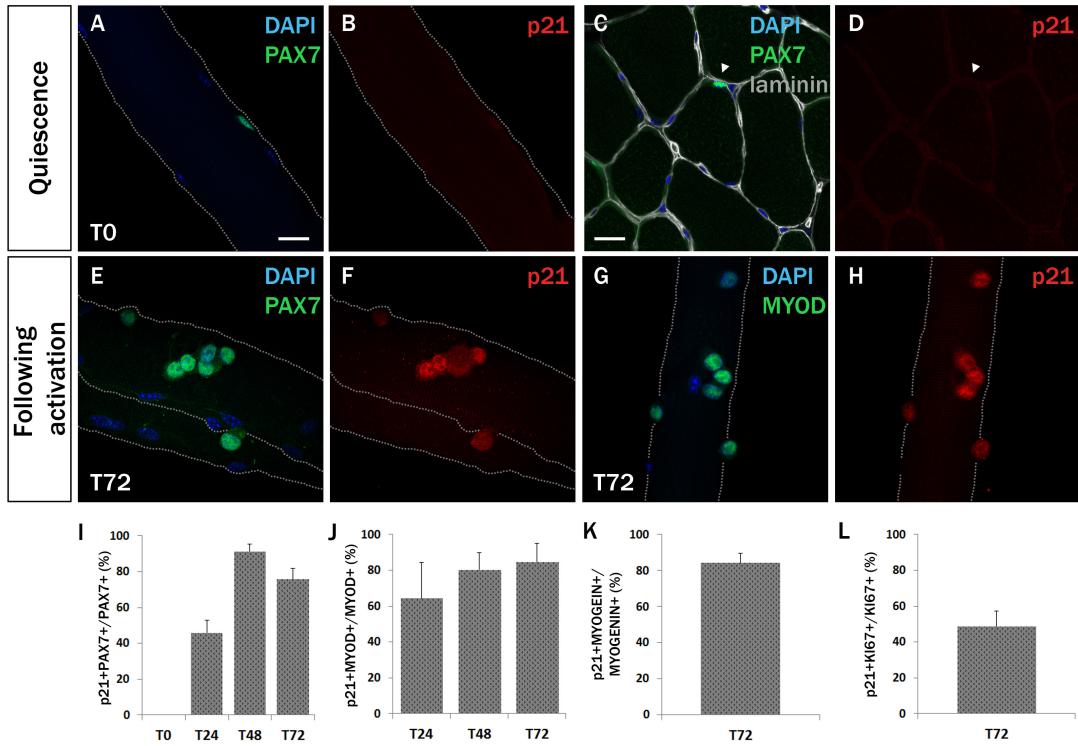


Figure 1

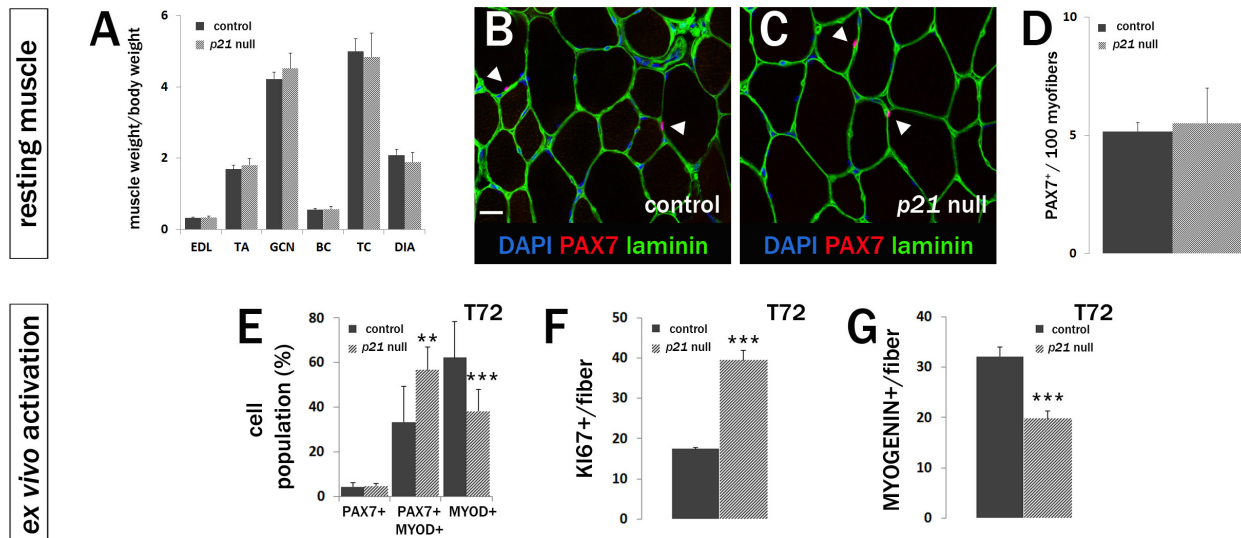


Figure 2

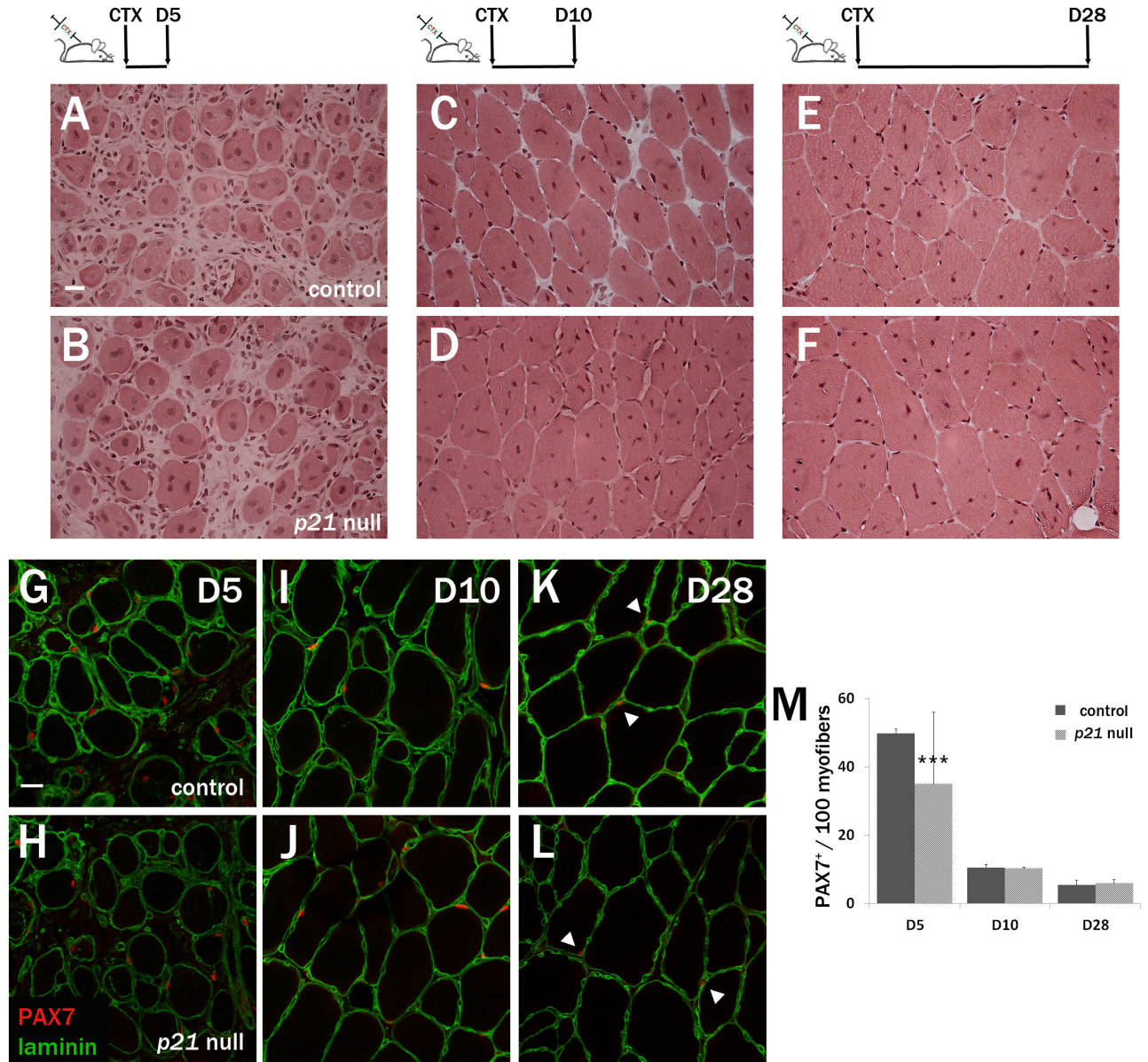


Figure 3

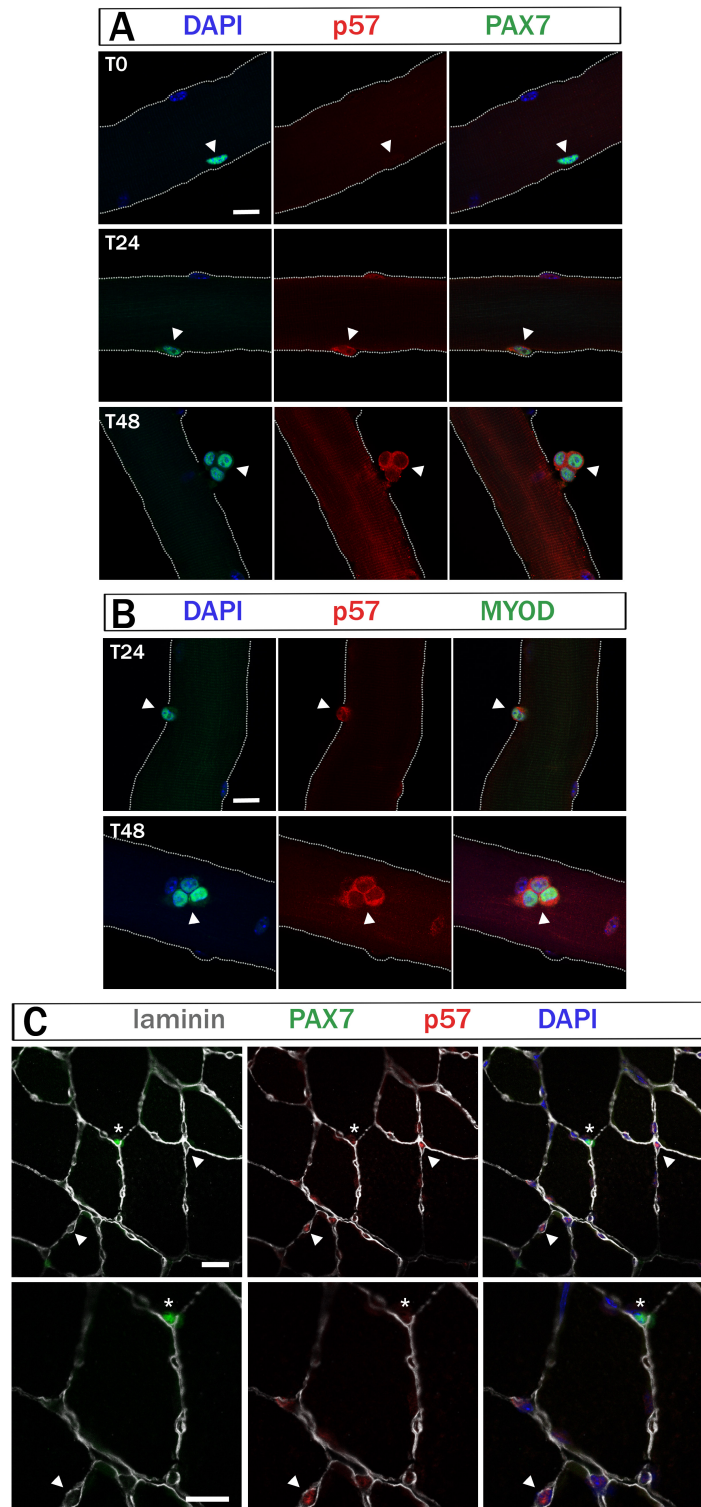


Figure 4

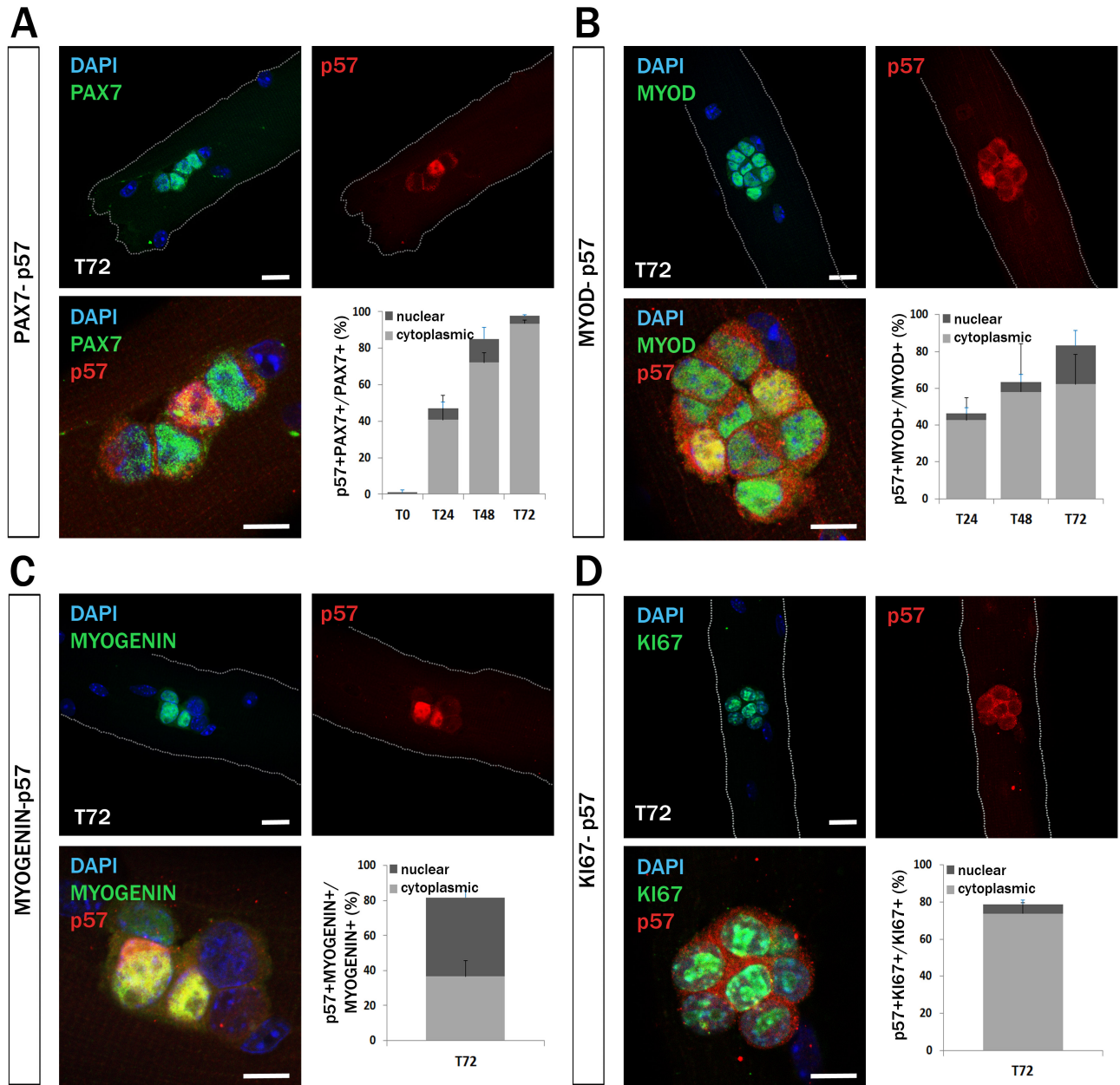


Figure 5

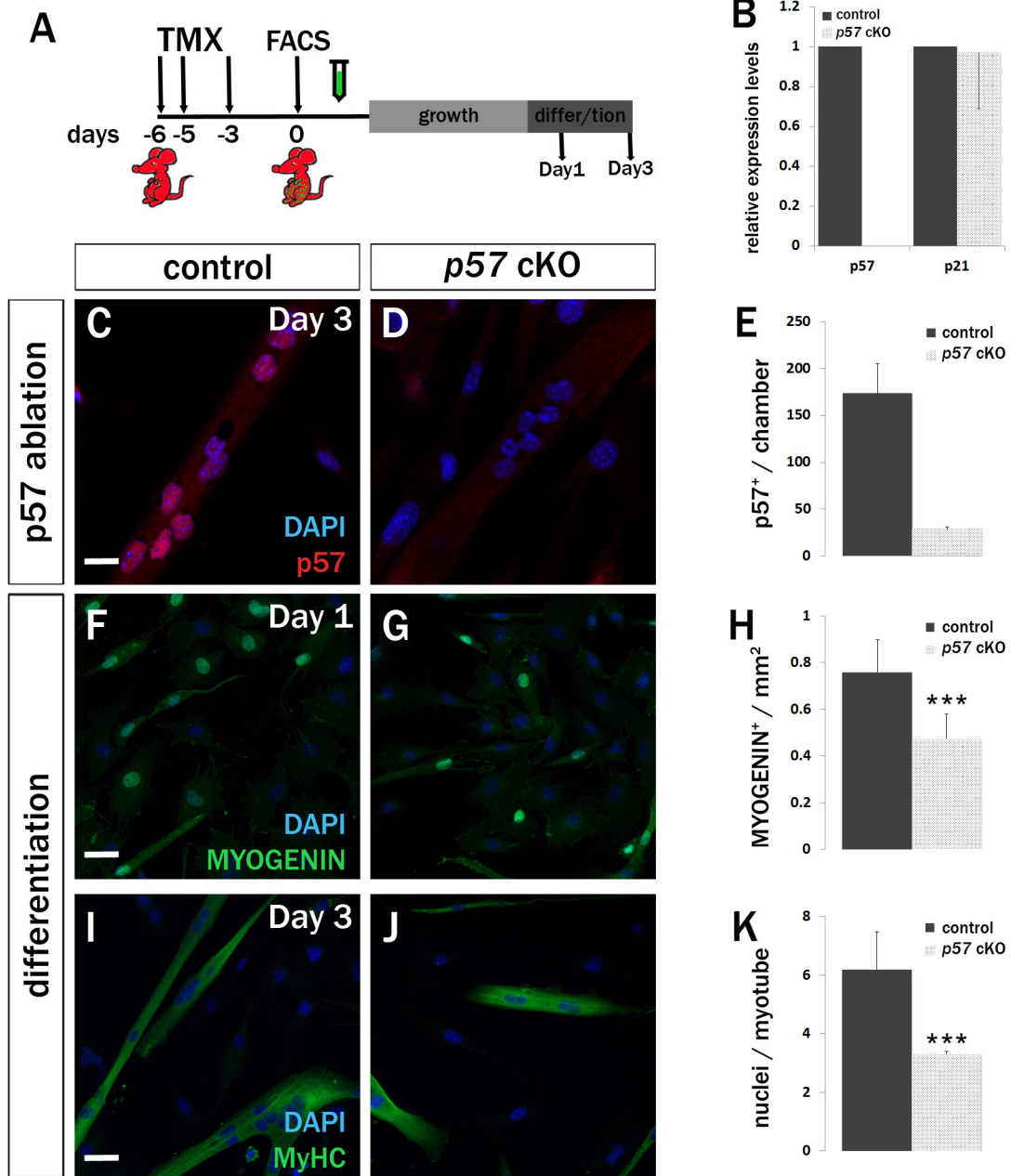


Figure 6

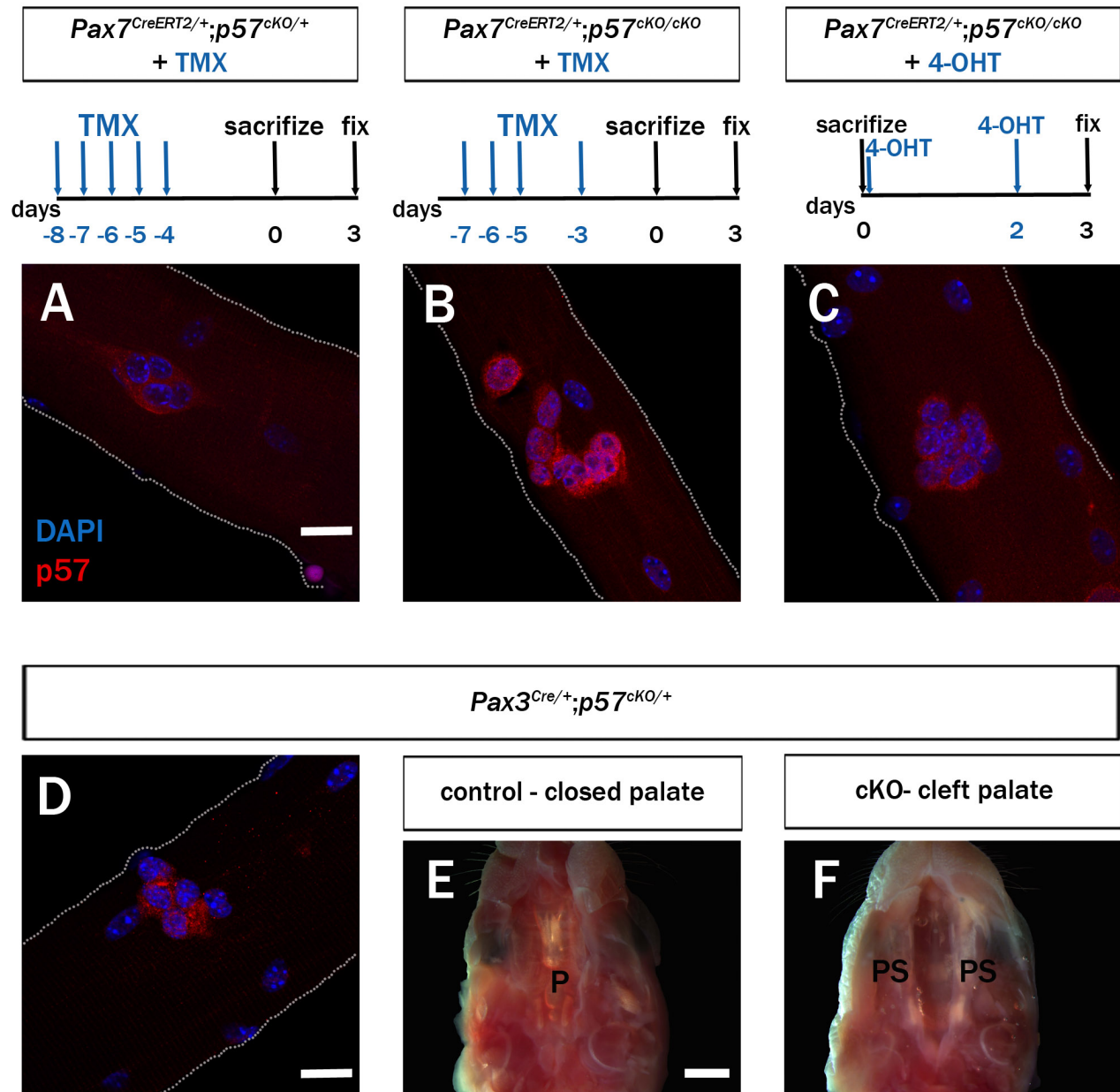


Figure S1

DISCUSSION

CDKIs, MRFs and Notch signaling interplay in cell cycle exit during development

The balance between proliferation and differentiation ensures normal tissue development during embryogenesis. Unregulated proliferation or premature differentiation can have deleterious defects. Thus, much effort is placed on understanding the mechanisms which define the equilibrium between sufficient propagation and differentiation, supporting the formation of morphologically and functionally correct structures. In the developing skeletal muscle, previous studies have identified key signaling pathways and factors controlling these procedures and we showed how they are coordinated in the co-existing and interacting proliferating progenitor and differentiating myoblast populations of the nascent muscle masses [Zalc et al., 2014].

One of the hallmarks of differentiation is the acquisition of a post-mitotic state, leading to the general assumption that growth arrest and differentiation are interconnected. During skeletal muscle formation in the embryo and fetus, cell cycle exit occurs at the transition from committed progenitors to determined myoblasts. Indeed, at the course of differentiation from Pax3/7+ progenitors to Myogenin+ myoblasts we observed progressively increasing expression of cyclin-dependent kinase inhibitors (CDKIs). This is consistent with the observed Hes/Hey repressive activity on Myogenic Regulator Factor (MRF)-mediated transactivation of the identified p57MRE, as our data and previous studies suggest high Notch and high MRF activity in muscle progenitors and in committed/differentiating cells, respectively. Monitoring the cell cycle status upon differentiation impairment (using MyoD mutant embryos) and vice-versa (using p21/p57 double knock-outs) showed that cell cycle exit and differentiation are synchronous, yet uncoupled [Zalc et al., 2014]. In the absence of MyoD, limb myogenesis is delayed and Myf5 is maintained at high levels [Kablar et al., 1997; Kablar et al., 1998]. We found that Myf5+Pax3/7- cells expressed p57 at E12.5 MyoD^{-/-} forelimbs [Zalc et al., 2014]. Given our model whereby MyoD binds and activates p57 via the p57MRE sequence, we assume that Myf5 operates similarly. Pax7+Myf5- undifferentiated progenitors were shown to represent a slow-cycling pool, while Pax7+Myf5+ cells correspond to a committed fast-cycling population [Picard & Marcelle, 2013]. Future studies are expected to address the subtle cell cycle regulation of those populations and the role of Myf5 in cell proliferation changes.

Apart from cell cycle regulation, p57 functions in muscle through a positive feedback loop with MyoD. Evidence from myogenic cell lines and zebrafish implicate it in MyoD stabilization (see below). It is not known whether the same mechanism operates in murine embryonic myogenesis. However, it could be suggested that growth arrest and differentiation are initially independent but subsequently synergize to drive terminal differentiation. Accordingly, p21/p57 double knock-out mice show impaired Mef2c expression [Zhang et al., 1999b], implicating p21 and p57 in the regulation of terminal differentiation.

In the model of coordinated regulation that we propose, the interplay of Notch-mediated p57 repression in Pax3/7 progenitors and the MRF-mediated p57 activation in myoblasts is essential for growth arrest. A similar Notch-based interplay of progenitor and committed cells has recently been proposed for the airway epithelium [Pardo-Saganta et al., 2015]. During embryonic and postnatal myogenesis, Notch signaling is crucial for sustaining progenitors and stem cells, while myogenic differentiation requires Notch downregulation [Mourikis & Tajbakhsh, 2014]. Further studies are required to precise the molecular network among PAX/MRF regulation, cell growth arrest, and Notch switch. Even though p57 and p21 are absent from cycling cells of E11.5 forelimbs under normal conditions [Zalc et al., 2014], we observed cycling Pax3+p57+ or Pax3+p21+ progenitors in *Pax3^{Cre/+}; Rbpj^{flox/flox}* E11.5 forelimbs [Zalc et al., 2014]. Although the latter might correspond to a transitory state, it is also possible that Notch regulates both cell cycle progression and arrest through a complex mechanism. As *Rbpj* ablation does not cause a complete growth arrest or differentiation of Pax3 progenitors [Zalc et al., 2014], other pathways might be involved. Indeed, in postnatal myogenesis the balance of Notch and Wnt or Notch and TGF- β /pSmad3 orchestrates proliferative expansion and differentiation [Brack et al., 2008; Carlson et al., 2008]. The opposing cell cycle effects of Notch and TGF- β are imposed through the control of CDKs, including p21 [Carlson et al., 2008].

Our study in embryonic myogenesis and growth arrest suggests a model for Notch/MRF crosstalk through p57 regulation. Whether the same mechanism applies in postnatal myogenesis remains to be shown, but a few lines of evidence render it a possible scenario. Firstly, p57MRE was identified by a

MyoD CHIP-seq in C2C12 cells, raising the possibility that it can be reused in adult muscle cells. Secondly, Notch has a well-established role in maintenance as well as homing of satellite cells [Bjornson et al., 2012; Bröhl et al., 2012; Mourikis et al., 2012b; Wen et al., 2012]. Thirdly, p57 has an emerging role stem cell homeostasis, including hematopoietic [Matsumoto et al., 2011; Zou et al., 2011], neural [Furutachi et al., 2013], and lung [Zacharek et al., 2011] stem cells. Fourthly, p57 is transcriptionally repressed by Notch effectors in several systems, apart from the embryonic muscle [Zalc et al., 2014], including neural plate cells [Park et al., 2005], pancreatic progenitors [Georgia et al., 2006], pituitary [Monahan et al., 2009], and thyroid gland [Carre et al., 2011]. It would be interesting to speculate that CDKI modulation represents a general safeguard mechanism for fine tuning of stem/progenitor-based prenatal growth and postnatal regeneration avoiding structure malformation and/or cancer.

Conditional p57 ablation for postnatal studies

Developmental data implicate p57 in the majority of the emerging tissues and organs [Zhang et al., 1997; Yan et al., 1997; Susaki et al., 2009], while by regulating cell cycle arrest, apoptosis, and cell mobility, p57 is a putative tumor suppressor [Kavanagh & Joseph, 2011]. However, *in vivo* p57 studies in the adult organism require the development of genetic tools that bypass the perinatal lethality of p57-deficient mice.

We generated a mouse model that allows conditional ablation of the coding region of p57 by the loxP/Cre system [Mademtzoglou et al., submitted-1]. We demonstrated that ubiquitous Cre expression results in p57 recombination and loss. Cre-induced p57 absence reproduced the phenotypes observed in p57-null mice, confirming its indispensable role during embryonic development [Zhang et al., 1997; Yan et al., 1997; Susaki et al., 2009; Mademtzoglou et al., submitted-1]. p57 expression has been associated with key differentiation steps of various organs [Matsuoka et al., 1995; Westbury et al., 2001]. Its loss caused proliferation, apoptosis, and differentiation defects in various tissues, such as skeleton, cranio-facial structures, sensory organs,

gastro-intestinal tract, and reproductive system [Yan et al., 1997; Zhang et al., 1997; Takahashi et al., 2000; Susaki et al., 2009; Mademtzoglou et al., submitted-1]. p57 ablation in a spatially- and temporally- defined manner during development will allow the characterization of the cell population(s) and time window, in which p57 function is crucial for individual structures. Furthermore, postnatal-specific p57 abrogation overcomes perinatal lethality and facilitates investigation of stem cell function and cell fate decisions.

The generated mice also contain a β -galactosidase reporter. We showed that this reporter faithfully recapitulates the expression profile that was described for p57, including widespread expression during development and highly localized in the adult [Matsuoka et al., 1995; Yan et al., 1997; Zhang et al., 1997; Westbury et al., 2001; Mademtzoglou et al., submitted-1]. Since early studies on p57 expression relied on northern and western analysis of entire organs, the new mouse provides a fast reporter-based way for screening of p57-expressing cell populations to define focused functional analyses.

Differences in genetic backgrounds in humans or model organisms affect responses to various factors, including disease susceptibility [Chow, 2016]. In our study, $p57^{FL-ILZ/+}$ mice showed improved survival in mixed 129S2 X C57Bl/6N or 129S2 X C57Bl/6J background compared to C57Bl/6J. Similarly, the effects of excessive p57 on embryonic growth retardation were responsive to genetic background [Andrews et al., 2007]. Strain background can also influence p57 imprinting, with minor expression of the paternal allele being detectable in certain crosses [Park & Chung, 2001].

CDKIs in the control of satellite cells

Regenerative myogenesis in adult skeletal muscle is of vital importance for recovery from injuries, but can be compromised by degenerative or disease states, including aging, that affect the functional capacity of satellite cells, namely the muscle stem cells. Central in their function preservation is the entry into a reversibly dormant state (i.e. quiescence) whenever they are not contributing to muscle formation, meaning when postnatal growth or post-injury repair ceases. Thus, unraveling their cell cycle transitions and (re)-acquisition of quiescence has gained significant interest in recent years.

Focusing on cell cycle exit signals, we hypothesized that p21 and p57 might drive adult satellite cells out of the cell cycle to promote myogenic differentiation or self-renewal of the quiescent pool. They have been shown to redundantly control embryonic muscle cell cycle status and differentiation [Zhang et al., 1999b], while the adult regenerative program is postulated to recapitulate features of embryonic myogenesis [Tajbakhsh, 2009]. In homeostasis, we found no p21 protein in quiescent satellite cells or any other muscle cell type [Mademtzoglou et al., submitted-2], complementing previous reports for negligible p21 mRNA [Macleod et al., 1995] or protein [Franklin & Xiong, 1996] in adult muscle. On the contrary, skeletal muscle is one of the tissues with high levels of p57 transcript in adult life [Matsuoka et al., 1995; Park & Chung, 2001] and we found p57 protein to be abundant in interstitial cells [Mademtzoglou et al., submitted-2], giving insight to its intramuscular localization. However, it was missing from the satellite cell compartment in muscle sections and in freshly isolated single myofibers [Mademtzoglou et al., submitted-2], in line with a previous study on mRNA analysis of FACS-sorted postnatal satellite cell populations at different ages and dormancy states [Chakkalakal et al., 2014]. The only report on p57 presence in quiescent satellite cells [Fukada et al., 2007] is based on FACS-sorting with a novel antibody that the same group previously described [Fukada et al., 2004]; nevertheless, this antibody is also immuno-recognizing a fraction of bone marrow cells [Fukada et al., 2004], while p57 has a well-established role and presence in the hematopoietic lineage [Matsumoto et al., 2011; Zou et al., 2011]. Furthermore, p57 immunostaining in Fukada et al. [2007] was performed with an antibody that recognizes a sequence in the carboxy-terminal region of p57, which

might cross-react with the respective homologous domain of p27 [Matsuoka et al., 1995; Galea et al., 2008; Pateras et al., 2009]. Combining our and previous observations [Fukada et al., 2004; 2007; Chakkalakal et al., 2014; Mademtzoglou et al., submitted-2], we conclude that quiescent satellite cells are p57 negative. Instead, they express p27, the last CDKI of the family encompassing p21 and p57 [Chakkalakal et al., 2014; our unpublished observation]. However, this observation does not preclude p21 and p57 participation in satellite cell cycle dynamics; in the pituitary p57 was found to promote cell cycle exit, while p27 prevented cell cycle re-entry [Bilodeau et al., 2009].

Upon satellite cell activation p21 and p57 rapidly became upregulated. Given their role in growth arrest during embryonic myogenesis [Zhang et al., 1999b], we did not expect p21 and p57 at early post-activation stages when myoblasts are dividing and no differentiation or self-renewal occurs [Zammit et al., 2004]. Their presence in actively proliferating cells seems intuitively contradictory to their role as cell cycle inhibiting molecules. However, p21 has been associated with cell cycle progression by promoting the assembly of cyclin-CDK complexes and their nuclear localization, whereas it induces growth arrest when it stoichiometrically exceeds these complexes [Xiong et al., 1993; Firpo et al., 1994; Michieli et al., 1994; Nourse et al., 1994; Zhang et al., 1994; Harper et al., 1995; Liu et al., 1996; LaBaer et al., 1997]. Thus, p21 was proposed to “titrate” proliferation by impeding it at zero/high amounts but supporting it in intermediate concentrations. In MyoD-converted fibroblasts p21 is involved in nuclear accumulation and activity of cyclin-Cdk complexes [Peschiaroli et al., 2002], possibly extending the previous observation in the myogenic context. Furthermore, experiments with C2C12, a widely used mouse myoblast cell line [Yaffe & Saxel, 1977] that can differentiate into myocytes, showed that the interaction of p21 with Cdk4 was higher in myocytes than myoblasts, leading to Cdk4 inhibition [Wang & Walsh, 1996b]. Whether these findings can be generalized to our experimental system remains to be shown. Of note, genetic ablation of p21 led to an increase in Ki67+ myoblasts in our *ex vivo* myofiber culture [Mademtzoglou et al., submitted-2], favoring a stronger contribution of p21 to the inhibition than promotion of Cdk activity and cell cycle progression.

We also found p57 in activated, proliferating cells, but mostly restricted to their cytoplasm, while in differentiating populations it became progressively nuclear. The molecular events regulating this shuttling remain to be identified. Cytoplasmic p57 was found to associate with LIM-kinase1 and regulate cell motility [Vlachos & Joseph, 2009; Chow et al., 2011; Guo et al., 2015]. We asked whether a same mechanism is found in muscle, given that one of the early manifestations of satellite cell activation is their mobilization [Siegel et al., 2009; Marg et al., 2014; Webster et al., 2016]. However, I was not able to detect *LIM-kinase1* expression in single myofiber cultures. Future studies with scratch assays in primary myoblast culture might show if it is associated with mobility with other, yet uncharacterized, partners. A second possibility is that p57 shuttling is related to cell cycle dynamics and differentiation progression. Such regulation on the cellular level, has previously been described for ERK in muscle progenitors [Michailovici et al., 2014]. Post-translational modifications of p57 protein might restrain it to the cytoplasm to allow cell cycle to progress. Alternatively, p57 might help the assembly of recently produced cyclin and CDK molecules in the cytoplasm, as p21 and p27 do (see above). In their absence, p57 was shown to resume their role in stabilizing cyclin-CDK complexes in mouse embryonic fibroblasts [Cerqueira et al., 2014]. Finally, it could be involved in the relocalization of Cdk or cyclins, as is the case for fibroblasts with ectopic MyoD and hematopoietic stem cells, respectively [Figliola & Maione, 2004; Zou et al., 2011]. Further studies are required to better understand the p57 nucleo-cytoplasmic translocation in activated and differentiating myoblasts. The fact that it persisted in the cytoplasm of half of the differentiating myoblasts [Mademtzoglou et al., submitted-2] complicates our speculation, but is consistent with previous data on p57 cytoplasmic distribution in myotubes forming from MyoD-converted fibroblasts [Figliola & Maione, 2004]. On the contrary, p21 did not undergo such nucleo-cytoplasmic shuttling [Mademtzoglou et al., submitted-2]. Similarly, when fibroblasts were induced to myogenic fate by MyoD, p21 had strictly nuclear presence in the forming myotubes [Figliola & Maione, 2004]. Moreover, p21, but not p57 or p27, appeared in the nucleus of Ki67+ proliferating b-cells, when proliferation was induced by adenoviral expression of cyclins and Cdks [Fiaschi-Taesch et al., 2013]. In general, under normal conditions, p21 is not expected to persist in the cytoplasm, as its presence there is linked to degradation [Hwang et al., 2009] or oncogenesis [Besson et al., 2004; Besson et al., 2008].

Upregulation of p21 and p57 upon satellite cell activation might be related to MyoD, which is also absent from quiescent satellite cells but appears as early as three hours post-activation *ex vivo* [Zammit et al., 2004; Zhang et al., 2010] or six hours post-injury *in vivo* [Grounds et al., 1992]. Both p21 and p57 are induced by MyoD, while later they enhance its activity and stabilization [Halevy et al., 1995; Reynaud et al., 1999; Reynaud et al., 2000b; Tintignac et al., 2004; Osborn et al., 2011; Battistelli et al., 2014; Zalc et al., 2014]. Interestingly, the p57-MyoD binding engages the p57 helix domain that was found indispensable for Cdk/cylin inhibition [Hashimoto et al., 1998; Reynaud et al., 2000b], possibly explaining p57 presence in cycling cells. MyoD is expressed in activated/proliferating myoblasts [Zammit et al., 2004; Liu et al., 2012] and has been proposed to be required for the transition from quiescence to the cell cycle, acting through the replication-related factor Cdc6 [Zhang et al., 2010]. MyoD deficient myoblasts divide in a perturbed way, with divisions taking longer and being less frequent [Megeny et al., 1996]. Curiously, despite MyoD presence in myoblasts, they continue to proliferate and do not proceed to differentiation for several days *in vitro* or *in vivo* [Tajbakhsh, 2009]. It is suggested that additional factors, including Myogenin, are needed to initiate/enhance transcription of at least a subset of MyoD targets [Blais et al., 2005; Cao et al., 2006]. This might also explain why exogenous MyoD induces growth arrest in non-myogenic cell lines [Crescenzi et al., 1990; Sorrentino et al., 1990], but not in myogenic ones or satellite cell-derived myoblasts.

p21 has been implicated at the early post-activation steps of satellite cells. Using the *ex vivo* system of floating myofibers my work shows that p21 is present in myoblasts soon after satellite cells are activated [Mademtzoglou et al., submitted-2], while during muscle regeneration, it was shown to peak at 3-4 days post-injury and then decline [Yan et al., 2003; our unpublished observation]. Conversely, resting muscle and quiescent satellite cells are devoid of p21 [Mademtzoglou et al., submitted-2], while muscle development in p21 mutants could to be rescued by p57 [Zhang et al., 1999b]. Thus, we hypothesized that under homeostatic conditions adult muscle would not be affected by p21 loss, while we expected deficits in satellite cell activation/differentiation and possibly muscle regeneration after injury. Indeed, resting muscles of p21 knock-out mice were indistinguishable from control littermates muscles in terms of muscle weight, histology, and satellite

cell compartment [Mademtzoglou et al., submitted-2], in line with previous observations [Deng et al., 1995; Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015]. However, p21 null mice had significantly less satellite cells at an early regeneration time point (i.e. D5), although starting with similar pre-regeneration satellite cell amounts [Mademtzoglou et al., submitted-2]. Moreover, regeneration was not overtly affected, as evidenced by the restoration of the satellite cell pool as well as muscle architecture (e.g. myofiber formation, myofiber morphology, central nucleation, interstitium minimization to pre-injury levels) [Mademtzoglou et al., submitted-2]. Similarly, satellite cell numbers [Chakkalakal et al., 2014] and muscle structure [Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015] one month post-injury revealed successful regeneration in previous studies, despite occasional defects at earlier time points [Hawke et al., 2003; Chinzei et al., 2015]. Furthermore, p21 absence did not compromise the long-term regeneration potential, as evidenced by repeated injuries and muscle recovery [Chakkalakal et al., 2014].

To get a better insight into the early activation/differentiation phase, we used single myofiber cultures. Although this system bears the disadvantages of an *ex vivo* situation, it might reflect a more physiological activation compared to the complete degeneration that is experimentally performed but rarely occurring in nature. When we cultured isolated myofibers from p21 mutant and control mice, activated satellite cells showed increased proliferation and decreased myogenic differentiation [Mademtzoglou et al., submitted-2]. This is indicative of an early defect, in line with our *in vivo* observations and previous data [Hawke et al., 2003; Chakkalakal et al., 2014; Chinzei et al., 2015]. AAV-mediated acute p21 knock-down triggered proliferation of quiescent cells (including satellite cells) and increased cellularity; nevertheless, when the delivered siRNA was used in primary myoblast culture it did not affect differentiation [Biferi et al., 2015]. The difference with previous studies was attributed to the effect of acute versus chronic p21 loss. Future experiments, including conditional genetic ablation of p21 with the loxP-Cre system would allow testing this hypothesis.

In the absence of p57 there was a differentiation deficit *in vitro* and myoblasts did not advance through the myogenic program. p57 correlates with differentiation in various embryonic tissues [Westbury et al., 2001]. It is highly and rapidly upregulated upon *in vitro* myogenic differentiation

[Reynaud et al., 1999], while it has even been considered as synonymous to it [Mounier et al., 2011]. However, the consequences of its abrogation on adult muscle have not been estimated, partially because of the perinatal lethality of p57 mutant mice, which we overpassed with the new conditional knock-out we generated. With the caveat that our *in vitro* system lacks the structural, neurogenic, and metabolic fidelity of the muscle tissue [Grounds, 2014], our data suggest that p57 is involved in adult myogenic differentiation. Although we wished to verify these findings in the *in vivo* context, our analysis was hindered by inefficient recombination [Mademtzoglou et al., submitted-2]. Previous studies on satellite cells suggest that the cell cycle state (quiescence versus activation/proliferation) may interfere with recombination resistance/success [Lepper et al., 2009; Günther et al., 2013; von Maltzahn et al., 2013]. Given these findings, a more appropriate scheme for tamoxifen-inducible Cre-mediated p57 recombination might include administration of tamoxifen during muscle regeneration. It will be interesting to monitor how p57 influences the post-regeneration re-establishment of the satellite cell compartment, given its emerging importance in the quiescence and maintenance of other stem cells, such as hematopoietic [Matsumoto et al., 2011; Zou et al., 2011], neural [Furutachi et al., 2013], and lung stem cells [Zacharek et al., 2011].

Overall, we conclude that p21 and p57 are essential at the early steps following satellite cell activation. Their presence seems compatible with activation/proliferation and possibly represents an early activation event. Their loss profoundly affects *ex vivo* myogenic differentiation. Our data so far indicate that p21 and p57 function in distinct ways during adult and embryonic myogenesis, in terms of early versus continuous myogenesis support. It remains to be established whether the preliminary differentiation/regeneration defects are rescued by p21 and p57 in p57 and p21 mutants, respectively. Satellite cell-specific double p21/p57 knock-out may elucidate their relative contributions and putative redundancies in the adult. A better understanding of cell cycle regulation in satellite cells is imperative to define the molecular events underlying their long-term preservation - through (re)-entry into quiescence- and their rapid response to regenerative needs -through prompt and tightly controlled activation. Defective stem cell cycle dynamics and continuous activation/proliferation can lead to DNA damage accumulation, apoptosis, pool exhaustion, and inability to support homeostatic or regenerative demands.

ANNEX

Further contribution to projects of the partner labs

Gene expression profiling of muscle stem cells identifies novel regulators of postnatal myogenesis

Alonso-Martin S, Rochat A, Mademtzoglou D, Morais J, De Reynies A, Auradé F, Chang T, Zammit PS, Relaix F

Skeletal muscle growth and regeneration require a population of muscle stem cells, the satellite cells, located in close contact to the myofiber. These cells are specified during fetal and early postnatal development in mice from a Pax3/7 population of embryonic progenitor cells. As little is known about the genetic control of their formation and maintenance, we performed a genome-wide chronological expression profile identifying the dynamic transcriptomic changes involved in establishment of muscle stem cells through life, and acquisition of muscle stem cell properties. We have identified multiple genes and pathways associated with satellite cell formation, including set of genes specifically induced (EphA1, EphA2, EfnA1, EphB1, Zbtb4, Zbtb20) or inhibited (EphA3, EphA4, EphA7, EfnA2, EfnA3, EfnA4, EfnA5, EphB2, EphB3, EphB4, EfnBs, Zfp354c, Zcchc5, Hmga2) in adult stem cells. Ephrin receptors and ephrins ligands have been implicated in cell migration and guidance in many tissues including skeletal muscle. Here we show that Ephrin receptors and ephrins ligands are also involved in regulating the adult myogenic program. Strikingly, impairment of EPHB1 function in satellite cells leads to increased differentiation at the expense of self-renewal in isolated myofiber cultures. In addition, we identified new transcription factors, including several zinc finger proteins. ZFP354C and ZCCHC5 decreased self-renewal capacity when overexpressed, whereas ZBTB4 increased it, and ZBTB20 induced myogenic progression. The architectural and transcriptional regulator HMGA2 was involved in satellite cell activation. Together, our study shows that transcriptome profiling coupled with myofiber culture analysis, provides an efficient system to identify and validate candidate genes implicated in establishment/maintenance of muscle stem cells. Furthermore, tour de force transcriptomic profiling provides a wealth of data to inform for future stem cell-based muscle therapies.

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Gene Expression Profiling of Muscle Stem Cells Identifies Novel Regulators of Postnatal Myogenesis

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Skeletal muscle growth and regeneration require a population of muscle stem cells, the satellite cells, located in close contact to the myofiber. These cells are specified during fetal and early postnatal development in mice from a Pax3/7 population of embryonic progenitor cells. As little is known about the genetic control of their formation and maintenance, we performed a genome-wide chronological expression profile identifying the dynamic transcriptomic changes involved in establishment of muscle stem cells through life, and acquisition of muscle stem cell properties. We have identified multiple genes and pathways associated with satellite cell formation, including set of genes specifically induced (*EphA1*, *EphA2*, *EfnA1*, *EphB1*, *Zbtb4*, *Zbtb20*) or inhibited (*EphA3*, *EphA4*, *EphA7*, *EfnA2*, *EfnA3*, *EfnA4*, *EfnA5*, *EphB2*, *EphB3*, *EphB4*, *EfnBs*, *Zfp354c*, *Zcchc5*, *Hmga2*) in adult stem cells. Ephrin receptors and ephrins ligands have been implicated in cell migration and guidance in many tissues including skeletal muscle. Here we show that Ephrin receptors and ephrins ligands are also involved in regulating the adult myogenic program. Strikingly, impairment of EPHB1 function in satellite cells leads to increased differentiation at the expense of self-renewal in isolated myofiber cultures. In addition, we identified new transcription factors, including several zinc finger proteins. ZFP354C and ZCCHC5 decreased self-renewal capacity when overexpressed, whereas ZBTB4 increased it, and ZBTB20 induced myogenic progression. The architectural and transcriptional regulator HMGA2 was involved in satellite cell activation. Together, our study shows that transcriptome profiling coupled with myofiber culture analysis, provides an efficient system to identify and validate candidate genes implicated in establishment/maintenance of muscle stem cells. Furthermore, tour de force transcriptomic profiling provides a wealth of data to inform for future stem cell-based muscle therapies.

Keywords: skeletal muscle, myogenesis, satellite cells, ephrins, zinc fingers

Q14 115 INTRODUCTION

Q6 116 During vertebrate development, successive phases of embryonic
117 and fetal myogenesis leads to formation and growth of
118 skeletal muscles (Relaix et al., 2005; Relaix, 2006; Buckingham
119 and Relaix, 2007). Skeletal muscle cells of trunk and limbs
120 in mouse originate from the early somites, which appear
121 at mid-gestation from undifferentiated presomitic mesoderm
122 (Tajbakhsh and Buckingham, 2000). Following several steps
123 of somite maturation, a population of muscle progenitor cells
124 (MPC) that express the paired-box/homeobox transcription
125 factors *Pax3* and *Pax7* emerge in the central region of the
126 developing somite. Similar cell populations are also found in
127 head muscles, though using a different set of transcriptional
128 regulators (Sambasivan et al., 2011). MPC will both self-renew
129 and give rise to all skeletal muscles via activation of a family
130 of four muscle-specific bHLH transcription factors (*Myf5*, *Mrf4*,
131 *MyoD*, and *Myog*: myogenin) that induce the myogenic program
132 (Bismuth and Relaix, 2010; Murphy and Kardon, 2011). Around
133 birth, while all MPC maintain the expression of *Pax7*, *Pax3*
134 expression in only maintained in a subset of muscles (Relaix
135 et al., 2006) (unpublished observations). MPC become in close
136 contact with the muscle fibers in response to different signals,
137 such as those from the Notch pathway (Seale et al., 2000;
138 Zammit et al., 2006a; Tajbakhsh, 2009; Brohl et al., 2012). During
139 establishment of this anatomical niche, emerging satellite cells
140 acquire stem cell-specific characteristics, including self-renewal
141 capacity (Mauro, 1961; Zammit et al., 2006a; Relaix and Marcelle,
142 2009). During postnatal muscle growth, satellite cells supply
143 myonuclei to maturing myofibers up to postnatal day 21 (P21)
144 before becoming mitotically quiescent (Lepper et al., 2009; White
145 et al., 2010). Adult satellite cells can be activated from their
146 mitotically quiescent state upon injury (Wang and Rudnicki,
147 2011; Relaix and Zammit, 2012), to proliferate, and co-express
148 MYOD and PAX7. They then differentiate via activation of *Myog*
149 (and down-regulation of *Pax7*) to repair damaged myofibers,
150 while a subpopulation of satellite cells will self-renew to restore
151 the pool of quiescent satellite cells by down-regulation of *MyoD*
152 (Zammit et al., 2004; Rudnicki et al., 2008; Relaix and Zammit,
153 2012).
154

155 Understanding regulation of myogenic progression from
156 MPCs to muscle stem cells is central to building a comprehensive
157 model of satellite cell function. Many transcriptional networks
158 that control embryogenesis are also important for myogenesis,
159 such as Notch, BMP or WNT proteins (Linker et al., 2003; Ono
160 et al., 2011; Brohl et al., 2012). Furthermore, a balance between
161 extrinsic cues and intracellular signaling pathways, such as IGF,
162 FGF, Notch, and TGF- β , is required to preserve stem cell function
163 (Brack et al., 2008; Kuang et al., 2008; Brack and Rando, 2012;
164 Dumont et al., 2015).

165 We have characterized the dynamics of skeletal muscle
166 progenitor and postnatal stem cells from embryonic development
167 to adult life, hence deciphering the intrinsic molecular pathways
168 involved in specification and regulation of these muscle
169 stem cells. Using this large microarray analysis of myogenic
170 progenitoris and stem cells during development and adult
171 myogenesis, we identified and evaluated several new candidate

172 factors mediating satellite cell specification and function, with
173 a focus here on EPHB1 and several transcriptional regulators,
174 including four zinc finger transcription regulators (*Zfp354c*,
175 *Zcchc5*, *Zbtb4*, and *Zbtb20*) and HMG2, a transcriptional co-
176 regulator belonging to the HMGI family of small high-mobility-
177 group (HMG) proteins (Zhou et al., 1995).
178

Eph Receptors and Ephrin Ligands

179 Eph/ephrin signaling has been shown to regulate muscle satellite
180 cell motility and patterning (Stark et al., 2011), but has
181 not been linked with regulation of the myogenic program.
182 Eph receptors belong to a large family of receptor tyrosine
183 kinases (RTK) involved in cell contact-dependent signaling and
184 patterning (Pitulescu and Adams, 2010). EPHs are classified
185 as EphAs or EphBs based on their binding affinity for the
186 ephrin ligands, ephrin-A (EFNA) or ephrin-B (EFNB) (Figures
187 S1A,B). EFNAs are GPI (glycosylphosphatidylinositol)-anchored
188 and lack a cytoplasmic domain while EFNBs are attached to the
189 membrane by a single transmembrane domain containing a short
190 cytoplasmic PDZ-binding motif (Pasquale, 2005). Interestingly,
191 both Eph receptors and ephrin ligands are competent to
192 signal following interaction (forward and reverse signaling,
193 respectively), and both *trans* and *cis* signaling have been
194 described (Arvanitis and Davy, 2008; Pitulescu and Adams,
195 2010). In addition, Eph/ephrin signaling is often part of a
196 complex signaling network of regulatory pathways, for instance
197 with adhesion molecules, other cell surface receptors or channels
198 and pores (Arvanitis and Davy, 2008).
199

200 Eph/ephrin interaction leads to a large set of developmental
201 processes and biological responses, including adhesion and
202 repulsion, increased or reduced motility, cell plasticity,
203 permeability and morphogenesis, and cell fate specification
204 (Palmer and Klein, 2003; Arvanitis and Davy, 2008). Eph/ephrins
205 are also implicated in regulation of stem cell niches and cancer
206 (Genander and Frisen, 2010; Murai and Pasquale, 2010; Pasquale,
207 2010).
208

Zinc Finger Transcription Factors

209 Zinc finger proteins belong to a large family of transcription
210 regulators subdivided in seven categories. There are about 800
211 zinc finger transcription factors in the human genome, with a
212 third of those containing a KRAB (Krüppel Associated Box)
213 domain, such as ZFP354C (see below) or related sequences
214 as ZBTB4 or ZBTB20 (Lupo et al., 2013). KRAB is the
215 most widespread family of transcription factors in the human
216 genome, but is also found in yeast (*S. cerevisiae*) and worm (*C.*
217 *elegans*) (Ganss and Jheon, 2004). The KRAB protein domain
218 is a powerful repression region that acts as a transcriptional
219 repressor, allowing the binding to co-repressor proteins (Urrutia,
220 2003). KRAB-containing proteins involved in cell proliferation,
221 differentiation, apoptosis, and tumor formation have been
222 described (Urrutia, 2003; Tian et al., 2006; Li et al., 2008).
223

224 *Zfp354c* (*Kid3*, *AJ18*) belongs to the *Kid* family of genes. The
225 corresponding proteins *Kid1*, *Kid2*, and *Kid3*, share a very similar
226 structure: a KRAB domain and 11–13 C2H2 motifs (Figure S1C),
227 these last zinc finger motifs consisting in two cysteine and two
228 histidine residues bonded tetrahedrally to a Zinc ion (Ganss and

229 Jheon, 2004). ZFP354c has been previously described as abundant
230 in the brain (Watson et al., 2000), but its expression has not been
231 tested in skeletal muscle. Interestingly, KRAB/C2H2 zinc finger
232 protein ZFP354C participates in the BMP (bone morphogenic
233 protein) signaling pathway (Jheon et al., 2003), a key regulator
234 of skeletal muscle development and stem cell function (Amthor
235 et al., 1998; Wang et al., 2010; Ono et al., 2011; Sartori et al., 2013).
236 Given the important role of BMP signaling in skeletal muscle
237 biology, ZFP354C is a good candidate as possible regulator of
238 myogenesis.

239 Zinc finger and BTB domain-containing protein 4 (*Zbtb4*,
240 *KALSO-L1*, *Znf903*) is a transcriptional repressor of specificity
241 protein (Sp) transcription factors (Sreevalsan and Safe, 2013),
242 that binds methylated DNA to repress transcription (Filion
243 et al., 2006; Weber et al., 2008). Despite its broad distribution,
244 ZBTB4 seems to be particularly expressed in the brain. In
245 addition, examination of publicly available microarray data sets
246 demonstrated an inverse relationship in the prognostic value
247 and expression of ZBTB4 and the histone methyltransferase
248 EZH2 in tumors from breast cancer patients (Yang et al., 2014).
249 Indeed, polycomb group protein EZH2 controls self-renewal and
250 safeguards the transcriptional identity of skeletal muscle stem
251 cells (Juan et al., 2011).

252 Zinc finger and BTB domain-containing protein 20 (*Zbtb20*,
253 *DPZF*, *Hof*, *Zfp288*) is a member of a subfamily of zinc
254 finger proteins containing C2H2 Krüppel-type zinc fingers
255 and BTB/POZ domains (Mitchellmore, 2002). ZBTB20
256 can function as a transcriptional repressor and plays an
257 essential role in the specification of pyramidal neurons in the
258 developing hippocampus (Nielsen et al., 2007), and promotes
259 astrocytogenesis during neocortical development (Nagao et al.,
260 2016). ZBTB20 is also a regulator of terminal differentiation
261 of hypertrophic chondrocytes (Zhou et al., 2015). This factor
262 has been recently described to be involved in liver regeneration
263 (Weng et al., 2014), and promoting cell proliferation and tumor
264 growth through repression of FOXO1 (Zhao et al., 2014; Kan
265 et al., 2016). *Zbtb20* null mice exhibit severe postnatal growth
266 retardation, metabolic dysfunction and lethality, suggesting that
267 ZBTB20 plays non-redundant roles in multiple organ systems
268 (Sutherland et al., 2009; Cao et al., 2016).

269 Zinc finger, CCHC domain-containing 5 (*Zcchc5*, *Mar3*, *Zhc5*)
270 belongs to the family of the gag-like retrotransposon genes
271 (glycosaminoglycans) exclusively found in mammals, and is
272 considered an ortholog of Ty3/gypsy group. *Zcchc5* is located
273 on the X chromosome, within the dystrophin (*Dmd*) locus
274 (X21.1) in human. The retrotransposition capacity of these genes
275 seems to have been lost, despite retaining an intact reading
276 frame (Brandt et al., 2005). Thus, the retrotransposons of this
277 family are considered as neogenes with new functions, but their
278 impact and regulation is still poorly understood. *Zcchc5* encodes a
279 nuclear protein containing a CX₂CX₄HX₄C DNA-binding motif,
280 also called CCHC domain, allowing DNA binding to regulate
281 transcription. Furthermore, the proteins of the family of genes
282 *Mart*, which includes *Zcchc5*, have been implicated in the control
283 of cell proliferation and apoptosis in cell lines of liver cancer
284 whereas some become up-regulated in regenerating mouse liver
285 (Okabe et al., 2003; Brandt et al., 2005). Interestingly, *Zcchc5* is

286 expressed in skeletal muscles of the limbs (Diez-Roux et al., 2011)
287 (www.eurexpress.org).

288 Architectural Factor HMGA2 (HMGI-C)

289 HMGA2, also called HMGI-C, is a transcriptional co-regulator
290 belonging to the HMGI family of small high-mobility-group
291 (HMG) proteins containing AT-hook DNA binding domains
292 (Zhou et al., 1995). HMGI proteins modulate gene expression
293 by altering chromatin architecture and/or by recruiting other
294 proteins to the transcription regulatory complex (Thanos and
295 Maniatis, 1992; Zhou and Chada, 1998; Pfannkuche et al., 2009).
296 *Hmga2* is highly expressed during embryonic development
297 and down-regulated in most adult tissues (Zhou et al., 1995;
298 Pfannkuche et al., 2009; Ashar et al., 2010). HMGA2 plays
299 an important role in maintaining adult stem/progenitor cells,
300 notably in maintaining neural stem/progenitor cells (Nishino
301 et al., 2008). *Hmga2* is also highly expressed in proliferating
302 skeletal myoblasts during myogenesis, modulating satellite
303 cell activation and proliferation both *in vivo* and *in vitro*
304 (Li et al., 2012). *Hmga2* knockout mice exhibit impaired
305 muscle development and reduced myoblast proliferation, while
306 overexpression of *Hmga2* promotes myoblast growth preventing
307 myoblast differentiation (Li et al., 2012). Thus, HMGA2 is a
308 key regulator of satellite cell activation and skeletal muscle
309 development.
310
311

312 METHODS

313 Mice

314 *Pax3^{GFP/+}* mice (Relaix et al., 2005) were used to isolate
315 MPC by fluorescent activated cell sorting (FACS) of the
316 GFP+ cells. *Pax3^{Cre/+}* mutant mice were kindly provided by
317 Jonathan A. Epstein (Engleka et al., 2005). *R26^{mT-mG}* mice
318 were obtained from The Jackson Laboratory (Stock No: 007576)
319 (Muzumdar et al., 2007). For myofiber cultures C57BL/6J
320 (Janvier[®]) male mice (8 weeks old) were used. For lineage tracing
321 experiments, *Pax3^{Cre/+}* mice were crossed with *R26^{mT-mG}* to
322 obtain *Pax3^{Cre/+}; R26^{mT-mG}* double mutant mice.
323

324 All animals were maintained inside a barrier facility, and
325 all *in vivo* experiments were performed in accordance with the
326 French and European Community guidelines for the care and
327 use of laboratory animals (Project No: 01427.03 approved by
328 MESR and File No: 15-018 from the Ethical Committee of
329 Anses/ENVA/UPEC).
330

331 Fluorescent Activated Cell Sorting

332 Trunk muscle samples (intercostal, pectoral and abdominal)
333 were isolated from the trunk as indicated in **Figure 1A**, at
334 different stages during development and after birth. Muscle
335 were minced and digested in 0.1% Trypsin (Life Technologies[®])
336 and 0.1% Collagenase D (Roche[®]) in DMEM High Glucose
337 without phenol red (Life Technologies[®]). Digested muscles after
338 filtration were cell-sorted by flow cytometry using a FACS Aria II,
339 using FITC channel to recover the GFP+ cells from *Pax3^{GFP/+}*
340 mice. GFP+ cells were stained using propidium iodide to exclude
341 dead cells (Figure S2A).
342

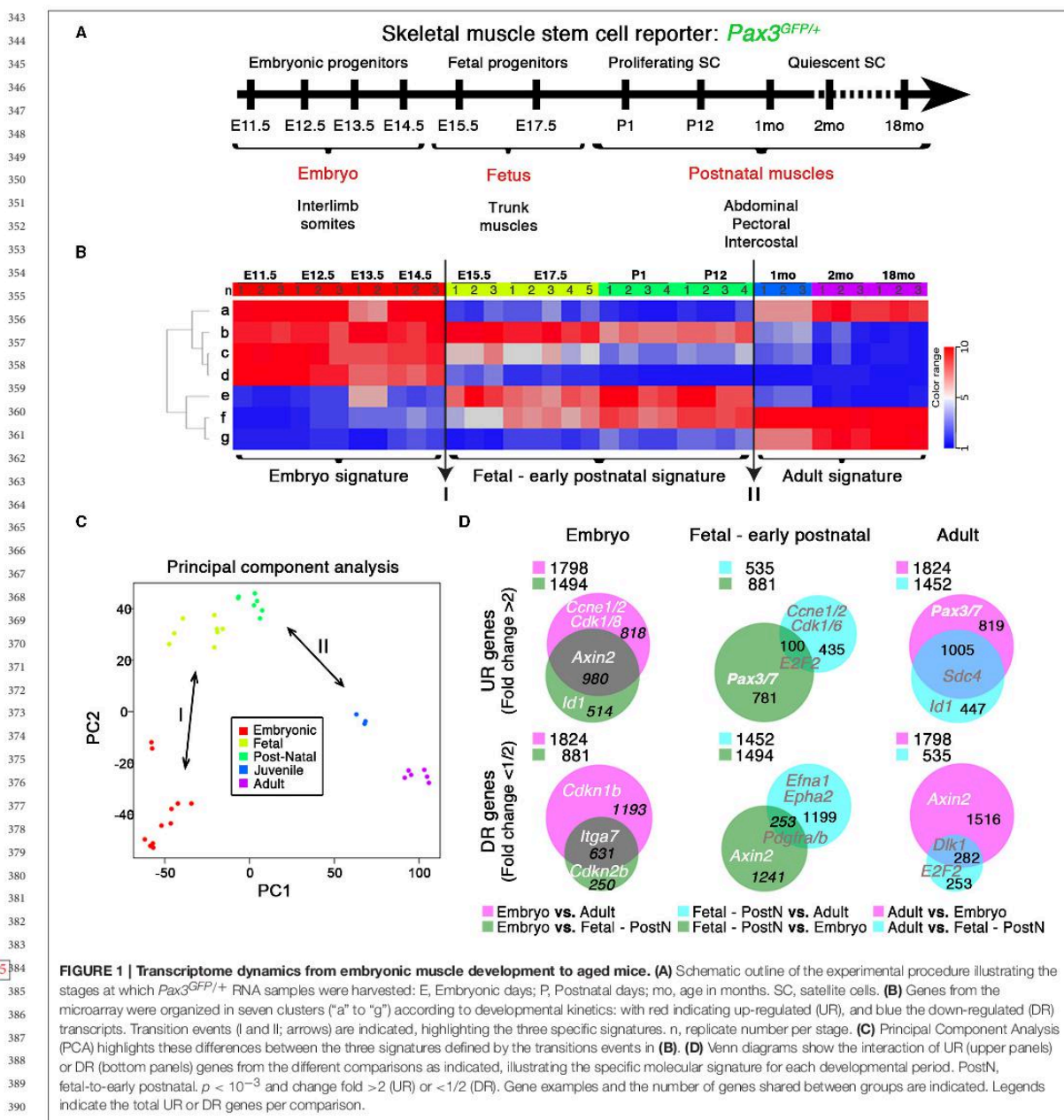


FIGURE 1 | Transcriptome dynamics from embryonic muscle development to aged mice. (A) Schematic outline of the experimental procedure illustrating the stages at which *Pax3^{GFP/+}* RNA samples were harvested: E, Embryonic days; P, Postnatal days; mo, age in months. SC, satellite cells. **(B)** Genes from the microarray were organized in seven clusters ("a" to "g") according to developmental kinetics: with red indicating up-regulated (UR), and blue the down-regulated (DR) transcripts. Transition events (I and II; arrows) are indicated, highlighting the three specific signatures. n, replicate number per stage. **(C)** Principal Component Analysis (PCA) highlights these differences between the three signatures defined by the transitions events in **(B)**. **(D)** Venn diagrams show the interaction of UR (upper panels) or DR (bottom panels) genes from the different comparisons as indicated, illustrating the specific molecular signature for each developmental period. PostN, fetal-to-early postnatal. $p < 10^{-3}$ and change fold >2 (UR) or <1/2 (DR). Gene examples and the number of genes shared between groups are indicated. Legends indicate the total UR or DR genes per comparison.

qPCR Analysis

RNA from trunk muscles was isolated through the RNeasy Fibrous Tissue kit (Qiagen®). For C2C12, total RNA extraction was performed using the RNeasy mini kit from Qiagen®. Total mRNA content was transcribed into coding DNA (cDNA) according to Transcriptor First Strand cDNA Synthesis kit

(Roche®) protocols. Quantitative analyzes were performed using the SYBR-Green kit (Roche®). qPCR was performed on biological duplicates (by sorting two different embryo series) with technical duplicates. The results obtained were analyzed by calculating the $2^{-\Delta\Delta Ct}$. *Hprt1* was used as reference gene.

457 Oligonucleotides of the following genes were selected, tested
458 and verified according to their efficiencies and specificities:

459 *EphB1*: FWD 5' - CCGTGGATGACTGGCTAAGT - 3'
460 REV 5' - TACCGATGGTACTGGTTCA - 3'
461 *Zbtb4*: FWD 5' - CGCTTCTCCATGTTGGCTAT - 3'
462 REV 5' - GTGAGCAGGGAAGTGGTGT - 3'
463 *Zbtb20*: FWD 5' - AATGCGAAAAGGGAAGCAGTA - 3'
464 REV 5' - ACAGGACCCGTGGAGTAATG - 3'

466 RNA Preparation, cDNA Synthesis, and 467 Microarray Hybridization

468 Microarray processing was performed by PartnerChip (Evry,
469 France), according to NuGEN (<http://www.nugen.com/nugen/index.cfm/products/pl/>) and Affymetrix (<http://www.affymetrix.com/support/technical/manuals.affx>) protocols. Briefly, total
470 RNA from FACS-sorted trunk muscle GFP+ cells was extracted
471 from independent experiments according to the RNasy[®]
472 Micro Kit (QIAGEN) RNA extraction protocol. RNA samples
473 were cleaned using Qiagen RNeasy mini-columns and their
474 quality assessed by spectrophotometry (Nanodrop ND-1000).
475 Total RNA was analyzed on Agilent microarrays (Bioanalyzer,
476 2100) to assess integrity of ribosomal RNA (28S and 18S
477 peaks). Synthesis and amplification of cDNA was performed
478 following NuGEN Ovation Pico WTA System protocol, and
479 100 ng of total RNA were used for first strand cDNA
480 synthesis. Second strand was synthesized following the Rib-
481 SPIA technology developed by NUGEN. Five micro gram of
482 single-stranded cDNA was fragmented and a biotin-labeled
483 nucleotide was attached to the 3' end of each fragment
484 (Encore Biotin Module, NuGEN). High-density oligonucleotide
485 arrays containing 45,000 sets of oligonucleotide probes (25 m)
486 that cover all 30,000 genes encoded by the murine genome
487 (Affymetrix Mouse Genome 430 2.0 Arrays, Ref 900495) were
488 used for gene expression detection. Hybridization during 16 h
489 at 45°C in a rotary oven (Affymetrix), washing and staining
490 (GeneChip[®] Fluidics Station 450) and scanning (GeneChip
491 Scanner 3000) were carried out according to NuGEN and
492 Affymetrix protocols. Expression Console software (Affymetrix)
493 was used for image analysis and to determine probe signal
494 levels. The quality and statistical analysis of the data were finally
495 made using the GeneSpring GX11 analysis software (Agilent
496 Technologies).

501 Expression Microarray Analysis 502 Pre-Treatment

503 Expression profiles of 36 samples (Pax3GFP+ cells at different
504 stages during development and after birth) were obtained
505 using Affymetrix Mouse Genome 430 2.0 Arrays. Expression
506 profiles were normalized in batch using RMA algorithm (affy
507 R package) yielding a (probe sets, samples) matrix. As the 36
508 samples were obtained by merging two series including 15 and
509 21 samples, Combat algorithm (Johnson WE—Biostatistics—
510 2007) was used to normalize the corresponding batch effect.
511 Expression profiles were aggregated by Gene Symbol (mean
512 across probe sets) using Affymetrix csv annotation file (na32
513 version).

514 Unsupervised Analysis

515 The gene expression matrix (GEO) was then row-mean-centered.
516 The resulting matrix was used for unsupervised classification of
517 the genes. Genes ($n = 21678$) were partitioned in ten clusters
518 using the kmeans classification algorithm. The biggest cluster
519 ($n = 8896$) contained genes showing almost no variation across
520 all samples: it was eliminated from further analysis. Three clusters
521 were found to be highly correlated (centroids correlation >0.95)
522 and were merged in a unique gene cluster (cluster g, **Figure 1B**).
523 We thus remained with seven clusters. For each sample, the mean
524 expression of all the genes of each cluster was calculated, yielding
525 a (seven clusters, 36 samples)-matrix shown in **Figure 1B**.

527 Supervised Analysis

528 Moderate T -tests (as implemented in limma R package) were
529 used to identify differentially expressed genes.

531 Pathways Analysis

532 To analyze the pathway enrichment, hypergeometric tests were
533 used, taking as “pathways” the terms (and related murine
534 genes) from the Gene Ontology (GO) (<http://www.geneontology.org>)
535 and the murine KEGG pathways (www.genome.jp/kegg).
536 Pathways enrichment in the seven gene clusters: in each of
537 the seven gene clusters, the pathway analysis was performed
538 using (i) all the genes included in the cluster, (ii) genes
539 selected based on their coefficient of variation and median-
540 absolute-deviation (different thresholds were used): the minimal
541 (hypergeometric test) p -value obtained from these different
542 (sub-) lists was retained. Pathways enrichment analysis of
543 differentially expressed genes: given a comparison between
544 two groups of samples, yielding a p -value and a fold change
545 for each gene, several lists of differentially expressed genes
546 were selected* and the minimal (hypergeometric test) p -value
547 obtained from these different lists was retained. (*) Lists of
548 differentially expressed genes: genes yielding a (moderate T -
549 test) $p > 1e-5$ were removed from the analysis; remaining
550 genes were ordered based on the fold change; the n genes
551 with highest (respectively lowest) fold change were selected
552 as a separate list; the $n/2$ genes with highest fold change
553 and the $n/2$ genes with lowest fold change were merged in
554 another list; this operation was performed for several values
555 of n (200, 300, 400, 500, 750, and 1000). Principal component
556 analysis (PCA) of the expression profiles was performed using
557 R software. Venn diagrams and pathway interaction schemes
558 were generated applying BioVenn (<http://www.cmbi.ru.nl/cdd/biovenn/>)
559 and GOrilla—REViGO (<http://cbl-gorilla.cs.technion.ac.il/>)
560 software packages, respectively. Pathway analysis was
561 completed employing DAVID Bioinformatics Resources 6.7
562 (<http://david.abcc.ncifcrf.gov/>).

564 Comparative Analysis of Microarray Data with 565 Published Available Datasets

566 Data normalization was performed with frozenRMA and
567 corrected for batch effect using Combat algorithm (Johnson
568 WE—Biostatistics—2007). Combined data series were the 36
569 samples from our study and those from published datasets
570 GSE50821 (Sinha et al., 2014) and GSE47177 (Liu et al., 2013).

571 The three Affymetrix series were used to compared adult
572 vs. old expression profiles (supervised meta-analysis young[2
573 months] vs. old[\geq 18 months]). This analysis showed that
574 the combination of the three sets found 32% significantly
575 deregulated genes and in the same direction of deregulation (32%
576 = proportion of the combined test under H1 = Test Stouffer).

577 Accession Numbers

578 The complete microarray data set, including the RMA data
579 used to produce intensity maps, have been deposited in NCBI's
580 Gene Expression Omnibus, and are accessible through GEO
581 Series accession number GSExxx (<http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSExxx>). To be requested to the editor
582 during the revision process.

583 Cloning (GMO Project No: 371)

584 To target activated satellite cells and not myofibers in our *ex vivo*
585 assays, a replication-deficient retrovirus, MIGR (*pMSCV-IRES-eGFP*),
586 has been used to transduce proliferating cells and overexpress
587 either dominant negative (DN) *EphB1*, *Zbtb4*, and *Zbtb20*
588 or full-length cDNA for *Zfp354c*, *Zcchc5*, *Zbtb4*, *Zbtb20*,
589 and *Hmga2* (Pear et al., 1998; Zammit et al., 2006b). The virus
590 is composed, besides the 5' and 3' LTR of the MSCV virus, the
591 latter being mutated to prevent replication, and the phi integrase,
592 of a multicloning site followed by an IRES-eGFP sequence to
593 track infected cells by fluorescence. This tracking cassette was
594 later modified into MISSINCK by substituting eGFP with an
595 insulin signal sequence-Cyan Fluorescent Protein (CFP)-KDEL
596 sequence in order to restrict fluorescent tracker expression to the
597 endoplasmic reticulum and Golgi.

598 Isolated Myofiber Cultures

599 Isolation of Muscular Fibers

600 Culture of single fibers was performed according to previously
601 described strategies (Moyle and Zammit, 2014). Briefly, dissected
602 EDL muscles were digested in a filtered solution of 0.2%
603 collagenase (SIGMA-ALDRICH®) in DMEM High Glucose/1%
604 L-Glutamine/1% Penicillin/Streptomycin (Life Technologies®)
605 (isolation medium). After 2 h of connective tissue digestion,
606 EDLs were mechanically dissociated fiber by fiber. Quiescent
607 satellite cells on the isolated myofibers were activated by a
608 solution of 10% horse serum/0.5% chicken embryo extract in
609 filtered isolation medium. Contracted fibers were removed.

610 Retrovirus Production and Infection

611 Retroviral particles (see *Cloning*) were produced in HEK293T
612 line cells by transfection using FuGENE® with a helper virus,
613 which contains the necessary elements to obtain the correct
614 encapsidation and active retrovirus (*phi integrase*, *gag*, *pol*
615 and *env* (VSV-g) genes). We collected the supernatants after
616 transfection at T = 72 h and T = 84 h, which displayed the highest
617 retroviral particle titers.

618 After 24 h of activation, myofiber-attached satellite cells were
619 infected with the retroviral particles diluted 1/10. 48 h afterwards
620 (T = 72 h), fibers were fixed to proceed with immunofluorescence
621 analysis.

622 C2C12 Cell Culture for Muscle Differentiation and Infection

623 Myogenic differentiation was induced according to previously
624 reported protocols (McMahon et al., 1994). Murine C2C12 cells
625 were cultured in 10% fetal bovine serum (Bio West®) in High
626 Glucose DMEM (Life Technologies®) for proliferation assay
627 (GM). Differentiation was induced by switching into medium
628 supplemented with 2% horse serum (Promega®) in High
629 Glucose DMEM (DM), generating multinucleated myotubes
630 surrounded by mononuclear reserve cells.

631 For retroviral infection, 10⁴ C2C12 cells were plated in GM
632 and incubated with undiluted retroviral supernatant containing
633 4 μ g/mL polybrene (SIGMA-ALDRICH®) for 3–4 h. Retroviral
634 medium was then removed, and the cells washed and incubated
635 in either proliferation (for PH3, KI67, EdU, and MYOD analysis)
636 or differentiation (for MYOG analysis) medium.

637 Immunostaining

638 Satellite Cells on Myofibers and Cryosections

639 Myofibers were fixed in 4% paraformaldehyde for 10 min, treated
640 with 0.5% triton and blocked in 10% goat serum/10% swine
641 serum (Moyle and Zammit, 2014). The following antibodies were
642 used: EPHB1 (Rabbit Abcam® ab5414) 1/100, PAX7 monoclonal
643 (DSHB®, PAX7-c) 1/50, MYOD monoclonal (DAKO®, clone
644 5.8A, M3512) 1/60, MYOG (Rabbit Santa Cruz Biotech®, sc-760)
645 1/50, MYOG monoclonal (DSHB®, F5D) 1/50, CD-31 (PECAM-
646 1) (Rat BD Pharmingen®, 550274) 1/500, and GFP (Rabbit Life
647 Technologies®) 1/500, or (Chicken Abcam® ab13970) 1/200.
648 Secondary antibodies employed to reveal the staining were Alexa
649 594 goat anti-mouse IgG (H+L), Alexa 488 goat anti-rabbit
650 IgG (H+L) (Life Technologies®), and DyLight-405 donkey anti-
651 chicken IgY (IgG) (H+L), and Cy5-Goat Anti-Rabbit IgG (H+L)
652 (Jackson ImmunoResearch®). Nuclei were counterstained with
653 DAPI.

654 C2C12 Cultured Myoblasts

655 The following antibodies were used: MYOD, MYOG, and GFP
656 (as above), PH3 (Rabbit Merck-Millipore®, 06-570) 1/50, Ki67
657 (BD Pharmingen®, 556003) 1/150, HA (Rabbit Sigma-Aldrich®,
658 H6908) 1/400, and GFP monoclonal (Sigma®) 1/50. Secondary
659 antibodies included Alexa 488 goat anti-mouse IgG (H+L),
660 Alexa 594 goat anti-mouse IgG (H+L), Alexa 488 goat anti-
661 rabbit IgG (H+L), Alexa 594 goat anti-rabbit IgG (H+L) (Life
662 Technologies®). EdU reaction was performed with Click-iT®
663 EdU Alexa Fluor® 647 Imaging Kit (ThermoFisher Scientific®).
664 Nuclei were counterstained with DAPI.

665 Imaging and Statistics

666 Analysis was carried out using a Leica TCS SPE confocal
667 microscope. Images were processed with either Adobe Photoshop
668 CS5 software (Adobe Systems) or ImageJ (version 1.47v; National
669 Institutes of Health, USA, <http://imagej.nih.gov/ij>).

670 Infected satellite cells in myofiber cultures were directly
671 counted under a Leica fluorescent microscope at 40x and 100x
672 magnification.

673 Mean \pm standard error (SEM) was given. The single (*),
674 double (**) and triple (***) asterisks represent *p*-values *p* < 0.05,
675 676 677 678 679 680 681 682 683 684

685 $p < 0.01$, and $p < 0.001$ respectively by Student's unpaired *t*-test
686 or Mann-Whitney *U*-test. All experiments have been performed
687 on at least three independent experiments for each condition.

688 Supplementary Movies were performed using a DSD2
689 Workstation with Imaris software (ANDOR).

690
691

692 RESULTS

693 Expression Dynamics of Skeletal Muscle 694 Stem Cells

695 *Pax3* is expressed in fetal progenitors and satellite cells of trunk
696 hypaxial muscles (Relaix et al., 2005; Relaix, 2006; Calhabeu et al.,
697 2013). We used a *Pax3* reporter mouse to perform a chronological
698 global profiling in embryonic, fetal and postnatal MPC and
699 satellite cells expressing *Pax3* (Figure 1A; Relaix et al., 2005).

700 Prospective isolation of *Pax3*-GFP myogenic progenitors and
701 stem cells was performed as previously described (Figure S2A;
702 Montarras et al., 2005; Lagna et al., 2010), taking advantage of
703 the GFP coding sequence targeting one allele of *Pax3* (Relaix
704 et al., 2005). *Pax3* is expressed in muscle progenitors but also
705 in early migrating neural crest cells (Epstein et al., 1993). Neural
706 crest cells give rise to many derivatives, including the peripheral
707 nervous system, melanocytes, and a subpopulation of venous
708 endothelial cells (by E13.5) among other cell types (Engleka et al.,
709 2005; Stoller et al., 2008). To exclude a possible contamination of
710 satellite cells with endothelial cells, we performed *Pax3*-lineage
711 tracing using *Pax3^{Cre/+}; R26^{mTmG}* mice (Figure S2B). While
712 adult myogenic cells were mGFP+ (*Pax3*-Cre recombined),
713 all endothelial cells remained mTOMATO+ (not recombined)
714 (Figure S2B and Movie S1). Moreover, all CD31 (PECAM-
715 1) + endothelial cells were included within the mTOMATO+
716 population (Figure S2B and Movie S2). These results demonstrate
717 that the *Pax3* lineage does not contribute to skeletal muscle
718 endothelial population, and that skeletal muscle expression of
719 PAX3 is specific to muscle stem cells.

720 Since *Pax3* is expressed in a subset of the *Pax7*-expressing
721 satellite cells, we compared our gene expression data with
722 previously published datasets of adult muscle stem cells where
723 markers different from PAX3 were used to isolate satellite
724 cells (Figure S3A; Liu et al., 2013; Sinha et al., 2014). *Pax3*-
725 expressing satellite cells were not significantly divergent from
726 previously reported datasets, while embryonic and fetal/early
727 postnatal datasets showed different specific profiles (Figure S3A).
728 Moreover, we compared available data from adult (3–8 month-
729 old) and old satellite cells (18–24 month-old) with our data. We
730 identified a similar variation in all datasets, demonstrating that
731 *Pax3*-expressing satellite cells do not define a subpopulation of
732 satellite cells. Our data therefore are likely representative of the
733 whole satellite population.

734 Expression profiles from 11 developmental stages were
735 normalized, generating a GEO showing the kinetics of each
736 transcript over time (Figures 1A,B). Transcript variations were
737 divided into seven clusters based on general expression profiles
738 (Figure 1B and Figure S3B), which were determined to be
739 functionally homogeneous and easily aggregated in defined
740 GO pathways (Figure S3B, Pathways). Furthermore, this *in*
741

742 *silico* analysis of the transcriptome through categorization of
743 expression trends (Figure 1B and Figure S3B, Pathways) and
744 specific molecular signatures (Figures 1B–D), yielded known
745 myogenic and related factors (Figure S3B, Genes) (Kuang et al.,
746 2008; Abou-Khalil et al., 2009; Boldrin et al., 2012; Conboy and
747 Rando, 2012). Strikingly, two transition events were revealed: (I)
748 from embryonic to fetal myogenesis (Messina and Cossu, 2009),
749 hypothesized to mark the early onset of satellite cell formation
750 (Kassar-Duchossoy et al., 2005); and (II) the acquisition of
751 quiescence in satellite cells around 3 weeks of age (Figures 1B,C;
752 Lepper et al., 2009; White et al., 2010). These transitions define
753 the three major developmental states: embryonic progenitors
754 (E11.5–E14.5), fetal-to-early postnatal (E15.5–P12) and adult
755 quiescent satellite cells (1–18 months), each with a specific
756 molecular signature (Figures 1B–D). Pairwise comparison
757 between different signatures of up-regulated (UR) and down-
758 regulated (DR) transcripts revealed the genes and pathways
759 defining each developmental period, provided in Figure S4
760 and Tables S1–S3 (UR), and Tables S4–S6 (DR), respectively.
761 Importantly, this *in silico* analysis also provides new markers for
762 muscle progenitor/stem cell maturation in both UR (extracellular
763 matrix formation, anatomical structure development, immune
764 and inflammatory responses) and DR (cell cycle and DNA repair
765 transcripts or developmental processes) pathways.

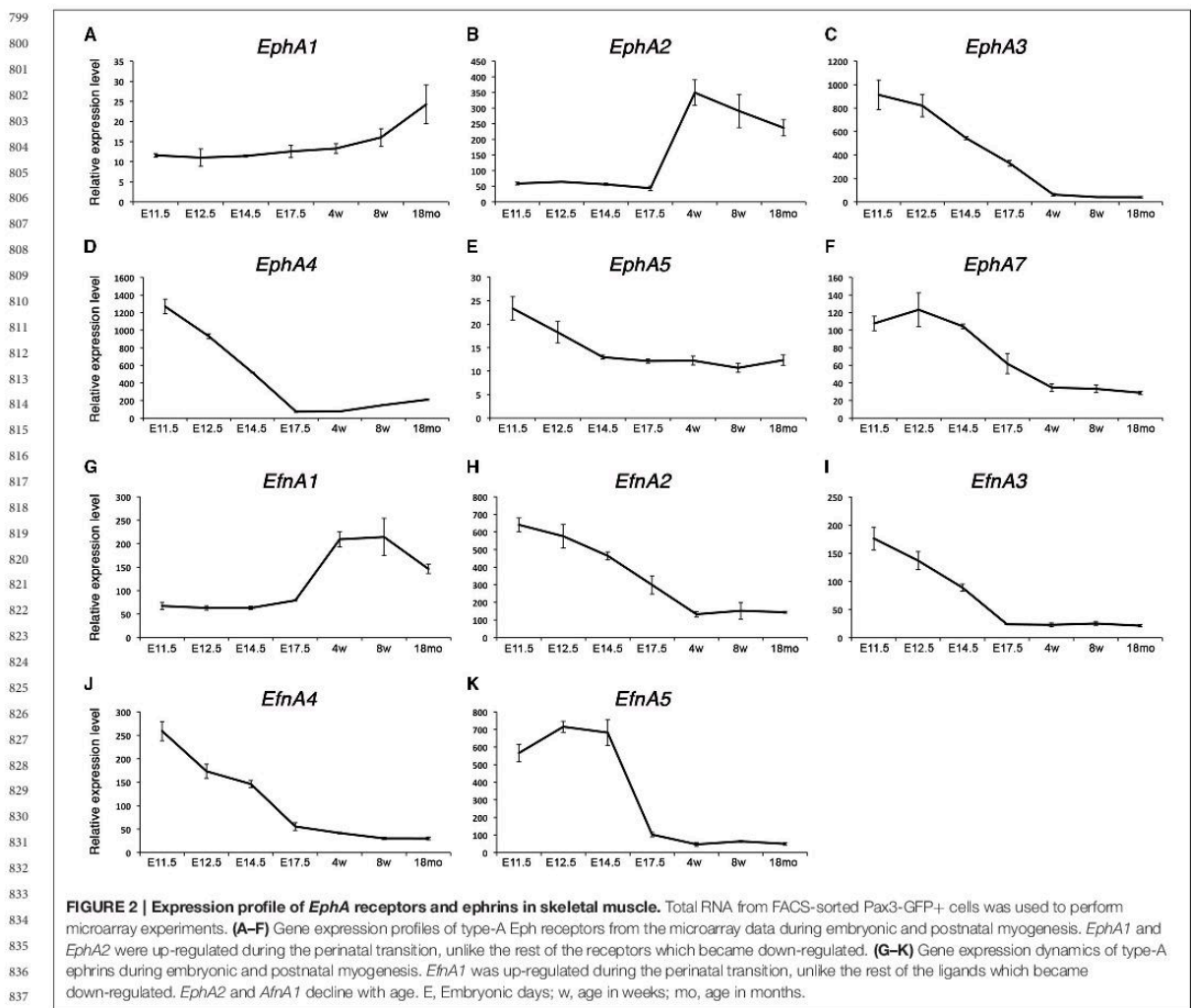
766 The dynamics of our transcriptional profiling reveal that each
767 stage of development is molecularly defined in a more progressive
768 manner than previously recognized.

769 Type A-Ephrins and Eph Receptors 770 Expression during Myogenesis

771 We have identified a set of transcripts specifically associated with
772 the embryonic and fetal stages of development or the satellite
773 cell lineage. Interestingly, Ephrin family members showed a
774 very dynamic behavior throughout development and postnatal
775 myogenesis, including *EphAs* and *EfnAs* (Figure 2). We could
776 distinguish two distinct behaviors: first, a set of *EphA* transcripts
777 that are up-regulated during the acquisition of muscle stem
778 cell properties (*EphA1* and *EphA2*, Figures 2A,B); second, an
779 independent set that is down-regulated over the same period
780 (*EphA3*, *EphA4*, *EphA5*, and *EphA7*, Figures 2C–F). *EPHA4*
781 has been reported to bind both EFNA and EFN B ligands
782 subtypes (Singla et al., 2010). This receptor was expressed in
783 the developing embryo, and repressed during postnatal growth
784 (Figure 2D). We found that *EphA4* is strongly expressed during
785 early embryonic development (E11.5) and ceases its expression
786 at the late fetal stage. In our transcriptome data, *EfnA2*, *EfnA3*,
787 *EfnA4*, and *EfnA5* ligands expression were also down-regulated
788 during fetal development, being no longer expressed during
789 aging (Figures 2H–K). Interestingly, only *EfnA1* became up-
790 regulated during the perinatal transition that characterizes the
791 emergence of satellite cells (Figure 2G).

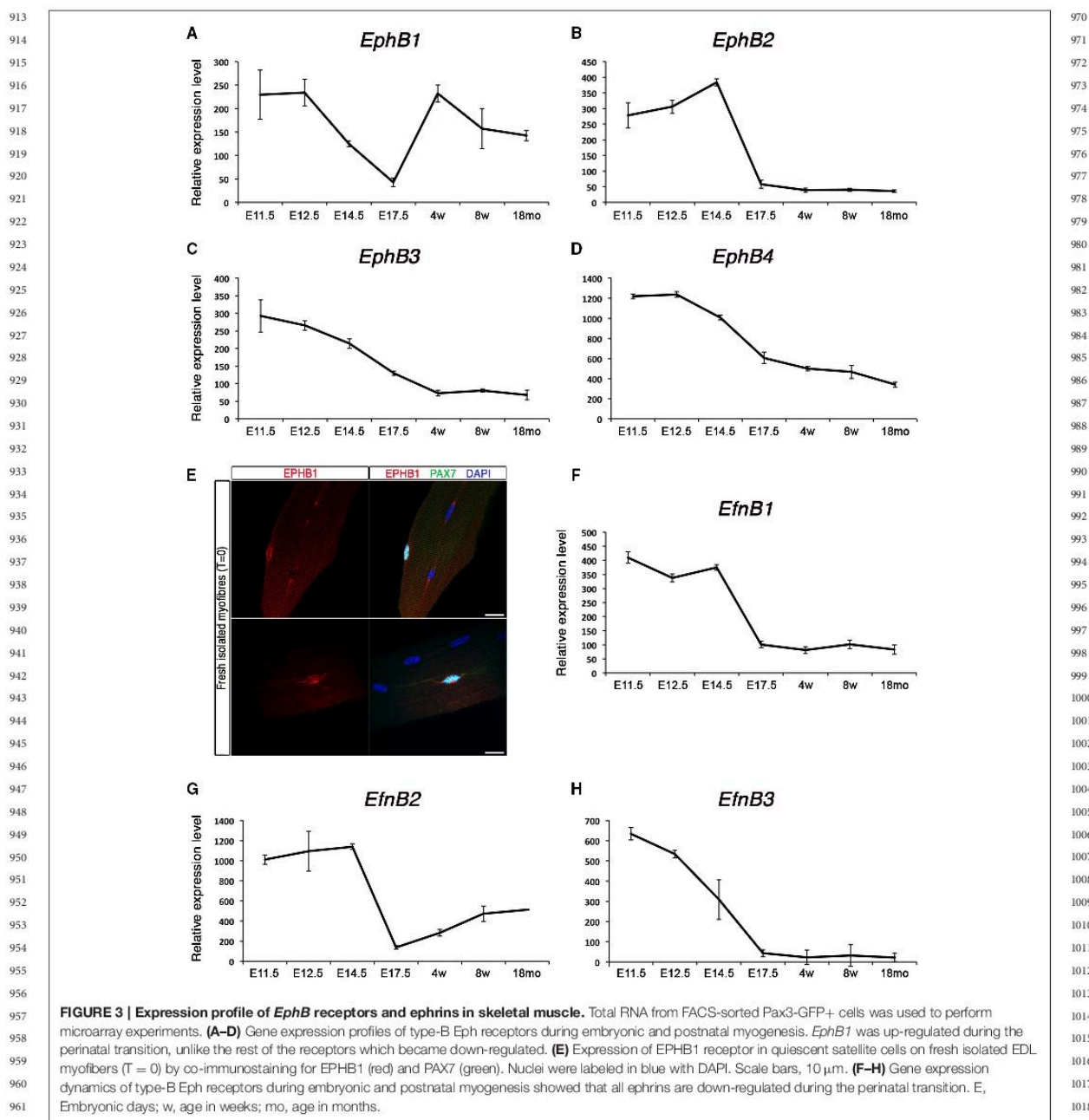
792 Type B-Ephrins and Eph Receptors 793 Expression during Myogenesis

794 Expression of *EphBs* and *EfnBs* at different stages is shown in
795 Figure 3. Among those, the transmembrane receptor *EphB1*
796
797
798



840 presents a unique dynamic expression profile: initially expressed
 841 early during myogenic development, then down-regulated
 842 during the fetal stage, and finally re-expressed in postnatal
 843 satellite cells (Figure 3A). By contrast, *EphB2*, *EphB3*, and *EphB4*
 844 are highly expressed during early development and progressively
 845 repressed as development proceeds (Figures 3B–D). We
 846 confirmed that *EphB1* was first expressed during the early
 847 stages of embryonic muscle development (Figure S5A), and
 848 down-regulated in the fetal stages. While it was weakly expressed
 849 in the early immature satellite cells (i.e., P2–P4), it was strongly
 850 up-regulated by P14, with expression then maintained, albeit at a
 851 lower level, in adult satellite cells. Interestingly, aged satellite cells
 852 (18 months old) show a marked decrease in *EphB1* expression
 853 (Figure S1A), corresponding to the timing when satellite cells
 854 start losing their regenerative capacity (Sousa-Victor et al., 2014).
 855

856 We used immunostaining on cultured floating myofibers to
 857 characterize expression of EPHB1 in muscle stem cells. This
 858 culture system recapitulates satellite cell activation, self-renewal
 859 and differentiation, similar to the situation observed during
 860 muscle regeneration in the adult (Zammit et al., 2004). After
 861 72 h, satellite cells were activated and proliferating (PAX7+ and
 862 MYOD+); some cells activated myogenin (MYOG+) and down-
 863 regulated PAX7, thus differentiating, and other cells will adopt
 864 a divergent fate, withdrawing from cell cycle and maintaining
 865 the expression of PAX7 while down-regulating MYOD (Zammit
 866 et al., 2004). Co-immunostaining of EPHB1 with PAX7, a
 867 specific marker of satellite cells, was observed on isolated fibers
 868 (Figure 3E), in 80% of the cells. However, expression was also
 869 observed in PAX7, MYOD, and MYOG positive myogenic cells
 870 at T = 72 (Figure S5B), demonstrating that EPHB1 was not
 871
 872



restricted to quiescent satellite cells, but maintained during the different steps of satellite cell activation and differentiation.

Finally, the kinetics of the ligands for type B-ephrins behaved similarly to most of the type A, being down-regulated during the perinatal transition to the emergence of satellite cells (Figures 3F–H).

EPHB1 Regulates Myogenesis in C2C12 Cells

C2C12 myoblasts are a classic model to analyze skeletal muscle differentiation (McMahon et al., 1994). Proliferating C2C12 cells were maintained in mitogen-rich medium, but differentiation was induced by switching into a serum poor-medium, thereby

1027 inducing MYOG expression and fusion into myotubes. Under
1028 long-term differentiation conditions, a reserve cell population
1029 emerges that shares some molecular and cellular features with
1030 quiescent satellite cells: for example, reserve cells express PAX7,
1031 are mitotically quiescent and aligned to the myotubes without
1032 fusing (Yoshida et al., 1998; Olguin and Olwin, 2004; Shefer et al.,
1033 2006).

1034 EPHB1 is expressed in both quiescent and activated satellite
1035 cells (Figures 3A,E and Figure S5). The extracellular region of
1036 the Eph receptor contains a globular ligand-binding domain,
1037 a cysteine-rich region (EGF-like motif), and two fibronectin-
1038 type III repeats (Figure S1). The intracellular region contains a
1039 tyrosine kinase domain, a SAM (Sterile Alpha Motif) protein-
1040 protein interaction domain and a C-terminal PDZ-binding motif
1041 (Figure S1A). To assess EPHB1 function in myogenic cells, we
1042 generated a dominant negative form of this receptor (EphB1DN)
1043 by removing the intracellular domain of the protein (Figure
1044 S1B) (Vindis et al., 2003, 2004; Haldimann et al., 2009; Oda-
1045 Ishii et al., 2010). Binding of ephrins to Eph receptors induces
1046 heterotetramers to initiate the signal cascade, which then will
1047 oligomerize and assemble in large signaling clusters (Pitulescu
1048 and Adams, 2010). EphB1 truncated receptor (EphB1DN) is
1049 therefore able to bind ephrin ligands, but cannot forward signal
1050 (Haldimann et al., 2009; Oda-Ishii et al., 2010). We induced
1051 expression of EphB1DN or control constructs using retroviral-
1052 mediated delivery in the C2C12 myoblastic cell line (Figure 4).
1053 EphB1DN was cloned into a modified retroviral vector carrying
1054 either an IRES-GFP or CFP to identify transduced cells and
1055 packaged using standard methods (Pear et al., 1998; Zammit
1056 et al., 2006b). These retroviral constructs were tested in C2C12
1057 and transduction of more than 90% of the cells was observed
1058 (Figure S6). Co-staining with EPHB1 antibody showed the
1059 expression of the receptor in C2C12 cells (Figure S6A). As our
1060 antibody is directed against the last 10 residues of the intracellular
1061 domain, a C-terminal 3HA-tagged version of EphB1DN was
1062 generated. Figure S6B shows a similar localization to the one of
1063 EPHB1 in transduced cells.

1064 We then assayed whether expression of EphB1DN would
1065 impact on proliferation of C2C12 cells using an antibody
1066 detecting the phosphorylated form of histone H3 at serine
1067 10 (PH3) (Figures 4A,B), and validated by KI67 and EdU
1068 incorporation (Figures S7A,B). By 24 h after infection with
1069 the EphB1DN-encoding retrovirus, C2C12 cells exhibited a
1070 significant increase in the mitotic index, suggesting either a
1071 decreased cell cycle time or a decreased myogenic commitment
1072 toward differentiation. To further characterize the role of EPHB1
1073 during myogenic differentiation, we analyzed expression of
1074 MYOD (Figures 4C,D) and MYOG (Figures 4E,F) in C2C12
1075 cells 48 and 72 h, respectively, after infection, and found an
1076 increased number of cells expressing these myogenic markers.
1077 We concluded that EphB1DN leads to increased proliferation
1078 and differentiation of C2C12 cells, suggesting a regulatory role
1079 for EPHB1 in satellite cell quiescence.

1081 EPHB1 Is Required for Satellite Cell 1082 Function and Renewal

1083 We next infected primary satellite cells on floating muscle
fibers to assay the consequence of expressing EphB1DN in

1084 activated satellite cells, and assayed self-renewal, proliferation
1085 and differentiation (Figure 5). 48 h after infection (72 h
1086 post isolation), the number of PAX7+ cells was reduced
1087 (Figures 5A,B). Consistently, we observed an increase in the
1088 MYOD+ (activated/proliferating and differentiating) population
1089 (Figures 5C,D). The number of MYOG+ (differentiating) cells
1090 was also increased (Figures 5E,F). Together, these results suggest
1091 that EPHB1 is involved in the maintenance of the pool of these
1092 adult stem cells, both by promoting self-renewal and by reducing
1093 activation and differentiation. To appropriately assess self-
1094 renewal of satellite cells, Pax7/MyoD double immunostaining
1095 was performed, taking advantage of a retrovirus with a CFP
1096 reporter expression restricted to the endoplasmic reticulum and
1097 Golgi (Figure 5G and Figures S6, S7). We confirmed that the
1098 decrease in the self-renewing satellite cell population (Pax7)
1099 correlated to an increase in differentiation (Figure 5G).

1100 Expression of Zinc Finger Containing 1101 Proteins during Myogenesis

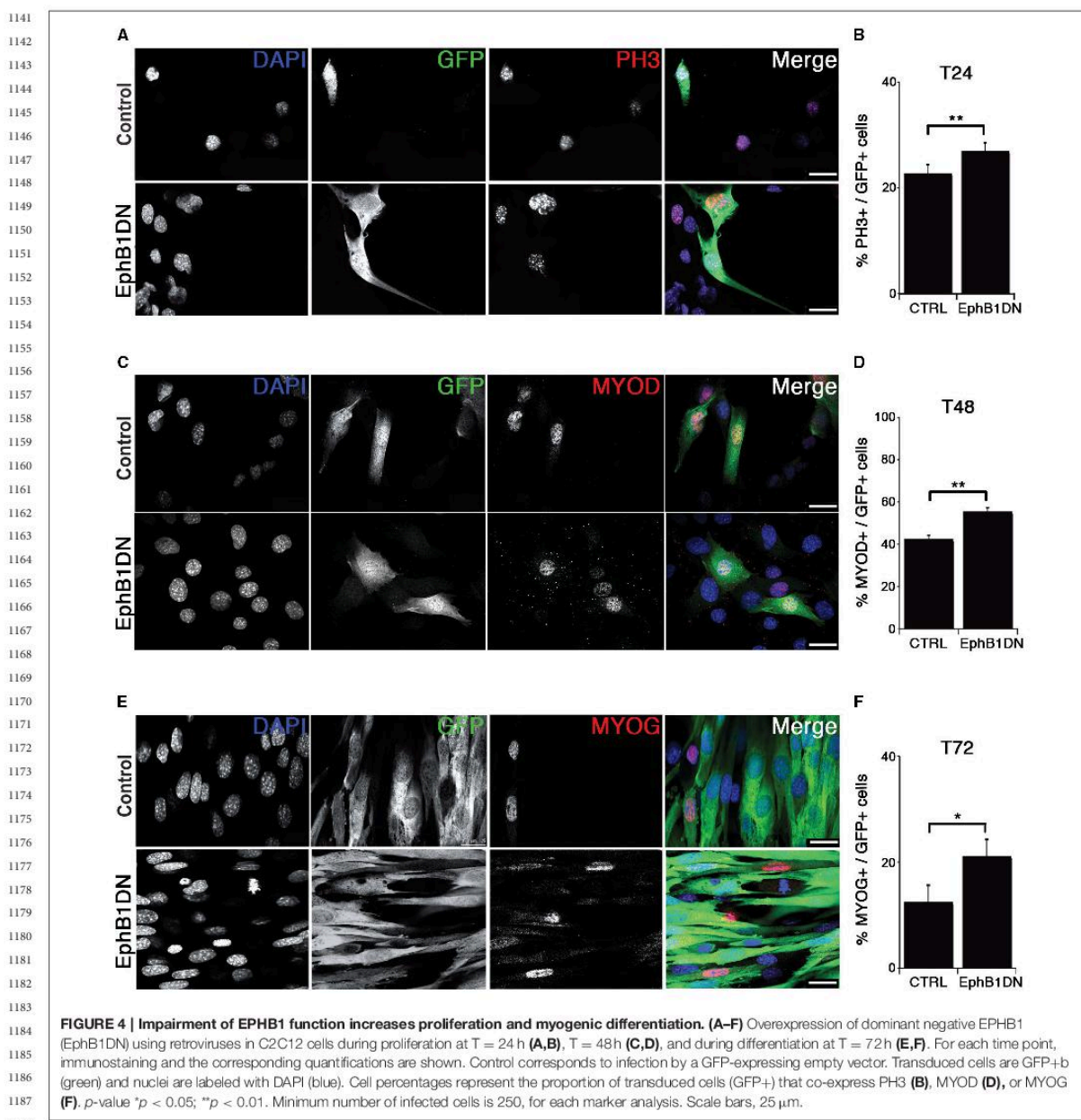
1102 Candidate genes coding for ZFP354c and ZCCHC5 zinc finger
1103 containing proteins were repressed during the emergence of
1104 satellite cells around birth (Figures 6A,B). Down-regulation
1105 of these factors was observed in muscle progenitors at the
1106 fetal stage overlapping with the emergence of satellite cells.
1107 These two zinc finger containing-proteins were not expressed
1108 in adult and aged satellite cells. While *Zfp354c* was highly
1109 expressed during early myogenesis and gradually repressed from
1110 fetal stages (Figure 6A), *Zcchc5* was not expressed during early
1111 embryonic myogenesis (Figure 6B), but appeared during early
1112 establishment/formation of the satellite cell pool, before being
1113 completely down-regulated during acquisition of satellite cell
1114 quiescence. According to the known functions of these factors,
1115 we can hypothesize their possible involvement during MPCs
1116 proliferation (*Zfp354c*), or for a correct determination of the
1117 MPC fate to become the muscle stem cells (*Zcchc5*).

1118 By contrast, two other zinc finger containing proteins *Zbtb4*
1119 and *Zbtb20*, were not expressed during development but were
1120 induced during establishment of satellite cells and acquisition of
1121 quiescence (Figures 6C,D). Moreover, high expression of these
1122 zinc finger containing-proteins was maintained in adult and
1123 aged satellite cells, implicating a possible function in maintaining
1124 quiescence of muscle stem cells. Strikingly, these factors are
1125 induced during cardiotoxin-induced muscle regeneration *in vivo*
1126 (Figure S8A).

1128 Effect of Zinc Finger Containing Proteins in 1129 Postnatal Satellite Cells

1130 We manipulated expression of *Zfp354c*, *Zcchc5*, *Zbtb4*, and
1131 *Zbtb20* using retroviral-mediated delivery in single myofiber
1132 cultures as above. We generated vectors carrying either a full-
1133 length transcript for overexpression, or dominant negative forms
1134 to analyze function.

1135 Overexpression in satellite cells of either *Zfp354c* or *Zcchc5*
1136 maintained expression in satellite cells that no longer expressed
1137 the endogenous gene (Figures 6E–J). Notably, overexpression
1138 of *Zfp354c* led to a decreased number of PAX7+ satellite
1139 cells compared to control (Figure 6E) with no apparent effect
1140 during activation (MYOD+) and differentiation (MYOG+)

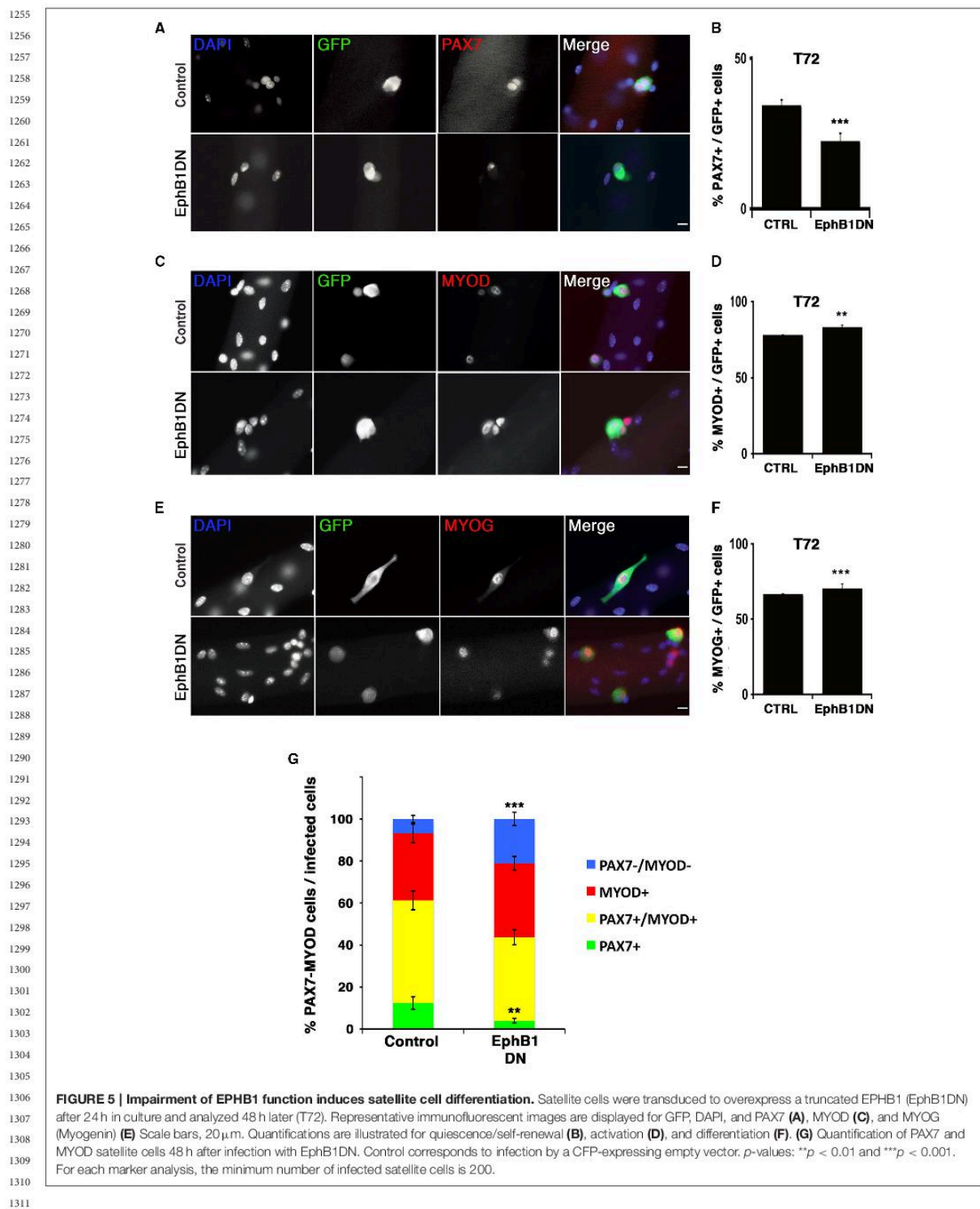


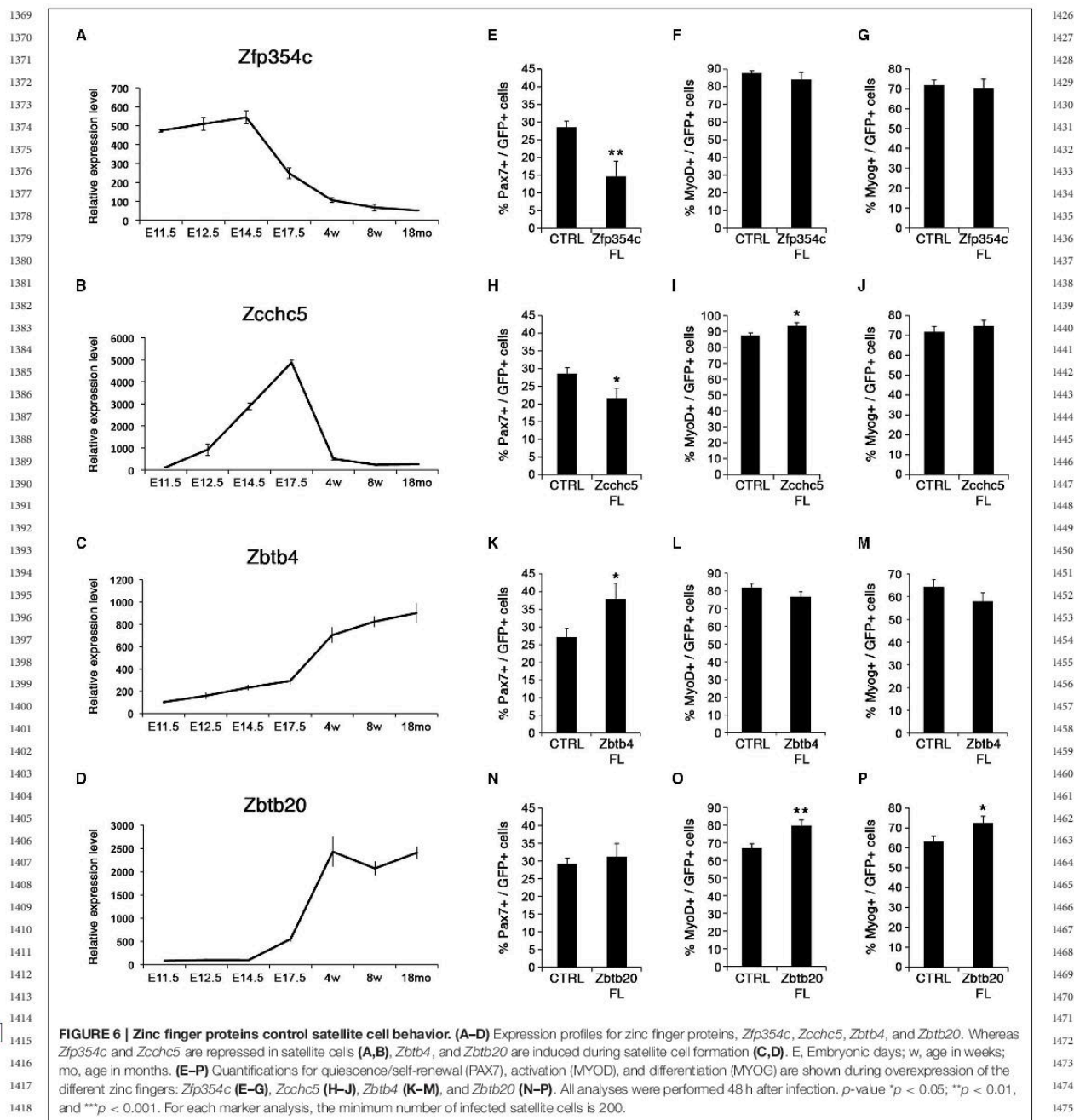
1190 (Figures 6F,G). These results demonstrate that overexpression of
1191 *Zfp354c* resulted in a reduction of self-renewal capacity of the
1192 satellite cells.

1193 Overexpression of *Zchc5* in satellite cells, as *Zfp354c*, resulted
1194 in a decreased of the PAX7+ population relative to control
1195 (Figure 6H). Strikingly, the proportion of MYOD+ satellite cells
1196 increased without affecting MYOG-expressing differentiated
1197

1247 cells (Figures 6I,J). These results showed that overexpression
1248 of *Zchc5* induced decreased self-renewal promoting the
1249 proliferation of satellite cells. Our functional data is consistent
1250 with a specific requirement of *Zchc5* function during the growth
1251 phase where production of MPC is needed.

1252 We next overexpressed the other two BTB-containing zinc
1253 finger factors, *Zbtb4* and *Zbtb20* (Figures 6K–P). *Zbtb4* increased
1254





PAX7⁺ satellite cells (Figure 6K), whereas *Zbtb20* promoted myogenic progression by increasing the activated/proliferating (MYOD⁺; Figure 6O) and differentiating (MYOG⁺; Figure 6P) populations. These results suggested that these transcriptional

repressors might be required for specification/maintenance of the muscle stem cell pool. Strikingly, inhibiting function by expression of ZBTB4 dominant negative constructs, missing the POZ DNA-binding domain, displayed an increase in

1483 satellite cell differentiation (MYOG+) without affecting the
 1484 activated/proliferating population (MYOD+) (Figures S8B,C).
 1485 On the other hand, ZBTB20 could behave with a previously
 1486 described phenotype in the brain of *Zbtb20* transgenic mice
 1487 (Nielsen et al., 2007), where overexpression of ZBTB20 represses
 1488 cell fate transitions in newborn pyramidal neurons. Moreover,
 1489 overexpression of ZBTB20 has been recently described as a
 1490 prognostic marker by promoting tumor growth of human
 1491 hepatocellular carcinoma (Kan et al., 2016). Thus, ZBTB20 could
 1492 be regulating muscle regeneration during satellite cell activation
 1493 as suggested in Figure S8A.

1495 **Hmga2 Must be Repressed for Appropriate** 1496 **Satellite Cell Function**

1497 *Hmga2* was highly expressed during early development
 1498 (Figure 7A), when MPCs expand to populate the future skeletal
 1499 muscle of the body. As development proceeds, *Hmga2* was no
 1500 longer expressed, and was not detected in the emerging satellite
 1501 cells prior to birth. Nishino and collaborators have described a
 1502 similar behavior where *Hmga2* is highly expressed in fetal neural
 1503 stem cells and declining with age (Nishino et al., 2008).

1504 We analyzed the effect of overexpressing *Hmga2* in satellite
 1505 cells (Figures 7B–E). A retroviral construct carrying full-length
 1506 cDNA including the coding sequence for the basic and acidic
 1507 region of the protein was generated (Figure 7B), and satellite cells
 1508 infected using single myofiber culture. *Hmga2* overexpression led
 1509 to a strong reduction in the pool of satellite cells expressing PAX7
 1510 (Figures 7C,D), with an increase on the activated/proliferating
 1511 MYOD+ muscle stem cells (Figure 7E); data consistent with the
 1512 work from Li and colleagues describing HMGA2 as a regulator
 1513 of myoblast proliferation by direct interaction with the RNA-
 1514 binding protein IGF2BP2 (Li et al., 2012).

1517 **DISCUSSION**

1518
 1519 PAX3 and PAX7 are key upstream regulators of skeletal
 1520 myogenesis (Relaix et al., 2005; Buckingham and Relaix, 2015).
 1521 Postnatally, while PAX7 labels all satellite cells (Seale et al.,
 1522 2000), PAX3 is maintained in a subset of these adult muscle
 1523 stem cells (Relaix et al., 2006). A complex balance between
 1524 extrinsic cues and intrinsic regulatory mechanisms is needed
 1525 to tightly control satellite cell determination and function. For
 1526 example, defects in satellite cell regulation or in their niche,
 1527 such as during postnatal growth or in degenerative conditions
 1528 and aging, can impair muscle regeneration with possible fatal
 1529 consequences (Dumont et al., 2015). Hence, identifying and
 1530 manipulating muscle progenitor stem cells, and understanding
 1531 the mechanisms underlying cell fate decision and self-renewal
 1532 (Relaix, 2006; Boutet et al., 2007) are essential for development
 1533 of stem cell-based therapeutic strategies.

1534 We have developed a FACs-based chronological
 1535 transcriptome profile of myogenic stem cells, sampled from
 1536 embryonic and fetal progenitors, to postnatal, adult, and
 1537 aging satellite cells. This provides a comprehensive description
 1538 of gene expression changes throughout life of muscle stem
 1539 cells and identifies two important transition events, which

1540 delimit three developmental periods of muscle stem cells with
 1541 specific molecular signatures: (1) embryonic, (2) fetal to early
 1542 proliferating postnatal progenitors, and (3) quiescent adult
 1543 muscle stem cells (Buckingham and Relaix, 2007; Braun and
 1544 Gautel, 2011). The intersection between specifically expressed
 1545 genes and functional pathways defines a molecular signature
 1546 unique to each developmental period. As such, our study is
 1547 instrumental for a better understanding of both myogenesis and
 1548 the establishment and maintenance of quiescent adult stem cells.

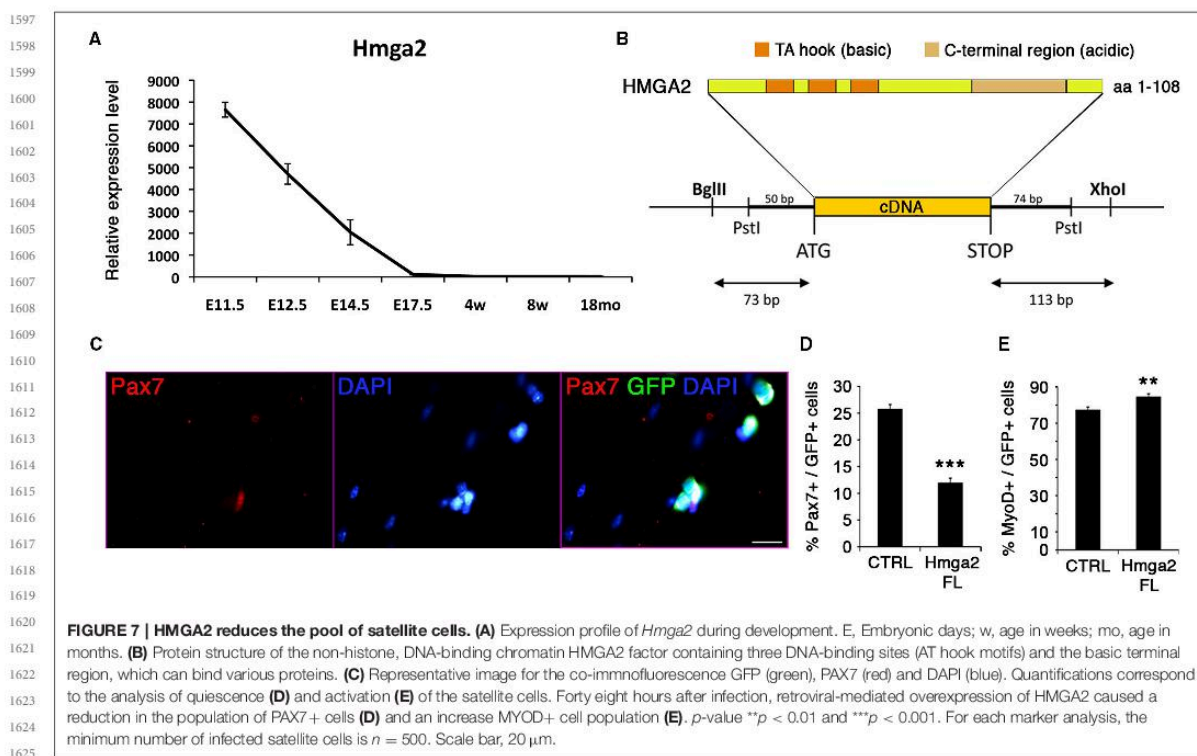
1549 The dynamics of our transcriptional profiling reveal that
 1550 cellular processes characterizing muscle stem cells, including
 1551 transition from the fetal lineage to postnatal stem cells,
 1552 establishment of quiescence and formation of a functional
 1553 niche, are defined molecularly in a more progressive manner,
 1554 highlighting that establishment of the satellite cell lineage is more
 1555 gradual than previously recognized. For example, cell division
 1556 processes (i.e., cyclins such as *Ccne1/2* or cyclin-dependent
 1557 kinases such as *Cdk1*) were gradually down-regulated throughout
 1558 the second transition, corresponding to the establishment of
 1559 satellite cell quiescence and consistent with analysis of fetal
 1560 progenitor cell proliferation (Picard and Marcelle, 2013). At
 1561 the same time, known satellite cell markers such as *Sdc4*
 1562 (Syndecan 4), *Igta7* (Integrin Alpha-7) or *Cav1* (Caveolin 1) were
 1563 progressively up-regulated (Cornelison, 2001; Gnocchi et al.,
 1564 2009).

1565 From this large-scale myogenesis transcriptome, we
 1566 functionally characterized a set of genes to provide novel
 1567 intrinsic factors that regulate satellite cell behavior (Figure 8A).

1569 **Eph/Ephrin Pathway and Myogenesis**

1570 EPHB1 is not only involved in motility and guidance in skeletal
 1571 muscle cells as previously shown (Stark et al., 2011), but also
 1572 acts as a novel regulator of myogenesis. Our findings point to a
 1573 function during self-renewal of satellite cells, since a dominant
 1574 negative form of EPHB1 led to increased proliferation and
 1575 differentiation in C2C12 myogenic cells and satellite cells in the
 1576 myofiber experimental model. The increase in cell differentiation
 1577 is achieved at the expense of self-renewal of the satellite cell
 1578 population (Figure 8B). Identifying the molecular regulators of
 1579 satellite cell renewal is important since it was recently shown
 1580 that targeted depletion of the satellite cell pool leads to complete
 1581 impairment of muscle regeneration following injury (Relaix and
 1582 Zammit, 2012).

1583 Eph/ephrin signaling takes place via direct cell-cell
 1584 interaction; either as *trans* or *cis* signaling (Arvanitis and
 1585 Davy, 2008; Pitulescu and Adams, 2010). This interaction could
 1586 take place with the muscle fiber, between satellite cells, or via
 1587 interactions with other cell types in the microenvironment
 1588 (i.e., macrophages and/or microvascular cells). The satellite
 1589 cell population is heterogeneous, with specific markers labeling
 1590 subpopulations of the satellite cell pool and different myogenic
 1591 behaviors *in vivo* or *ex vivo* (Relaix et al., 2006; Kuang et al.,
 1592 2007; Rudnicki et al., 2008; Ono et al., 2010; Rocheteau et al.,
 1593 2012). Whether arising through lineage or stochastic events,
 1594 more “stem” satellite cells likely correspond to independently
 1595 identified label-retaining satellite cells during growth and after
 1596 injury (Shinin et al., 2006; Rocheteau et al., 2012; Chakkalakal



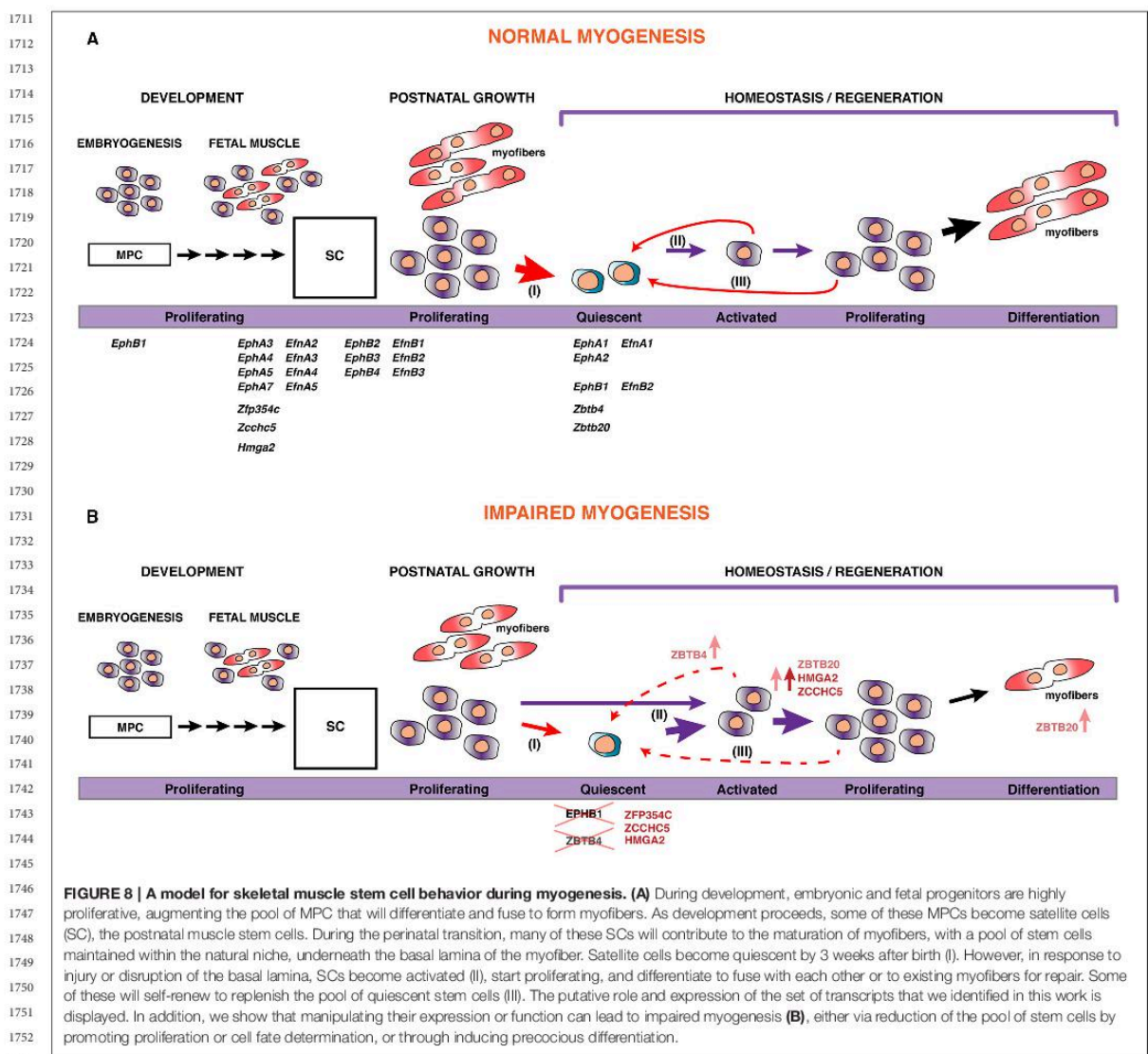
et al., 2014), or displaying different rates of cell division (Ono et al., 2012). Interestingly, satellite cells can asymmetrically divide and it will be of interest to evaluate if interaction between fibers and/or the satellite cells via the Eph/ephrin signaling plays a role in these cell fate decisions. Finally, our results are consistent with the work from Chumley and colleagues, showing that proliferative neuronal progenitor cells increase in *EphB1* mutant mice (Chumley et al., 2007), thereby demonstrating an important role of EPHB1 in maintenance of neuronal progenitors in the quiescent state.

Eph/ephrin signaling has also been shown to play a role in regulating other stem cell niches, for instance in the dental (Stokowski et al., 2007) or osteochondral (Arthur et al., 2011) system. Using an ephrin “stripe” assay revealed that satellite cells respond to a subset of ephrins with repulsive behavior *in vitro* (Stark et al., 2011). Our finding that EPHB1 is also regulating myogenesis suggests that this guidance signaling might impact multiple aspects of muscle regeneration, including escape from the niche, directed migration to sites of injury, cell-cell interactions among satellite cell progeny, and differentiation and patterning of regenerated muscle.

Identification of Novel Zinc Finger Proteins Regulating Myogenesis

We identified a set of zinc finger containing proteins with a dynamic expression profile during myogenesis. We have shown

that overexpression of *Zfp354c* decreased self-renewal of satellite cells (summarized in Figure 8). In the skeletal system, the highest *Zfp354c* expression is in proliferating bone cells compared to mature and differentiated chondrocytes. Interestingly, ZFP354C is induced as an early response to BMP-7 (Jheon et al., 2001). It has been shown that overexpression of *Zfp354c* affects osteoblast differentiation, a lineage that is also regulated by BMP signaling (Jheon et al., 2001). Moreover, overexpression of this gene results in a decrease in osteogenic differentiation by suppressing BMP-7 induced alkaline phosphatase activity, an early marker of osteogenesis (Jheon et al., 2003). Furthermore, BMP signaling prevents myogenic differentiation of satellite cells, and is also involved in regulation of satellite cells during proliferation or differentiation (Friedrichs et al., 2011; Ono et al., 2011). In essence, there is strong evidence of a functional interaction between ZFP354C and BMP7, though the precise relationship between the two proteins is not fully understood (Jheon et al., 2002). Future studies will be necessary to evaluate whether a functional interaction between ZFP354C and BMP7 regulates myogenesis and, in general, to identify the downstream gene regulatory networks for all four zinc finger proteins presented here, ZFP354C, ZCCHC5, ZBTB4, and ZBTB20, which are able to strongly repress transcription of target genes. These zinc fingers, thus, could be used as potentially powerful tools for regulation of muscle stem cell function.



HMGA2 Function and Its Role in Myogenesis and Satellite Cell Fate Decision

HMGA2 is a co-regulator of chromatin structure and pluripotency in stem cells (Pfannkuche et al., 2009). The role of HMGA2 in myoblast proliferation has been previously described in neonatal and regenerating muscle (Li et al., 2012). *Hmga2* is sharply induced during satellite cell activation. We found that *Hmga2* is highly expressed during early muscle development and progressively down-regulated in the fetal stages, while it is not expressed during growth or aging (Figure 7A). It has been shown that *Hmga2* knockout mice are smaller and show

defects in postnatal skeletal muscle (Zhou et al., 1995; Li et al., 2012). In addition, HMGA2/IGF2BP2 has been shown to be critical for myoblast proliferation and early myogenesis, but should be down-regulated in order for myoblasts to differentiate into multinucleated skeletal muscle. Indeed, when satellite cells are activated entering cell cycle, HMGA2 is up-regulated and activates the expression of IGF2BP2 (Li et al., 2012). Our transcriptome analysis shows that *Igf2bp2* behaves similarly to *Hmga2* before birth, but in contrast to *Hmga2*, *Igf2bp2* is induced in adult stem cells, including aged satellite cells (data not shown). This suggests that IGF2BP2 could be functionally independent of HMGA2 in adult and aged satellite cells.

In conclusion, understanding the molecular signals that control and regulate the muscle stem cell population is essential in order to open new therapies strategies for muscle diseases. Here we provide a set of potential new regulators of myogenesis that improves the understanding and knowledge of the intrinsic factors controlling muscle stem cell acquisition, establishment, maintenance and function in the adult, and could be targeted to modify the regenerative capacity of endogenous skeletal muscle stem cells.

AUTHOR CONTRIBUTIONS

SA, AR, and JM designed and performed experiments, and analyzed data. SA wrote the manuscript. AD analyzed bioinformatic data. DM, FA, and TC performed experiments. PZ designed experiments. FR oversaw the entire project, designed experiments, analyzed data and wrote the manuscript. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

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Online resources

1. https://embryology.med.unsw.edu.au/embryology/index.php/Mouse_Timeline_Detailed
(Timeline and hallmarks of mouse embryonic development)