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Skeletal muscle dedifferentiation during salamander limb regeneration

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35 Abstract

36 Salamanders can regenerate entire limbs throughout their life. A critical step during 37 limb regeneration is formation of a blastema, which gives rise to the new extremity. 38 Salamander limb regeneration has historically been tightly linked to the term 39 dedifferentiation, however, with refined research tools it is important to revisit the 40 definition of dedifferentiation in the context. To what extent do differentiated cells 41 revert their differentiated phenotypes? To what extent do progeny from differentiated 42 cells cross lineage boundaries during regeneration? How do cell cycle plasticity and 43 lineage plasticity relate to each other? What is the relationship between 44 dedifferentiation of specialized cells and tissue resident stem cells in terms of their 45 contribution to the new limb? Here we highlight these problems through the case of 46 skeletal muscle.

47

48 Tracking muscle cells in the salamander limb

Limb skeletal muscle fibers are formed by the fusion of somite-derived precursors. These multinucleate, elongated cells have a specialized cytoarchitecture built up by proteins, which make the fibers easily distinguishable from their precursor cells at the molecular level. A key feature of the myofibers in the context of the present review is the quiescent state of the myonuclei within the multinucleated syncytium, which is often referred to as the stable post-mitotic state [1,2].

55 Skeletal muscle has considerable regenerative capacity in all vertebrates, including 56 mammals. However the myonuclei in mammals do not resume proliferation after an 57 injury. Instead, a population of muscle stem cells, the so-called satellite cells, starts to 58 proliferate and subsequently differentiates into muscle to replenish lost fibers [3–5]. 59 Although satellite cells were first described in amphibians [6] *, their presence in 60 adult salamanders [7–9] was unequivocally confirmed more than 40 years later by the 61 isolation of single newt myofibers along with an attached population of cells expressing the canonical satellite cell marker, Pax7 [10]. This finding challenged the 62 63 traditional view that solely the myofiber itself, rather than a quiescent stem cell population are the progenitor cells during salamander limb regeneration [11], and also 64 65 highlighted the need to carry out cell type specific tracking experiments during limb 66 regeneration.

Limb regeneration starts with a rapid wound healing followed by formation of ablastema from which the new limb develops [12,13]. Pioneering histological analyses

69 suggested more than half century ago that myofibers undergo fragmentation, and 70 indicated the migration of mononucleate myofiber fragments into the salamander limb 71 blastema [14*,15]. Furthermore, myofiber fragmentation temporally coincides with 72 disorganization and histolysis of the stump tissues in general, and concomitant 73 production of blastema cells [16]. Cell cycle reentry by myonuclei was also suggested 74 but it is important to remember that the available tools at the time did not allow 75 discrimination among myonuclei, satellite cell nuclei and the nuclei of other 76 interstitial cells within muscle tissue [17]. The model of myofiber-dedifferentiation 77 gained further support from several studies on myotubes, which are the *in vitro* model 78 cell type for resident myofibers. Although myotubes lack striation, they do express a 79 range of terminal differentiation markers, and their nuclei are stably quiescent. 80 However, myotubes from the aquatic salamander, the newt, reenter the cell cycle and 81 replicate their DNA upon appropriate stimulation, which is a distinctive feature of 82 these cells compared to their mammalian counterparts [18*,19]. Furthermore, upon 83 implantation of myotubes into the blastema, could give rise to mononucleate progeny 84 in the blastema [20,21].

85 Although these studies collectively suggested a distinctive plasticity of 86 differentiated salamander muscle cells, genetically integrated, heritable labeling of 87 myonuclei was required to address whether and to what extent myofibers 88 dedifferentiate during limb regeneration. These experiments were performed in the 89 red spotted newt (Notophthalmus Viridescens) and the Mexican axolotl (Ambystoma 90 Mexicanum), and revealed unexpected differences between these two salamander 91 species [22] **. First, myofibers in newts gave rise to proliferating blastema progeny, 92 but no such cells were found in the axolotl limb blastema. Second, in sharp contrast to 93 the axolotl, the fraction of myofibers carrying the tracer was similar in pre-existing 94 and regenerated muscle in the new limb in newt. Third, the newt blastema was largely 95 devoid of PAX7⁺ cells, except for a few cells appearing during the first few days of 96 limb regeneration [10,23]. The axolotl limb blastema on the other hand contained a large number of PAX7⁺ cells. To what extent these differences at the molecular level 97 98 reflect differences in the cellular contribution of satellite cell progeny to the 99 regenerating limb will be discussed further down. Importantly, the dissimilarities 100 between the two species were independent of the developmental stages of the animals, 101 since myofiber-progeny did not contribute to the new limb in axolotls that were 102 experimentally induced to undergo metamorphosis, and PAX7⁺ cells were also 103 lacking in the blastemas of larval newts. On the other hand, a recent analysis in the 104 Japanase fire-bellied newt (Cynops pyrrhogaster) indicated that skeletal muscle 105 dedifferentiation only occurs in metamorphosed animals [24]. Remarkably, that work 106 also suggested that in larval stage the vast majority of blastema cells turn from being PAX7⁻ into PAX7⁺ between day12 to day15 after amputation. The possibility that 107 108 proliferating PAX7⁺ cells in the axolotl blastema are derived from myofibers, whose 109 nuclei upregulate Pax7 after amputation was raised [25], but the cell tracking 110 experiments do not provide support for such a process.

111

112 Satellite cell progeny vs dedifferentiated cells in the blastema

113 Does the lack of PAX7⁺ cells in the newt blastema mean that satellite cells do not significantly contribute to muscle (or to other tissues for that matter) in the 114 115 regenerating limb? At a first glance this appears as a logical conclusion, especially in 116 light of the contrasting observations in the axolotl [26]. However, it is important to 117 keep in mind that the tracing experiments in newts specifically targeted myofibers, 118 but not the satellite cells. Currently, it is perfectly possible that satellite cell progeny 119 contribute to the limb blastema also in newts but these progeny downregulate 120 expression of the Pax7 gene within the blastema. If this were the case, a major 121 difference between the newt and axolotl in terms of satellite cell contribution to the 122 blastema would be at the level of gene regulation rather than in the cell source per se 123 (Figure 1). In order to unequivocally determine the fate of satellite cells and to relate 124 the contribution from satellite cells to myofiber dedifferentiation, one would need to 125 trace satellite cell progeny during newt limb regeneration. So far this has not been 126 feasible due to lack of suitable cell type specific promoter constructs.

127 As a surrogate approach to *bona fide in vivo* tracing, satellite cells were previously 128 isolated and, following *in vitro* expansion, re-injected into to regenerating newt limb 129 [10,23]. Although in vitro expansion could lead to such epigenetic changes in the 130 cultured cells that naturally are not occurring, these experiments suggested that 131 satellite cell progeny have the capacity to contribute to the regenerate. In addition, the 132 experiments indicated that satellite cell progeny could not only give rise to muscle but 133 also to other cell types in newts - a plasticity, which might be reflected by 134 downregulation of Pax7 in the satellite cell progeny [23]. This scenario would 135 represent yet another difference between axolotls and newts. While axolotl muscle 136 tissue, and presumably the satellite cells within, were shown only to form muscle during limb regeneration [27] **, satellite cells may cross lineage boundaries in the
newt. Again, the distinctive difference in the newt compared to the axolotl in that case
would be the plasticity rather than the lack of contribution by satellite cells and their
progeny.

141

142 Cell cycle plasticity and lineage plasticity

The results of the myofiber tracing studies in newts refined our understanding ofmyofiber plasticity from at least two aspects.

145 First, they showed that cell cycle reentry is a post-fragmentation event occurring in 146 mononucleate myofiber progeny rather than in the myonuclei within the syncytium 147 before breaking up of the myofiber. This is in line with earlier experiments showing 148 that myotubes that were blocked to re-enter the cell cycle still could give rise to 149 mononucleate (obviously non proliferating) progeny upon implantation into the 150 blastema [21]. However they contrast other conclusions that some myonuclei did 151 enter S-phase in the syncytium during limb regeneration [28]. Further experiments are 152 required to resolve the discrepancy between the two studies. The mechanistic 153 separation of cell fragmentation from cell cycle reentry is also consistent with the 154 observations showing that, although without detectable proliferation, also axolotl limb 155 and tail blastemas harbored mononucleate myofiber-derived progeny [22,29]. This 156 indicates that fragmentation of myofibers may represent an alternative fate direction 157 of the muscle fiber - a question that we will discuss further.

158 Second, they provided no evidence for the myofiber progeny to cross lineage 159 boundaries, as the label introduced to intact muscle prior to limb removal was only 160 found in muscle fibers and not elsewhere in the new limb. How the muscle identity of 161 the myofiber progeny is maintained is not clear but myofiber derived mononucleate 162 progeny that had lost expression of terminal muscle differentiation marker myosin 163 heavy chain, still expressed the early myogenic factor Myf5 in the blastema [22]. It 164 will be important to determine whether Myf5 expression is a prerequisite for retaining 165 the myogenic commitment of myofiber progeny. Yet another open question is 166 whether myofiber progeny acquire muscle stem cell properties, which also requires 167 further investigations. So far we can conclude that dedifferentiated myofiber-derived 168 cells neither do acquire Pax7-expression nor are they found in satellite cell position in the regenerated muscle within the new limb, suggesting that they act as lineage 169 170 committed progenitors during regeneration.

171

172 Mechanisms of myogenic dedifferentiation

173 Three key features thus define dedifferentiation of skeletal muscle fibers during limb 174 regeneration: (1) Fragmentation of the syncytium into mononucleate cells, (2) loss of 175 terminally differentiated markers, but retention of at least one early myogenic 176 determinant and (3) proliferation of the fiber-derived mononucleate cells. As outlined 177 above, myofiber fragmentation does not depend on cycle reentry by the myonuclei, 178 and conversely, fragmentation of the muscle syncytium does not predestine the 179 derived mononucleate cells to proliferate. The underlying mechanisms of these two 180 processes should thus be possible to disentangle from each other.

Means to force myotubes of both salamander and mammalian origin to reenter the cell cycle has been extensively explored. Key gate-keepers that prevent myonuclei reentering the cell cycle or initiate myogenic dedifferentiation have been identified, such as the retinoblastoma (Rb) protein [18], MSX1 [30], p21 [31], p19ARF [32], and thoroughly discussed in an excellent recent review [26]. Here we focus myogenic dedifferentiation cues specifically studied in the context of salamander limb regeneration.

188 A series of experiments involving both culture based assays and cell tracking 189 approaches during limb regeneration showed that fragmenting muscle cells displayed 190 hallmarks of a programmed cell death (PCD) process, such as activation of caspase-3, 191 and that inhibition of caspase activity counteracted the derivation of mononucleate 192 cells from both cultured myotubes as well as myofibers in the limb [33] *. 193 Importantly, inducing a programmed cell death response by myotubes was sufficient 194 to cause cellularization of cultured myotubes but only a fraction of the derived 195 mononucleate cells could be rescued from dying by apoptosis inhibitors and induced 196 to proliferate. Although still not proven, the emerging model suggests that limb 197 amputation evokes myofibers to embark on a programmed cell death program, which 198 is manifested by fragmentation of the syncytium. However, the derived mononucleate 199 cells must be rescued from the full execution of the cell death program in order to 200 gain ability for resuming proliferation within the blastema. This idea is consistent 201 with the observations that axolotl myofibers also fragment into mononucleate cells 202 during appendage regeneration [22,29], but these cells cannot be traced further during 203 axolotl regeneration and presumably die.

204 At present it is unclear how the molecular components of the programmed cell 205 death program cause myofiber disassembly. An experimentally approachable 206 hypothesis is that caspases are involved in the disintegration of structural elements, 207 which are required for maintaining the integrity of syncytium. Noteworthy in the 208 context are the experiments showing that caspase activity is required for spermatid 209 individualization during sperm maturation in drosophila – a process during which 210 each spermatid becomes encapsulated by an independent plasma membrane [34]. 211 Caspases might also expel obstacles of subsequent proliferation that reside in the 212 chromatin structure.

213 What could be the reasons why, in contrast to the newt, myofiber derived 214 mononucleate cells do not contribute to the regenerate in the axolotl (formally only 215 proven in the limb)? Differences both in intrinsic cell properties as well as in extrinsic 216 cues that cells encounter in the limb might provide explanations but no such 217 differences have yet been identified. Assays on cultured newt myotubes indicated that 218 inhibition of p53 activity is necessary for cell cycle reentry [35*,36] and p53 219 knockdown was also required to render mammalian myotube-derived mononucleate 220 cells ability to resume proliferation [33]. However, p53 activity decreases also during 221 axolotl blastema formation, and p53 stabilization led to impairment of limb 222 regeneration [35]. Similarly, with a creative screening strategy using newt myotubes 223 the Tanaka lab recently identified a MARCKS (Myristoylated alanine-rich C-kinase 224 substrate)-like protein (MLP), which on one hand promotes proliferation of myofiber 225 derived mononucleate cells in newts, and on the other hand initiates regeneration of 226 both limbs and tails in the axolotl [37] **.

227

228 Future perspectives

229 Our understanding of how and to what extent skeletal muscle contributes to limb 230 regeneration has significantly increased during the past years. In this review we also 231 highlighted outstanding questions that still have not been addressed experimentally. 232 One such issue is to determine the relative contribution from dedifferentiating 233 myofibers and from satellite cells to the regenerating newt limb. Even if we have 234 gained more insight to myofiber dedifferentiation at the cellular level, we are still 235 short of insights into the underlying molecular mechanisms. One way forward is to 236 combine cell tracking approaches with genome wide expression analyses and

and genome editing technologies.

239 Figure legends

- 240 Figure 1. Contribution of skeletal muscle cells to the blastema formation during newt
- 241 *limb regeneration*. Myofiber dedifferentiation results in proliferating, Myf5⁺/PAX7⁻
- 242 mononuclear cells (black) in the blastema that give rise to the skeletal muscle in the
- 243 new limb. Lack of PAX7⁺ cells in the newt blastema indicates either a minimal
- 244 contribution of satellite cells (green) to the blastema formation or a down-regulation
- of *pax7* gene expression in the progeny of satellite cells.
- 246

247 Figure 2. Model of myofiber dedifferentiation during newt limb regeneration. Injury

evokes myofibers to activate caspases, which are involved in the disassembly of the

249 syncytium. The resulting fragments apoptotic fragments will either die or survive and

250 proliferate. The identity of the pro-survival and proliferation cues is largely unknown.

- 251 Although not proven in newts, downregulation of p53 activity is likely to play a role
- 252 in cell survival. The MLP promotes proliferation of myofiber progeny during newt
- limb regeneration.

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379	salamanders.		



