1	Regulation of stem cell identity by miR-200a during spinal cord regeneration					
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14	Key words: axolotl, spinal cord, stem cell, mesoderm, regeneration					
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16	Summary Statement: After spinal cord injury, miR-200 fine-tunes expression levels					
17	<i>brachyury</i> and β -catenin to direct spinal cord stem into cells of the mesodermal or					
18	ectodermal lineage.					
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20	Abstract					
21	AxolotIs are an important model organism for multiple types of regeneration, including					
22	functional spinal cord regeneration. Remarkably, axolotls can repair their spinal cord					
23	after a small lesion injury and can also regenerate their entire tail following amputation.					
24	Several classical signaling pathways that are used during development are reactivated					
25	during regeneration, but how this is regulated remains a mystery. We have previously					
26	identified miR-200a as a key factor that promotes successful spinal cord regeneration.					
27	Here, using RNA-seq analysis, we discovered that the inhibition of miR-200a results in					
28	an upregulation of the classical mesodermal marker brachyury in spinal cord cells after					
29	injury. However, these cells still express the neural stem cell marker sox2. In vivo					
30	lineage tracing allowed us to determine that these cells can give rise to cells of both the					
31	neural and mesoderm lineage. Additionally, we found that miR-200a can directly					
32	regulate brachyury via a seed sequence in the 3'UTR of the gene. Our data indicate that					
33	miR-200a represses mesodermal cell fate after a small lesion injury in the spinal cord					
34	when only glial cells and neurons need to be replaced.					

35

36 Introduction

37 Regeneration has been observed throughout the plant and animal kingdoms for 38 many years (Sanchez Alvarado and Tsonis, 2006). Among vertebrates, the Mexican 39 axolotl salamander has the remarkable ability to faithfully regenerate its spinal cord after 40 injury. This process has been most commonly studied in the context of tail amputation 41 (McHedlishvili et al., 2012, Monaghan et al., 2007, Piatt, 1955, Rodrigo Albors et al., 42 2015, Zhang et al., 2000, Zhang et al., 2002a), but the axolotl spinal cord also 43 regenerates after a more targeted transection injury (Butler and Ward, 1965, Butler and 44 Ward, 1967, Clarke et al., 1988, O'Hara et al., 1992, Zukor et al., 2011). These lines of 45 investigation have identified a population of Sox2⁺/GFAP⁺ glial cells that function as bona 46 fide neural stem cells (NSCs) in the axolotl spinal cord (Echeverri and Tanaka, 2002, Fei 47 et al., 2016, Fei et al., 2014a, McHedlishvili et al., 2012, Rodrigo Albors et al., 2015). 48 These Sox2⁺/GFAP⁺ NSCs proliferate after injury and differentiate into new glia and 49 neurons (Echeverri and Tanaka, 2002, McHedlishvili et al., 2012, Rodrigo Albors et al., 50 2015). Inhibition of NSC function by CRISPR/Cas9-mediated knockout of Sox2 results in 51 deficient regenerative outgrowth of the spinal cord after tail amputation (Fei et al., 2016, 52 Fei et al., 2014a).

53 Molecular signals required for the NSC response to injury have been identified 54 after both tail amputation and spinal cord transection. Sonic hedgehog, Wnt/PCP, and 55 Fgf signaling are indispensable for the pro-regenerative NSC response to tail amputation 56 (Rodrigo Albors et al., 2015, Schnapp et al., 2005, Zhang et al., 2000, Zhang et al., 57 2002a). During spinal cord regeneration after transection, the transcriptional complex 58 AP-1^{cFos/JunB} and MAP kinase signaling are critical regulators of the NSC response to 59 injury (Sabin et al., 2015a, Sabin et al., 2019). Additionally, microRNA (miRNA) signaling 60 is important to fine-tune the NSC response to injury after both tail amputation and spinal 61 cord transection (Diaz Quiroz et al., 2014, Gearhart et al., 2015, Lepp and Carlone, 62 2014, Sehm et al., 2009).

63 We previously demonstrated that miR-200a is an important regulator of the glial 64 cell response after spinal cord transection (Sabin et al., 2019). The function of miR-200a 65 has been most extensively studied during neurodevelopment and epithelial-to-66 mesenchymal transition (EMT) (Trumbach and Prakash, 2015, Zaravinos, 2015). miR-67 200a functions to inhibit EMT by directly repressing the expression of the transcription 68 factor β -catenin (Su et al., 2012, Zaravinos, 2015). This leads to maintained epithelial

69 polarity and decreased Wnt signaling. During neurodevelopment, miR-200 family

70 members regulate many processes: neuronal survival (Karres et al., 2007),

71 neuroepithelial progenitor proliferation, NSC identity and neuroblast transition (Morante

et al., 2013), neural progenitor identity and cell cycle dynamics (Peng et al., 2012), as

73 well as fine tunes signaling networks necessary for neurogenesis (Choi et al., 2008,

74 Vallejo et al., 2011) and gliogenesis (Buller et al., 2012).

75 Early experiments aimed at determining the potential of GFAP⁺/Sox2⁺ NSCs 76 prospectively labeled these cells with the Glial fibrillary acidic protein (GFAP) promoter 77 driving GFP expression and used live in vivo fluorescence imaging to follow GFP⁺ glial 78 cells after tail amputation. Most GFP⁺NSCs gave rise to new neurons and glia but a 79 small proportion of labeled cells left the spinal cord and contributed to muscle and 80 cartilage within the regenerated tail (Echeverri and Tanaka, 2002). Similar experiments 81 using grafting of GFP⁺ spinal cords into non-transgenic animals further confirmed that 82 spinal cord cells exited the spinal cord and contributed to cells of other lineages during 83 tail regeneration (McHedlishvili et al., 2007).

84 Recent reports have identified a population of progenitors, called 85 neuromesodermal progenitors (NMPs), which reside in the posterior of developing 86 vertebrate embryos (Henrique et al., 2015, Kimelman, 2016b). NMPs are competent to 87 contribute to both the mesoderm and spinal cord during embryonic development 88 (Garriock et al., 2015, Henrique et al., 2015, Tzouanacou et al., 2009). Extensive genetic 89 and biochemical analysis determined that NMPs can be defined by the co-expression of 90 low levels of the transcription factors Brachyury and Sox2 (Gouti et al., 2017, Koch et al., 91 2017a, Turner et al., 2014a, Wymeersch et al., 2016). Also, the relative level of Fgf and 92 Wnt signaling activity regulate NMP cell fate decisions (i.e. differentiation into 93 mesodermal progenitors or neural progenitors) (Bouldin et al., 2015, Garriock et al., 94 2015, Goto et al., 2017, Gouti et al., 2017, Gouti et al., 2015, Martin, 2016, Martin and 95 Kimelman, 2008, Turner et al., 2014a). Both Fgf and Wnt signaling are important 96 regulators of the NSC response to tail amputation, as inhibition of either Wnt or Fgf 97 blocks tail regeneration (Makanae et al., 2016, Ponomareva et al., 2015, Zhang et al., 98 2000, Albors et al., 2015). Although the role of individual For ligands in spinal cord 99 regeneration is relatively unknown, the expression of *wnt5* has been elegantly shown to 100 be essential for orientated cell division and outgrowth of the spinal cord during whole tail 101 regeneration (Albors et al., 2015). However, the activity of these pathways after spinal 102 cord transection has not been well characterized.

In this study we identify a role for miR-200a in stabilizing the NSC identity after spinal cord transection in axolotl, by repressing expression of the mesodermal marker *brachyury*. Furthermore, we uncovered other genes in the miR-200 pathway and provide evidence that depending on the injury context, such as spinal cord lesion repair or spinal cord outgrowth during tail regeneration, that miR-200a plays an important role in determining the identity of NSCs in the spinal cord during the regenerative process.

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110 **Results**

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112 Transcriptional profiling identifies conserved miR-200a targets in homeostatic 113 versus regenerating spinal cords

114 We previously identified miR-200a as a key microRNA (miRNA) that inhibits *c-jun* 115 expression in neural stem cells in the spinal cord after injury, and hence plays an 116 important role in preventing reactive gliosis and promoting a pro-regenerative 117 response(Sabin et al., 2019). Additional RNA sequencing (RNA-seq) on uninjured and 4 118 days post injury spinal cord tissue electroporated with a control or specific, antisense 119 miR-200a inhibitor identified other targets of miR-200a in axolotl spinal cord (Fig. 1A, 120 **Table S1**). During normal regeneration at 4 days post injury there were 1,163 genes that 121 are differentially expressed (Log2 fold change ≥ 2 -fold, $p \leq 0.05$) compared to uninjured 122 spinal cords. Inhibition of miR-200a in the uninjured spinal cord resulted in 6,235 123 transcripts with a greater than 2-fold differential expression compared to control 124 uninjured spinal cords. Interestingly, of the 6,235 differentially expressed genes, only 125 2.760 were up-regulated (Fig.1A, Fig. S1, Table S1). We used GOrilla analysis to 126 identify gene ontology (GO) terms for the subset of genes that were significantly up-127 regulated after miR-200a inhibition. GO terms involved with translation, RNA 128 metabolism, peptide metabolism, and translation initiation were significantly enriched in 129 this gene set ($p \le 10^{-24}$) (**Fig.S1**). Interestingly, the 3,475 genes that were significantly 130 down-regulated in uninjured spinal cords after miR-200a inhibition were enriched for GO 131 terms involved with organismal development, developmental process, cellular 132 differentiation, and signaling ($p \le 10^{-22}$) (**Fig. S1**).

Analysis of differentially expressed genes at 4 days post injury after miR-200a inhibition identified a total of 1,007 genes that were differentially expressed compared to control spinal cords. This is a much smaller gene set and suggests that there is a higher specificity to genes affected by miR-200a during spinal cord regeneration. A total of 797

137genes were up-regulated and 210 genes were down-regulated after miR-200a inhibition138(Fig. S1). Genes that were up-regulated were enriched for GO terms involved with139nucleic acid metabolism, specifically RNA metabolism, and protein localization ($p \le 10^{-6}$)140(Fig. S1). Interestingly, the top GO terms enriched in down-regulated genes were141involved with nervous system processes, specifically synaptic signaling and chemical142synaptic signaling, as well as nervous system development ($p \le 10^{-6}$).

143 Taking a more targeted gene-level approach, we generated a heat map of the 30 144 most significantly up-regulated and down-regulated genes in all four conditions (Fig.1C). 145 Consistent with the GO analysis, genes involved in RNA processing, nucleic acid 146 metabolism, and protein targeting were among the most up-regulated genes in our data 147 set (tdrd9, acap1, eme1, zfp324b). Similarly, genes involved with neurotransmitter 148 transport, neuronal polarization, neurotrophin signaling, and neuronal differentiation 149 were among the most down-regulated genes (slc6a6, brsk1, slc6a14, arhgap8, 150 *neurog1*). Surprisingly, the transcription factor *brachyury* (*T*) was among one of the most 151 up-regulated genes at 4 days post injury in response to miR-200a inhibition (Fig.1C). In 152 4 days post injury controls, *brachyury* was not up-regulated in response to injury, the 153 RNA seq transcripts per million (TPM) values on Control uninjured were 0.782 TPM, 154 versus 4 dpi post injury 0.92 TPM, show no significant increase (Table S1). However, 155 inhibition of miR-200a in uninjured spinal cords led to a 2-fold increase in brachyury 156 expression (2.236 TPM), while the combination of miR-200a inhibition during injury led to 157 a highly significant 7-fold increase in its mRNA levels (5.8 TPM, **Table S1**). 158 We used quantitative RT-PCR (qRT-PCR) to verify genes of interest revealed by

159 RNA-seq, this approach confirmed that *brachvurv* is detectable at very low levels in 160 uninjured and control 4 days post injury spinal cords but is significantly up-regulated 161 after miR-200a inhibition in 4 days post injury spinal cords (Fig.1D). This is an intriguing 162 finding as *brachyury* is considered a classical marker of mesodermal tissue and was 163 originally thought to be absent from in the nervous system. However, more recent 164 research has identified a bipotent cell population in development, in which some spinal 165 cord neural progenitor cells are developmentally derived from Sox2⁺/Brachyury⁺ 166 neuromesodermal progenitors (NMPs) (Garriock et al., 2015, Tzouanacou et al., 2009, 167 Wymeersch et al., 2016). In the axolotl, the bona fide stem cells that line the central 168 canal are identified by the expression of the glial cell marker GFAP and the neural stem 169 cell marker Sox2. These GFAP⁺/Sox2⁺ cells respond to the injury, divide, migrate and 170 repair a lesion in the spinal cord, or regenerate lost cells and tissues in the context of

171 whole tail regeneration (Sabin et al., 2015a, Fei et al., 2014b, Echeverri and Tanaka, 172 2002, Echeverri and Tanaka, 2003a, McHedlishvili et al., 2007, McHedlishvili et al., 173 2012). Given that NMPs and axolotl glial cells both express Sox2 and that sox2 is a miR-174 200a target during mouse brain development (Peng et al., 2012), we assayed sox2 175 transcript abundance. Interestingly, while sox2 is slightly up-regulated in control 4 days 176 post injury compared to uninjured spinal cords, sox2 expression did not increase in miR-177 200a inhibitor treated spinal cords. Instead, the sox2 transcript abundance remains near 178 uninjured homeostatic levels (Fig.1D). This observation suggests that axolotl sox2 is not 179 a direct target of miR-200a as it is in mammals (Pandey et al., 2015, Peng et al., 2012, 180 Wang et al., 2013).

181 To identify the cells that express *brachyury* in the 4 days post injury spinal cord 182 after miR-200a inhibition, in situ hybridization was used. In situ hybridization determined 183 that cells lining the central canal are *brachyury*⁺ after miR-200a inhibition (**Fig.1Evii**), and 184 importantly, this is the same population of cells that express sox2 (Fig.1Evi, viii). 185 Collectively, this data indicates that miR-200a inhibition leads to increased brachyury 186 expression in stem cells in the axolotl spinal cord. Although these progenitor cells have 187 been traditionally thought of as NSCs due to their expression of the classical NSC 188 marker sox2, they also express low levels of the mesodermal marker brachyury 189 (Fig.1D), suggesting that they are in fact a population of bipotent stem cells and may in

- 190 fact have broader differentiation potential.
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192 Inhibition of miR-200a leads to changes in cell fate after spinal cord injury

193 To determine if miR-200a inhibition and subsequent upregulation of the mesodermal

- 194 marker *brachyury* leads to changes in the number of NSC in the regenerating spinal
- 195 cord, we quantified the number of Sox2⁺ cells in control versus inhibitor treated animals
- 196 (Fig.2A, B). Previous work has shown that after spinal cord injury the cells that are
- 197 activated to partake in the regeneration process lie within 500µm of the injury site.
- 198 However, we detected no significant difference in the total number of Sox2⁺ cells 500µm
- rostral or caudal of the injury site (Fig.2A, B). During the normal regenerative process in
- 200 the spinal cord, the Sox2⁺ stem cells will replenish the GFAP⁺ cell population and
- 201 differentiate into new neurons (Albors et al., 2015, Echeverri and Tanaka, 2002,
- 202 Echeverri and Tanaka, 2003a, McHedlishvili et al., 2007, McHedlishvili et al., 2012). To
- 203 determine if the number of newborn neurons is affected by miR-200a inhibition we
- 204 quantified the number of NeuN and EDU⁺ cells 500µm rostral and caudal to the injury

205 site. We found that overall, significantly fewer NeuN⁺/EdU⁺ cells were found in the miR-206 200a inhibitor animals (Fig. 2C, D). Collectively, these data demonstrated that the total 207 number of the Sox2⁺ spinal cord stem cells is relatively the same in control versus miR-208 200a inhibited animals, but less NeuN⁺ cells are found in the inhibitor treated animals. 209 These findings suggest that either more cells remain in a progenitor-like state, or the 210 expression of *brachyury* in the miR-200a inhibitor treated animals changes the fate of 211 the cells. To address this question, we used *in vivo* cell tracking to determine the fate of 212 these cells during regeneration of the lesioned spinal cord. We have previously tracked 213 the fate of GFAP⁺ spinal cord stem cells during regeneration of the lesioned spinal cord 214 and found that these cells proliferate and migrate to replace the portion of injured neural 215 tube, and that this is a bidirectional process (Sabin et al., 2015a). The same technique 216 was used in these studies, the axolotl GFAP promoter driving expression of a 217 fluorescent protein was injected into the lumen of the spinal cord, the animals were 218 electroporated to label small groups of cells. The miR-200a inhibitor was injected into 219 animals with fluorescently labelled cells and then the spinal cord ablation was performed 220 (Sabin et al., 2019). The animals were imaged every 3-days over a two-week time 221 period. In the control labelled animals, we found the cells behaved as we had previously 222 described, the labeled cells proliferated and partook in repair of the neural tube, 223 replenished the endogenous stem cell population, and differentiated to replace lost 224 neurons (Fig.3 A-D), consistent with our previous work (Sabin et al., 2015). In contrast, 225 in the miR-200a inhibitor treated animals although the cells proliferated and partook in 226 repair of the lesioned spinal cord, we also discovered that the cells exited from the spinal 227 cord and differentiated into muscle cells. The labelled cells which started in the spinal 228 cord where always found in the muscle layer adjacent or directly above the spinal cord 229 (Fig.3I, J). In all miR-200a inhibitor treated animals we observed at least 1 muscle fiber 230 being formed in all animals (n=25), although in some animal's multiple fibers were seen. 231 Additionally, in inhibitor and control animals' some cells differentiated into neurons and 232 remained within the neural tube to give rise to new glial cells (data not shown). This data 233 suggests that miR-200a represses brachyury in sox2⁺ spinal cord stem cells, maintaining 234 the cells in neural primed state. Inhibition of miR-200a in these cells results in the co-235 expression of neural (sox2) and mesoderm (brachyury) markers, converting the cells into 236 a bipotent progenitor population capable of making both neural and mesodermal cells. 237

238 Molecular regulation of progenitor cells by miR-200a

239 Our data strongly indicate that inhibition of miR-200a leads to the expression of 240 brachyury in stem cells within the axolotl spinal cord (Fig. 1D,). However, the signaling 241 pathway(s) upstream of brachyury expression were not known. As a first step we first 242 tested whether miR-200a could directly repress brachyury expression. The axolot 243 brachyury 3' untranslated region (UTR) contains three miR-200a seed sequences, this 244 indicates that miR-200a could directly regulate *brachyury* expression. Consistent with 245 this hypothesis, co-transfection of B35 neural cells with a *brachyury* 3' UTR luciferase 246 reporter and miR-200a mimic led to decreased luciferase activity compared to the 247 control mimic (Fig. S2A). This finding confirmed that miR-200a directly represses 248 brachyury expression in axolotl spinal cord stem cells in homeostatic conditions and 249 during normal regeneration.

250 During normal spinal cord regeneration in the context of a tail amputation model 251 it has been found that wnt genes are re-expressed in the caudal 500µm of the 252 outgrowing spinal cord and are necessary for this outgrowth (Albors et al., 2015). Further 253 studies have shown that inhibition of all Wnt or Fgf signaling after tail amputation 254 abolished regenerative outgrowth, suggesting both are necessary for spinal cord and tail 255 regeneration (Ponomareva et al., 2015). As both Fgf and Wnt signaling regulate cell fate 256 decisions of brachyury⁺/sox2⁺ NMPs during development, we first tested whether Fgf 257 signaling could be affected by miR-200a inhibition during regeneration. We assayed for 258 expression of fgf8 and fgf10 by qRT-PCR on isolated spinal cord tissue. Fgf8 expression 259 was slightly down-regulated at 4 days post injury after miR-200a inhibition compared to 260 uninjured spinal cords (Fig.4A), while fgf10 expression was significantly up-regulated 261 after miR-200a inhibition in 4 days post injury spinal cords compared to controls 262 (Fig.4A). This finding is consistent with the idea that miR-200a inhibition could lead to an 263 increase in fgf ligand expression in regenerating spinal cords. However, given that Wnt 264 signaling directly regulates Brachyury expression (Arnold et al., 2000, Yamaguchi et al., 265 1999) and NMP cell fate decisions (Bouldin et al., 2015, Garriock et al., 2015, Martin, 266 2016, Martin and Kimelman, 2008), we wanted to further examine the role of Wnt 267 signaling.

The expression levels of *wnt3a*, *wnt5a*, *wnt8a* were quantified using qRT-PCR (**Fig. 4B**). Both *wnt3a* and *wnt8* transcript levels were not significantly altered in the inhibitor treated animals compared to controls. However, we did detect a significant difference in *wnt5a* levels. In 4 day post injury controls, *wnt5a* was up-regulated after injury, although this change in expression was not found in the miR-200a inhibitor

273 treated animals. To further verify the qRT-PCR results for faf and wnt genes we 274 performed fluorescent in situs for faf10 and wnt5a in control and miR-200a inhibitor 275 treated regenerating animals. This confirmed that indeed fgf10 transcript levels are up-276 regulated in cells within the spinal cord in comparison to the control regenerating 277 animals (Fig.4C, D). Additionally, wnt5a transcript levels were down-regulated in cells 278 within the spinal cord in comparison to controls (Fig.4C, D). Although here we see only 279 changes in wnt5a expression, a Wnt which is known to play an important role in 280 regeneration (Albors et al., 2015), there are many additional Wnt ligands, therefore Wnt 281 signaling activity could still be affected by miR-200a inhibition. To establish a baseline 282 for Wnt signaling activity after spinal cord injury we assayed *lef1* expression, which is a 283 direct transcriptional target downstream of Wnt signaling (Filali et al., 2002). Lef1 284 expression was significantly up-regulated in control 4 days post injury compared to 285 uninjured spinal cords (Fig. S3A), indicating a potential increase in Wnt signaling after 286 injury. Remarkably, *lef1* expression was significantly up-regulated even further after miR-287 200a inhibition in 4 days post injury compared to control regenerating spinal cords (Fig. 288 **S3A**). Collectively, these data indicate that miR-200a inhibition could result in increased 289 Wnt signaling, potentially independent of changes in *wnt* ligand expression.

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291 miR-200a modulates Wnt signaling activity by directly targeting β-catenin

292 While miR-200a inhibition could lead to increased Wnt signaling, it was not clear 293 how this was occurring. During tumor progression, miR-200a inhibits the epithelial-to -294 mesenchymal transition subsequently blocking tumor cell metastasis (Su et al., 2012, 295 Zaravinos, 2015). This is partially achieved through the direct repression of β -catenin by 296 miR-200a, resulting in decreased Wnt signaling (Su et al., 2012). We did not observe a 297 significant up-regulation of specific wnt ligand expression after miR-200a inhibition in the 298 spinal cord cells (Fig. 4B). However, as determined by *lef1* expression, miR-200a 299 inhibition could lead to increased Wnt signaling (Fig. S3A). Therefore, we hypothesized 300 that miR-200a might regulate Wnt signaling by targeting β -catenin. To test our 301 hypothesis, we first assayed for changes in β -catenin transcript abundance (ctnnb1). 302 qRT-PCR analysis confirmed that after injury in control 4 days post injury spinal cords, 303 there is an increase in *ctnnb1* abundance compared to uninjured spinal cords, similar to 304 what we observed for *lef1* (Fig. S3A). There is a slight increase of *ctnnb1* transcript 305 levels after miR-200a inhibition compared to control 4 days post injury spinal cords (Fig. 306 **S3A**), indicating β -catenin could be a direct target of miR-200a in axolotl.

307 To determine whether miR-200a could target axolotl β -catenin, we cloned the 308 ctnnb1 3' UTR and identified two miR-200a seed sequences. We subcloned the ctnnb1 309 3' UTR into a luciferase reporter and co-transfected cells with a control mimic or miR-310 200a specific mimic. There was decreased luciferase activity in miR-200a mimic 311 transfected cells compared to control, suggesting that miR-200a could regulate β -catenin 312 expression (Fig. S3B). To confirm that the decrease in luciferase activity is due to direct 313 regulation by miR-200a, we mutated both seed sequences in the *ctnnb1* 3' UTR and 314 repeated the luciferase experiments. Mutation of the miR-200a seed sequences 315 completely alleviated the repression, confirming that axolotl β -catenin is a direct target of 316 miR-200a, similar to mammals (Fig. S3B).

Taken together, these data are consistent with the idea that miR-200a could modulate Wnt signaling through the direct regulation of β -catenin transcript levels. Inhibition of miR-200a leads to increased *lef1* expression, which is indicative of increased Wnt signaling. Increased levels of Wnt signaling may contribute to the increased *brachyury* expression and changes in *fgf10* levels in axolotl stem cells after spinal cord lesion.

323

The role of spinal cord stem cells in spinal cord injury versus tail regeneration 324 325 We have shown that when a lesion occurs in the axolotl spinal cord the glial cells 326 adjacent to the injury site respond to the injury cue and proceed to behave like NSCs; 327 they divide, migrate, self-renew and replace lost neurons. However, previous work has 328 shown that during spinal cord regeneration after amputation, rather than injury, these 329 glial cells can transdifferentiate and give rise to cells of both the ectodermal and 330 mesodermal lineage (Echeverri and Tanaka, 2002, McHedlishvili et al., 2007). We next 331 examined the expression of *brachyury* in the context of whole tail regeneration and 332 discovered that brachyury is expressed in the $sox2^+$ stem cells of the spinal cord 500µm 333 adjacent to the injury site at 7-days post amputation (Fig. 5). To determine if this is an 334 attribute of the larval animals only, we also examined regenerating tail tissue from 2-year 335 old adult animals and found that in response to tail amputation that these progenitor cells 336 in the adult spinal cord indeed co-express *brachyury* and *sox2* during tail regeneration in 337 adult (Fig. 5) and larval animals (Fig.S4). These data suggest that during the 338 regenerative process the cells lining the central canal determine what tissue types need 339 to be restored. When only a small portion of the neural tube must be regenerated 340 following injury, the progenitor cells adopt a neural stem cell state to successfully

341 regenerate the spinal cord. During whole tail regeneration following amputation when

342 multiple tissue lineages must be regenerated, these cells within the spinal cord become

bipotent progenitors capable of making mesoderm and ectoderm (**Fig.6**).

344 Collectively, these experiments have shed light on the context dependent nature of

345 miRNA signaling during spinal cord lesion repair versus tail amputation and have

- 346 identified new signaling pathways that regulate progenitor cell fate during axolotl spinal
- 347 cord regeneration.
- 348

349 **Discussion**

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351 The current study has identified miR-200a as a regulator of stem cell fate in the 352 regenerating axolotl spinal cord. GO term analysis of genes down-regulated in the 353 uninjured and 4 days post injury spinal cord after miR-200a inhibition showed that these 354 genes were involved with nervous system development, organismal development, 355 synaptic signaling, and cellular differentiation (Fig.1, Fig.S1). Specifically, genes 356 involved with neuronal differentiation (*neurog1, neuroD4*) and neuronal processes like 357 synaptic transmission (CHRNB1, GABRA4) and neurotransmitter uptake (SLC6A6, 358 SLC18A3, SLC6A14) were down-regulated (Fig.1, Fig. S1). This suggests that miR-359 200a normally functions to promote NSC identity. This is consistent with multiple reports 360 across various species that inhibition of miR-200a and other miR-200 family members 361 results in the loss of neural progenitor identity and precocious neuronal or glial 362 differentiation (Buller et al., 2012, Choi et al., 2008, Morante et al., 2013, Peng et al., 363 2012. Trumbach and Prakash. 2015. Valleio et al., 2011). However, we have found that 364 in the axolotl spinal cord that even in uninjured conditions in larval or adult axolotls, the 365 cells lining the central canal in fact express low levels *brachyury* and *sox2*, the classical 366 markers of mesoderm and neural stem cells (Fig.1D, Fig.5). These cells may represent 367 a bipotent progenitor cell population and our data suggests that increased levels of 368 brachyury are necessary for a progenitor to make the decision to exit the spinal cord and 369 become a cell type of mesodermal origin (Fig.6).

During embryonic development in multiple species a small population of cells that co-express Sox2 and Brachyury have been identified and are now called neuromesodermal progenitor cells (Gouti et al., 2014, Henrique et al., 2015, Jurberg et al., 2013, Kimelman, 2016b, Taniguchi et al., 2017, Tsakiridis et al., 2014, Tsakiridis and Wilson, 2015, Turner et al., 2014b, Tzouanacou et al., 2009). Neuromesodermal

375 progenitor cell commitment to the neural lineage is partially determined by the relative 376 levels of Sox2 compared to Brachyury, given that the two transcription factors function to 377 antagonize one another (Koch et al., 2017a) and by the respective levels of fgf versus 378 wnt that the progenitor cells encounter (refs). During axolotl development to date a 379 definitive population of neuromesodermal progenitors has not been defined, however 380 work published by Taniguichi et al has shown that a posterior region of the axolotl neural 381 plate is positive for *brachyury* and *sox2* and this region gives rise to mesoderm during 382 development (Taniguchi et al., 2017). This finding is consistent with the idea that the 383 axolotl may also have a bipotent progenitor pool of cells established during early 384 development, however more work is needed, especially lineage tracing to establish if 385 these cells behave similar to what is seen in other species like chick, mouse and 386 zebrafish. Our results here showing that by gRT-PCR and RNAscope in situs that 387 brachyury and sox2 are detected in the progenitor cells of the spinal cord in both larval 388 and adult axolotis would suggest that axolotis retain a population of cells in the spinal 389 cord throughout life that are bipotent. Work from McHedlishvili et al previously showed 390 that adult axolotl retains expression of embryonic markers of dorsal/ventral patterning 391 like pax7, pax6 and shh genes that are not expressed in adult mammalian spinal cord 392 (McHedlishvili et al., 2007). They additionally showed that like earlier lineage tracing 393 work in axolotl, cells from the spinal cord do in fact migrate out and form a range of other 394 cell types including blood vessels, skin, cartilage, and muscle cells. Overall, these 395 bodies of work indicate that the cells in the axolotl spinal cord retain a multi-potent 396 progenitor cell state and are capable to respond to injury cues which direct them towards 397 different cell fates as needed. Very early work on tail regeneration in salamanders had 398 already hinted that the terminal vesicle structure formed at the growing end of the spinal 399 cord during tail regeneration is an area of epithelial to mesenchymal transition where 400 cells delaminate from the neural tube and exit to contribute to regeneration of 401 surrounding tissues of other developmental lineages (Benraiss et al., 1997, Egar and 402 Singer, 1972, O'Hara et al., 1992). Our data now gives molecular insights into the 403 identity of these cells. We found that miR-200a inhibited cells increase levels of 404 brachyury and then these cells during regeneration of a spinal cord lesion will form 405 muscle which is not observed in control regenerating lesions. However, we have not 406 observed these cells to form cartilage, skin, fin mesenchyme of any other cell type. We 407 cannot rule out that they have this potential, but the lineage tracing is limited as the 408 fluorescent protein expression is driven by the GFAP promoter, we expect this promoter

409 is turned off as the cells differentiate and as we can only image very 3 days, we may in

410 fact miss some differentiation events. We observed in all animals where miR-200a is

411 inhibited that at least one muscle fiber is formed from the labelled cells, however, again

412 we may be missing some differentiation events due to the limitations of our labelling

413 technique.

414 It is still not clear whether Brachyury directly regulates Sox2 levels in the regenerating

415 axolotl spinal cord or whether it is via an indirect mechanism. Work from other labs in

416 other research organisms has indicated that Brachyury and Sox2 can have a mutually

417 repressive relationship (Kimelman, 2016a, Koch et al., 2017b, Martin, 2016). We have

418 shown that miR-200a directly regulates *brachyury* and β -catenin via seed sequences in

419 the 3'UTR of these genes. When miR-200a is inhibited in the spinal cord cells, *brachyury*

420 is expressed at higher levels in these cells but *fgf* and *wnt* levels are also perturbed.

421 Work in development on NMP's has shown that feedback loops exist between

422 brachyury, fgf and wnt genes, and hence a complex signaling network may exist that is

423 driven by specific levels of certain regulators in these cells at particular times. An

424 additional level of complexity is the fact that Wnt is a secreted protein and although we

425 see downregulation of it within the progenitor cells in the spinal cord, we also see that

426 cells outside the spinal cord express *wnt* (**Fig.4**) and therefore the progenitor cells may

427 also be influenced by external gradients of Wnt protein.

428 During development, Wnt and Fgf signaling tightly regulate neuromesodermal cell fate

429 decisions (Goto et al., 2017, Gouti et al., 2017, Gouti et al., 2015, Martin, 2016) and both

430 genes are known to play important roles in regeneration (Sun et al., 2002, Wilson et al.,

431 2000, Zhang et al., 2000, Zhang et al., 2002b, Caubit et al., 1997, Ghosh et al., 2008,

Lin G, 2008, Stoick-Cooper et al., 2007, Tanaka and Weidinger, 2008, Wehner et al.,

433 2017, Zakany and Duboule, 1993). Canonical Wnt signaling is crucial for radial glial cell

434 proliferation during neural tube development (Shtutman et al., 1999) and spinal cord

435 regeneration in zebrafish (Briona et al., 2015). Therefore, it is not surprising to see a

436 potential increase in Wnt signaling during spinal cord regeneration in axolotl. However, it

437 is interesting that miR-200a does not regulate expression of *wnt* ligands, but instead

438 regulates β -catenin levels (**Fig. S3**). This is reminiscent to the role of miR-200a in

inhibiting EMT by repressing β -catenin and canonical Wnt signaling (Su et al., 2012,

440 Zaravinos, 2015). The increase in *ctnnb1* levels after miR-200a inhibition is not

441 statistically significant, however slight changes in transcript abundance can have

442 profound effects on protein levels (Schwanhausser et al., 2011). Therefore, a modest

443 increase in transcript abundance could represent a biologically significant increase in β -

444 *catenin* protein levels.

445 The signals that inform injured cells what tissue must be replaced remain a mystery. 446 Here we show that glial cells in the spinal cord appear to sense the difference between a 447 lesion of the spinal cord that primarily needs replacement of neural stem cell and 448 neurons, versus regeneration in the context of whole tail regeneration where cells of 449 multiple developmental germ layer origin must be regenerated. Interestingly we find that 450 cells of both the larval and adult tail regenerate bipotent progenitors that express 451 brachyury and sox2 in response to tail amputation, suggesting that the presence of 452 these bipotent progenitors is not only a hallmark of embryonic development, but rather a 453 stem cell population that is maintained in the animals specifically for regeneration. In the 454 future it will be important to determine if all cells in the spinal cord have this potential or 455 whether there are sub-populations of stem cells present in the axolotl spinal cord. 456 457 458 459 460 461 Materials and Methods 462 463 Animal Handling and Spinal Cord Injury All axolotls used in these experiments were obtained and bred at the University 464 465 of Minnesota or the Marine Biological Laboratory in accordance with IACUAC 466 regulations. Prior to all *in vivo* experiments, animals (3-5cm) were anesthetized in 0.01% 467 p-amino benzocaine (Sigma). Spinal cord ablations were performed as previously 468 described (Diaz Quiroz et al., 2014, Sabin et al., 2015a). Briefly, a 26-gage needle was 469 used to clear away skin and muscle to expose the spinal cord 6-10 muscle bundles 470 caudle to the cloaca. Then, using the needle, a segment of spinal cord 1 muscle bundle 471 thick, approximately 500µm, was removed. Animals were placed in cups and monitored 472 for the duration of the experiments. 473 474 Immunohistochemistry and EdU 475 Tissue was harvested and fixed in fresh 4% paraformaldehyde (Sigma) overnight

476 at 4°C. Then tails were washed three times in phosphate buffered saline + 0.1% Tween

477 20 (PBSTw). Next the tails were incubated in a 50:50 solution of PBSTw and 30%

478 sucrose. Finally, tails were transferred to 30% sucrose solution and allowed to

479 equilibrate overnight at 4°C. The next day samples were embedded for cross-sectioning

480 in TissueTek (Sakura) and stored at -20°C.

481 For EdU staining, animals were injected intraperitoneal with EdU at a 482 concentration of $0.5 \,\mu g/\mu L$ in PBS+1% Fast Green at 5 and 7-days post injury then 483 harvested at 14-days post injury. The tissue was processed for sectioning as described 484 above and stained using the Click-iT EdU Imaging Kit (Invitrogen) according to the 485 manufacturer's instructions.

486 After staining for EdU, samples were processed for immunohistochemical 487 analysis using either anti-Sox2 (Abcam) or anti-NeuN (Chemicon) primary antibodies as 488 previously described (Sabin et al., 2015a, Sabin et al., 2019). Briefly, slides were 489 subjected to a boiling citrate antigen retrieval step and then washed with PBSTw 3 times 490 for 5 minutes each. Samples were blocked (PBS+0.1% Triton-X+2% bovine serum 491 albumin +2% goat serum) for an hour at room then incubated overnight at 4°C in primary 492 antibodies diluted (1:100) in blocking buffer. The next day, slides were washed 4 times 493 with PBSTw and then incubated with secondary antibody (Invitrogen) diluted in blocking 494 buffer (1:200) for 2 hours at room temp and cell nuclei were counterstained with 4',6-495 diamidino-2-phenylindole (DAPI) (1:10,000). After secondary incubation the slides were 496 washed four times with PBSTw and mounted in Prolong Anti-fade mounting solution 497 (Invitrogen). All samples were imaged using an inverted Leica DMI 6000B fluorescent 498 microscope.

499 All images were generated using Fiji and cells were counted with the Cell 500 Counter plugin.

501

502

Quantitative Reverse Transcriptase Polymerase Chain Reaction

503 Injured spinal cords 500µm rostral and 300µm caudal to the lesion from 7-10 504 control or miR-200a inhibitor electroporated animals were micro dissected and pooled 505 for each biological replicate. Total RNA was isolated using Trizol (Invitrogen) according 506 to the manufacturer's instructions. Subsequent cDNA was synthesized from 1µg of 507 DNasel (NEB) treated RNA using either High Capacity cDNA Reverse Transcription kit 508 (Applied Biosystems) or miScript II RT kit (Qiagen). The gRT-PCR was carried out using 509 Light Cycler 480 SYBR Green I Master (Roche). MicroRNA gRT-PCR was carried out

- 510 with custom designed primers to conserved miRNAs (Qiagen) and custom primers from
- 511 IDT were used to quantify axolotl mRNAs:
- 512
- 513 18S_F: CGGCTTAATTTGACTCAACACG
- 514 *18S_*R: TTAGCATGCCAGAGTCTCGTTC
- 515 brachyury_F: GAAGTATGTCAACGGGGAAT
- 516 brachyury_R: TTGTTGGTGAGCTTGACTTT
- 517 sox2_F: TTGTGCAAAATGTGTTTCCA
- 518 sox2_R: CATGTTGCTTCGCTTTAGAA
- 519 wnt3a_F: AAGACATGCTGGTGGTCTCA
- 520 *wnt3a_*R: CCCGTACGCATTCTTGACAG
- 521 wnt5a_F: ACCCTGTTCAAATCCCGGAG
- 522 wnt5a_R: GGTCTTTGCCCCTTCTCCAA
- 523 wnt8a_F: TTGCTGTCAAATCAACCATG
- 524 wnt8a_R: TGCCTATATCCCTGAACTCT
- 525 ctnnb1_F: ACCTTACAGATCAAAGCCAG
- 526 ctnnb1_R: GGACAAGTGTTCCAAGAAGA
- 527 *lef1_*F: GTCCCACAACTCCTACCACA
- 528 *lef1_*R: TAGGGGTCGCTGTTCACATT
- 529 *fgf8_*F: TTTGTCCTCTGCATGCAAGC
- 530 *fgf8_*R: GTCTCGGCTCCTTTAATGCG
- 531 *fgf10_*F: AAACTGAAGGAGCGGATGGA
- 532 fgf10_R: TCGATCTGCATGGGAAGGAA
- 533
- 534 Brachyury and Sox2 Probe Synthesis
- 535 Approximately 500bp fragments of axolotl *brachyury* and *sox2* were PCR
- 536 amplified using OneStep PCR Kit (Qiagen) from RNA extracted from axolotl embryos at
- 537 various developmental stages using the following primers:
- 538
- 539 brachyury ISH For: CCCCAACGCCATGTACTCTT
- 540 *brachyury* ISH Rev: GGCCAAGCGATATAGGTGCT
- 541
- 542 sox2 ISH For: TGGCAATCAGGAAGAAGTC
- 543 sox2 ISH Rev: GCAAATGACAGAGCCGAACT

544

545 The resulting PCR fragments were gel purified using the Monarch Gel 546 Purification Kit (New England Biolabs) and TA cloned into pGEM-T Easy (Promega) then 547 transformed into DH5α competent *E. coli* (Invitrogen). Blue/White positive selection was 548 used to pick clones and recovered plasmids were sent for sequencing. Positive clones 549 were digested with the appropriate enzyme to linearize the plasmid and anti-sense 550 ribonucleoprobe synthesis was carried out using Sp6 or T7 RNA polymerase (New 551 England Biolabs)+DIG labeled UTP (Roche). Subsequent probes were cleaned up using 552 the RNA Clean Up kit (Qiagen) and resuspended in hybridization buffer.

553

554 Fluorescent In situ Hybridization

555 All RNAscope® in situ hybridization procedures were performed according to the 556 manufacturer's instructions (Advanced Cell Diagnostics). In brief, cryosections were 557 incubated in PBS for 10 minutes to remove the OCT, and then baked at 60°C for 30 558 minutes. The slides were next post-fixed in 4% paraformaldehyde for 15 minutes at 4°C, 559 and then dehydrated in a graded series of ethanol dilutions before being incubated in 560 absolute ethanol for 5 minutes. After briefly air-drying the slides for 5 minutes, sections 561 were next treated with hydrogen peroxide to quench endogenous peroxidase activity for 562 10 minutes at room temperature. Next, samples were briefly washed in deionized water, 563 then incubated in target retrieval buffer at 90°C for 5 minutes. Following target retrieval, 564 the slides were rinsed in deionized water for 15 seconds and treated with absolute 565 ethanol for 3 minutes. Slides were next permeabilized in protease III for 30 minutes 566 before hybridization with RNAscope® probes at 40°C for 2 hours. Following 567 hybridization, sections were placed in 5x SSC overnight. The next day, sections were 568 incubated in Amp1 and Amp2 at 40°C for 30 minutes each, followed by Amp3 for 15 569 minutes. Next, slides were treated with HRP-C1 to detect brachyury or fgf10, followed by 570 a 30-minute incubation in Opal-690 fluorescent dye. After treatment with HRP blocking 571 buffer, samples were next incubated in HRP-C2 to detect either sox2 or wnt5a, followed 572 by a 30-minute incubation in Opal-570 dye. After an additional treatment with HRP 573 blocking buffer, slides were counterstained with DAPI and imaged using a Zeiss 780 574 Confocal Microscope.

575

576 Lineage Tracing

577	Cells of the uninjured spinal cord were transfected with a construct containing a GFP or					
578	tdTomato fluorescent protein under the control of the axolotl GFAP promoter. The cells					
579	were injected and electroporated as previously described (Echeverri and Tanaka, 2003a,					
580	Sabin et al., 2015a). One day after electroporation the animals were screened for					
581	fluorescent cells. Positive animals were then injected with a control inhibitor or miR-200a					
582	inhibitor and then a spinal cord lesion performed as described in (Sabin et al., 2019).					
583	Animals were imaged every 3 days until the lesion site was no longer visible and the					
584	animals regained motor and sensory function, typically 12-14 days post injury.					
585						
586	Cloning 3' Untranslated Regions for miRNA Luciferase Assays					
587	For 3' UTR luciferase experiments, primers were designed to amplify the					
588	brachyury and β -catenin 3' UTR based off sequences obtained from axolotl-omics.org.					
589	All the 3' UTRs were amplified with a 5' Spel and 3' HindIII restriction site.					
590						
591	brachyury 3' UTR For 1 AGCACTAGTATGTGAAATGAGACTTCTAC					
592	brachyury 3' UTR Rev 1 TGCAAGCTTCTTATTCTTCCCATTTAACTTAAA					
593						
594	ctnnb1 3' UTR For 1 ATAACTAGTTTGTGTAATTTTTCTTAGCTGTCATAT					
595	ctnnb1 3' UTR Rev 1 ATCAAGCTTAATTGCTTTATAGTCTCTGCAGAT					
596						
597	ctnnb1 3' UTR SDM1 For AGTGCCTGATGAATTCAACCAAGCTGAG					
598	ctnnb1 3' UTR SDM1 Rev CTCAGCTTGGTTGAATTCATCAGGCACT					
599						
600	ctnnb1 3' UTR SDM2 For ATTTAATGGTGTAGGAATTCAATAGTATAA					
601	ctnnb1 3' UTR SDM2 Rev TTATACTATTGAATTCCTACACCATTAAAT					
602						
603	The PCR fragments and pMiR Report (Life Technologies) were digested with Spel and					
604	HindIII (NEB) and the fragments were ligated over night at 4° C with T4 DNA Ligase					
605	(NEB) and heat shock transformed into DH5 α competent <i>E. coli</i> (Promega).					
606						
607	Mutation of miR-200 sites in Brachyury 3' UTR					
608	To mutate the 3 miR-200a and 3 miR-200b sites in the axolotl Brachyury 3' UTR,					
609	we used the QuikChange Lightening Multi Site-Directed Mutagenesis kit (Agilent) as per					
610	the manufacturer's instructions.					

611					
612	miR-200a1 SDM: gactgctttctatggacactttttaatttctgaagataagctcccacccg				
613	miR-200a2 SDM: cacacataaatcttttcgtgctgaacaaattatgatccatgaaaccagtgcatcattt				
614	miR-200a3 SDM: tccaatgtgtgtaatcctctcaattatcgcctctgcgtgtagaatgtc				
615					
616	miR-200b1 SDM: atgcattacaatgcattgttttctggacggcaatgaaagctgtgatgaaatatttaagat				
617	miR-200b2 SDM: caccataagagacaataaatgcaccggaatactgtgatatttgatgcctgcac				
618	miR-200b3 SDM: gaatcattaccatgtatttatcaggccggaatattcaaaatgtgacttcctctgtga				
619					
620	<u>3' UTR luciferase experiments</u>				
621	B35 neuroblastoma cells were plated in a 96 well plate (Celltreat Scientific				
622	Products) at a concentration of 2.0 x 10^5 cells/mL and allowed to adhere overnight. The				
623	next day cells were co-transfected with the appropriate Luciferase 3' UTR reporter				
624	plasmid, β -Galactosidase control, and 100nM of miR-200a, miR-200b, or control mimic				
625	(Qiagen) per well using Lipofectamine 3000 (Invitrogen). After 48 hours luciferase				
626	activity was determined using Dual Light Luciferase & β -Galactosidase Reporter Gene				
627	Assay System (Thermo) according to the manufacturer's protocol.				
628					
629	Pie Chart and Venn Diagram Generation				
630	Pie charts were generated using previously published data (Sabin et al., 2019) to				
631	represent the total number of differentially expressed genes in a given comparison using				
632	Excel. Venn Diagrams were generated with Venny (v2.1.0) (Oliveros 2007-2015) and				
633	saved as .csv files to be modified in Adobe Illustrator.				
634					
635	<u>Gene Ontology (GO) Analysis</u>				
636	GO terms were determined using GOrilla (Eden et al., 2009). We used 2				
637	unranked list of genes: a background list (all differentially expressed genes in our data				
638	set) and a target list (genes that were differentially regulated in a given comparison).				
639	Using this approach, GOrilla generated a list of enriched Biological Process GO terms				
640	and we selected the top 9-13 terms with the lowest p-value and generated				
641	representative bar graphs using Excel.				
642					
643	Calculation of the proportion and distribution of neural stem cells and newborn neurons				

644 The number of Sox2⁺ neural stem cells were counted in control and miR-200a 645 inhibitor spinal cords 2-weeks post injury. The proportion of Sox2⁺ neural stem cells was 646 calculated by dividing the total number of Sox2⁺ neural stem cells by the total number of 647 DAPI⁺ spinal cord cells times 100. To analyze regenerative neurogenesis, control or 648 miR-200a inhibitor animals were injected with EdU at 5 and 7-days post injury, then tails 649 were harvested for cryosectioning at 14 days post injury. The proportion of newborn 650 neurons was determined by dividing the number of NeuN⁺/EdU⁺ double positive neurons 651 by the total number of NeuN⁺ neurons times 100.

To visualize whether the proportion of Sox2⁺ neural stem cells changed rostral and caudal to the injury site in control and miR-200a inhibitor spinal cords, we quantified the average number of Sox2⁺ neural stem cells at a defined distance rostral and caudal from the lesion. We divided that number by the average number of DAPI⁺ spinal cord cells at that same distance and then binned the vales from 3 adjacent serial sections encompassing a region of 80µm. We used the same method for graphing the distribution of newborn neurons relative to the injury site.

659

660 <u>Statistical Analyses</u>

661 All results are presented as mean +/- s.d. unless otherwise stated. Analyses 662 were performed using Microsoft Excel or GraphPad Prism v8. Data set means were 663 compared using ANOVA for three or more tests with a Tukey test (for multiple 664 comparisons) or Dunnett test (to compare to a control mean). When two groups were 665 compared an unpaired t-test was used. When multiple comparisons were made using a 666 t-test, an adjusted p-value was determined using the two stage Benjamin, Krieger, and 667 Yekutieli procedue with a false discovery rate <5%. Differences between groups was 668 considered significant at three different levels (p-values of *≤0.05, **≤0.01 and 669 ***≤0.001) and are indicated in the figure legends.

670

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- 673

674 **Competing Interests**

- The authors declare no competing financial interests.
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- 681

682 Data Availability

- 683 The authors declare that all data supporting the findings of this study are available within
- the article and its Supplementary Information files or from the corresponding author upon
- reasonable request. The RNA seq data has been deposited in the public GEO database
- 686 with the accession number GSE122939.
- 687
- 688
- 689

690 Figure Legends

691 Figure 1. miR-200a inhibition during spinal cord injury leads to brachyury expression 692 axolotl in spinal cord stem cells. (A) RNA-Sequencing analysis identified a large subset 693 of differentially regulated genes following injury. The Vehn diagram compares the 694 number of overlapping differentially expressed genes between uninjured, 4 days post 695 injury control and miR-200a inhibitor treated samples. (B) Pie-chart showing the relative 696 portion of all transcripts that are differentially regulated. Regeneration specific transcripts 697 are defined as differentially expressed transcripts with a log2fc > 1 or < 1 and padi < 0.05698 between 4dpi control animals and 4dpi animals treated with mir-200a inhibitor that were 699 not differentially expressed in uninjured animals. (C) Log2fold change 700 heat map demonstrates the 30 most up- and down-regulated genes in uninjured and 4 701 days post injury control versus miR-200a inhibitor treated spinal cords. This analysis 702 revealed that the transcription factor brachyury (T) is dramatically up-regulated after 703 miR-200a inhibition. (D) qRT-PCR analysis confirmed that miR-200a inhibition led to 704 increased brachyury expression and blocked the up-regulation of the neural stem cell 705 marker sox2 in 4 days post injury (dpi) spinal cords (n=3). (E) Fluorescent In situ 706 hybridization confirmed the gRT-PCR analysis and revealed miR-200a inhibition leads to 707 brachyury expression (n=5) in spinal cord cells and a failure to up-regulate sox2 708 expression (n=3). * $p\leq0.05$, ** $p\leq0.01$, *** $p\leq0.001$, N.S. is not significant. Error bars 709 represent \pm S.T.D. Scale bar= 50 μ m.

710

711 Figure 2: Chronic miR-200a inhibition affects the birth of new neurons. Inhibition of miR-712 200a for 2 weeks does not affect the (A) proportion or (B) distribution of Sox2⁺ stem cells 713 in the spinal cord throughout the regeneration zone (n=6). Inhibition of miR-200a for 2 714 weeks results in an smaller proportion of new born neurons compared to controls (C). 715 (D) An increased proportion of new born neurons reside closer to the injury site while 716 chronic miR-200a inhibition blocked this increase in new born neurons close to the injury 717 site. (n=6). *p≤0.05, **p≤0.01, *** p≤0.001, N.S. is not significant. Error bars represent 718 ±S.T.D.

719

Figure 3: miR-200a inhibited spinal cord cells form muscle during spinal cord lesion repair. The cells lining the central canal of the spinal cord were labelled using GFAP promoter driving GFP or tdTomato by injection and electroporation. Control cells were followed over a 14 day period and cells gave rise to new glial cells or neurons only (A-F) (n=20). Cells which were injected with the miR-200a inhibitor were followed in parallel over the same time period and were found to exit the spinal cord and give rise to muscle cells (G-L). (n=25). Scale bar = $50\mu m$.

727

Figure 4: miR-200a inhibition affects the expression of Wnt and FGF signaling ligands.
(A) qRT-PCR analysis revealed fg10, but not *fgf8*, was significantly up-regulated after
miR-200a inhibition (n=3). (B) qRT-PCR analysis showed that miR-200a inhibition
differentially affects the expression of *wnt5a*, but not additional Wnt ligands (*wnt3a*, *wnt8a*; n=3). (C) Fluorescent *in situ* hybridization in control (C) and miR-200a inhibitor
treated (D) animals confirmed the qRT-PCR analysis, and demonstrates an increase in

fgf10 expression and a down-regulation of *wnt5a* within stem cells in the spinal cord (n=6). *p≤0.05, **p≤0.01, *** p≤0.001, N.S. is not significant. Error bars represent ±S.T.D. Scale bar= $50\mu m$.

737

Figure 5. Spinal cord amputation leads to *brachyury* expression in spinal cord stem
cells. (Ai-iV) Fluorescent *in situ* hybridization revealed that *sox2* is abundant within the
uninjured adult spinal cord, while *brachyury* is absent (n=2). (Bi-iv) At 7 days post
amputation, *brachyury* was localized to spinal cord stem cells that share an overlapping
expression pattern with *sox2* (n=2). Scale bar= 50µm.

743

744 Figure 6. A proposed model for the role that miR-200a plays in different injury 745 paradigms. When a lesion occurs in the spinal cord miR-200a levels remain high, which 746 inhibits *brachyury* expression and modifies levels of β -catenin, potentially stabilizing a 747 neural stem cell identity in the cells adjacent to the injury site. After spinal cord injury 748 these cells replace neurons and glial only. In contrast, when the tail is amputated 749 progenitor cells respond to injury cues and replace multiple cell types of different 750 developmental origins. These cells in the spinal cord then up-regulate brachyury in the 751 sox2⁺ stem cells of the spinal cord and direct these cells to proliferate and form cells of 752 both ectodermal and mesodermal origin.

753

754 **Supplementary Figure 1:** miR-200a affects expression of common and unique gene 755 sets in the uninjured and regenerating spinal cord. (A, B, C) Pie chart representation of 756 the proportion of up-regulated (Red) or down-regulated (Blue) genes in (B) uninjured 757 control compared to uninjured miR-200a inhibitor electroporated spinal cords or (A) 4 758 days post injury control compared to 4 days post injury miR-200a inhibitor electroporated 759 spinal cords. (C). Pie chart illustrating the number of overlapping versus individual genes 760 that are differentially regulated. (B-F) Gene Ontology terms enriched in gene specifically 761 in (B) control regenerating or (E) genes common to all data sets or specific to uninjured 762 tissue (F).

763

Supplementary Figure 2: Multiple miR-200 members directly regulate the *brachyury* 3' UTR. (A) Co-transfection of B35 cells with a *brachyury* 3' luciferase reporter and a miR-200a (A) or miR-200b (B) mimic results in decreased luciferase activity compared to the control mimic (n=5). Mutation of all miR-200a seed sequences in the *brachyury* 3' UTR alleviates this repression suggesting it is a direct target of miR-200 in axolotl (n=4). *** $p \le 0.001$, N.S. is not significant. Error bars represent ±S.T.D.

770

Supplementary Figure 3: miR-200a may regulate the expression of Wnt signaling
 components. (A) qRT-PCR analysis revealed that miR-200a inhibition leads to increased

773	expression of Wnt signaling transcriptional components <i>lef1</i> and β -catenin. (B) Co-				
774	transfection of B35 cells with a β -catenin 3' UTR luciferase reporter and a miR-200a				
775	mimic leads to decreased luciferase activity compared to controls. Mutation of both miR-				
776	200a seed sequences in the β -catenin 3' UTR alleviates this repression (n=3). *p≤0.05,				
777	*** p≤0.001, N.S. is not significant. Error bars represent ±S.T.D.				
778					
779	Supplementary Figure 4: brachyury is expressed in regenerating spinal cord stem cells				
780	following amputation in larval animals. At 4 days post amputation (dpa) both sox2 (ii)				
781	and <i>brachyury</i> (iii) are abundant in the spinal cord stem cells and share an overlapping				
782	expression pattern (iv)(n=6). Scale bar= 50mm.				
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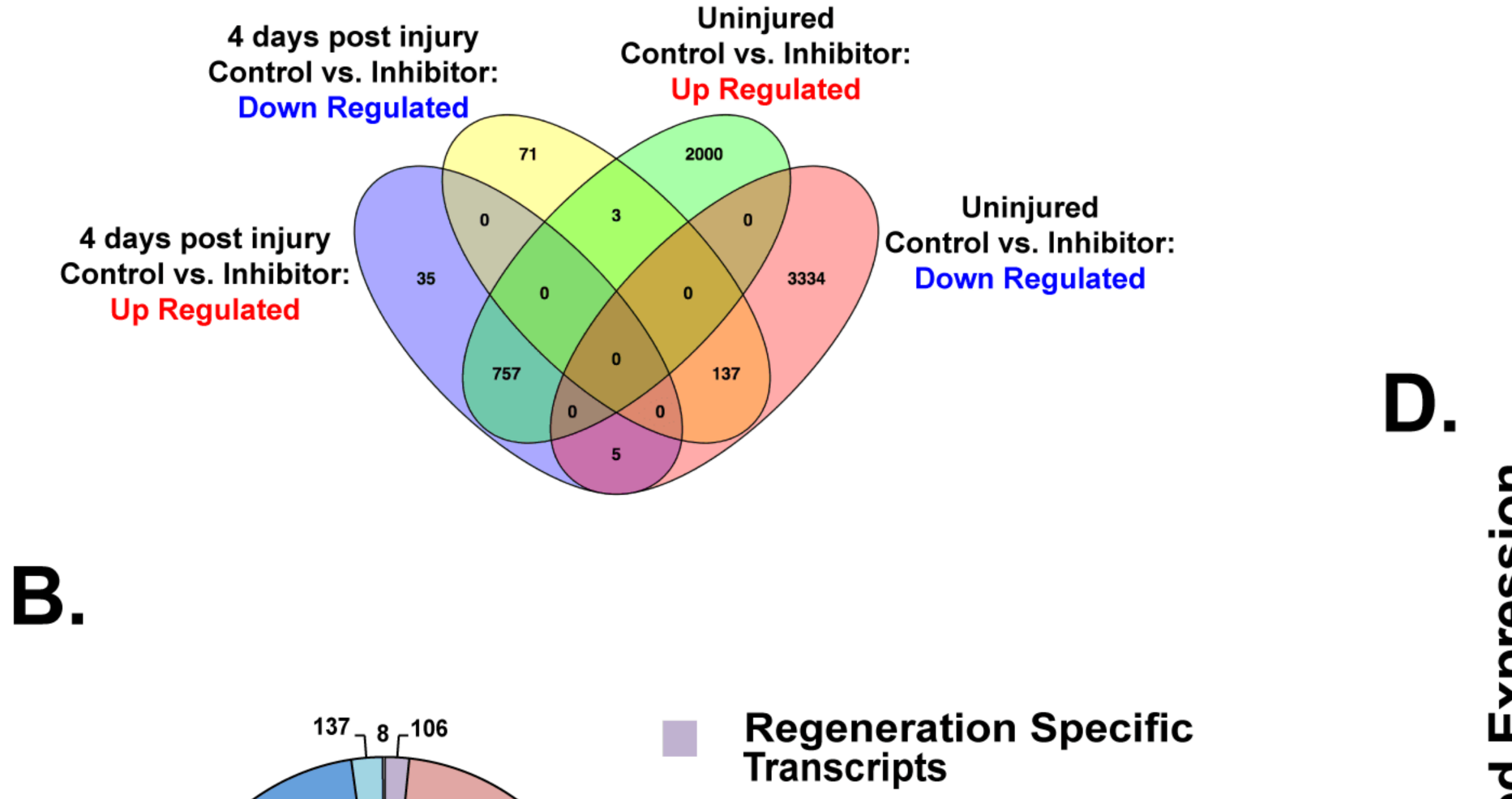
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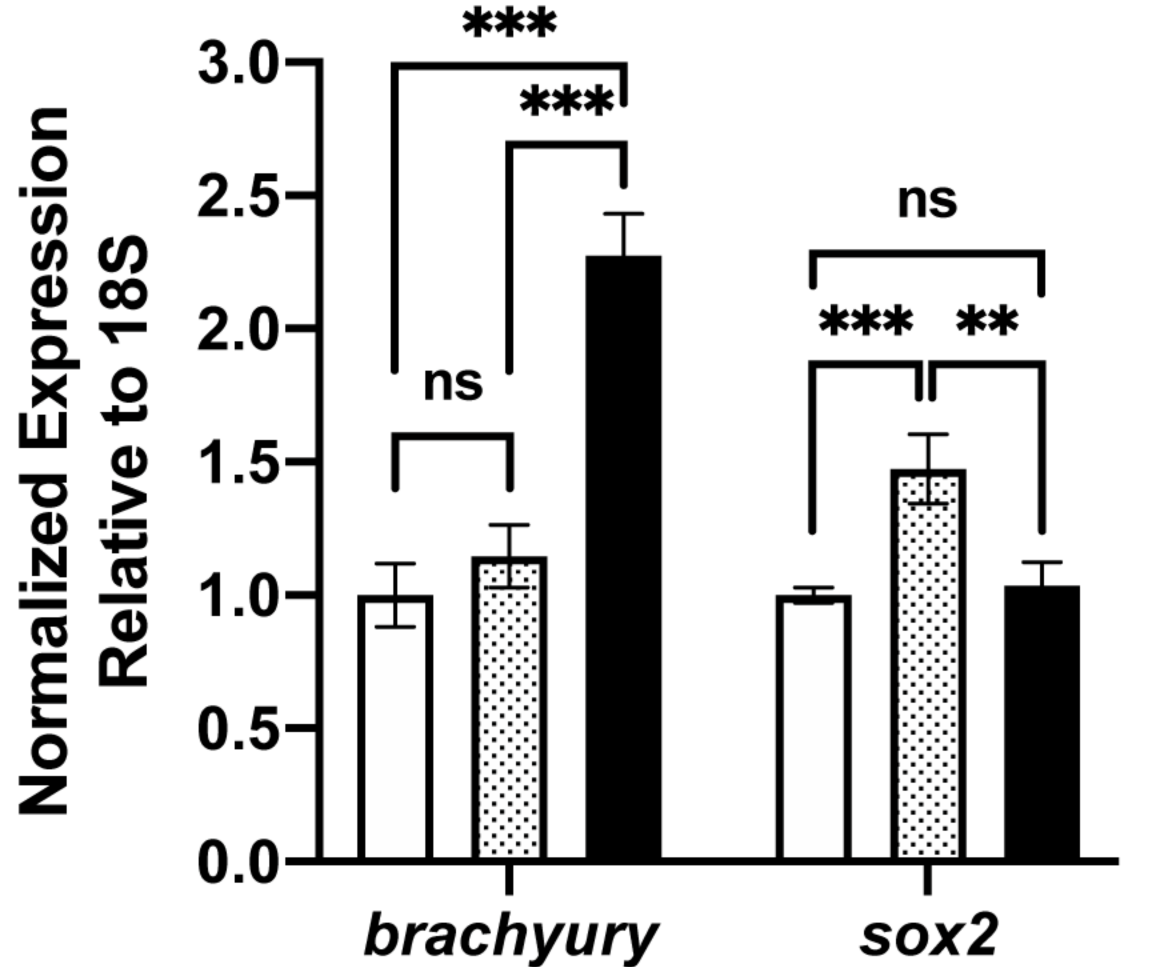
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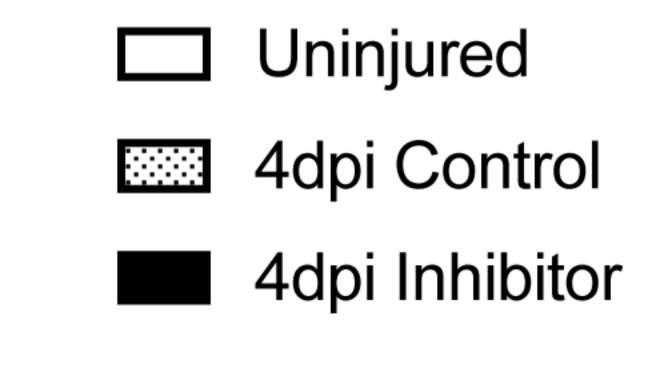
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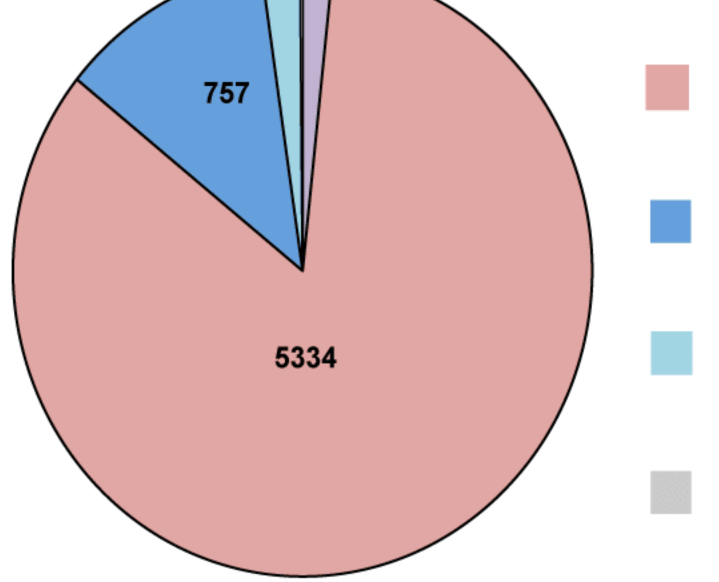


C.











- Common Up Regulated Transcripts
- Common Down Regulated Transcripts

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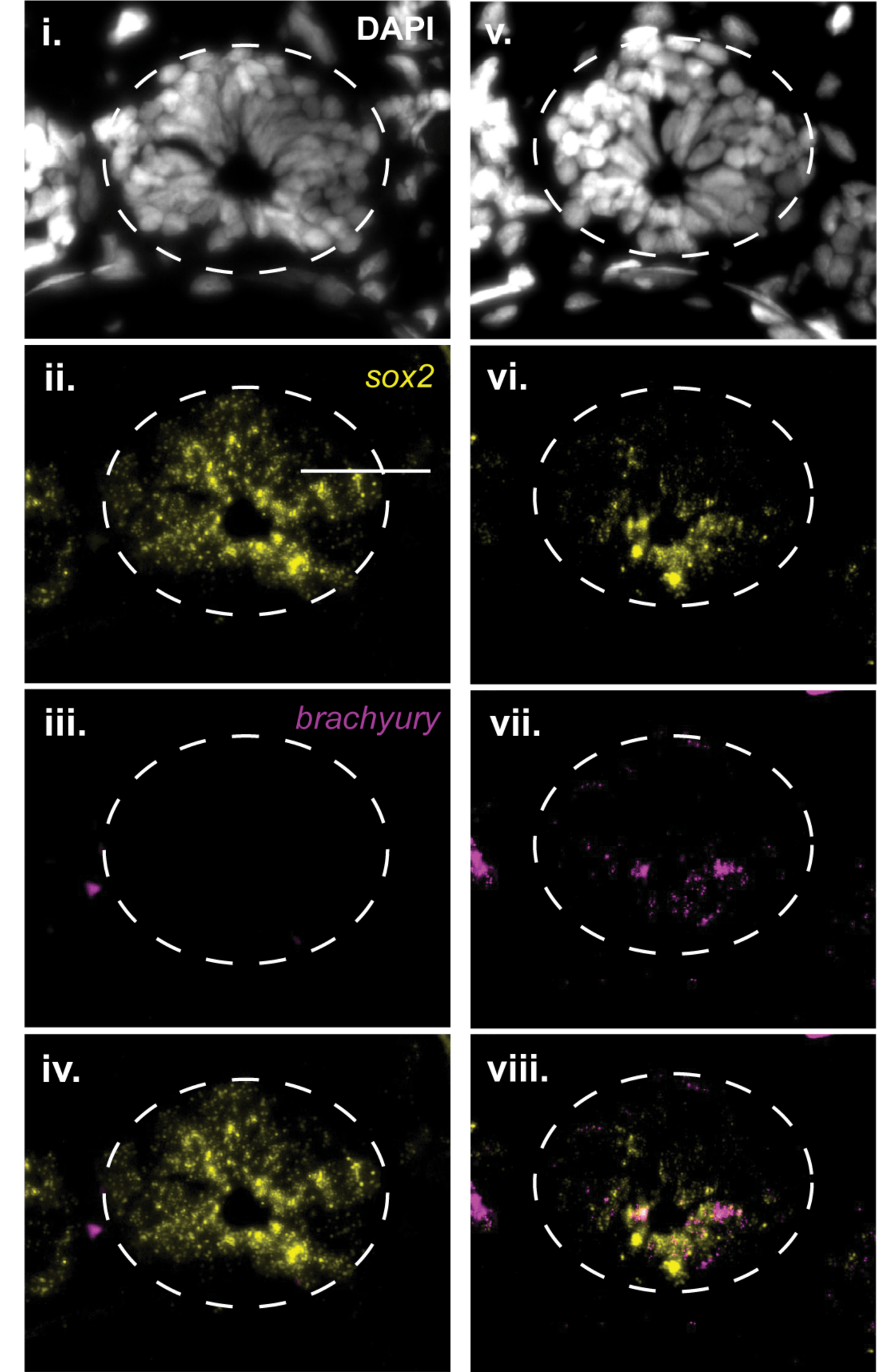
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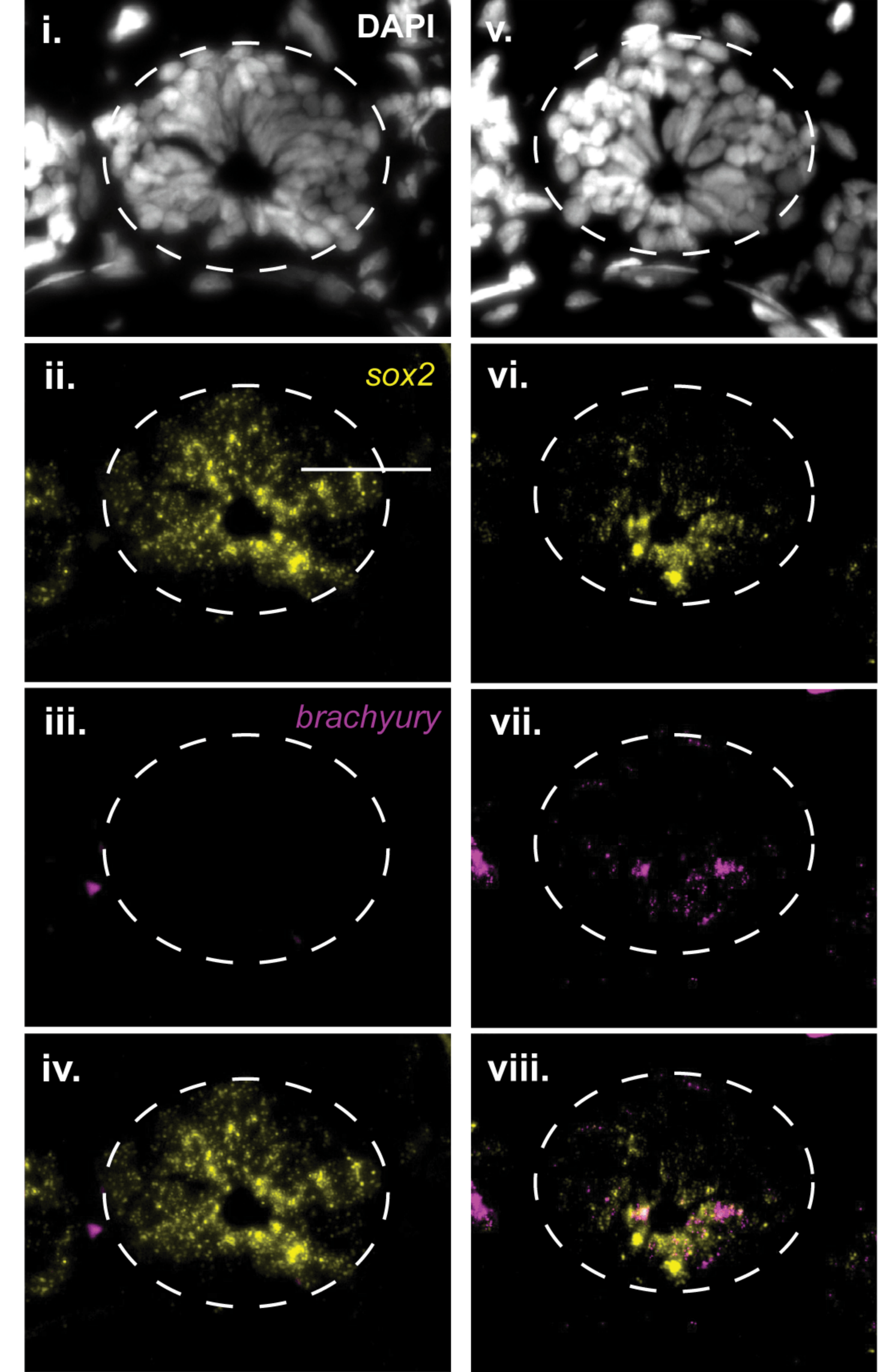
Other

4dpi Control

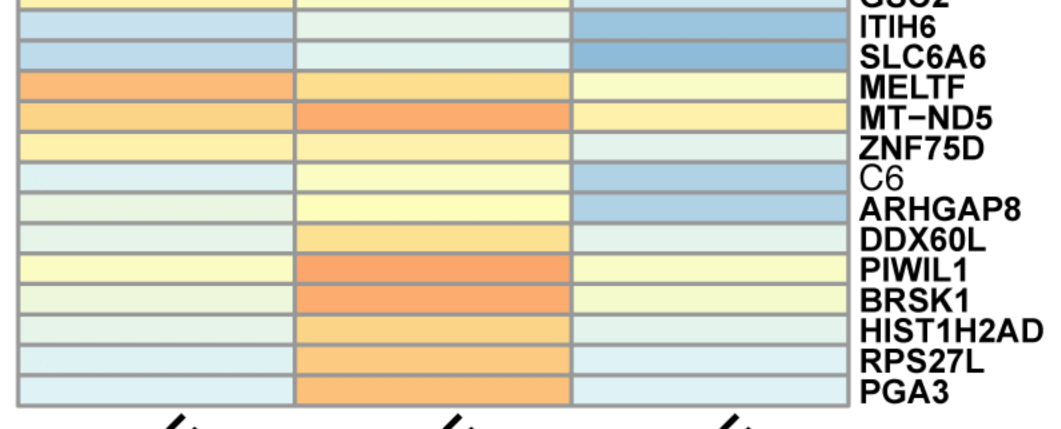
Ε.

4dpi Inhibitor





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			MRPL27
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			C3orf49
			C1QTNF2
			JMJD7-PLA2G4B
			 OTOS
			ZNF324B
			TYRP1
			HSPA1A
			CLEC3A
			HSPA1L
			T
			HSPA6
			FNDC7
			SSC5D
			PZP
			HAO1
			MATN3
			TCHP
			PRG2
			SLC13A1
			G6PC
			CCDC152
			MC5R
			 SCNN1A
			NEUROG1
			WFDC5
			APOA1
			PBXIP1
			 OASL
			MT-ND1
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			MOGAT1
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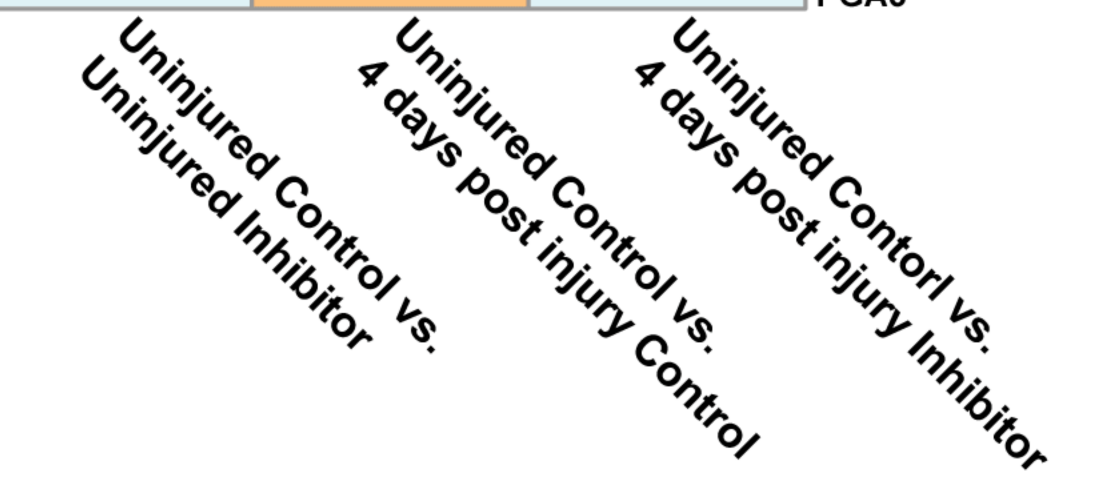
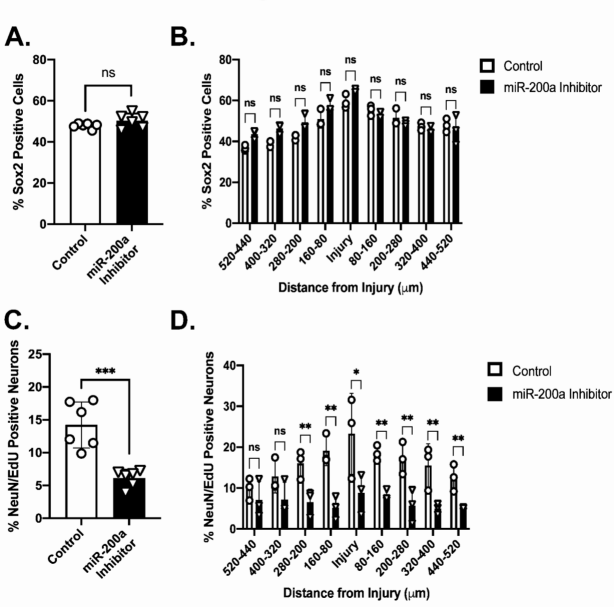
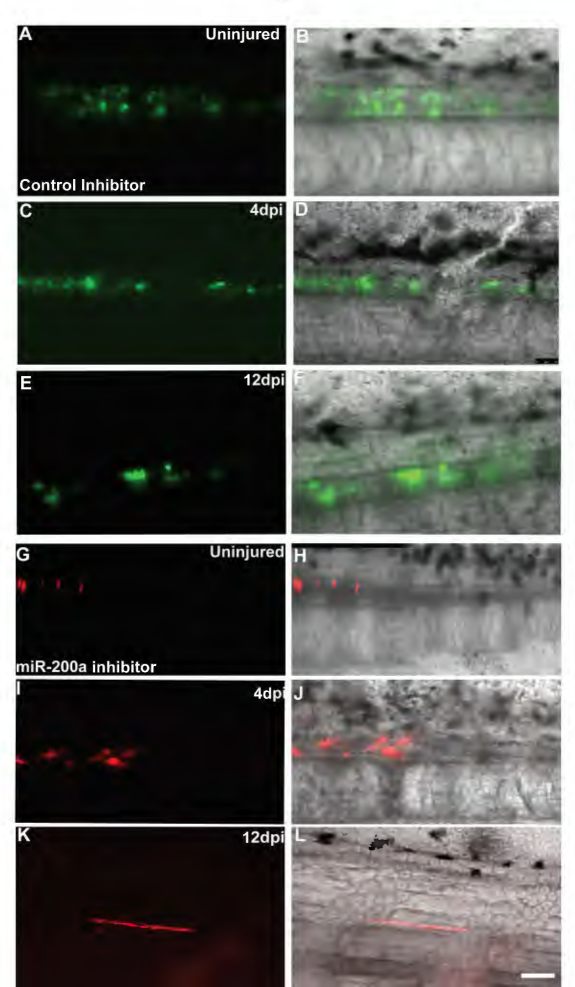


Figure 2





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