1 2	Multiple sources of Shh are critical for the generation and scaling of ventral spinal cord oligodendrocyte precursor populations
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4	Running Title: Shh sources scale spinal OPC numbers
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32 Abstract:

33	Graded Sonic Hedgehog (Shh) signaling emanating from notochord and floorplate patterns
34	the early neural tube. Soon thereafter, Shh signaling strength within the ventricular zone
35	becomes dis-contiguous and discontinuous along the ventral to dorsal axis suggesting a
36	distribution of Shh that cannot be achieved by diffusion alone. Here we discover that
37	sequential activation of Shh expression by ventricular zone derivatives is critical for
38	counteracting a precocious exhaustion of the Olig2 precursor cell population of the pMN
39	domain at the end of motor neuron genesis and during the subsequent phase of ventral
40	oligodendrocyte precursor production. Selective expression of Shh by motor neurons of the
41	lateral motor column at the beginning of oligodendrogenesis ensures a more yielding pMN
42	domain at limb levels compared to thoracic levels. Thus, patterned expression of Shh by
43	ventricular zone derivatives including earlier born neurons contributes to the scaling of the
44	spinal cord along the anterior - posterior axis by regulating the activity of a select
45	ventricular zone precursor domain at later stages of development.
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55 *Introduction*:

The "Bauplan", or body plan, of vertebrates can be recognized among members of the same 56 species and across phyla despite significant differences in the absolute size of individuals of the 57 same species or the form of different species. Chiefly responsible for our ability to recognize the 58 common features of a body plan is the proportionate differentiation and growth of the constituent 59 parts of an organism irrespective of absolute size. How developmental processes maintain a 60 constant ratio of physical pattern features with changing size, a property known as scale 61 62 invariance, is not completely understood (Huang and Umulis 2018). 63 Successful development is dependent on highly stereotyped series of inductive events that result 64 in the determination of progressively increasing numbers of cell fates within rapidly growing 65 tissues. Ensuring scale invariance could be achieved by mechanisms that are distinct from cell 66 fate determination processes and could act at different developmental stages (Barkai and Ben-Zvi 67 2009, Umulis and Othmer 2013). Alternatively, patterning mechanisms themselves could be 68 modified to generate a size-invariant output (Kicheva and Briscoe 2015). Cell fate determination 69 70 is governed by a handful of cell signaling factors termed morphogens that are secreted from spatial fix-points in the developing embryo and form gradients of activity across a developmental 71 field (Lander 2007). Morphogen signaling induces distinct transcriptional programs in naïve 72 73 precursor cells dependent on signaling thresholds that are determined by signal strength and duration (Kicheva, Bollenbach et al. 2014). Experimental and theoretical studies revealed several 74 mechanisms by which morphogen signaling can be scaled to embryo size (Gregor, Bialek et al. 75 2005, Houchmandzadeh, Wieschaus et al. 2005, Howard and ten Wolde 2005, McHale, Rappel 76

et al. 2006, Ben-Zvi and Barkai 2010, Cheung, Miles et al. 2014, Uygur, Young et al. 2016). The
commonality of these studies is that they define mechanisms of scale invariance across embryos
during early stages of development. However, scaling occurs within the same embryo and
throughout development as different body parts reveal proportionate changes in size regardless
of when they are specified during development. How is proportionality ensured across
developmental stages within the same embryo?

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We have explored this question in the context of the switch from neurogenesis to gliogenesis in 84 85 the developing neural tube. The spinal cord is a tube-like structure with enlargements at segmental levels that provide control of, and receive proprioceptive information from, limbs. The 86 larger spinal cord at brachial (forelimb) and lumbar (hindlimb) levels compared to thoracic levels 87 is chiefly the result of the presence of a greater number of motor neurons (MNs) that sub serve 88 limb musculature, and correspondingly larger numbers of interneurons and glial cells (Bjugn and 89 Gundersen 1993, Dasen 2017). Since the basic patterning and structure of the spinal cord is 90 highly similar along the neuraxis, the difference in size between limb levels and thoracic 91 segments allows comparative studies that might reveal mechanisms involved in scale invariance. 92 93 MNs are among the very first neurons in the developing neural tube whose fate becomes specified while most interneurons and all types of glia develop later (Jessell 2000). Whether and 94 how the earlier production of neurons plays a role in the proportionate differentiation of 95 96 subsequent cell types is not well understood. MNs and oligodendrocyte precursor cells (OPCs) emerge subsequently from the same domain, the "pMN" domain, within the ventricular zone of 97 98 the developing spinal tube (Bergles and Richardson 2015, Traiffort, Zakaria et al. 2016) 99 suggesting that cell-autonomous as well as cell non-autonomous mechanisms could be involved

in the sequential and proportionate differentiation of MNs and OPCs (Traiffort, Zakaria et al.
2016). The elucidation of the molecular control mechanisms that determine the precise "switch"
from neurogenesis to gliogenesis of the pMN domain and the segment-specific yield of the pMN
domain have been hampered by a scarcity of cell type selective- and temporally- specific genetic
tools to dissect MN and OPC production within the pMN domain. Despite these difficulties,
studies have implicated the morphogen Sonic Hedgehog (Shh) in regulating both, MN and OPC
production (Briscoe and Ericson 1999, Traiffort, Zakaria et al. 2016).

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Shh plays a critical role in MN and OPC differentiation at several developmental stages. First, 108 graded Shh signaling originating from the notochord induces distinct transcriptional programs in 109 overlying naïve neural ectoderm cells in a concentration-dependent manner that leads to the 110 111 establishment of molecularly distinct precursor domains along the ventral to dorsal axis of the developing neural tube (Dessaud, McMahon et al. 2008, Alaynick, Jessell et al. 2011, Yu, 112 McGlynn et al. 2013), The transcription factors activated by Shh are responsible for determining 113 the cell fates in the derivatives of these precursor domains. Thus, Olig2 expression marks MN 114 precursors, (pMN) (Mizuguchi, Sugimori et al. 2001, Novitch, Chen et al. 2001), Nkx2.2 115 116 expression marks the more ventral V3 interneuron progenitors (p3), and Dbx1 is expressed in the p0 domain located at the equator of the neural tube leading to interneurons situated dorsal to 117 MNs (Briscoe, Pierani et al. 2000). Subsequently, persistent Shh signaling originating from the 118 119 medial floorplate (MFP) located at the ventral midline of the developing neural tube is critical 120 for maintaining the identities of each of these domains throughout neurogenesis (Dessaud, Ribes et al. 2010). The ventral neural tube expands rapidly during neurogenesis and Shh signaling 121 strength measured by expression levels of Shh target genes declines progressively within the 122

123 precursor domains (Balaskas, Ribeiro et al. 2012, Kicheva, Bollenbach et al. 2014). Surprisingly, the beginning of oligodendrogenesis is marked by increased Shh signaling strength within the 124 pMN domain (Danesin, Agius et al. 2006). How this dynamic increase in Shh signaling is 125 126 achieved subsequently to a decline in Shh signaling strength and whether it is of functional 127 importance for OPC specification and production is an area of active research (Traiffort, Zakaria et al. 2016). Since the Shh gradient emanating from the MFP displays a constant decay length 128 and does not scale with congruent and rapid growth during early development (Cohen, Kicheva 129 et al. 2015), its influence on pMN domain activity must likely wane during neurogenesis. 130 131 Accordingly, multiple adaptation mechanisms have been proposed to underlie temporally dynamic Shh signaling at the time of initiation of OPC production (Traiffort, Zakaria et al. 132 2016). One such mechanism is the accumulative storage of Shh in the extracellular matrix 133 134 followed by Sulfatase 1 dependent release leading to a greater concentration of Shh than could be achieved by continuous production and diffusion (Danesin, Agius et al. 2006, Touahri, 135 Escalas et al. 2012, Al Oustah, Danesin et al. 2014). While the genetic ablation of Sulfatase 1 136 curtails the production of OPCs, the current experiments cannot exclude the possibility that other 137 signaling factors than Shh are released from the extracellular matrix by Sulfatase 1 and then 138 139 participate in the regulation of OPC production. Another mechanism might be the sequential production of Shh by previously specified ventricular zone derivatives (VZD). Prominent 140 sources of VZD_{Shh} are the lateral floorplate (LFP_{Shh}) (Charrier, Lapointe et al. 2002, Park, Shin et 141 142 al. 2004, Al Oustah, Danesin et al. 2014), which is constituted by cells emigrating from the p3 domain, and MNs (MN_{shh}) (Oppenheim, Homma et al. 1999, Akazawa, Tsuzuki et al. 2004). The 143 144 selective ablation of Shh from LFP or MNs without impacting earlier patterning of the spinal

- cord has not been achieved and causal evidence for the involvement of LFP_{Shh} and MN_{Shh} in the
 regulation of OPC generation is therefore lacking.
- 147 Here we produced a series of mouse lines with different degrees and tissue selectivity of
- 148 conditional Shh gene ablation from the MFP, LFP, and MNs during spinal cord development.
- 149 We find that VZD_{Shh} is critical for the maintenance of the Olig2 expressing cell population in the
- pMN domain during the phase of oligodendrocyte precursor cell (OPC) production. While early
- neural tube patterning and MN development can proceed successfully in the absence of these
- 152 Shh sources, the pMN domain exhausts of Olig2 expressing cells during neurogenesis leading to
- a subsequent and spinal level specific reduction in OPC production. LFP_{Shh} is needed for OPC
- 154 production throughout the spinal neuraxis. In contrast, MN_{Shh}, which occurs earlier at limb levels
- than thoracic levels, is critical for maintaining a larger pMN domain at limb levels compared to
- thoracic segments throughout OPC production. Our data provides causal evidence for the critical
- 157 involvement of sequential and spinal level restricted expression of Shh by VZD for scaling of
- 158 OPC production along the neuraxis.
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170 *Results:*

171 Shh expression in the ventral spinal cord

To examine the contributions of different sources of Shh in the ventral spinal cord to 172 pMN domain activity at the time of the transition from MN to OPC production we first expanded 173 on previous descriptions of Shh expression during spinal cord development. We used a gene 174 expression tracer and conditional loss of function allele of Shh (Shh-nLZ^{C/C}, abbreviated Shh^{C/C}) 175 in which a bicistronic mRNA is transcribed from the un-recombined Shh locus that encodes Shh 176 and nuclear targeted LacZ (Gonzalez-Reyes, Verbitsky et al. 2012). The allele allows the 177 quantification of the numbers of cells that express Shh and the determination of Shh ablation 178 179 efficiencies in response to Cre activity with single-cell resolution. This approach reveals developmental stage- and spinal level- specific patterns of Shh expression and relative 180 contributions to ventral Shh production from at least 4 distinct cell populations. At brachial 181 levels at E12.5 we find Shh in all cells of the MFP (defined by co-expression with FoxA2 and 182 situated at the ventral midline stacking 4-5 cells high along the ventral to dorsal axis), LFP 183 184 (defined by reduced levels of FoxA2 and Shh compared to MFP, situated immediately dorsal to the MFP and stacking about 3-4 cells high along the ventral to dorsal axis), LFP* (defined by co-185 expression of Nkx2.2 and Shh and situated in part at the lateral edges of the LFP and in part as 186 187 isolated cells flanking the p3 domain), and in 30% of all motor neurons (MNs) of the lateral motor column (LMC, defined by co-expression with Hb9 and lateral position in the ventral 188 horns) (Fig. 1). In contrast, at thoracic and lumbar levels at E12.5, we find Shh expression only 189 190 in MFP, LFP*, and LFP (Fig. 1). Based on numbers of nLacZ+ cells present at E12.5 we

estimate that about 57% of Shh production occurs by MFP, 30% by the collective LFP and 13%
by MNs at brachial levels while about 50% of Shh production occurs each by MFP and LFP at
thoracic and lumbar levels (Fig. 1B). The onset of Shh expression in these tissues is sequential
and overlaps with both MN and OPC production (Fig. 1C), suggesting the possibility to dissect
each source's Shh contribution to pMN activity.

At E13.5 lumbar LMC MNs begin to express Shh resulting in a pattern of expression that is qualitatively similar to brachial levels at E12.5 (**Fig. S2A**). By E14.5 medial motor column (MMC) MNs at all spinal segments begin to express Shh (**Fig. S1A**). At P20 Shh expression occurs in MNs, V0 cholinergic neurons, and remaining FP cells at all spinal cord segments (**Fig. S1B and S2C**). We observe a similar temporal and segmental pattern of Shh expression in the developing chick spinal cord (**Fig. S1C**).

To investigate the role of Shh signaling from these Shh sources onto the pMN domain, we generated a series of mouse lines with conditional and in part overlapping ablation of Shh. We used ChAT-Cre (ChAT_{Shh}-^{/-}) to target MNs, Nestin-Cre (Nestin_{Shh}-^{/-}) to target all Shh expressing VZD (MNs, LFP, and LFP*), and Olig2-Cre (Olig2_{Shh}-^{/-}), to target MNs, LFP, and MFP.

At E12.5 we did not observe Shh recombination in MFP cells in $ChAT_{Shh}^{-/-}$ mutants. Nestin_{Shh}^{-/-} mutants had a few MFP cells which displayed Cre activity, but this ablation was insignificant (**Fig. 1D and 1G**). However, in Olig2_{Shh}^{-/-} mutants the Shh recombination efficiency among MFP cells in brachial and thoracic spinal segments revealed a significant ~44% and ~39% loss resp. (**Fig. 1D and 1G**). We found no recombination of Shh in the LFP or LFP* in ChAT_{Shh}^{-/-} at brachial or thoracic segments (**Fig. 1D, E, H**). Consistent with the previous reported expression of Nestin-Cre in all VZD between E10.5 to E12.5 (Kramer et al.,

214	2006), and transient expression of Olig2-Cre in all VZD of the pMN domain and ventral to it
215	(Ribes and Briscoe, 2009), we find near complete (~80%, and ~90% resp.) ablation of Shh from
216	the LFP and LFP* in both $Nestin_{Shh}^{-/-}$ and $Olig2_{Shh}^{-/-}$ in brachial segments (Fig. 1D, E, H). At
217	thoracic segments LFP recombination was less efficient at ~60% and ~72% for $Nestin_{Shh}$ ^{-/-} and
218	$Olig2_{Shh}^{-/-}$ resp. LFP* recombination was ~77% and ~94% for $Nestin_{Shh}^{-/-}$ and $Olig2_{Shh}^{-/-}$ resp.
219	(Fig. 1D, E, H). Cre efficiency in MNs was about 80% in $ChAT_{Shh}^{-/-}$ and $Nestin_{Shh}^{-/-}$ mutants,
220	and 98% for $Olig2_{Shh}^{-/-}$ (Fig. 1F and 1G). The location of Shh expression and the varied degrees
221	of ablation of Shh at E12.5 are schematically summarized in Fig. 1J.
222	Using a conditional reporter allele (Fig. 2A), we find that Olig2-Cre but not Nestin-Cre is
223	active in the MFP prior to notochord (NC) regression at E10.5 resulting in the ablation of Shh
224	from $46 \pm 5.1\%$ and $8.5 \pm 2.8\%$ resp. of FoxA2+ cells (Fig. 2B and 2C). These results reveal that
225	the degree of ablation of Shh from MFP in $Nestin_{Shh}^{-/-}$ and $Olig2_{Shh}^{-/-}$ animals is established at
226	the time of MN production and remains fixed. The drastic loss of Shh expression in the MFP
227	already at E10.5 in Olig2 _{Shh} -/- animals prompted us to ascertain a possible patterning defect along
228	the ventral midline. Consistent with previous reports that Shh expression by the notochord (NC)
229	is sufficient for the establishment of precursor domains in the ventral spinal cord (Dessaud,
230	Ribes et al. 2010, Yu, McGlynn et al. 2013) we find that the relative location and size of the p3-
231	(Nkx2.2), pMN- (Olig2) and p0- (Dbx1) domains, and the location of the ventral border of the
232	Pax6 expression domain are indistinguishable between $Olig2_{Shh}^{-/-}$ and control at E10.5 (Fig. 2D
233	and 2E).
234	Together, the varied degrees of quantifiable and source selective ablation of Shh from the

ventral spinal cord and the preservation of early ventral tube patterning indicated that this set of

recombinant mouse lines might be informative in the investigation of the effect of VZD sourcesof Shh onto the pMN domain during MN and OPC generation.

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239 Shh signaling from MFP, but not VZD sources influences MN generation

We investigated whether ablating Shh from VZD sources would impact the generation of
MNs from Olig2 precursors of the pMN domain. We analyzed columnar pattern, relative

242 distribution of MNs among columns, and absolute numbers of MNs of different columnar

243 identity at brachial and thoracic levels. We first visualized MN columnar organization by

immunostainings for MMC (Hb9+ Lhx3+), LMC_M (Hb9+ Isl1/2+), and LMC_L (Hb9+, Isl1/2-,

245 Lhx3-) at brachial (Fig. 3A) and MMC (Hb9+, Is11-), HMC (Hb9+, Is11/2+), and PGC (Hb9-,

246 nNos+) at thoracic (Fig. 3E) and found no apparent differences in the staining pattern either

among ChAT_{shh}^{-/-}, Nestin_{Shh}^{-/-}, Olig2_{Shh}^{+/-}, or Olig2_{Shh}^{-/-} compared to Shh^{C/C} controls. Further

supporting unaffected MN positioning and patterning, we find inconspicuous ventral root

formation in $Olig2_{Shh}^{-/-}$ compared to $Shh^{C/C}$ controls at E10.5 and E12.5. (Fig. S3).

Quantification of MN numbers revealed no differences in ChAT_{Shh}^{-/-} and Nestin_{Shh}^{-/-} 250 compared to Shh^{C/C} controls at brachial and thoracic levels. In contrast, in Olig $2_{\text{Shh}^{+/-}}$ and 251 Olig2_{shh}^{-/-} we find a 22% and 38%, resp. reduction in the numbers of total MNs at brachial (Fig. 252 **3B**), and 22% and 32% at thoracic levels compared to Shh^{C/C} controls (Fig. 3F). Since the 253 production of late born MNs could be affected to a greater extent than early born MNs by the 254 255 ablation of Shh from previously born VZD, we compared the relative size of the MN columns and the numbers of late born MNs just emerging from the ventricular zone at E12.5. We found 256 that MNs attained columnar identities in normal relative proportions in all genotypes (Fig. 3D 257 258 and 3H). However, quantification of migrating late born Hb9+ MNs showed a dose-dependent

259	reduction of ~10% in $Olig2_{Shh}^{+/-}$ and 47% in $Olig2_{Shh}^{-/-}$ in brachial, and 23% and 47% resp. in
260	thoracic segments, suggesting that the deficit in MN generation that we observe in $\text{Olig2}_{\text{Shh}^{+/-}}$ and
261	$Olig2_{Shh}^{-/-}$ mice is greatest towards the end of MN production (Fig. 3I). We did not observe a
262	decrease in numbers of late born migrating MNs in ChAT _{Shh} ^{-/-} or Nestin _{Shh} ^{-/-} . We next examined
263	if the earlier deficits in MN generation in Olig2 _{Shh} ^{-/-} mice could be overcome by reduced
264	apoptosis during the phase of programmed cell death. To this end, we found reduced levels of
265	Caspase3+ Hb9+ brachial MNs in Olig2 _{Shh} -/- at E12.5 indicating that reduced cell death of MNs
266	at least in part will compensate for a reduced rate of MN production in Olig2 _{Shh} -/- animals (Fig.
267	3J).

We associated the degree of reduction in the numbers of MNs with the tissue specific 268 efficiency of Shh ablation in ChAT_{Shh}^{-/-}, Nestin_{Shh}^{-/-}, and Olig2_{Shh}^{-/-} (Fig. 3K). The 80% efficient 269 270 ablation of Shh from MNs at brachial levels as well as the near complete ablation of Shh from the LFP in Nestin_{Shb}^{-/-}, has no effect on MN numbers at E12.5. In contrast, the near complete 271 ablation of Shh from MNs, LFP and about 50% of MFP in Olig2_{Shh}^{-/-} results in a Shh dose-272 dependent reduction of MN numbers at all levels and in all MN columns (Fig. 3K, orange line). 273 Together these results reveal that MN differentiation can proceed in the absence of Shh signaling 274 from the ventricular zone derivatives MN and LFP, but the efficacy of MN generation becomes 275 progressively impacted by dose-dependent reductions in Shh signaling from the MFP. 276

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Shh from VZD sources in addition to MFP is critical for pMN domain maintenance during the onset of gliogenesis in a spinal level specific manner.

280 Previous studies did not evaluate the contribution from individual Shh sources for the

281 maintenance of established precursor domains after NC retraction. We therefore next

investigated ventricular zone organization and precursor domain maintenance in $ChAT_{Shh}^{-/-}$, Nestin_{Shh}^{-/-}, and Olig2_{Shh}^{-/-}. We find that the relative location and distance to each other of the p3, pMN, and p0 precursor domains are preserved in $ChAT_{Shh}^{-/-}$, Nestin_{Shh}^{-/-}, and Olig2_{Shh}^{-/-} at E12.5 (**Fig. 4A**). However, we observe a Shh source and dose-dependent, spinal level-specific decline in the numbers of Olig2+ cells in the pMN domain of $ChAT_{Shh}^{-/-}$, Olig2_{Shh}^{+/-}, Nestin_{Shh}^{-/-}, and

287 $\text{Olig2}_{\text{Shh}^{-/-}}$ compared to $\text{Shh}^{\text{C/C}}$ or Olig2-Cre control (Fig. 4B and 4C).

In ChAT_{Shh}^{-/-} we find a 26% decrease in pMN/Olig2+ cells (pMN_{Olig2}^+) at brachial levels 288 (but not at thoracic or lumbar) consistent with expression of Shh in MNs at brachial but not yet 289 in MNs at thoracic and lumbar levels which will begin to express Shh about one day later (Fig. 290 **4**C). Upon more detailed analysis, we find the effect of MN Shh to become progressively more 291 pronounced during the rapid enlargement of the pMN that occurs at brachial segments between 292 E12.5 to E12.75 (Fig. 4D and 4E). The numbers of pMN_{Olig2}^+ cells in Nestin_{Shh}^{-/-} displays an 293 anterior-posterior progressive decrease of 40% at brachial, 51% at thoracic, and 59% at lumbar 294 segments (Fig. 4B and 4C). The most severely affected genotype is Olig2_{Shh}^{-/-} with a decrease in 295 pMN_{Olig2}^+ cells of 66% at brachial, 61% at thoracic, and 72% at lumbar (Fig. 4B and 4C). 296 We next associated the degree of reduction in the numbers of pMN_{Olig2}^+ cells at brachial 297

and thoracic levels with the time of onset and tissue specific efficiency of Shh ablation in ChAT_{Shh}^{-/-}, Olig2_{Shh}^{-/-}, and Nestin_{Shh}^{-/-} (**Fig. 4F**). In controls, we find about 19% more pMN_{Olig2}⁺ cells at brachial than thoracic segments. The 80% efficient ablation of Shh from MNs in ChAT_{Shh}^{-/-} reduces the numbers of pMN_{Olig2}⁺ cells at brachial levels to those present in controls at thoracic levels while the ablation of Shh from MNs has no effect on the numbers of pMN_{Olig2}⁺ cells at thoracic levels (**Fig. 4F, red line**). Nestin_{Shh}^{-/-} and Olig2_{Shh}^{-/-} exhibit near complete ablation of Shh from MNs and LFP, but significantly different ablation efficiencies in the MFP,

resulting in a reduction of pMN_{Olig2}^+ cells at brachial levels that scales with the degree of Shh ablation from the MFP. At thoracic levels, however, we find the same magnitude in the reduction of the numbers of pMN_{Olig2}^+ cells in $Nestin_{Shh}^{-/-}$ and $Olig2_{Shh}^{-/-}$ despite of a much greater total reduction in numbers of Shh producing cells by Olig2-Cre compared to Nestin-Cre. This observation suggests that Shh produced by LFP cells, though few in number, has a disproportionate significance compared to MFP derived Shh for pMN_{Olig2}^+ cells at thoracic segments.

Next, we determined if the decreased numbers of pMN_{Olig2}^+ cells is the result of 312 313 diminished initial specification or a failure of maintenance. We lineage traced Olig2 cells in Olig2_{Shh}^{-/-} mutants, Olig2_{Shh}^{+/-} controls, and Olig2-Cre controls using the R26mT/mG reporter 314 allele from which myristylated GFP is expressed in all derivatives of Olig2 expressing cells and 315 316 immunostained for Olig2 at E12.5 (Fig. 5A). We observe GFP expression in the ventral spinal cord forming a dorsal boundary at a similar relative distance to the MFP in mutants and controls 317 but a decline in pMN_{Olig2}⁺ GFP+ double positive cells in mutants compared to controls. Notably, 318 the absence of Olig2 expressing cells within the GFP labeled area in mutants is most pronounced 319 in the dorsal half of the pMN domain. Together, these results demonstrate that ongoing Shh 320 signaling originating from VZD in addition to MFP are critical for the selective maintenance of 321 pMN_{Olig2}⁺ cells once the influence of notochord Shh has waned and the pMN switches to 322 gliogenesis of OPCs. 323

Providing further evidence for decreased pMN domain activity at the beginning of OPC production at E12.5 we find a \sim 3-fold reduction in the size of p* precursor domain in Olig2_{Shh}-/compared to controls (**Fig. 5B and 5C**). The p* domain forms at the ventral border of the pMN domain and is marked by Olig2+/Nkx2.2+ cells (Agius et al., 2004). Interestingly, we found

328	LFP* cells (which we define as Nkx2.2+ nLacZ+ cells) in direct contact with Nkx2.2+ Olig2+
329	double positive cells of the p* domain in Shh ^{C/C} controls, highlighting a cyto-architectural
330	arrangement that could underpin the disproportionate importance of LFP compared to MFP_{Shh}
331	for the maintenance of the pMN_{Olig2}^+ cell population (Fig. 5B).
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333	Diminishment of pMN_{Olig2}^+ cells during the phase of ventral oligodendrogenesis results in
334	reduced OPC production.
335	Reduced Shh signaling originating from MNs, LFP, or MFP leaves the pMN domain
336	impoverished of pMN_{Olig2}^+ cells at the beginning of oligodendrogenesis (Fig. 4). Nevertheless,
337	the pMN_{Olig2}^+ cell population could recover during OPC production by increased precursor
338	recruitment from dorsal ventricular precursor domains (Ravanelli and Appel 2015), proliferation
339	of remaining pMN_{Olig2}^+ precursor cells, or increased differentiation and amplification of OPC
340	fated cells that have left the pMN domain. We therefore first visualized the size of pMN_{Olig2}^+
341	population at the end of ventral oligodendrogenesis at E14.5. We find a moderate reduction in
342	the numbers of pMN_{Olig2}^+ cells in $ChAT_{Shh}^{-/-}$ and an almost complete absence of pMN_{Olig2}^+ cells
343	in Nestin _{Shh} -/- and Olig2 _{Shh} -/- compared to Shh ^{C/C} controls suggesting that increased recruitment of
344	precursors to the pMN domain does not occur in mutants (Fig. S4 and Fig. 6A). We then
345	determined whether pMN_{Olig2}^+ cells and/or migrating OPCs in $Olig2_{Shh}^{-/-}$ increase their rate of
346	proliferation during the phase of OPC production. We injected EdU into pregnant dams at E11.5,
347	E12.5, and E13.5 and quantified the numbers of pMN_{Olig2}^+ cells 24h later. We found comparable
348	broad EdU+ labeling throughout the ventricular zone of Olig2 _{Shh} ^{-/-} mutants and controls,
349	suggesting overall progenitor proliferation is not affected in mutants (Fig. 6A). Within the pMN
350	domain we observe a ~25% decrease in the numbers of pMN_{Olig2}^+ cells in Shh ^{C/C} controls over

351	the course of OPC production from E12.5 to E14.5 (Fig. 6B). In contrast, numbers of pMN_{Olig2}^+
352	cells in Olig2 _{Shh} -/- mutants decline to near undetectable levels during the same period indicating a
353	precocious exhaustion of the pMN_{Olig2}^+ cell population during OPC generation (Fig. 6B). We
354	next determined the rate of proliferation of pMN_{Olig2}^+ cells in controls and mutants. In controls,
355	we find a similar proliferative rate of about 47% of pMN_{Olig2}^+ cells at E12.5, E13.5 and E14.5. In
356	contrast, in $Olig2_{Shh}^{-/-}$ we find the rate of proliferation to be reduced to 37% at E12.5 and E13.5,
357	followed by a further decrease to 16% by E14.5 (Fig. 6C). These results indicate that the
358	precocious exhaustion of the pMN_{Olig2}^+ cell population is associated with a reduced proliferation
359	rate of precursor cells during OPC production which is compounded by reduced numbers of
360	pMN_{Olig2}^{+} cells that are present at the beginning of OPC production.
361	We next tested whether OPC production in Olig2 _{Shh} ^{-/-} recovers through increased
362	amplification of precursors once they have emerged from the pMN. We analyzed numbers of
363	EdU+ Olig2+ cells in the mantle zone (Fig. 6D). While there are very few Olig2+ cells in the
364	mantle zone of mutants and controls at E12.5, with ongoing expansion of these cells, we find a 3-
365	fold and 15-fold reduction of Olig2+ cells at E13.5 and E14.5, resp.in mutants compared to
366	controls (Fig. 6E). The rate of proliferation among Olig2+ cells in the mantle zone in $Olig2_{Shh}^{-/-}$
367	is similar at E13.5 and E14.5 compared to controls suggesting that OPCs in mutants do not
368	amplify at an increased rate compared to controls (Fig. 6F). The cells that do emerge from the
369	pMN in Olig2 _{Shh} -/- disperse as rapidly as their control counterparts, resulting in a ventral spinal
370	cord that is populated with nascent OPCs with a 15-fold lower density compared to controls (Fig.
371	S5). Additionally, we examined OPC proliferation at E14.5 in posterior thoracic and lumbar
372	segments and found no detectable increase in proliferation rate in any of the pMN, mantle, or
373	white matter (WM) areas (Fig. S6). These results reveal that the yield of the pMN domain during

374	OPC production largely determines the numbers of OPCs that settle in white and grey matter.
375	Thus the scaled production of OPCs and oligodendrocyte along the anterior posterior extent of
376	the spinal cord must at least in part be determined by the patterned expression of VZD_{Shh} that we
377	find to be critical for the maintenance of the pMN_{Olig2+} precursor population.
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392 Discussion:

Our study provides causal evidence that the sequential and patterned expression of Shh by 393 previously specified cell types in the ventral spinal cord is critical for scaled oligodendrocyte 394 395 precursor specification along the anterior posterior axis of the developing spinal cord. We took advantage of neural tube development as an established model system to reveal that Shh-396 397 expressing signaling centers become established among post mitotic derivatives of ventricular 398 zone differentiation (VZD_{Shh}) in a temporally- and spatially-patterned manner as development proceeds. We find that VZD_{Shh} signaling is critical for the production of OPCs in proportionate 399 numbers to previously specified neurons. Our data provides genetic evidence in support of the 400 hypothesis that extra-midline sources of Shh are critical for switching neurogenesis to 401 402 gliogenesis and add a novel mechanism that contributes to pattern scaling along the anterior to 403 posterior axis. Our observations do not rule out contributions of other well-established cellautonomous and cell non-autonomous mechanisms that adapt morphogen function and allow 404 pattern scaling (Houchmandzadeh, Wieschaus et al. 2005, Ben-Zvi and Barkai 2010, 405 406 Hamaratoglu, de Lachapelle et al. 2011, Ben-Zvi, Fainsod et al. 2014, Uygur, Young et al. 2016), but provide a mechanism by which Shh mediated signaling from the midline in the ventral neural 407 408 tube becomes progressively augmented such that Shh signaling remains a relevant instructive 409 signal despite rapid growth.

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Sequential expression and function of VZD_{Shh}.

411 Despite its complexity, development is orchestrated by the actions of just a handful of
412 morphogens. The temporal and spatial segregation of developmental fields that are patterned by

413 morphogens can allow the same morphogen to play an instructive role repeatedly and at multiple anatomic regions specifying vastly different cell types and tissues. However, in the developing 414 spinal cord all cell types are descendants of the same developmental field, the ventricular zone, 415 yet the same morphogen, Shh, is involved in the sequential specification of multiple cell types 416 417 (Dessaud, McMahon et al. 2008). The number of source tissues of Shh in the ventral spinal cord 418 expands concomitantly with development, suggesting that "moving the Shh source" might play an important role (Danesin and Soula, 2017). We reassessed Shh expression in mice using a 419 nuclear targeted LacZ based gene expression tracer allele and verified the principle findings in 420 421 chick by RNA in situ. The advantage of the gene expression tracer allele is that the identity, location and numbers of Shh expression can be determined with single-cell resolution. Ablating 422 Shh and nLacZ by Nestin-Cre, which is expressed in the ventricular zone, reveals that one half to 423 424 one third of all Shh expressing cells in the ventral spinal cord are ventricular zone descendants at the time that Shh signaling strength within the pMN domain rises coincident with the switch 425 from neurogenesis to gliogenesis (Dessaud, Yang et al. 2007, Balaskas, Ribeiro et al. 2012, 426 427 Touahri, Escalas et al. 2012, Kicheva, Bollenbach et al. 2014, Kicheva and Briscoe 2015). While the gene expression tracer allele cannot be used to draw conclusions about the relative Shh 428 429 signaling strength in the ventricular zone that is contributed by VZD_{Shh}, the anatomic arrangement of these sources and the genetic ablation of Shh from these sources suggests a 430 disproportionate effect on the activity of the pMN domain. The LFP proper extends the midline 431 432 source of Shh by several cell diameters more dorsally as previously observed (Charrier et al., 2002). A subset of LFP cells, which we designate here as LFP* cells, co-express Nkx2.2 and Shh 433 (Al Oustah, Danesin et al. 2014) and are in direct contact with cells that co-express the Shh high-434 435 threshold genes Nkx2.2+ Olig2+ and form the p* domain at the beginning of oligodendrogenesis

436 (Fu, Qi et al. 2002, Traiffort, Zakaria et al. 2016). At the same developmental stage, we detect expression of Shh in the nascent lateral, but not medial, MN column in mice and chick. Shh can 437 be released from the axonal as well as the dendritic compartment of neurons ((Beug, Parks et al. 438 439 2011). Hence nascent MNs could secrete Shh via dendrites or possibly trailing cellular processes close to the pMN domain. Together, the spatial and temporal pattern of VZD_{shh} expression that 440 we observe is consistent with the results from our selective gene ablation studies which 441 demonstrate that these sources of Shh influence pMN domain activity, a speculation put forward 442 previously in regard of the function of LFP_{Shh} (Danesin and Soula 2017). 443 444 A striking feature of the phenotype of Shh ablation from VZD is the temporal and qualitative segregation of the effects on MN and OPC production. VZD_{shh} has no detectable effect on MN 445 production or the extent of programmed cell death of MNs in our paradigms. In contrast, ablation 446 of VZD_{Shh} results in severely reduced OPC production: (1) During the period of OPC production 447 the initial 2-fold diminishment of pMN_{Olig2}⁺ cells increases to a more than 10-fold reduction in 448 pMNOlig2+ cells compared to controls. (2) During the same period, the numbers of Olig2+ cells 449 that emerge from the pMN domain drop 10-fold. (3) The proliferation rate of these nascent OPCs 450 451 is equal to or lower compared to their control counterparts. Together these observations reveal 452 that the numbers of OPCs that emigrate from the pMN domain and settle in the white and grey matter are determined mainly by the size of the pMN_{Olig2+} precursor cell population. 453 The relative contribution of VZD_{Shh} to regulating the size of the pMN_{Olig2+} precursor cell 454 population is spinal level specific. For example, the ablation of VZD_{Shh} results in a pMN_{Olig2}^+ 455 loss of about 40% at brachial levels. Half of that effect at brachial levels can be attributed to the 456 457 expression of MN_{shh} since the ablation of Shh from cholinergic neurons by ChAT-Cre alone results in an almost 26% reduction in the size of the pMN_{Olig2}^+ population (Fig. 3.2B). The 458

remaining size of the pMN_{Olig2}^+ population at brachial levels in the ChAT_{Shh}^{-/-} spinal cord is 459 similar to the average size of the pMN_{Olig2}⁺ domain at thoracic levels in control spinal cords. 460 Since thoracic MNs do not express Shh until the end of OPC production at E14.5, our 461 observations reveal that the increased size of the pMN domain at brachial levels is dependent on 462 MN_{shh}. Thus, the selective expression of MN_{shh} at brachial and lumbar levels provides a 463 mechanism for ensuring a proportionate increase in the production of OPCs at brachial and 464 lumbar - compared to thoracic- levels that is matched to the increased numbers of MNs at limb 465 levels. 466

467

Potential mechanisms of actions of VZD_{Shh}.

Does Shh from different sources have distinct functions in the ventricular zone of the 468 ventral spinal cord? Three observations support origin specific functions of Shh in the ventral 469 470 spinal cord in the pMN domain: (1) MFP_{shh} but not VZD_{shh} influences MN production (Fig. 3). Consistent, ablation of Shh in the MFP by Olig2-Cre is complete at the beginning of MN 471 production while ablation of Shh in VZD by Nestin-Cre only begins towards the end of MN 472 generation (Fig. 1). In further support of a critical role of MFP_{shh} in MN generation and 473 consistent with previous findings (Yu, McGlynn et al. 2013), we find a Shh gene dose-dependent 474 reduction of the numbers of MNs of similar magnitude among early and late forming MN 475 columns as well as late born MNs still in transit at E12.5. Hence, based on the timing of Shh 476 expression and the consistent deficit in MN production throughout the period of MN generation, 477 478 the reduced rate of MN production must be associated with the reduction of midline-derived Shh rather than VZD-derived Shh. Strikingly, however, whether Shh expression is ablated from all 479 VZDs (Nestin_{Shh}^{-/-}) or partially from the MFP in addition to LFP and MNs (Olig2_{Shh}^{-/-}), the pMN 480 domain is left with similar strongly reduced numbers of pMN_{Olig2}⁺ cells at the end of MN 481

482 production (Fig. 2). Since ventral precursor domains form normally and a full complement of pMN_{Olig2}⁺ cells is induced in Olig2_{Shh}^{-/-} (Fig. 1), these results point to distinct functions of Shh 483 derived from the MFP and VZDs in pMN domain activity: Our data indicates that MFP_{Shh} 484 determines the rate of MN production while VZD_{Shh} is critical to counteract the exhaustion of the 485 pMN_{Olig2}^{+} population during MN production. (2) We find that the high threshold, Shh dependent 486 and p* domain defining co-expression of Nkx2.2 and Olig2 (Fu, Qi et al. 2002) occurs in cells 487 that are in close proximity to LFP* cells. Ablation of VZD_{Shh} results in a 3-fold reduction in the 488 numbers of Nkx2.2/Olig2 expressing cells at E12.5 (Fig. 4). Additional ablation of Shh from 489 50% of the MFP in Olig2_{shh}^{-/-} did not increase the severity of this phenotype (Fig. 4). Thus, 490 consistent with the anatomic juxtaposition of LFP cells to p* domain cells our gene ablation 491 studies demonstrate that VZD_{Shh} is critical for the production of the Nkx2.2 expressing 492 subpopulation of ventral oligodendrocyte lineage cells. (3) The dorsal most aspects of the pMN 493 domain is almost completely devoid of Olig2+ expressing cells at the end of MN production at 494 E12.5 in Nestin_{Shh}^{-/-} and Olig2_{Shh}^{-/-} (Fig. 2). Given that early pioneer OPCs elaborate cellular 495 contacts selectively with MNs (Osterstock, Le Bras et al. 2018) it seems plausible that the LFP 496 and MFP most distant pMN areas are served selectively by Shh from MN_{Shh}. In this scenario Shh 497 signaling from different sources might subdivide the pMN domain into sub-regions from which 498 distinct oligodendrocyte subtypes might emerge (Dimou and Simons 2017, Ravanelli, Kearns et 499 al. 2018) Dependent on the mode of delivery, VZD_{Shh} could also exhibit distinct modes of 500 501 action: for example, LFP_{Shh} could result in maintenance of Olig2 expression in pMN precursor cells while MN_{Shh} could attract precursor cells to migrate into the pMN domain from the dorsal 502 503 ventricular zone as is observed in zebrafish (Ravanelli and Appel, 2015). 504 Together, our data provides genetic evidence in support of the idea that morphogen signaling centers

are established sequentially in the developing ventral spinal cord. These new "organizer tissues" express

506	Shh which acts together with Shh produced by the medial floor plate to influence SVZ activity as the
507	spinal cord grows. These Shh sources are critically involved in the switch of neurogenesis to
508	oligodendrogenesis in the pMN domain. Endowing VZDs with morphogenic activity makes subsequent
509	differentiation conditional to the completion of previous developmental milestones in the ventral spinal
510	cord. Further, expression of the morphogen linked to the numbers and types of VZDs produced,
511	provides a mechanism that could ensure scale invariance in regard of neuron and glia production along
512	the anterior-posterior axis of the spinal cord.
513	
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518	University of New York.
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- Materials and Methods Transgenic Mice All animal experiments were approved by the Institutional Animal Use Care Committee at CUNY. The following mouse strains were used and genotyped as described previously: Shh-nLZ^{L/+} animals (Gonzalez-Reves et al., 2012), Chat-Cre (Rossi et al., 2011), Olig2-Cre (Dessaud 2007), Nestin-Cre (Tronche et al., 1999), Rosa26^{mT/mG} (Muzumdar et al., 2007). Mice were maintained on a C57BL/6 background. Noon on the day of the plug was considered E0.5. Mice were kept on a 12 hr dark/light cycle and the day of birth designated P1. For E12.25, E12.50, and E12.75 pMN analysis, pregnant dams were sacrificed at E12.5 according to plug date and embryos were binned into three groups (E12.25, E12.50, and E12.75), based on how many Olig2+ cells have migrated out of the pMN domain. Embryo sections which had an average of less than 5 cells migrate out of the pMN were considered 12.25, 5-20 cells migrating out – E12.50, and >20 cells migrating out - E12.75. In vivo EdU Assay

553	Pregnant dams received EdU (5-ethynyl-20 -deoxyuridine, Invitrogen) dissolved in PBS by		
554	intraperitoneal injection (50 mg/kg) and sacrificed after 24 hours. Tissue sections were stained		
555	using the Click-iT Plus EdU Alexa Fluor 647 Imaging Kit (Thermofisher).		
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558	Tissue Processing		
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560	All mice were sacrificed using an overdose of anesthetic, subjected to transcardial perfusion with		
561	4% (w/v) paraformaldehyde (PFA) in 0.1 M PBS pH 7.4. Spinal cords and embryos were		
562	dissected, postfixed in 4% PFA for 1 hr at 4°C, cryoprotected with 30% (w/v) sucrose in 0.1M		
563	PBS for 24–48 hr, embedded and frozen in OCT medium, and stored at -80°C. Tissues were		
564	sectioned at 20 µm and collected onto glass slides.		
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566	Immunocytochemistry and Microscopy		
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568	20um thick spinal cord cryosections were air dried for 30 min. Then sections were washed with		
569	PBS for 10 mins and with 0.3% [v/v] Triton X-100 in PBS for 20 min. Sections were then pre-		
570	treated with blocking solution (10% $[v/v]$ horse serum and 0.3% $[v/v]$ Triton X-100 in PBS) for		
571	90 mins and incubated with primary antibodies overnight at 4C. The next day, following 3 PBS		
572	washes the sections were incubated with secondary antibodies for 2 hr at room temperature. A		
573	list of all antibodies and compounds used is provided in a table. For cell counts, at least three		
574	sections per animal from at least three mice were examined, unless otherwise noted. Images were		
575	acquired using a Zeiss LSM880 confocal microscope.		

577 Statistical analysis was performed using Prism 7 (Graphpad Software Inc.) Analysis of multiple

578 groups was made using one-way ANOVA followed by the Tukey or Dunnett's post hoc analysis

tests. For 2-groups analyses, unpaired Student's t test was used. The data are presented

580 graphically as: (p < 0.05), (p < 0.01), and (p < 0.001).

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REAGENT	SOURCE	IDENTIFIER		
Antibodies				
Chicken polyclonal anti beta Galactosidase	Abcam	Cat# ab9361, RRID:AB_307210		
Goat polyclonal anti beta Galactosidase	Biogenesis	Cat# 4600-1409, RRID:AB_2314510		
Goat polyclonal anti Choline acetyltransferase	Millipore	Cat# AB144P, RRID:AB_2079751		
Rabbit polyclonal anti Olig2	Millipore	Cat#AB9610, RRID:AB_570666		
Mouse monoclonal anti Nkx2.2	DSHB	Cat# 74.5A5, RRID:AB_531794		
Rabbit polyclonal anti nNos	Immunostar	Cat# 24431, RRID:AB_572255		
Mouse monoclonal Ankyrin G (AnkG) (463)	Santa Cruz	Cat# sc-12719, RRID:AB_626674		
Mouse monoclonal Lim3 (Lhx3)	DSHB	Cat# 67.4E12, RRID:AB_2135805		
Rabbit polyclonal Lim1	Gift Jessell Lab	N/A		
Guinea Pig polyclonal Hb9	Gift Jessell Lab	N/A		
Rabbit polyclonal FoxA2	Gift Jessell Lab	N/A		
Rabbit polyclonal Pax6	Gift Jessell Lab	N/A		
Rabbit polyclonal Dbx1	Gift Jessell Lab	N/A		

Rabbit polyclonal Isl1/1	Gift Jessell Lab	N/A
Rabbit polyclonal anti Nkx2.2	Gift Jessell Lab	N/A
Donkey anti rabbit Alexa 488	Jackson	Cat# 711-545-152,
	ImmunoResearch	RRID:AB_2313584
Donkey anti mouse Alexa 488	Jackson	Cat# 715-545-150,
	ImmunoResearch	RRID:AB_2340846
Donkey anti guinea pig Alexa 488	Jackson	Cat# 706-545-148,
	ImmunoResearch	RRID:AB_2340472
Donkey anti goat Alexa 488	Jackson	Cat# 705-545-147,
	ImmunoResearch	RRID:AB_2336933
Donkey anti rabbit Cy3	Jackson	Cat# 711-165-152,
	ImmunoResearch	RRID:AB_2307443
Donkey anti mouse Cy3	Jackson	Cat# 715-165-150,
	ImmunoResearch	RRID:AB_2340813
Donkey anti guinea pig Cy3	Jackson	Cat# 706-165-148,
	ImmunoResearch	RRID:AB_2340460
Donkey anti rat Cy3	Jackson	Cat# 712-165-153,
	ImmunoResearch	RRID:AB_2340667
Donkey anti rabbit Alexa 594	Jackson	Cat# 711-585-152,
	ImmunoResearch	RRID:AB_2340621
Donkey anti mouse Alexa 594	Jackson	Cat# 715-585-150,
	ImmunoResearch	RRID:AB_2340854
Donkey anti guinea pig Alexa 594	Jackson	Cat# 706-585-148,
	ImmunoResearch	RRID:AB_2340474
Donkey anti rat Alexa 594	Jackson	Cat# 712-585-153,
	ImmunoResearch	RRID:AB_2340689
Donkey anti chicken Cy5	Jackson	Cat# 703-175-155,
	ImmunoResearch	RRID:AB_2340365
Donkey anti goat Cy5	Jackson	Cat# 705-175-147,
	ImmunoResearch	RRID:AB_2340415
Donkey anti mouse Cy5	Jackson	Cat# 715-175-150,
	ImmunoResearch	RRID:AB_2340819
Donkey anti guinea pig Alexa Cy5	Jackson	Cat# 706-175-148,
	ImmunoResearch	RRID:AB_2340462

Chemicals			
X-Gal	Roche	10745740001; CAS 7240-90-6	
DAPI	Sigma Aldrich	CAS 28718-90-3	
Commercial A	ssays		
Click-iT EdU	Thermofisher	C10340	
Mouse Stra	ins		
Mouse: Shh ^{tm1Ahk}	Gonzalez-Reyes L, et al., 2012	MGI:5440762	
Mouse: B6;129S6-Chat ^{tm1(cre)Lowl} /J	The Jackson Laboratory	JAX: 006410	
Mouse: B6.Cg-Tg(Nes-cre)1Kln/J	The Jackson Laboratory	JAX: 003771	
Mouse: Olig2 ^{tm1(cre)Tmj}	Dessaud E, et al., 2007	MGI:3774124	
Mouse: Gt(ROSA)26Sor ^{tm4(ACTB-tdTomato,-EGFP)Luo} /J	The Jackson Laboratory	JAX: 007576	

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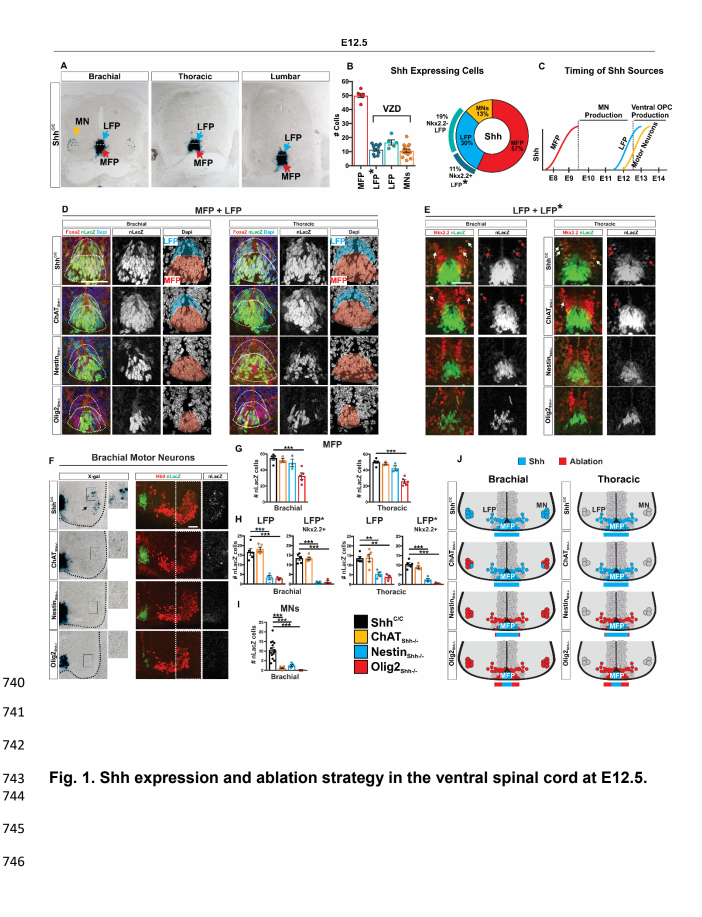
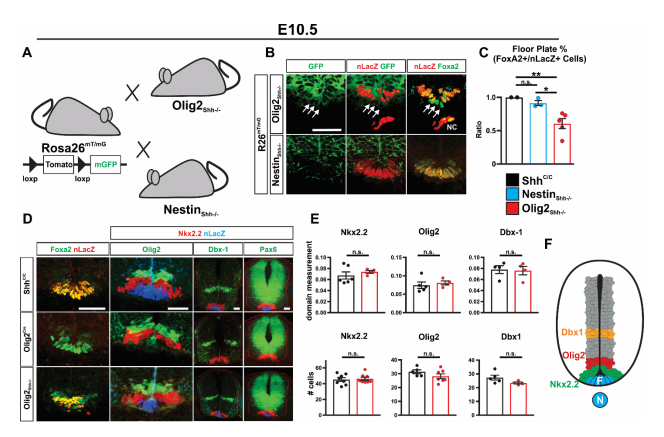


Fig. 1. Shh expression and ablation strategy in the ventral spinal cord at E12.5. 747 748 (A) X-gal staining of Shh expressing cells of E12.5 control Shh^{C/C} spinal cord sections. 749 750 Three sources identified: Medial floor plate (MFP), lateral floor plate (LFP), and motor neurons (MN). 751 (B) Numbers of nLacZ expressing cells in MFP (FoxA2+) n=6, LFP (FoxA2+ Nkx2.2-) 752 n=5, LFP* (Nkx2.2+) n=11, and among MNs (Hb9+) n=14. Ventricular zone derived 753 754 (VDZ): LFP, LFP*, and MNs. Breakdown percentages of each Shh source relative to total. Means ± SEM are shown. 755 (C) Overlap in MN and ventral OPC generation from pMN domain with the timing of Shh 756 757 expression from all sources. (**D**) Immunostaining for nLacZ and FoxA2 on brachial and thoracic segments. 758 Identification of MFP (FoxA2+ nLacZ+) and LFP (FoxA2+ nLacZ+). LFP cells are 759 identified as dorsal to MFP cells, oriented in tangent to MFP, and expressing lower 760 levels of nLacZ. 761 (E) Immunostaining for nLacZ and Nkx2.2 on brachial and thoracic segments. 762 Identification of LFP (Nkx2.2- nLacZ+) and LFP* (Nkx2.2+ nLacZ+). Arrows point to 763 migrating LFP* Nkx2.2+ nLacZ+ cells. 764 (F) X-gal staining and immunostaining for nLacZ and Hb9 on brachial segments. 765 (G) Quantification of MFP nLacZ recombination at brachial and thoracic segments for 766 each Shh source per genotype. Brachial and thoracic segments Shh^{C/C} n=5, ChAT_{Shh}^{-/-} 767 n=3, Nestinshh^{-/-} n=3, Olig2shh^{-/-} n=5. Means ± SEM are shown. One-way ANOVA, 768 Dunnett's multiple comparison post hoc test. *p<0.05, **p<0.01, ***p<0.001. 769 (H) Quantification of LFP and LFP* nLacZ recombination at brachial and thoracic 770 segments for each Shh source per genotype. Brachial and thoracic segments Shh^{C/C} 771 n=5, ChAT_{shh}^{-/-} n=4-5, Nestin_{shh}^{-/-} n=4, Olig2_{shh}^{-/-} n=3. Means ± SEM are shown. One-772

- way ANOVA, Dunnett's multiple comparison post hoc test. *p<0.05, **p<0.01,
- 774 ***p<0.001.
- (I) Quantification of nLacZ recombination in brachial MNs. Shh^{C/C} n=14, ChAT_{Shh}^{-/-} n=8,
- ⁷⁷⁶ Nestin_{Shh}^{-/-} n=5, Olig2_{Shh}^{-/-} n=5. Means \pm SEM are shown. One-way ANOVA, Dunnett's multiple comparison post hoc test. *p<0.05, **p<0.01, ***p<0.001.
- (J) Schematic of Shh expressing cells at E12.5 brachial and thoracic segments and
- strategy of Cre ablation for each genotype. Scale bars, 50 μm.
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784 Fig. 2. Early spinal cord patterning is unaffected in Olig2_{Shh}^{-/-}.

(A) Detection of Cre activity in $Olig2_{Shh}^{-/-}$ and $Nestin_{Shh}^{-/-}$ mutants at E10.5 using the conditional R26mT/mG allele.

(B) GFP co-labeling of Cre activity in addition to a loss of nLacZ from Foxa2+ cells

demonstrates Shh ablation in MFP of Olig2_{Shh}^{-/-} but not Nestin_{Shh}^{-/-} embryos. Arrows
 indicate MFP FoxA2+ cells that have lost nLacZ expression. NC. notochord.

indicate MFP FoxA2+ cells that have lost nLacZ expression. NC, notochord.
 (C) Quantification of recombination frequency of the ratio of Foxa2+ nLacZ+ double

positive cells. Shh^{C/C} n=2, Nestin_{shh}-/- n=3, Olig2_{shh}-/- n=5. Means \pm SEM are shown.

792 One-way ANOVA, Tukey post hoc test. *p<0.05, **p<0.01.

(**D**) Immunostaining of Shh-sensitive domains, Olig2, Nkx2.2, Dbx1, and Pax6 are

- unaffected at E10.5 in Olig2_{Shh^{-/-}}, despite MFP recombination.
- (E) Quantification of relative domain sizes and numbers of Nkx2.2, Olig2, Dbx1. Domain

measurement Shh^{C/C} n=4-6, Olig2_{Shh}^{-/-} n=4. Means ± SEM are shown. Cell counts

- 797 Shh^{C/C} n=5-8, Olig2_{Shh^{-/-}} n=4-8. Data were analyzed by Student's t test. *p<0.05, 798 **p<0.01.
- (**F**) Scheme highlighting position of p3 (Nkx2.2), p0 (Dbx-1), and pMN (Olig2) domains relative to the MFP at E10.5.
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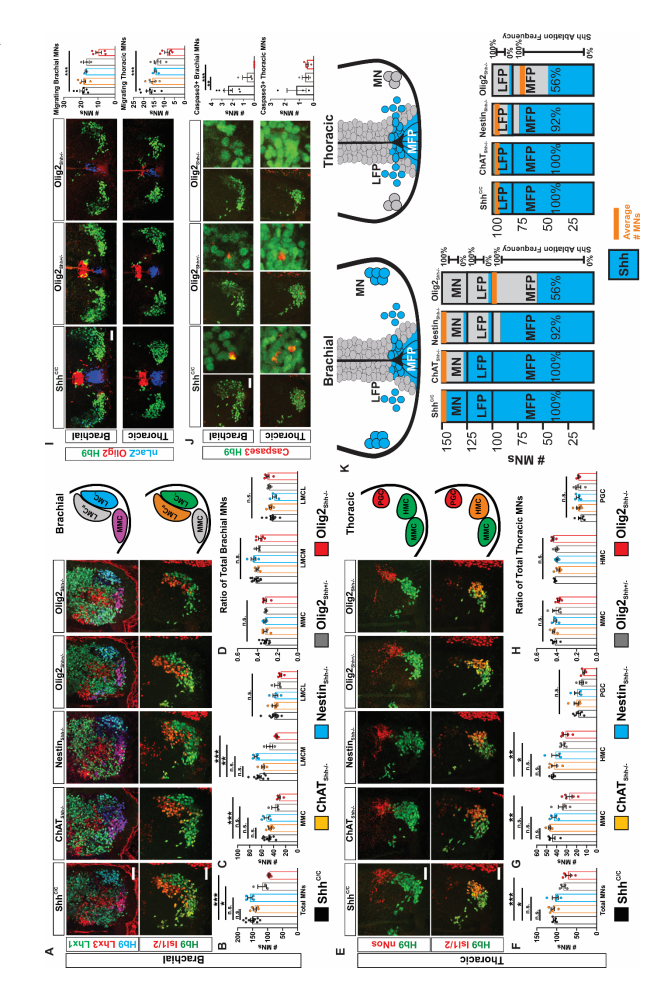


Fig. 3. Shh signaling from MFP, but not VZD influences MN generation.

(A) E12.5 brachial sections immunostained with Hb9, Lhx3, and Lhx1 to distinguish

807 MMC and LMCL columns, and Hb9 and Isl1/2 to distinguish LMCM and LMCL columns.

(**B**) Quantification of total brachial MNs. Shh^{C/C} n=11, ChAT_{Shh}-/- n=5, Nestin_{Shh}-/- n=4,

809 $\text{Olig2}_{\text{Shh}^{+/-}}$ n=3, $\text{Olig2}_{\text{Shh}^{-/-}}$ n=5. Means ± SEM are shown. One-way ANOVA, Dunnett's

multiple comparison post hoc test. p<0.05, p<0.01, p<0.001.

(C) Quantification of total numbers of MMC, LMCM, and LMCL MNs. Means ± SEM are

shown. One-way ANOVA, Dunnett's multiple comparison post hoc test. *p<0.05,
 p<0.01, *p<0.001.

(D) Ratio of each motor column to total brachial MNs. Means ± SEM are shown. One-

815 way ANOVA, Dunnett's multiple comparison post hoc test. NS, not significant, P>0.5.

(E) E12.5 thoracic sections immunostained with Hb9 and nNos to distinguish MMC and

817 HMC from PGC column, and Hb9 and Isl1/2 to distinguish MMC and HMC columns.

- (F) Quantification of total thoracic MNs. Shh^{C/C} n=4-5, ChAT_{Shh}-/- n=5, Nestin_{Shh}-/- n=4,
- 819 Olig2_{Shh}^{+/-} n=4, Olig2_{Shh}^{-/-} n=5. Means \pm SEM are shown. One-way ANOVA, Dunnett's

820 multiple comparison post hoc test. *p<0.05, **p<0.01, ***p<0.001.

(G) Quantification of total numbers of MMC, HMC, and PGC MNs. Means ± SEM are

shown. One-way ANOVA, Dunnett's multiple comparison post hoc test. *p<0.05,
 p<0.01, *p<0.001.

(H) Ratio of each column to total thoracic MNs. Means ± SEM are shown. One-way

ANOVA, Dunnett's multiple comparison post hoc test. NS, not significant, P>0.5.

(I) Immunostaining and quantification of late born migrating Hb9+ MNs at E12.5 brachial

- and thoracic segments. Shh^{C/C} n=6-7, ChAT_{Shh}^{-/-} n=5, Nestin_{Shh}^{-/-} n=3-4, Olig2_{Shh}^{+/-} n=3-
- 4, Olig2_{Shh}^{-/-} n=5. Means ± SEM are shown. One-way ANOVA, Dunnett's multiple comparison post hoc test. ***p<0.001.
- (J) Immunostaining and quantification of Hb9+ Caspase3+ apoptotic MNs at E12.5
- brachial and thoracic segments. Shh^{C/C} n=3-6, Olig2_{Shh}^{+/-} n=3, Olig2_{Shh}^{-/-} n=4-5. Means ±
- 832 SEM are shown. One-way ANOVA, Dunnett's multiple comparison post hoc test. 833 **p<0.01, ***p<0.001.
- (**K**) Schematic representing Shh ablation within the genotypes and associated numbers
- of average MNs for brachial and thoracic sections. Scale bars, 50 μm.
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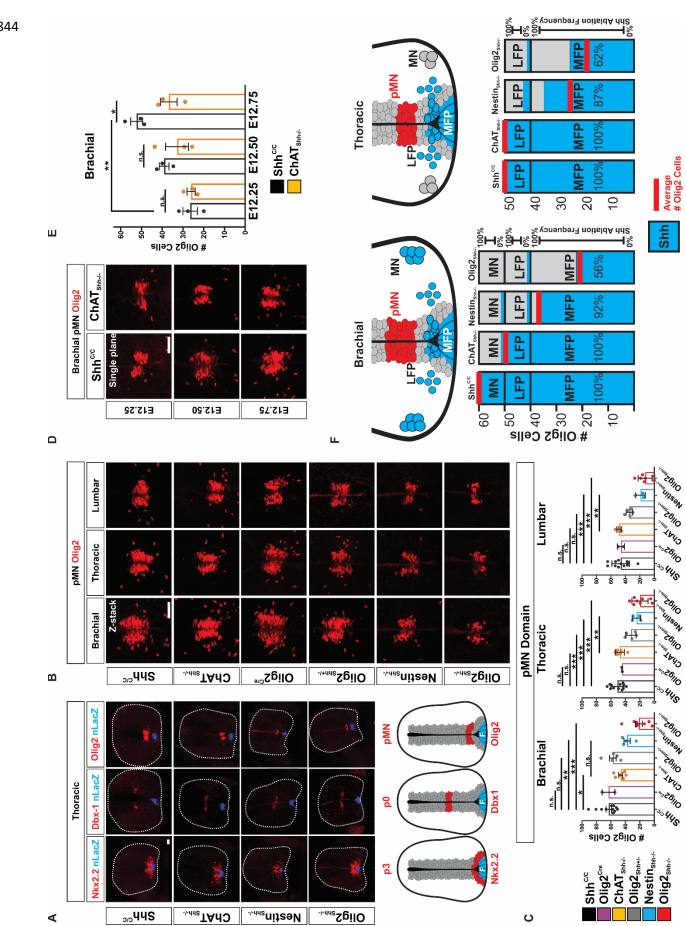
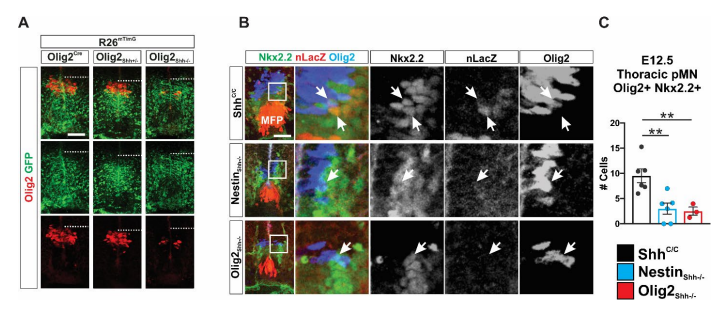


Fig. 4. Shh from VZD in addition to MFP is critical for pMN domain maintenance in a spinal level specific manner.

- (A) Immunostaining of nLacZ with Nkx2.2, Dbx-1, and Olig2 on E12.5 thoracic sections.
- (B) Immunostaining for Olig2 in the pMN domain along the AP axis on brachial,
- thoracic, and lumbar sections for Shh^{C/C}, ChAT_{Shh}-/-, Nestin_{Shh}-/-, Olig2-Cre, Olig2_{Shh}+/-, and Olig2_{Shh}-/-.
- (C) Quantification of numbers of Olig2 cells in the pMN domain for brachial, thoracic,
- and lumbar sections. Means ± SEM are shown. Shh^{C/C} n=12, Olig2-Cre n=2, ChAT_{Shh}-/-
- n=6, Olig2_{shh}^{+/-} n=5, Nestin_{shh}^{-/-} n=4, Olig2_{shh}^{-/-} n=8. One-way ANOVA, Dunnett's or
- Tukey's multiple comparison post hoc test. *p<0.05, **p<0.01, ***p<0.001.
- (D) Expansion of pMN domain at onset of gliogenesis at brachial segments between
 E12.25-E12.75 is reduced in ChAT_{Shh}-/-.
- (E) Quantification of Olig2 cells in the pMN domain on brachial segments. Means ±
- 858 SEM are shown. E12.25 Shh^{C/C} n=3, ChAT_{Shh}-/- n=3, E12.50 Shh^{C/C} n=3, ChAT_{Shh}-/- n=3,
- E12.75 Shh^{C/C} n=3, ChAT_{Shh}-/- n=3. Data were analyzed by Student's t test. *p<0.05.
- (**F**) Schematic representing Shh ablation within the genotypes and associated numbers
- of average pMN Olig2 cells for brachial and thoracic sections. Blue columns indicate
- percent of Shh expressing cells normalized to Shh^{C/C} controls for MFP, LFP, and MNs.
- Grey areas indicate Cre ablation. Red bars indicate average numbers of Olig2 cells
 within the pMN. Scale bars, 50 μm.
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Fig. 5. pMN_{Olig2+} expression and p* domain formation requires Shh from MFP and VZD during onset of gliogenesis.

(A) E12.5 Lineage tracing reveals correct establishment of the pMN in Olig2shh^{-/-} as

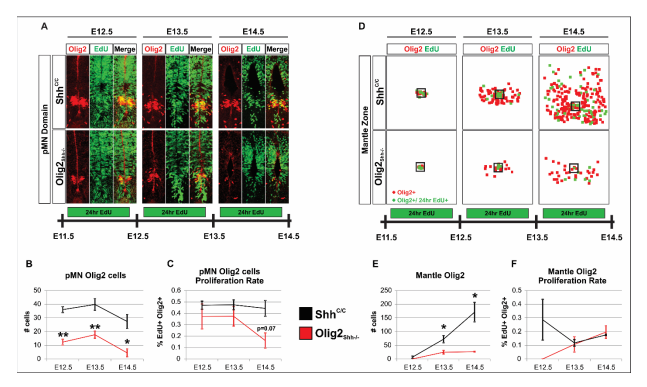
indicated by dorsal boundary of R26mT/mG expression, however failure of maintenance
 of Olig2 in pMN as detected by immunolabeling. Scale bars, 50 µm.

(B) p* domain is found in contact with LFP* in Shh^{C/C} controls but not in Nestin_{Shh^{-/-}} and Olig2_{Shh^{-/-}} as identified by immunolabeling of Olig2+ Nkx2.2 (p*) and nLacZ+ Nkx2.2+ (LFP*).

(C) Quantification of p* domain in Shh^{C/C}, Nestin_{Shh}-/-, and Olig2_{Shh}-/-. Means \pm SEM are shown. Shh^{C/C} n=6, Nestin_{Shh}-/- n=6, Olig2_{Shh}-/- n=3. One-way ANOVA, Dunnett's multiple comparison post hoc test. **p<0.01.

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898 Fig. 6. Exhaustion of pMN_{Olig2+} cells results in diminished OPC production.

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900 (A) pMN domain proliferation in lumbar sections labeled by EdU incorporation in 24hr

901 intervals for E12.5- E14.5 in Shh^{C/C} and Olig2_{Shh^{-/-}}.

(**B**) Total Olig2 cells in pMN at E12.5-E14.5. Means ± SEM are shown. Shh^{C/C} n=3-4,

Olig2_{Shh}^{-/-} n=3-4. Data were analyzed by Student's t test. *p<0.05, **p<0.01.

904 (**C**) Proliferation rate of Olig2 cells in pMN E12.5, E13.5, and E14.5. Means \pm SEM are 905 shown. Shh^{C/C} n=3-4, Olig2_{Shh^{-/-}} n=3-4. Data were analyzed by Student's t test.

906 (D) Tracing of migrating Olig2 cells on representative lumbar sections with proliferation

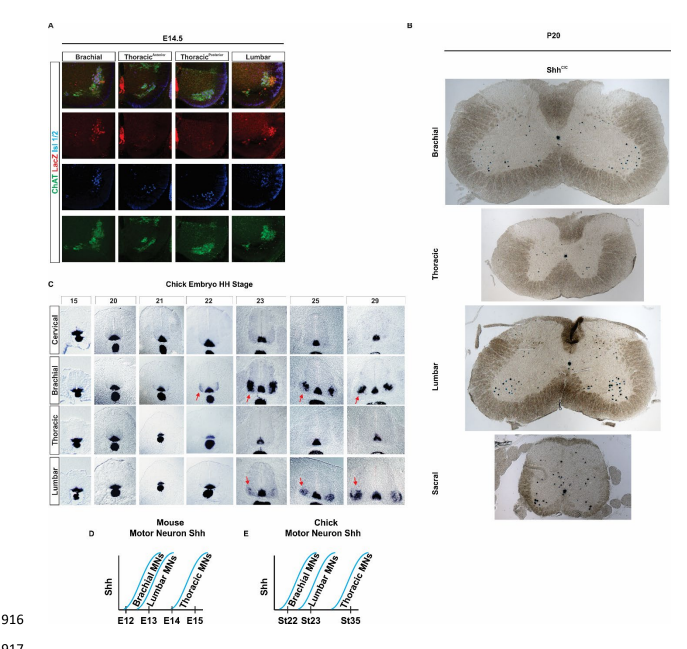
907 labeled by EdU incorporation in 24hr intervals between E12.5- E14.5. Box indicates

pMN domain, Olig2+ cells in the mantle zone (red), Olig2+ EdU+ cells (green).

(E) Total Olig2 cells in mantle zone at E12.5, E13.5, and E14.5. Olig2 cell numbers in

910 Olig2_{Shh}-/- mantle remain reduced. Means ± SEM are shown. Shh^{C/C} n=3-4, Olig2_{Shh}-/-

- n=3-4. Data were analyzed by Student's t test. *p<0.05.
- 912 (F) Proliferation rate of Olig2 cells in mantle zone at E12.5, E13.5, and E14.5. Olig2
- cells that have migrated out of the pMN in Olig2_{Shh}-/- mutants proliferate at the same rate
- as controls. Means \pm SEM are shown. Shh^{C/C} n=3-4, Olig2_{Shh^{-/-}} n=3-4.
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918 Fig. S1. Shh expression in MNs.

(A) Shh expression by MNs along AP axis at E14.5. Immunostaining colocalization of 919

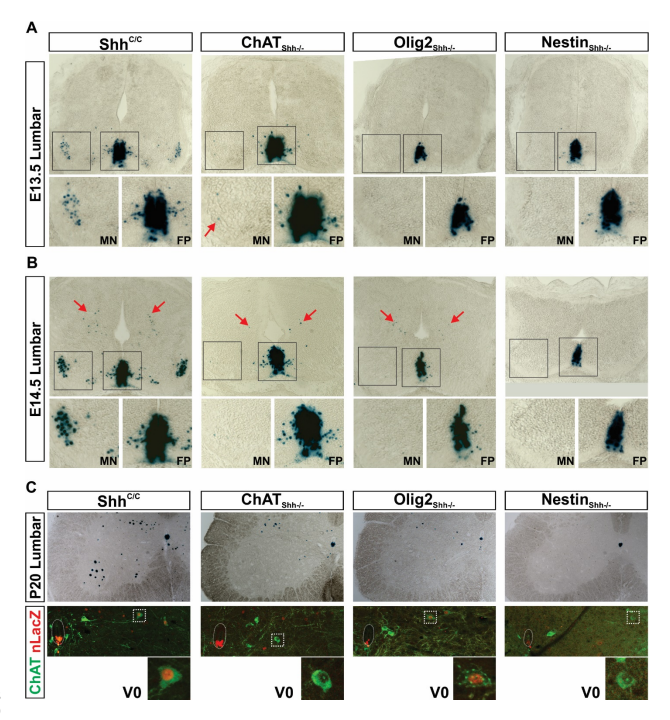
nLacZ with MN markers IsI1/2 and ChAT. Revealing that thoracic MNs express Shh 920 much later than brachial and lumbar. 921

(B) X-gal staining revealing Shh expression pattern throughout AP axis of P20 control 922

- Shh^{C/C} spinal cords at brachial, thoracic, lumbar, and sacral segments. 923
- (C) In situ hybridization for Shh in developing chick neural tube demonstrating 924

comparable timing and pattern of Shh expression by MNs as in mouse. 925

(**D** and **E**) Timeline of Shh expression by MNs in (**D**) mouse and (**E**) chick. 926



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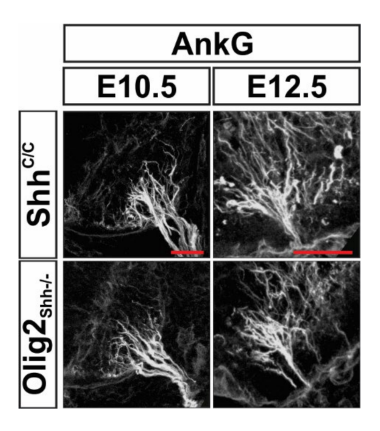
Fig. S2. Ablation of Shh from MFP and VZD sources 930

(A and B) X-gal staining revealing ablation of Shh at lumbar spinal cord at (A) E13.5, 931

and (B) E14.5. Arrows in B point to a dorsal Shh source located near the ventricular 932 zone appearing at E14.5. 933

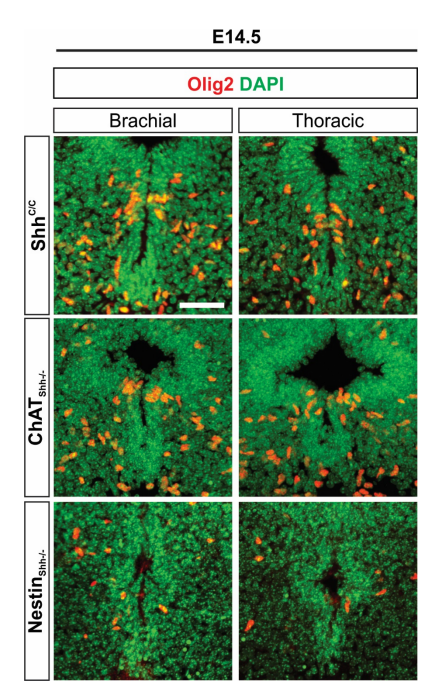
(C) X-gal staining of P20 spinal cords revealing loss of Shh expression from MNs and 934 V0 cholinergic interneurons in ChAT_{Shh}^{-/-}, MNs and descendants of neurons ventral to

- 935 the Olig2 domain in Olig2shh^{-/-}, and all neuronal sources in Nestinshh^{-/-}.
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940 Fig. S3. AnkG staining at E10.5 and E12.5 revealing correct MN axon fasciculation

941 and exit from ventral horns in Olig2_{Shh}^{-/-}.





- 955 Fig. S4. Immunostaining of Olig2 and DAPI on brachial and thoracic segments
- reveals the continued depletion of Olig2 cells from the pMN of Nestinshh^{-/-} but not
 ChAT_{shh}^{-/-} at E14.5.

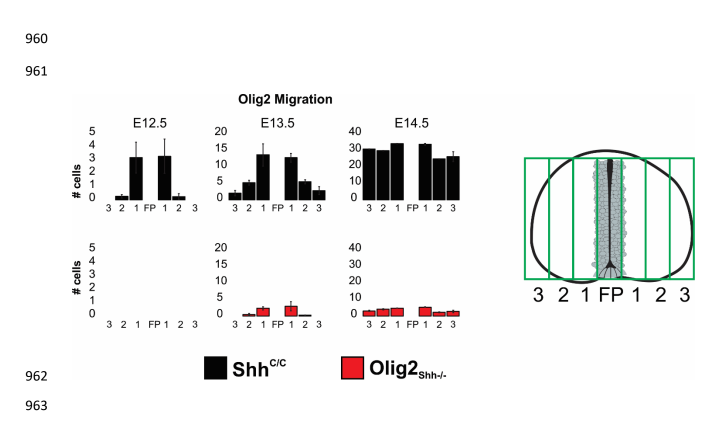


Fig. S5. Analysis of migrating Olig2 cell dispersion in Olig2_{Shh}-/- mutant and control embryos.

- Lumbar spinal cord sections from E12.5-E14.5 were binned into 6 zones excluding the ventricular zone, and numbers of Olig2 cells in each zone were quantified. Means ±
- SEM are shown. Shh^{C/C} (n = 3-4 embryos), Olig2_{Shh^{-/-}} (n = 3 embryos).
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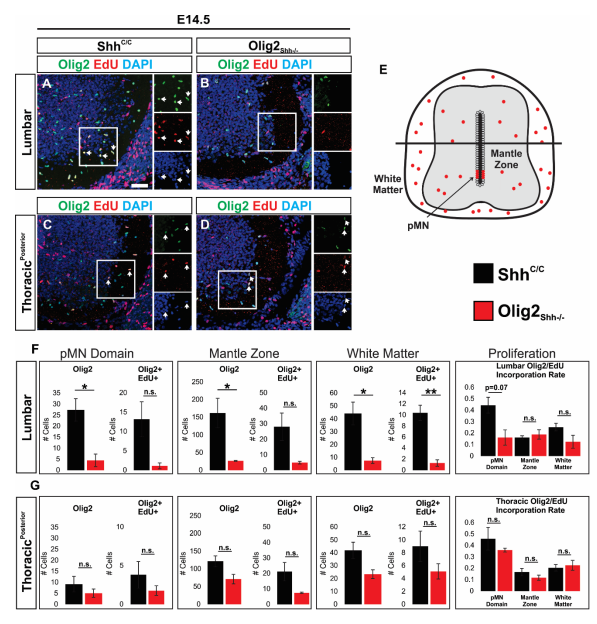
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980 Fig. S6. OPCs are reduced and do not increase proliferation rate.

981 (A-D) Immunostaining on E14.5 sections for Olig2, EdU, and DAPI at (A and B) lumbar

and (**C and D**) posterior thoracic. Arrows indicate co-expression of Olig2 and EdU.

983 (E) Schematic depicting areas analyzed.

984 (**F and G**) 24hr pulse chase with a single EdU injection to label proliferating Olig2 cells.

Total Olig2 cells and Olig2+ Edu+ double positive cells are reduced in the pMN, mantle zone, and white matter at both (**F**) lumbar and (**G**) posterior thoracic segments in

- Olig2_{sh}^{-/-} compared to control. Shh^{C/C} (n = 3 embryos), Olig2_{sh}^{-/-} (n = 3 embryos).
- Means \pm SEM are shown. Data were analyzed by Student's t test. *p < 0.05, **p<0.01.

989 Scale bars, 50 µm.

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