Asymmetric activation of Dll4-Notch signaling by Foxn4 and proneural factors activates BMP/TGFβ signaling to specify V2b interneurons in the spinal cord

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ABSTRACT

During development of the ventral spinal cord, the V2 interneurons emerge from p2 progenitors and diversify into two major subtypes, V2a and V2b, that play key roles in locomotor coordination. Dll4-mediated Notch activation in a subset of p2 precursors constitutes the crucial first step towards generating neuronal diversity in this domain. The mechanism behind the asymmetric Notch activation and downstream signaling events are, however, unknown at present. We show here that the Ascl1 and Neurog basic helix-loop-helix (bHLH) proneural factors are expressed in a mosaic pattern in p2 progenitors and that Foxn4 is required for setting and maintaining this expression mosaic. By binding directly to a conserved Dll4 enhancer, Foxn4 and Ascl1 activate Dll4 expression, whereas Neurog proteins prevent this effect, thereby resulting in asymmetric activation of Dll4 expression in V2 precursors expressing different combinations of proneural and Foxn4 transcription factors. Lineage tracing using the Cre-LoxP system reveals selective expression of Dll4 in V2a precursors, whereas Dll4 expression is initially excluded from V2b precursors. We provide evidence that BMP/TGFβ signaling is activated in V2b precursors and that Dll4-mediated Notch signaling is responsible for this activation. Using a gain-of-function approach and by inhibiting BMP/TGFβ signal transduction with pathway antagonists and RNAi knockdown, we further demonstrate that BMP/TGFβ signaling is both necessary and sufficient for V2b fate specification. Our data together thus suggest that the mosaic expression of Foxn4 and proneural factors may serve as the trigger to initiate asymmetric Dll4-Notch and subsequent BMP/TGFβ signaling events required for neuronal diversity in the V2 domain.

KEY WORDS: V2 interneuron, Spinal cord, Foxn4, Ascl1, bHLH proneural factor, Dll4-Notch, BMP/TGFβ, Chick, Mouse

INTRODUCTION

Neuronal diversity during spinal cord (SC) development is initially generated by activities of two competing signaling pathways: sonic hedgehog (Shh) ventrally and bone morphogenetic proteins (BMPs)/Wnt dorsally (Marti et al., 1995; Ericson et al., 1997; Liem et al., 1997; Edlund and Jessell, 1999; Jessell, 2000; Muroyama et al., 2002). Additional pathways subsequently get involved (Sockananth and Jessell, 1998; Novitch et al., 2003; Mizuguchi et al., 2006; Wildner et al., 2006; Del Barrio et al., 2007; Peng et al., 2007). Themed on the traditional paradigm, Hh signals are localized to the ventral SC and Gli repressor forms restrict activity in the dorsal SC (Jacob and Briscoe, 2003; Meyer and Roelink, 2003; Matise and Wang, 2011). Similarly, Wnt ligands are mostly restricted to dorsal regions, whereas inhibitors such as secreted Frizzled related proteins (sFRPs) are expressed in the ventral SC (Wodarz and Nusse, 1998; Kim et al., 2001; Kawano and Kypfa, 2003). BMP/TGFβ signaling, however, does not phenocopy this model. For instance, though implicated in dorsal fate specification, expression of Tgfβ2 is observed in notochord and floor plate (Garcia-Campmany and Marti, 2007). Additionally, expression of BMP/TGFβ signaling mediators Smad3, Smad4 and receptor-activated Smad1 and Smad5 is observed in almost all dorsoventral progenitor domains (Chesnutt et al., 2004; Garcia-Campmany and Marti, 2007; Hazen et al., 2012). This suggests undeciphered instructive roles for this pathway during ventral neurogenesis.

The V2 interneurons (INs) emerging from the p2 progenitor domain diversify into two major subtypes: V2a INs expressing Chx10 (Vsx2 – Mouse Genome Informatics) and V2b INs expressing Gata2 and Gata3 (Ericson et al., 1997; Zhou et al., 2000). The winged helix/forkhead transcription factor (TF) Foxn4 is essential for Ascl1 and Dll4 expression in this domain (Li et al., 2005; Del Barrio et al., 2007). Notably, although Foxn4 is initially expressed in all p2 progenitors, Dll4 transcription is observed only in a subset of INs (Del Barrio et al., 2007; Peng et al., 2007). It has been speculated that Dll4+ precursors give rise to V2a INs, whereas the neighboring Dll4– precursors, which receive the Dll4 ligand and activate Notch pathway, differentiate into V2b INs (Peng et al., 2007). The restriction of Dll4 expression to a subset of progenitors is the crucial step for generating asymmetry in immature postmitotic V2 precursors, which in turn is crucial for generating diversity. The mechanism behind this restriction, however, is presently unknown.

Notch ligands are regulated by proneural basic helix-loop-helix (bHLH) class of TFs (Bertrand et al., 2002; Castro et al., 2006; Henke et al., 2009). p2 progenitors express proneural TFs Ascl1, Neurog1 and Neurog2 as they initiate differentiation before onset of Dll4 expression. However, to date, no study has addressed the specific roles of these proneural genes in regulating Dll4 expression in V2 domain. Here, we provide evidence that Ascl1, Neurog1 and Neurog2 are expressed in a mosaic, balanced pattern in p2 progenitors and that Foxn4 is required for setting and maintaining this expression dynamic. The readout of this mosaic expression pattern results in asymmetric activation of Dll4 expression in V2 precursors expressing different combinations of proneural and Foxn4 TFs. One mechanism leading to this differential outcome
**RESULTS**

**Mosaic expression pattern of proneural factors Ascl1, Neurog1 and Neurog2 in p2 progenitors dictates V2 subtype specification**

Although earlier studies have analyzed expression of proneural bHLH TFs Ascl1, Neurog1 and Neurog2 in the developing SC (Parras et al., 2002), no study has addressed the specific roles of these proneural factors in generating V2 subtype diversity. As a first step to characterize the function of these proneural factors in V2 fate specification, we carried out detailed immunostaining expression analysis of Ascl1, Neurog1 and Neurog2 in the ventral mouse and chick SCs. At embryonic day (E) 10.5, Ascl1 shows a distinct expression pattern in the ventral SC that previous studies have mapped to p2 progenitors (Fig. 1A). The broader Neurog1 and Neurog2 expression in the ventral neural tube also overlaps with the p2 domain (Fig. 1B,C). A similar expression pattern for Neurog2 in the ventral mouse SC that previous studies have mapped to p2 IN progenitors (Fig. 1A). The broader Neurog1 expression pattern in the ventral mouse and chick SCs. At embryonic day (E) 10.5, Ascl1 shows a distinct expression pattern in the ventral SC that previous studies have mapped to p2 progenitors (Fig. 1A). The broader Neurog1 and Neurog2 expression in the ventral neural tube also overlaps with the p2 domain (Fig. 1B,C). A similar expression pattern for Neurog2 in the ventral mouse SC that previous studies have mapped to p2 IN progenitors (Fig. 1A). The broader Neurog1 expression pattern in the ventral mouse and chick SCs.

Ascl1, Neurog1 and Neurog2 are expressed in distinct patterns along the dorsoventral axis of the developing mouse spinal cord at E10.5. The bracket in B indicates the dorsal domain of Neurog1 expression. (E) Chicken Neurog2 displays an expression pattern similar to that of mouse Neurog2 at stage 22. (D) Magnified view of the p2 region marked in D. (F) Schematic illustration of the salt and pepper expression pattern showing progenitors that express only Ascl1 or Neurog1 and those that express both Ascl1 and Neurog1. (G-J) Co-expression analysis of Chx10 or Gata2 with Neurog1 in mouse (G,H) and with Neurog2 in chick (I,J) spinal cords. No overlap is observed except for occasional Neurog2 and Chx10 co-expressing neurons (arrows in I). Scale bar: 60 μm for A-E; 30 μm for G,H; 10 μm for I,J.

**Ascl1 and Neurog factors bind to a Dll4 enhancer but only Ascl1 can activate it**

Dll4-Notch signaling has been demonstrated to determine the V2b versus V2a interneuron fate in the V2 domain (Del Barrio et al., 2007; Peng et al., 2007). It is therefore possible that proneural factors and Foxn4 may control the V2 fates by directly regulating Dll4 expression. To determine this, we tested whether Foxn4, Ascl1 and Neurog2 were able to activate gene expression through an evolutionarily conserved Dll4 enhancer CR1 found to be active in the retina (Luo et al., 2012) (Fig. 3A). In the chick spinal cord, CR1 was able to drive DsRed reporter expression in a pattern that mimics that of the mouse Dll4 along the dorsoventral axis, predominantly in the V2 domain (supplementary material Fig. S1). Co-transfection of the CR1-DsRed reporter construct with a Foxn4-GFP expression plasmid led to robust DsRed expression in the electroporated side of the spinal cord (Fig. 3D-E’), whereas co-transfection with a control GFP expression vector induced a low level of DsRed expression, presumably resulting from endogenously expressed Foxn4 and other factors (Fig. 3B-C’).

To test the roles of proneural factors, the enhancer construct was co-electroporated with Ascl1- and Neurog-GFP expression plasmids. Similar to Foxn4, Ascl1 efficiently induced DsRed expression (Fig. 3F-G’); however, maximal reporter expression level was achieved only by cotransfection with both Ascl1 and Foxn4 expression plasmids (Fig. 3L-M’), indicating a synergistic effect. Neurog1 and Neurog2, on the other hand, were unable to drive the reporter expression (compare Fig. 3B-C’ with 3H-I’; data not shown). Furthermore, co-transfection of Neurog1 with Ascl1,
enhancer by Ascl1 and Foxn4, respectively (Fig. 3A,X-AA motifs nearly abolished DsRed reporter activation from the CR1 enhancer but have differential effects on Dll4 gene expression.

**Dll4 is selectively expressed in V2a precursors to non-cell-autonomously specify the V2b fate**

Dll4-Notch signaling has been shown to determine the V2b versus V2a fate choices adopted by p2 progenitors (Del Barrio et al., 2007; Peng et al., 2007), suggesting that Dll4 expression may exhibit cell-type specificity. To test this possibility, we produced a bacterial artificial chromosome (BAC) Dll4-Cre transgenic line (unpublished) to map the progeny of Dll4-expressing precursors by crossing with the R26R-YFP reporter line (Srinivas et al., 2001). For comparison we included Foxn4-Cre; R26R-YFP embryos in the analysis (Li et al., 2010). Whereas YFP-expressing progenies of both Cre lines arise from the p2 domain that expresses the Notch intracellular domain (NICD) (Fig. 4A,C), Dll4 expression is initiated later than that of Foxn4 (Fig. 4A,B). Analysis of Cre expression in E11.5 Foxn4-Cre embryos showed extensive overlap with that of the V2a IN marker Chx10 and V2b-specific Gata2 (Fig. 4D,E). Cre expression in Dll4-Cre mice, however, showed a different pattern. Whereas there was extensive overlap between Cre and Chx10 in V2a neurons, there was no colocalization between Cre and Gata2 in V2b nascent neurons (Fig. 4F,G). Similar analysis using YFP staining revealed near complete overlap of YFP with Chx10 and Foxn4 but repressed by Neurog proneural factors through the CR1 enhancer.

To narrow down the region in CR1 that can be activated by Ascl1, we tested a series of deletion reporter constructs by co-electroporation with the Ascl1-GFP expression plasmid (Fig. 3A,S-T″). However, no reporter expression was seen with the CR1a, CR1b and CR1g constructs (Fig. 3A,R-R″,U-V″). However, no reporter expression was seen with the CR1a, CR1b and CR1g constructs (Fig. 3A,S-T″,W-W″). These results thus define a 26-base sequence between CR1f and CR1g that can be activated by Ascl1 (Fig. 3A). Interestingly, this region falls within the previously defined Foxn4 critical region and contains both an E-box and a Foxn4 binding motif that are highly conserved among many vertebrate species (supplementary material Fig. S2A,B) (Luo et al., 2012). Mutating the E-box and cluster of Foxn4 binding motifs nearly abolished DsRed reporter activation from the CR1 enhancer by Ascl1 and Foxn4, respectively (Fig. 3A,X-AA′′).

We performed an electrophoretic mobility shift assay (EMSA) to test whether Ascl1 and Neurog proteins were able to directly bind to the minimal enhancer region using an oligo probe containing the critical E-box (Probe 2) and a control upstream probe that lacks an E-box (Probe 1) (supplementary material Fig. S2B). Although unable to form any complex with Probe 1, in the presence of the ubiquitous bHLH factor E12, in vitro translated Ascl1 formed a strong complex with Probe 2, which could be abrogated by excess wild-type but not mutant cold probes (supplementary material Fig. S2C, lanes 1-12, and S2D). Similarly, Neurog1 formed a specific complex with Probe 2 in the presence of E12 and the DNA-protein complex remained when both Ascl1 and Neurog1 were present (supplementary material Fig. S2C, lanes 13-18, and S2D). In addition, we carried out chromatin immunoprecipitation (ChIP) assays to show that both Ascl1 and Neurog1 were able to occupy in vivo the critical E-box region of the CR1 enhancer in cell culture and mouse embryonic neural tubes, whereas no enrichment was shown for control DNA in Dll4 3′ UTR (Fig. 3BB,CC; supplementary material Fig. S2E). Thus, Ascl1 and Neurog factors may bind to the same E-box in the Dll4 enhancer but have differential effects on Dll4 gene expression.

**Fig. 2. Foxn4-mediated equilibrium between Ascl1 and Neurog expression is essential for generating V2 interneuron diversity.**

(A-C) There is almost complete overlap between Ascl1 and Foxn4 expression in the p2 domain at E10.5-E12.5. (D-H) In spinal cords of E11.5 Foxn4 mutant embryos, there is complete loss of Ascl1 (D,D′) and Dll4 expression (G,G′) in the p2 domain, whereas normal expression of Neurog1 and Neurog2 is observed (E-F′). Arrows in G,G′ indicate Dll4 expression in blood vessels. Schematic representation in H,H′ shows that Neurog expression is insufficient to induce Dll4 expression in the absence of Ascl1 and Foxn4. Scale bars: in F′, 11.8 μm for A-C; in F′′, 20.7 μm for D-F′; in G′, 10 μm for G,G′.

Foxn4, or both Ascl1 and Foxn4 resulted in greatly reduced reporter expression (Fig. 3J-K′,N-Q′), indicating that Dll4 expression is activated by Ascl1 and Foxn4 but repressed by Neurog proneural factors through the CR1 enhancer.

Because misexpressed Dll4 promoted V2b differentiation (Peng et al., 2007), our data imply that Foxn4- and Ascl1-activated Dll4 non-cell-autonomously activates Notch signaling to specify the V2b fate. Consistent with this and a previous report (Peng et al., 2007), overexpression of the constitutively active NICD robustly induced Foxn4 and Gata2 expression, but markedly reduced Chx10 V2a cells in the chick SC (Fig. 4N-P). The induced Gata2 cells appeared to be V2b precursors, as they were mostly located in the ventricular zone, postmitotic, yet negative for the mature neuron marker TuJ1 (Tubb3 – Mouse Genome Informatics)
Fig. 3. Direct activation ofDll4 expression by Foxn4 and Ascl1, but not Neurog1, through a phylogenetically conserved enhancer. (A) Schematics of truncated CR1 DsRed reporter constructs. Corresponding relative activities by Ascl1 are indicated to the right (ND, not determined). The green and red ovals in the CR1 fragment indicate the E-box motif CANNTG, with the red being the critical one. The four cyan ovals show the clustered ACGC Foxn4 binding motifs. In the CR1cmE construct, the E-box is mutagenized to ATGATG, and in CR1cm(1-3), the first three Foxn4 binding motifs are mutated to AAAA. The vertical dashed lines outline the critical 26-bp region containing the critical E-box and a ACGC motif. (B-Q) DsRed expression in spinal cords co-electroporated with the CR1 reporter construct and indicated expression plasmids. Reporter activity was visualized in whole-mount spinal cords (B,B′,D,D′,F,F′,H,H′,J,J′,L,L′,N,N′,P,P′) or their cross sections (C,C′,E,E′,G,G′,I,I′,K,K′,M,M′,O,O′,Q,Q′). GFP alone showed only minimal endogenous enhancer activation (B,C,B′,D,D′). Both Foxn4 and Ascl1 could strongly induce DsRed expression (D,G′). With Neurog1 barely any induction of DsRed expression was observed (H,I′). This repression was maintained even when Neurog1 was co-electroporated with Ascl1 (J,K′). Foxn4 and Ascl1 together, by contrast, caused synergistic increase of enhancer activation (L,M′), and electroporation of all three factors together could rescue Neurog1 inhibition to some extent (N,O′). Notably, Neurog1 and Foxn4 co-electroporation also resulted in significant reduction of ectopic enhancer activation observed with Foxn4 alone (P-Q′). (R-W′) Ascl1 regulation of truncated CR1 DsRed reporter constructs. As visualized in whole-mount embryos (R,R′,U,U′,V,V′) and transverse sections (R″,U″,V″), CR1d and CR1f exhibited high levels of DsRed expression, albeit less than full-length CR1. By contrast, CR1a, CR1b and CR1g had little or no activity (S,T′,W,W′). (X-AA′) E-box mutation nearly abolished DsRed reporter activation from CR1c by Ascl1 (X-Y′) while mutating Foxn4 binding motifs abrogated activation by Foxn4 (Z-AA′). (BB,CC) ChIP assay showing enrichment of the critical E-box region by anti-Ascl1 and anti-Neurog1 antibodies on chromatin of neural tubes pooled from E10.5, E11.5 and E12.5 wild-type embryos (BB,CC). Control DNA region fromDll4 3′ UTR was not significantly enriched (BB). Statistical significance was determined by Student’s t-test: *P<0.05. Scale bars: in Q′, 70 μm for B,B′,D,D′,F,F′,H,H′,J,J′,L,L′,N,N′,P,P′; in Q″, 30 μm for C,C′,E,E′,G,G′,I,I′,K,K′,M,M′,O,O′,Q,Q′; in W, 54 μm for R-W″; in V″, 30 μm for R″,U″,V″; in AA′, 20 μm for X-AA′.
Msx1 is expressed in the ventral spinal cord (Fig. 5C-E). By double-immunolabeling, we determined that Msx1 is expressed specifically in the V2 domain, as it is excluded from the Isl1/2+ motoneurons located ventrally, the Jag+ V1 INs located dorsally, and from the Pax6+ progenitors (Fig. 5F,G,K-M). In the V2 domain, Msx1 is co-expressed with Gata2 mostly in the more medially located V2b progenitors (Fig. 5F,G,K-M). In the V2 domain, Msx1 is co-expressed with Chx10, but excluded from Gata2 expressing cells. (H-K) In E11.5 Foxn4-Cre; R26R-YFP spinal cords, both Chx10- and Gata2-immunoreactive subtypes were immunoreactive for YFP. Similar tracing analysis with Dll4-Cre; R26R-YFP mice revealed extensive overlap between YFP and Chx10 expression (J) but showed a transient exclusion of YFP from Gata2-immunoreactive V2b precursors at early stages of differentiation (K, highlighted by the dashed line). (L,M) Schematic illustrating the Cre expression patterns in E11.5 Foxn4-Cre and Dll4-Cre spinal cords. Dotted vertical lines in A-M mark the transition zone between proliferating progenitors and differentiating neurons. (N-R) Overexpression of activated Notch receptor (NICD) in the chick spinal cord caused widespread induction of Foxn4 and Gata2 expression (N-O′) but downregulation of Chx10 expression (P,P′). NICD-expressing cells highly colocalized with those labeled by bromodeoxyuridine (BrdU) and failed to express Tuj1 (Q-R′). Scale bars: in C, 20 μm for A-C; in K, 12 μm for D-K; in R, 20 μm for N-R′.

**BMP signaling promotes V2b interneuron generation**

During our analysis of p2 progenitor markers, we observed Msx expression in this domain using an antibody that detects all three Msx proteins (Msx1, 2 and 3). The ventral Msx expression is first seen in mice at E10.5 (Fig. 5A), while in the chick spinal cord it is observed between stages 17-18 when differentiation is initiated (Fig. 5B). To characterize expression patterns of individual murine Msx genes, we performed in situ hybridization, which showed that only Msx1 is expressed in the ventral spinal cord (Fig. 5C-E). By double-immunolabeling, we determined that Msx1 is expressed specifically in the V2 domain, as it is excluded from the Isl1/2+ motoneurons located ventrally, the Jag+ V1 INs located dorsally, and from the Pax6+ progenitors (Fig. 5F,G,K-M). In the V2 domain, Msx1 is co-expressed with Gata2 mostly in the more medially located V2b precursors but excluded from Chx10+ V2a cells (Fig. 5H,I,N). This expression pattern is similar to p21 (Fig. 5J), which marks nascent INs as they undergo a transition from late G1 to early G0 phase (Misra et al., 2008).

As Msx genes are known BMP signaling targets (Liu et al., 2004), our observations raised the possibility that BMP signaling may have a role in V2b IN specification. To test this possibility, we transfected a construct encoding the activated BMP receptor 1b (ABMPR) into chick SCs. Exogenous activation of BMP receptors was confirmed by ectopic induction of phosphorylated (p) Smad (pSmad1,5,8) and Msx (Fig. 5P-Q′,X). We found that misexpressed ABMPR increased the number of Gata3+ cells by approximately tenfold while reducing the number of Chx10+ cells by >80% (Fig. 5T-U′,X). Interestingly, ectopic induction of Gata3+ cells was largely limited to the ventral spinal cord (supplementary material Fig. S3A-B). ABMPR overexpression also significantly promoted Nkx2.2+ cells but decreased Lhx3+ cells, Evx1/2+ V0 cells and Isl1/2+ motoneurons (Fig. 5R,S′,V-W′,X). Together, these results suggest that activation of BMP signaling is sufficient to induce V2b and inhibit V2a IN differentiation.

**Inhibition of BMP/TGFβ signaling causes loss of V2b interneurons**

To determine whether BMP/TGFβ signaling is necessary for V2b neuron generation, we overexpressed BMP (noggin, Smad7 and chordin) and TGFβ (Ski and Smad7) inhibitors in chick SCs. Electroporation of all of these inhibitors caused great to near complete depletion of Gata3+ neurons (Fig. 6B,B′,E,E′,G,G′,I-L). As V2a INs could be inhibited both by activating and inhibiting the BMP/TGFβ signaling pathway, these effects are possibly secondary to ectopic induction of V2b cells.

To investigate directly the role of BMP/TGFβ pathway in V2b fate specification, we examined Smad3 knockout mice and...
conditional Smad4 knockout animals (Datto et al., 1999; Chu et al., 2004). Neither model showed any overt V2 phenotype at early stages examined (data not shown), suggesting redundancy between signaling components. We therefore analyzed the effect of knocking down both Smad3 and Smad4 simultaneously by RNAi constructs. Knocking down both Smads caused some general inhibition of differentiation, especially in the dorsal SC, as observed with Isl1/2 and Lhx1/5 staining (Fig. 6M,M′,O,O′). Notably for Isl1/2 the inhibition was restricted to the dorsal dI3 population and the ventral motoneurons were not inhibited (Fig. 7M). With respect to V2 fates, simultaneous Smad inhibition caused a 68% reduction of Gata3-expressing V2b INs but no significant change in Chx10+ V2a INs (Fig. 6Q,Q′,S,S′,V). Co-electroporation of scrambled Smad3 and Smad4 RNAi constructs did not cause significant alteration in either Gata3+ or Chx10+ INs (Fig. 6R,R′,T,T′,W). Consistent with a role of these Smads in V2 fate specification, Smad4 expression was observed in progenitors throughout the spinal cord (Fig. 6U). Smad3 has already been
shown to be expressed in the V2 domain (Martí et al., 1995). Together, these data demonstrate a requirement of BMP/TGF\(\beta\) signal transduction in V2b IN specification.

**Dll4-initiated Notch signaling acts upstream of BMP/TGF\(\beta\) signaling in V2b precursors**

Given that both Dll4-Notch and BMP/TGF\(\beta\) signaling are involved in V2b fate specification, it is important to decipher the hierarchy and interactions of these two signal transduction pathways. Overexpression of NICD caused widespread ectopic expression of pSmad and BMP target Msx1/2 in the chick SC, including the V2 domain (Fig. 7B-C'). Consistent with activation of Notch signaling by NICD, robust Dll4 and Gata3 expression was induced but Ascl1 and Chx10 expression was strongly suppressed (Fig. 7A,A',D-F'). By contrast, activation of the BMP pathway by ABMPR could not induce Dll4 expression (data not shown). Moreover, ABMPR-activated Gata3 induction was not suppressed by the dominant-negative Maml1, a Notch signaling inhibitor (supplementary material Fig. S3) (Peng et al., 2007).

To further delineate the effects of the two pathways, we co-transfected BMP inhibitor chordin with NICD. In the co-electroporated embryos, Dll4 induction and Ascl1 and Chx10 inhibition were still observed, but pSmad and Msx1/2 upregulation was completely blocked and Gata3 induction greatly diminished (Fig. 7G-L'). These results...
suggest that the activation of BMP signaling in V2b precursors depends on and occurs downstream ofDll4-Notch signaling during V2 IN development.

DISCUSSION

Asymmetric Dll4 activation by proneural and Foxn4 TFs as a basis of V2 subtype diversity

Although widely expressed, proneural TFs Ascl1, Neurog1 and Neurog2 have predominantly cross-repressive patterns with minimal overlap (Gowan et al., 2001; Parras et al., 2002; Helms and Johnson, 2003; Nakada et al., 2004; Helms et al., 2005; Ge et al., 2006; Battiste et al., 2007; Sugimori et al., 2007; Osório et al., 2010; Quiñones et al., 2010). This repression is validated in the developing SC also, where Ascl1, Neurog1 and Atoh1 are expressed in discrete cross-repressive domains (Gowan et al., 2001; Parras et al., 2002; Nakada et al., 2004; Helms et al., 2005; Kele et al., 2006).

Functionally, proneural factors have a crucial role in neuronal differentiation, a function largely dependent on activating Notch ligands. Of these, Dll1 expression is regulated by an enhancer having two distinct subdomains: one that requires Ascl1 binding for activity and the other that binds Neurog2 but is only partly required for SC expression (Bertrand et al., 2002). Dll3 enhancer, by contrast, has one critical E-box subdomain that can bind Ascl1, Neurog1 and Ascl1/Neurog2 heterodimers. Dll3 expression is mostly dependent on Ascl1, but not Neurog1 (Henke et al., 2009).

Differential regulation of Delta ligands is dictated largely by the expression pattern of proneural factors. V2 domain is unique, as it has progenitors expressing Dll4, Ascl1 and Neurog in exclusive as well as overlapping patterns. It thereby provides a context to analyze whether these factors have distinct roles based on exclusive expression patterns, or based on intrinsic properties. In this study, we show that although both Ascl1 and Neurog factors can bind to a conserved Dll4 enhancer, only Ascl1 can ectopically activate it. A balanced mosaic expression of the proneural factors is therefore crucial for initiating asymmetric expression of Dll4 in p2 progenitors. Although our studies focused on one conserved enhancer, it is conceivable that other Dll4 enhancers might be regulated in a different manner. Overall, mutually exclusive expression patterns might facilitate the regular function of proneural factors by heterodimerization with ubiquitously expressed bHLH E-proteins, whereas overlapping expression might be a mechanism adopted to create a mixture of functional and nonfunctional heterodimers with E-proteins and each other, thereby creating a basis for asymmetric activation of Notch ligand expression.

The generation of neural diversity by bHLH TFs is dependent on inputs from other regionally expressed factors (Powell and Jarman, 2008). One such example is the synergistic activation of Dll1 by simultaneous binding to its enhancer of Ascl1 and regionally expressed Brn1 (Pou3f1 – Mouse Genome Informatics) and Brn2 (Pou3f2 – Mouse Genome Informatics) (Castro et al., 2006). In the V2 domain Foxn4 is required for Ascl1 expression and V2b IN specification (Li et al., 2005; Del Barrio et al., 2007). Interestingly, the CR1 Dll4 enhancer contains a functional E-box and Foxn4 binding motifs located in close proximity (Fig. 8). Foxn4 may therefore operate at two levels for Dll4 regulation, one by directly binding and activating the enhancer and the other by regulating expression of Ascl1, which in turn can also bind to and activate the Dll4 enhancer. The juxtaposed location of the highly conserved binding motifs suggests a mechanism that requires proximity of these two TF binding sites for optimal Dll4 enhancer activity, perhaps dependent on direct interaction between Foxn4 and Ascl1. This may explain the synergistic effect of Foxn4 and Ascl1 on the Dll4 enhancer and the observed Foxn4 and Ascl1 co-expression in p2 progenitors at all stages of neurogenesis.

Because of the apparently complete colocalization between Foxn4 and Ascl1 in p2 progenitors, our data suggest the presence of four types of progenitors in the p2 domain: (1) those expressing Foxn4 and Ascl1; (2) those expressing Foxn4, Ascl1 and Neurog1 (or/and 2); (3) those expressing Neurog; and (4) those that express none of these TFs (Fig. 8A). We propose that this mosaic expression pattern of Foxn4 and proneural factors initiates asymmetric expression of Dll4 in p2 progenitors, which eventually leads to the specification of different V2 IN subtypes (Fig. 8). As expression of proneural genes has been shown to oscillate in neural progenitors (Kageyama et al., 2008; Shimojo et al., 2008; Kageyama et al., 2009), the mosaic expression pattern observed here may result in part from this oscillatory expression and may represent only a snapshot of the dynamic expression levels and asymmetry of Foxn4 and proneural factors in p2 neural progenitors. We found that although both Ascl1 and Neurog1 were able to bind the conserved E-box in the Dll4 enhancer, only Ascl1 could activate the enhancer, whereas Neurog
proteins inhibited the enhancer activation by Ascl1 and Foxn4 (Fig. 3; supplementary material Fig. S2), presumably by competition for common cofactors, competition for binding sites, or by disruption of pre-formed complexes. Thus, p2 progenitors that express both Foxn4 and Ascl1 are expected to express Dll4 strongly, whereas those expressing all three TFs will have a weak Dll4 expression, and those expressing Neurog alone or none of these transcription factors will lack Dll4 expression (Fig. 8A). The uneven expression of TFs coupled with lateral inhibition/cis-inhibition subsequently generates p2a progenitors with high levels of Dll4 ligand and neighboring p2b progenitors with high levels of activated Notch. The activated Notch in turn cell-autonomously activates BMP/TGFβ signaling essential for V2b fate specification.

Requirement of BMP/TGFβ signaling in V2b interneuron specification

Neurogenesis along the dorsoventral axis of the neural tube is initiated by extracellular inductive signals. BMP signals initiate patterning of the dorsal neural tube (Liem et al., 1997). Conversely, their downregulation is essential for specification of ventral fates (McMahon et al., 1998; Liem et al., 2000; Patten and Placzek, 2002; Timmer et al., 2002). Notably, Smad3 expression pattern supports this notion, as Smad3 is excluded specifically from motoneuron progenitors (García-Campmany and Martí, 2007). However, none of the previous studies looked specifically at the effect of BMP/TGFβ signaling on the V2 fate specification. We show in this study that overexpression of activated BMP receptor 1b could efficiently induce Gata2/3-expressing neurons. We further demonstrated the necessity of BMP/TGFβ signaling in V2b IN generation by overexpression of pathway inhibitors as well as simultaneous knockdown of Smad3 and Smad4 expression. Our study thus for the first time reveals an important instructive role for BMP/TGFβ signaling in ventral V2b IN fate specification.

BMP/TGFβ signaling acts downstream of Notch signaling in V2b interneuron generation

Our results show that Dll4 ligand expression represents the earliest step for initiating V2 subtype diversification. BMP/TGFβ signaling acts at a later step, initiated downstream in a subpopulation of p2 progenitors that are receiving Notch signaling. This conclusion is based on two observations. First, activation of Notch signaling induced pSmad expression that could be inhibited by chordin; however, chordin was unable to inhibit Notch-activated Dll4 expression. Second, activation of BMP signaling by dominant active ABMPR failed to induce Dll4 expression (data not shown) and co-expression of dominant-negative Maml1 did not prevent Gata3 induction. Nevertheless, there appeared to be a slight reduction in Gata3 induction compared with ABMPR overexpression alone. This might reflect sequestration of the endogenous Notch signaling operating in this region by dominant-negative Maml1. It is interesting to note that perturbations to Notch signaling lead to interconversion of V2 subtype fates (Li et al., 2005; Del Barrio et al., 2007; Peng et al., 2007). Inhibition of BMP/TGFβ signaling, by contrast, just inhibited V2b fates but no fate conversions were observed. This is true both when pathway inhibitors as well as RNAi knockdown was utilized. This indicates that BMP/TGFβ activation is a late event that comes into play after binary V2 fate decisions have been made. So although required for downstream activation of V2b specific markers, BMP/TGFβ signaling may not be involved in the binary fate decision per se.
MATERIALS AND METHODS

**Animals**

All experiments with mice were performed in accordance with animal protocols approved by Rutgers University. The C57BL/6j mice were purchased from the Jackson Laboratory and CD1 mice from the Charles River Laboratories. The Foxa4 knockout (Li et al., 2004), Foxa4-Cre (Li et al., 2010), Smad3 knockout (Datto et al., 1999), Smad4DNtop (Chu et al., 2004) and R26R-YFP (Streit et al., 2001) mice were generated previously and maintained by breeding with C57/BL6j mice. The stage of mouse embryos was determined by taking the morning when the copulation plug was seen as E0.5. All genotypes described were confirmed by polymerase chain reaction (PCR).

**In ovo electroporation and expression constructs**

Electroporation was performed on stage 11-12 chick embryos using a BTX square wave electroporator as described (Li et al., 2005). Transfected embryos were incubated for 48 hours and processed for immunohistochemistry (Li et al., 2005). For RNAi knockdown, embryos were collected 30 hours post-transfection. Following full-length cDNAs were subcloned into the pCIG vector (Megason and McMahon, 2002): Neurog1 (Ma et al., 1999) and dominant active BMPR-1b (ABMPR) (Timmer et al., 2002) into the pMWiii vector. The Neurog1 expression plasmid for dominant negative Maml1 was previously subcloned into pCIG. Neurog1 amino acids 1753–2185 (Gaiano et al., 2000) was also subcloned into pCIG. Bmp4 (Sela-Donenfeld and Kalcheim, 2002) was subcloned into the pCIG vector (Megason and McMahon, 2002): Neurog1 and dominant active BMPR-1b (ABMPR) (Timmer et al., 2002) into the pMWiii vector. The expression plasmid for dominant negative Maml1 was described previously (Peng et al., 2007). Transfection with the Bmp4 or ABMPR expression plasmids was visualized by co-electroporating the pCIG vector.

**shRNA plasmids**

Previously generated Smad3 shRNA and its control were used for Smad3 knockdown (García-Campmany and Martí, 2007). Smad4 shRNA plasmid was generated by inserting the already characterized region of Smad4 effective for gene silencing (Jazag et al., 2005) into the pBBS/u6 (pU6) RNAi vector (Sui et al., 2002): 5′-GATCCGGCCGCAATGTAAGAAGTCTTCAAGAGACGCTTCC-TACTAAGGCGCTCTTGGTTTTTG-3′ and 5′-AAATCAAAAAAGGAGCCGATATGAGGACTGTCTCTTGAACAGTCCTTCACTATGGCTGCCG-3′. Scrambled control sequences generated by Genscript were: 5′-GATCCGGAGTAAGCTACCGGTAAGGCTTCAAGAGAGCCTTACCGT-GAGTAGCTACCTCTCCTTTTGG-3′ and 5′-AAATCAAAAAAGGAGGTAAGCGTACCGTAAAGCCTACCTGAGATCCGTTACGCTC-3′. ChIP assays were also performed on chromatin DNA prepared from 293T cells co-transfected with the CR1 reporter construct and Ascl1, Neurog1 and E12 expression plasmids according to the instruction of the Simple Enzymatic Chromatin IP kit (Cell Signaling).

**Immunostaining, RNA in situ hybridization and BrdU labeling**

In ovo BrdU labeling, immunostaining and in situ hybridization were performed as described previously (Li et al., 2004; Mo et al., 2004; Misra et al., 2008). The following primary antibodies were used: rabbit anti-Foxn4 (Li et al., 2004); mouse anti-Ascl1, -Lhx3, -Isl1/2, -Evx1/2, -Nkx2.2, -Lhx1/5, and -Msx1/2 (Developmental Studies Hybridioma Bank); mouse anti-Neurog1 and rabbit anti-Neurog2 (Lo et al., 2004); mouse anti-Tuj1 and -Cre (Covance); rat anti-BrdU (Sigma); goat anti-Neurog1 and rabbit anti-Neurog2 (Cell Signaling); rabbit anti-Ascl1 and goat anti-Cdx2 (Abcam); goat anti-Neurog1 and -Neurog2 and -DesRed, mouse anti-Gata3 and -Smad4, and rabbit anti-Gata2 (Santa Cruz); sheep anti-Chx10 (Merck); guinea pig anti-Chx10 and -Gata2 (Peng et al., 2007); rabbit anti-NICD and -pSmad1/5,8 (Cell Signaling); goat anti-Dll4 (R&D); rabbit anti-Ascl1 (Gowan et al., 2004); and guinea pig anti-Lbx1 (Müller et al., 2002). Images were captured with a Nikon Eclipse 80i microscope. Probes used for RNA in situ hybridization were: chicken Foxn4 (Li et al., 2005) and mouse Msx1, Msx2 and Msx3 (Wang et al., 1996).

**Analysis of the CR1 Dll4 enhancer activity**

Dll4 enhancer alignment and reporter constructs were described previously (Luo et al., 2012). In addition, the critical E-box in the CR1 construct was mutated to ATGATG using the mutagenesis service of Genewiz. The enhancer reporter plasmids were co-transfected with various TF expression plasmids into the spinal cord of developing chick embryos as described above. Whole-mount images were taken with a fluorescence microscope, or the tissue was fixed and processed for cryosection and staining.

**Electrophoretic mobility shift assay**

EMSA was carried out as previously described (Liu et al., 2000). In vitro translated Ascl1, E12 and Neurog1 protein products were generated by a TNT T7 or Sp6 coupled reticulocyte lysate system (Promega) using Ascl1, E12 and Neurog1 expression plasmids (Henke et al., 2009). Competition was performed by adding excess amount of wild-type or mutant cold oligonucleotides to the reaction mixtures. The E-box was mutated from CAGATG to ATGATG in the mutant Probe 2 oligonucleotide (supplementary material Fig. S2B).

**Chromatin immunoprecipitation assay**

ChIP assays were performed on chromatin prepared from neural tube tissue pooled from E10.5, E11.5 and E12.5 wild-type mouse embryos using the Magna ChIP HiSens kit from Millipore. Isolated chromatin was sheared using a Diagenode Bioruptor for 30 minutes at 50% power with 30 seconds on/off cycles. Sixty micrograms of chromatin was incubated with 5 μg antibodies overnight. Antibodies used were rabbit anti-Ascl1 and goat anti-Neurog1 purchased from Bioss and Santa Cruz, respectively. ChIP enrichment was quantified by quantitative reverse transcription PCR (qRT-PCR) with Sybr Green mix from ABI. Percentage ChIP efficiency was calculated as described by Henke et al. (Henke et al., 2009). The following primers were used: for the critical E-box, 5′-CCACGCTGCGCTGCGACG-3′ and 5′-GGAGATTTCAGAACTGTTGCTGCC-3′, and for 3′ UTR, 5′- CCTCCCTTCACCCCATTTCTCC-3′ and 5′-TGTAAGAACGAGGCCAAGC-3′. ChIP assays were also performed on chromatin DNA prepared from 293T cells co-transfected with the CR1 reporter construct and Ascl1, Neurog1 and E12 expression plasmids according to the instruction of the Simple Enzymatic Chromatin IP kit (Cell Signaling).

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**Competing interests**

The authors declare no competing financial interests.

**Author contributions**

K.M., H.L., S.L., M.M. and M.X. designed and performed the experiments, and wrote the manuscript. All authors participated in the editing process.

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**Supplementary material**

Supplementary material available online at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.092536/-/DC1

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