Sequential actions of BMP receptors control neural precursor cell production and fate

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Bone morphogenetic proteins (BMPs) have diverse and sometimes paradoxical effects during embryonic development. To determine the mechanisms underlying BMP actions, we analyzed the expression and function of two BMP receptors, BMPR-IA and BMPR-IB, in neural precursor cells in vitro and in vivo. Neural precursor cells always express Bmpr-1a, but Bmpr-1b is not expressed until embryonic day 9 and is restricted to the dorsal neural tube surrounding the source of BMP ligands. BMPR-IA activation induces (and Sonic hedgehog prevents) expression of Bmpr-1b along with dorsal identity genes in precursor cells and promotes their proliferation. When BMPR-IB is activated, it limits precursor cell numbers by causing mitotic arrest. This results in apoptosis in early gestation embryos and terminal differentiation in mid-gestation embryos. Thus, BMP actions are first inducing (through BMPR-IA) and then terminating (through BMPR-IB), based on the accumulation of BMPR-IB relative to BMPR-IA. We describe a feed-forward mechanism to explain how the sequential actions of these receptors control the production and fate of dorsal precursor cells from neural stem cells.

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Animal development involves a complex progression of tissue induction and morphogenesis, expansion of precursor cell populations, and the death or terminal differentiation of these cells into specific functional types. Bone morphogenetic protein (BMP) signaling is repeatedly used in this dynamic process for both vertebrates and invertebrates. Bone morphogenetic proteins [BMPs] have diverse and sometimes paradoxical effects during embryonic development. To determine the mechanisms underlying BMP actions, we analyzed the expression and function of two BMP receptors, BMPR-IA and BMPR-IB, in neural precursor cells in vitro and in vivo.
Type I receptors also have BMP-binding properties but are primarily responsible for transducing the signal into the cell: BMPR-IA (Alk3), BMPR-IB (Alk6), and ActR-I (Alk2) are all known to transduce BMP signals [Kawabata et al. 1998]. The activated type I receptors in turn phosphorylate the DNA-binding proteins Smad1, Smad5, and Smad8, each of which can then heteromerize with Smad4 and translocate to the nucleus to regulate transcription of downstream target genes [Itoh et al. 2000].

The expression patterns of Bmpr-1a and Bmpr-1b in the CNS [Dewulf et al. 1995; Zhang et al. 1998] suggest that they have distinct roles in the transduction of BMP signals. In an effort to understand their role in neural development, we used multipotent CNS stem cells as a defined system to understand signals that control proliferation and fate choice [Johe et al. 1996; Studer et al. 1998]. We have also developed a vector using POU-regulated genomic elements that control expression of the intermediate filament gene nestin in CNS precursor cells [Josephson et al. 1998]. This vector, called pNERV, expresses mutant BMP receptors selectively in CNS stem cells in vitro and in transgenic mice. Using these tools, we show that most of the known actions of BMPs during neural precursor development can be attributed to the distinct actions of BMPR-IA and BMPR-IB. Furthermore, the activities of these receptors are linked sequentially, with BMPR-IA acting first and inducing the expression of Bmpr-1b. These results identify a feed-forward mechanism by which BMPs can control both the production and fate of precursor cells.

Results

Manipulating BMP receptor expression in neuroepithelial precursors

Extending on previous studies [Dewulf et al. 1995], we observed Bmpr-1a mRNA expression in all proliferative zones of the neural tube after gastrulation (Fig. 1A), regardless of location along the anterior-posterior or dorsal-ventral axis. In contrast, Bmpr-1b mRNA expression was not seen at embryonic day E8.75 or earlier but was first detected at E9.0 (Fig. 1B). It was expressed along the entire anterior-posterior axis but was restricted to the dorsal domain of the neural tube. Expression of Bmpr-1b (and to a lesser extent Bmpr-1a) was lower in the extreme dorsal-most domains corresponding to the roof plate and choroid plexus epithelium. Expression of both receptors was greatly reduced in postmitotic mantle layers in forebrain, hindbrain, and spinal cord (Fig. 1A,B), indicating that they are down-regulated after terminal differentiation.

To further examine the role of BMP receptors in nervous system development, we generated constructs expressing constitutively active (ca) or dominant-negative (dn) forms of BMP receptors [Zou et al. 1997] under the control of a new nestin promoter-based vector called pNERV [Fig. 1C; see Materials and Methods]. Expression of pNERV-driven constructs was strong in cultured CNS stem cells and was used for in vitro analysis of BMP receptor signaling. In vivo, transgene expression was exclusively localized to the proliferating neuroepithelium and neural crest (Fig. 1D,E). We observed very few position effects, possibly because the 5.4-kb nestin basal promoter contains A/T rich sequences similar to previously described matrix attachment regions [Charron et al. 1995]. In situ mRNA expression, epitope tag staining, and intensity of phenotype correlated well with transgene dose for each receptor at all ages analyzed (Fig. 1D,E). As expected, transgene expression under pNERV diminished once cells became postmitotic in older embryos. At all transgene doses, mice expressing caBmpr-1a showed qualitatively different gross phenotypes from those expressing caBmpr-1b (Table 1; Fig. 1D,E), suggesting that each receptor mediated distinct responses.

BMPR-IA activation increases precursor cell numbers

Embryos expressing caBmpr-1a could be readily distinguished from wild-type or caBmpr-1b embryos by a failure of the tail to elongate properly. This stubby, curly “pig-tail” was apparent by E12.5 and, along with an occasional cranial hematoma, was generally the only external sign of abnormality in these embryos (Fig. 2A,B). These animals died between E17.5 and birth, as no transgenics were found in newborn litters. Dissection of embryos revealed a pronounced gyrus-like appearance to the hindbrain and spinal neuroepithelium by E11.5 (Fig. 2C–L), leading to a dense packing of cells by E12.5 that obscured the ventricular lumen. This phenotype was observed in almost all caBmpr-1a transgenic embryos before E13 (Table 1). Because tissue gyriﬁcation is an indicator of increased proliferation, bromodeoxyuridine (BrdU) incorporation was assayed in wild-type versus transgenic embryos at two ages. At E11.5, the number of BrdU+ cells in the spinal cord of caBmpr-1a embryos was twofold higher (P < 0.01) than that of wild-type littermates (Fig. 2M). This proliferative effect was most pronounced in the dorsal neural tube. E14.0 caBmpr-1a embryos also showed twofold higher BrdU labeling than that of their wild-type littermates (Fig. 2N; P < 0.01), although the total number of proliferating cells was lower than that at E11.5. This suggested that BMPR-IA activation promotes proliferation in precursor cells throughout development.

Early dorsal patterning is induced by BMPR-IA activation

Inspection of the anterior neural tube revealed that the normal bilateral telencephalic vesicles (Fig. 3A,C) had been transformed in caBmpr-1a embryos into a single vesicle lacking a defined dorsal midline (Fig. 3B,D). This holoprosencephaly [Golden 1999] occurred with moderate and high transgene copy numbers (Table 1; Fig. 1D), was first seen at E11.5, and was severe at E15.5 (Fig. 3P). In the older animals, the entire alar domain rostral to the midbrain–hindbrain boundary was transformed into a cuboidal epithelium of single-cell thickness (Fig. 3N,P).
Figure 1. Bmpr expression in wild-type and transgenic embryos. (A) In situ hybridization for Bmpr-1a on wild-type embryos showing normal expression pattern during development; note ubiquitous expression in proliferating ventricular zone and lack of expression in differentiated mantle layer. Dorsal–ventral axis of neural tube is indicated by arrows. (B) In situ hybridization for Bmpr-1b on wild-type embryos showing normal expression pattern during development; note that expression is restricted to dorsal proliferating neuroepithelium and is not expressed in postmitotic mantle layer. All images in A and B are from adjacent sections except for E15.5 sections. (C) Diagram of construct [pNERV] used to control expression of mutant bone morphogenetic protein (BMP) receptors for all experiments. (D,E) Examples of transgene expression at E10.5 (moderate copy number, c.n.), E11.5 (lower c.n.), and E13.5–E14.0 (high c.n.). In situ mRNA hybridization for Bmpr-1a (D) or Bmpr-1b (E) shows ratio of receptor mRNA expression in wild-type versus transgenic embryos. Insets show Southern blot of pNERV transgene band just above endogenous nestin second intron band. Note distinct phenotypes caused by each transgene. Also note diminished transgene expression in older embryos. D, Dorsal; V, ventral; FB, forebrain; MB, midbrain; HB, hindbrain; SC, spinal cord; VZ, ventricular zone; and ML, mantle layer. [A,B] Bar, 500 µm for E10.5 and 500 µm for E12.5 and E15.5; [D,E] bar, 200 µm for E10.5 and E11.5 and 500 µm for E13.5 and E14.0.
and specifically express are derived from a less dorsal anlage in the neural tube (Fig. 3F,H). Its expression had expanded to include the entire alar domain of the forebrain and midbrain (Fig. 3E,G; Lim et al. 1997; Kaestner et al. 2000) indicated that its expression had expanded to include the entire alar domain along the entire rostrocaudal axis of the neural tube (Figs. 3Q, 4G).

This cuboidal morphology most closely resembled choroid plexus, which is the most dorsal CNS cell type generated in the anterior neural tube. In situ analysis of the choroid plexus marker Foxj1 (formerly known as Hfh4; Fig. 3E,G, Lim et al. 1997; Kaestner et al. 2000) indicated that its expression had expanded to include the entire alar domain of the forebrain and midbrain (Fig. 3F,H). The basal plate remained Foxj1 negative. In contrast to the choroid plexus, the cortex and ganglionic eminence are derived from a less dorsal anlage in the neural tube and specifically express Foxg1 (formerly known as Bf1; Fig. 3IK, Tao and Lai 1992, Kaestner et al. 2000). In caBmp-1a embryos, Foxg1 expression [Fig. 3J,L,Q] was replaced by expression of Foxj1 [Fig. 3F,H,Q]. As with Foxj1, the basal plate remained Foxg1 negative.

The hindbrain and spinal cord were also analyzed for induction of dorsal markers. The domain of Wnt1 expression, which is normally restricted to the roof plate and adjacent dorsal commissural precursors [Fig. 4A], was expanded to include the entire alar domain of the neural tube in caBmp-1a embryos [Fig. 4B]. Pax7, a dorsal precursor marker that in the mouse has a pattern of expression complementary to Wnt1 [Fig. 4C], was diminished or extinguished from the dorsal neural tube [Fig. 4D]. No ectopic Wnt1 and Pax7 expression was seen in the ventral neural tube [Fig. 4B,D], and ventral expression of Pax6 was unaffected in the caBmp-1a animals [Fig. 4E,F]. Thus, caBmp-1a expression dorsalized the alar domain along the entire rostrocaudal axis of the neural tube, but its effects were not seen in the ventral-most neural tube [Figs. 3Q, 4G].

Apoptosis is the early response to BMPR-IB activation

In contrast to caBmp-1a transgenics, embryos expressing caBmp-1b had obvious gross abnormalities in the rostral neural tube [Fig. 5A–H]. As early as E10.5, openings in the forebrain and midbrain neuroepithelium could be detected because of either failed closure of the neural tube or subsequent re-opening. Nearly all embryos age E11 or older showed exencephaly or at least partial neural tube opening, and almost all animals displayed a collapsed lumen in the caudal neural tube [Table 1; Fig. 6A,B]. No transgenics were recovered among newborn litters, indicating that the mice died at or before birth.

As the phenotype of these embryos strongly suggested a dysregulation of cell death [Haydar et al. 1999], we measured the expression of the proteolytically activated form of Caspase3 as an indicator of apoptosis [Srinivasan et al. 1998]. An increase in activated Caspase3 was seen in the ventricular neuroepithelium of caBmp-1b embryos at all ages measured between E9.5 and E12.5 compared with wild-type embryos [Fig. 5I–P]. This increase was widespread but most prominent in the forebrain and hindbrain. No enhanced cell proliferation was seen at any age analyzed [Figs. 6M, 7S]. In addition, dorsoventral identity was not altered, as measured by Pax6, Pax7, and Wnt1 expression [Fig. 6E–H, data not shown]. Staining of E11.5 embryos for the 160-kD subunit of neurofilament [NF160; Fig. 6I, J] showed no evidence of increased neuronal differentiation. Islet1 staining, which identifies early differentiating dorsal interneurons and ventral motorneurons [Jurata et al. 2000], was also unaffected [Fig. 6K, L]. Therefore, the principal early response to caBmp-1b was apoptosis.

Differentiation is a later response to BMPR-IB activation

By E13.5, cells in caBmp-1b embryos were disorganized and spilled from the dorsal and ventral roots of the spinal cord [Fig. 7A,B]. At high magnification [data not shown], individual cells were observed spanning the glia limitans, suggesting that a chemotropic response was causing cell migration. Among the closest cell types to the dorsal and ventral roots are the two postmitotic Islet1+ cell groups, one marking dorsal interneurons and the other marking ventral motorneurons [Jurata et al. 2000]. Antibody staining revealed that Islet1+ cells were ectopically located along the dorsal and ventral roots and were also scattered throughout the spinal cord [Fig. 7P,Q]. These cells could be distinguished from Islet1+ cells of the dorsal root ganglia by having less intense staining.

Although E11.5 caBmp-1b embryos showed no difference in BrdU labeling [Fig. 6M], by E13.5 the number of BrdU+ cells decreased, suggesting that terminal differentiation occurred [Fig. 7S]. The strongest phenotype showed a twofold decrease in BrdU+ cells, which correlated with a stronger migratory response and more severe exencephaly. Likewise, staining was reduced or eliminated for the pan-neural precursor marker nestin [Fig. 7E], as well as Pax7 [Fig. 7H], and Pax6 [Fig. 7K], indicating that both dorsal and ventral precursors were differentiating. Staining for NF160 showed wider and more scattered expression in transgenic embryos [Fig. 7N]. Large bundles of NF160+ fibers could be seen leaving both the dorsal and ventral roots, suggesting that large numbers of cells were differentiating into neurons and their processes exiting the spinal cord. An increase in the number of Islet1+ cells was also seen [Fig. 7Q], indicating that mid-gestation precursors prematurely differentiated into early neuronal types.

Evidence of a link between receptor actions

There was genetic evidence of a link between the BMPR-IA and BMPR-IB pathways based on certain phenotype

### Table 1. Summary of gross phenotypes from caBmpr-1a and caBmpr-1b transgenics

| Phenotype                        | Age range       | # expressing/# transgenic |
|---------------------------------|-----------------|---------------------------|
| Open fore/midbrain              | E10.0–E15.5     | 2/19                      |
| Complete exencephaly            | E10.0–E15.5     | 1/9                       |
| Collapsed neural tube           | E10.0–E15.5     | 0/19                      |
| Holoprosencephaly               | E11.0–E15.5     | 7/17                      |
| Gyrification                    | E11.0–E12.5     | 10/11                     |
| Cell migration/mixing           | E12.5–E15.5     | 5/7                       |

This table summarizes the gross phenotypes observed in transgenic embryos of the caBmpr-1a and caBmpr-1b alleles.

- **Open fore/midbrain**: Phenotype observed in embryos with a failure of neural tube closure, affecting the forebrain and midbrain regions.
- **Complete exencephaly**: Full separation of the forebrain from the spinal cord, with the cerebrum exposed externally.
- **Collapsed neural tube**: Partial or complete collapse of the neural tube, common in embryos with lethal phenotypes.
- **Holoprosencephaly**: Failure of the forebrain to develop properly, leading to a single nasal cavity.
- **Gyrification**: Development of convolutions in the cerebral cortex.
- **Cell migration/mixing**: Disruption of normal cell migration patterns, leading to embryonic lethality.

**Note**: The table entries reflect the number of embryos (expressing or transgenic) to the total number analyzed in each category. The data are compiled from observations made at specific embryonic ages (E10.0–E15.5), providing a comprehensive overview of the developmental abnormalities associated with these transgenic alleles.
similarities at early and mid-gestation. Activated Caspase3 was also increased in early caBmpr-1a embryos in the thinning forebrain neuroepithelium (data not shown), but unlike caBmpr-1b transgensics, these embryos had complete neural tube closure and normal craniofacial structure (Table 1; Fig. 2). This suggested that apoptosis occurred as a delayed response to BMPR-IA activation.

By E14.0, caBmpr-1a embryos also showed a migratory and differentiative response (Table 1; Fig. 7). However, comparison of the two transgensics at mid-gestation indicated that although caBmpr-1b showed only a differentiative response (Fig. 7), caBmpr-1a embryos still showed increased proliferation [Fig. 2N] as well as increased neuronal differentiation [Fig. 7O,R]. Thus, dorsal instruction and proliferation were seen only in caBmpr-1a embryos, but apoptosis and differentiation were seen in both transgensics. This suggested that BMPR-IB acts downstream of BMPR-IA.

Figure 2. Excess proliferation in early neural tube in caBmpr-1a transgenic embryos. (A,B) Whole mount images at E15.5 show wild type versus pNERV.caBmpr-1a embryos; note the slight distortion of the cranium [open arrow] and the truncated tail [arrow]. (C,D) Sagittal sections through E11.5 wild-type and caBmpr-1a embryos; note the extensive neuroepithelial gyrification in the transgenic embryos. (E,F) Higher (200x) magnification of hindbrain (box in C and D). (G,H) Bmpr-1a mRNA expression in wild-type and caBmpr-1a transgenic embryos shown in C and D. (I–L) Transverse sections showing hematoxylin and eosin and bromodeoxyuridine (BrdU) staining in E11.5 wt and caBmpr-1a transgenic embryos. (M,N) Bar graph indicating numbers of BrdU+ cells per unit spinal cord length in wild-type versus caBmpr-1a embryos at E11.5 and E14.0; P < 0.01. Bars: A–D,G,H, 1 mm; E,F,I–L, 200 μm.
BMPR-IB expression is induced by BMPR-IA activation and repressed by Shh

Two other sets of evidence suggested that the actions of BMPR-IA and BMPR-IB are linked. First, in situ hybridization revealed that although *Bmpr-1a* is expressed ubiquitously among neural precursors at all gestational ages (Fig. 1A), *Bmpr-1b* is expressed only after E8.75 and is restricted to dorsal neural precursors (Fig. 1B; Dewulf et al. 1995) surrounding the source of BMP ligands (Furuta et al. 1997). Second, the domain of *Bmpr-1b* expression expanded ventrally in some *caBmpr-1a* transgenic embryos [Fig. 8A]. The hypothesis most consistent with these observations is that activation of BMPR-IA induces or stabilizes *Bmpr-1b* expression in precursor cells, and subsequent BMPR-IB activation is responsible for the death or differentiation of these cells.

We tested this hypothesis by measuring BMP responses in dissociated stem cells from E12 rat ventral midbrain [VM; Fig. 8B; Studer et al. 1998], a region that does not normally express *Bmpr-1b* or dorsal identity genes. We measured the expression of *Bmpr-1b*, which is normally restricted to the dorsal two thirds of the neural tube, and the expression of *Msx1*, which is induced by BMP treatment (Bendall and Abate-Shen 2000) but restricted to the dorsal-most domain in the spinal cord and forebrain [Grove et al. 1998; Bang et al. 1999]. Treatment of VM stem cells with 5 ng/mL BMP2 resulted in an induction of both *Bmpr-1b* and *Msx1* expression within 2 h [Fig. 8C]. In contrast, *Bmpr-1a* expression was present in untreated VM stem cells and remained unchanged after BMP2 treatment [Fig. 8C], consistent with its ubiquitous expression in vivo. The induction of *Bmpr-1b* and *Msx1* was transient and began to decline after 8 to 24 h. Both *Bmpr-1b* and *Msx1* were induced by BMP2 in a dose-dependent manner [Fig. 8D]. The actions of BMP2 were inhibited by Sonic hedgehog (Shh), a ventralizing factor that is expressed in the notochord and floor plate [Briscoe and Ericson 1999]. Conversely, inhibiting Shh activity with cyclopamine [Cooper et al. 1998] promoted *Bmpr-1b* and *Msx1* induction [Fig. 8E]. Thus, BMPR-IA activation induces (and Shh represses) *Bmpr-1b* expression in the manner of a dorsal identity gene.

Bmpr-1b induction correlates with the onset of terminal differentiation in vitro

We then tested the effects of BMP signaling on stem cell proliferation and fate choice. We first measured BrdU
incorporation in E12 rat VM stem cells and found that low doses of BMP2 rapidly promoted proliferation (Fig. 9A), whereas higher doses had the opposite effect (Fig. 9B). To assay the instructive and differentiative effects of BMPs, we looked at the induction of neural crest, a dorsal precursor cell that emigrates from the neural tube to generate the peripheral nervous system, smooth muscle, and other cell types (Garcia-Castro and Bronner-Fraser 1999). Previous studies have shown that BMP treatment of cultured cortical, striatal, and spinal CNS stem cells efficiently changes their fate to neural crest (Mujtaba et al. 1998; Molne et al. 2000; T.G. Hazel, D.M. Panchision, M.H. Sailer, and R.D. McKay, unpubl.). We measured the generation of the postmitotic neural crest–derivative smooth muscle, as measured by the expression of smooth muscle α-actin (SMA; Shah et al. 1996; Tsai and McKay 2000). Low doses of BMP2 were insufficient to generate smooth muscle, whereas higher doses effectively generated smooth muscle (Fig. 9C). The BMP2 doses causing terminal differentiation [Fig. 9B,C] correlated with the doses that induced Bmpr-1b [Fig. 8D,E]. These results suggest that BMP2 causes proliferation through BMPR-IA until a threshold of Bmpr-1b induction is reached, after which cells undergo terminal differentiation.

**BMPR-IA and BMPR-IB act sequentially**

To test the hypothesis that the two BMP receptors act sequentially, we transfected cortical stem cells (Johe et al. 1996) with pNERV-driven mutant receptor constructs. We measured neural crest precursors by expression of p75NGFR (Morrison et al. 1999), differentiated neural crest progeny by expression of SMA, and mitotically arrested cells by expression of p21cip1, a cyclin-dependent kinase inhibitor (Jernvall et al. 1998). In non-transfected or mock-transfected conditions, cultures were composed of nestin+ stem cells, with no staining for these markers or markers of any other cell types [data not shown]. Stem cells transfected with caBmpr-1a initially expressed p75NGFR [Fig. 10A,E] but later expressed SMA [Fig. 10C,F] as well as p21cip1 [Fig. 10C,G]. This indicated that caBmpr-1a dorsalized cortical stem cells to neural crest precursors and then terminally differentiated these cells to smooth muscle. In contrast, stem cells transfected with caBmpr-1b expressed only p21cip1 [Fig. 10D,G], indicating that they had undergone mitotic arrest without prior dorsalization to the neural crest fate. Interestingly, the nuclei of p21+/SMA+ cells in caBmpr-1a transfected cultures had a characteristic large oval shape, whereas p21− nuclei of caBmpr-1b transfected cells were small and indistinguishable from the p21− nuclei of undifferentiated surrounding cells (Fig. 10C,D).

Stem cells cotransfected with both caBmpr-1a and dnBmpr-1b expressed the neural crest precursor marker p75NGFR but did not generate postmitotic smooth muscle [Fig. 10E–G], indicating that dnBmpr-1b blocked terminal differentiation. Stem cells cotransfected with both dnBmpr-1a and caBmpr-1b were indistinguishable from caBmpr-1b transfected cells alone [Fig. 10E–G], indicating that blocking BMPR-IA function did not prevent BMPR-IB from causing mitotic arrest. These results support the conclusion that initial instruction and proliferation are mediated by BMPR-IA, subsequent mitotic arrest is mediated by BMPR-IB, and these receptors are sequentially linked.

**Discussion**

**A model for BMP signaling during neural development**

In this study we show that the expression patterns of two type 1 BMP receptors reflect their distinct functions in CNS development. Bmpr-1a is expressed early and ubiquitously in precursor cells [Fig. 1A]. We found that BMPR-IA activation dorsalized and promoted proliferation of CNS precursor cells in vivo [Figs. 2–4] and in vitro
In contrast, Bmpr-1b is expressed only after E8.75 and is dorsally restricted [Fig. 1B]. We found that BMP2 treatment and BMPR-1A activation induced the expression of Bmpr-1b in ventral [Bmpr-1a⁺, Bmpr-1b⁻] precursor cells [Fig. 8], which may account for its delayed and restricted expression during embryogenesis. BMPR-IB activation caused apoptosis in early gestation embryos [Fig. 5] and accelerated differentiation of precursor cells to neurons at mid-gestation [Fig. 7]. Apoptosis and terminal differentiation were also seen in caBmpr-1a transgenic mice, suggesting that BMPR-IB functions downstream of BMPR-1A. This hypothesis was confirmed by in vitro experiments. First, ventral [Bmpr-1a⁺, Bmpr-1b⁻] precursor cells showed increased proliferation at low BMP2 doses that minimally induced Bmpr-1b expression [Figs. 8 and 9A]. Second, these same cells differentiated at higher doses that caused maximal Bmpr-1b induction [Figs. 8 and 9B,C]. Third, activating BMPR-IA while blocking BMPR-IB induced dorsal precursors that did not differentiate, yet activating BMPR-IB while blocking BMPR-1A yielded mitotically arrested cells that were not dorsalized [Fig. 10E–G].

Expression of caBmpr-1a transformed the anterior alar neural tube to Foxj1⁺ choroid plexus precursors [Fig. 3] and transformed the posterior alar neural tube to Wnt1⁺ roof plate or dorsal commissural precursors [Fig. 4]. This suggests that reports of BMP-induced holoprosencephaly (Golden et al. 1999) and Foxg1/Bf1 repression (Furuta et al. 1997) can be grouped with BMP-induced dorsalization of posterior neural precursors [Liem et al. 1995, 1997; Nguye et al. 1998, 2000; Barth et al. 1999; Nikaido et al. 1999] as inducing effects of BMPR-IA activation. Likewise, the apoptosis and terminal differentiation seen in acute BMP application studies in cortex (Furuta et al. 1999) as inducing effects of BMPR-IA activation. Although the noggin⁻/⁻/chordin⁻/⁻ double mutant has a phenotype more similar to early caBmpr-1a transgenics, including holoprosencephaly (Bachiller et al. 2000). Dissociated CNS stem cells can respond to BMPs by generating neural crest progeny (Munjaba et al. 1998; Molne et al. 2000; T.G. Hazel, D.M. Panchision, M.H. Sailer, and R.D. McKay, unpubl.). Our results show that this is caused by the dorsalizing actions of BMPR-IA followed by the differentiative actions of BMPR-IB [Fig. 10].
These results lead us to propose a model for BMP actions on neuroepithelial precursors during development (Fig. 11). An early, unpatterned precursor responds to BMPs through the actions of BMPR-IA, which is expressed ubiquitously among precursor cells from gastrulation onward. Activation of BMPR-IA induces the expression of dorsal identity genes, including Bmpr-1b, and promotes the proliferation of this dorsalized cell. As cell surface levels of accumulating BMPR-IB exceed those of BMPR-IA, BMPs then cause a termination response of cell death or cell differentiation. The in vitro results indicate that a single BMP ligand can drive this feed-forward, self-limiting process (Figs. 8 and 9) and suggest that the dose of BMPs determines the rate at which it occurs.

Alternative BMP pathways have opposing actions on cell number

Our study shows that BMPR-IA and BMPR-IB mediate antagonistic drives: an initial one that instructs precursor cells and expands their numbers, and a final one that limits the numbers of that cell type. Previous studies support a proliferative role for BMP ligands and BMPR-IA. Application of BMP7 causes proliferation in the ventral hindbrain [Arkell and Beddington 1997], consistent with the expression of Bmpr-1a but not Bmpr-1b in this region. Mice that lack Bmpr-1a die before gastrulation partially because of a proliferation defect; in these animals, total cell number in the epiblast is reduced by 20% to 30%, and primitive streak formation is not initiated [Mishina et al. 1995]. In chick limb bud, BMPR-IA activation delays chondrocyte differentiation whereas BMPR-IB activation promotes apoptosis or chondrocyte condensation [Zou et al. 1997]. Mice that lack functional Bmpr-1b in the limb have complex defects, including incomplete distal limb differentiation (Zou et al. 1997; Baur et al. 2000; Yi et al. 2000). Interestingly, the targeted Bmpr-1b−/− mouse has no obvious reported neural defects [Yi et al. 2000], suggesting that redundant mechanisms exist to promote terminal differentiation.

The proliferation of vertebrate limb muscle precursors is promoted by low doses of BMPs, acting through Pax3. At higher doses, BMPs promote repression of Pax3 and terminal differentiation [Amthor et al. 1998, 1999]. This is consistent with the dose-dependent BMP stimulation of proliferation and terminal differentiation we see in our assays (Fig. 9A–C). It also suggests that the increase in proliferation by 6 h [Fig. 9A] depends first on the induction of dorsal patterning genes (e.g., Pax, Fox) through BMPR-IA. We show that terminal differentiation correlates with the induction of Bmpr-1b [Figs. 8C–E, 9A–C]. Ectopic Bmpr-1b expression has been reported in chick limb buds treated with TGFβ3 ligands [Merino et al. 1998, 1999] or infected with caBmpr-1a [Zou et al. 1997]. Bmpr-1b repression is seen in limb buds treated with noggin [Pizette and Niswander 2000]. Although the functional significance of these results is not established, they suggest that the sequential activation of BMP receptors may regulate precursor cell identity and number in other tissues.

We also observed that Bmpr-1b (and to a lesser extent
Terminal differentiation in mid-gestation transgenic embryos. Lumbar level of E13.5 wt, E13.5 *caBmpr-1b* (moderate copy number, c.n.) and E14.0 *caBmpr-1a* (high c.n.) spinal cord. (A–C) Hematoxylin and eosin staining shows well-defined boundaries in wild-type embryos compared with loss of boundaries in both *caBmpr-1b* and *caBmpr-1a* embryos; note migratory response. (D–L) Staining with markers of precursor cells such as nestin, Pax7, and Pax6 show a reduction or elimination of staining in *caBmpr-1b* and *caBmpr-1a* embryos. (M–R) Staining with markers of differentiated cells such as NF160 and Islet1 shows an increase in number and/or area of expression in *caBmpr-1b* and *caBmpr-1a* embryos. Note the migration of Islet1+ cells and the spreading of NF160-positive fibers throughout the spinal cord and out of the dorsal and ventral roots. (S) Bar graph indicating numbers of BrdU+ cells per unit spinal cord length in E13.5 wild-type versus *caBmpr-1b* embryos, variability between transgenic animals reflects variable penetrance of the phenotype. Bar, 200 µm.
Bmpr-1a expression is paradoxically lower near the immediate source of BMPs (Fig. 1A,B). This may be because of accelerated apoptosis or terminal differentiation of cells in regions of highest BMP activity (Fig. 9), resulting in a loss or down-regulation of the receptors (Fig. 11). Alternatively, in Drosophila the highest Dpp levels negatively regulate expression of the receptor thick veins (tkv) to control the range of the Dpp gradient (Lecuit and Cohen 1998; Teleman and Cohen 2000; Entchev et al. 2000). If a similar mechanism operates in mammals, then Bmpr-1b expression would be both positively (Fig. 8) and negatively regulated by BMP ligands.

An intriguing question is how the distinct BMPR-IA and BMPR-IB signals are transduced inside the cell. There is no evidence to date that differential activation of Smad1/5/8 proteins can account for the distinct actions of BMPs. TGFβ family ligands have been reported to also activate MAP kinases of the p38, p46/54JNK, and p42/44ERK families, but the evidence is contradictory about whether these responses are either direct or specific for such effects as mitogenesis or apoptosis (Hu et al. 1999; Iwasaki et al. 1999; Monzen et al. 1999; Nakamura et al. 1999; Yamaguchi et al. 1999; Kimura et al. 2000). If a similar mechanism operates in mammals, then Bmpr-1b expression would be both positively (Fig. 8) and negatively regulated by BMP ligands.

Figure 8. Induction of Bmpr-1b expression by BMP2 and caBmpr-1a. (A) In situ hybridization for Bmpr-1b on transverse hindbrain sections of E11.5 wild-type and caBmpr-1a embryos; open arrow indicates ectopic Bmpr-1b expression in ventral hindbrain. (B) Diagram showing isolation and culturing of E12.5 rat ventral midbrain (VM) stem cells. (C) RT–PCR of E12.5 VM stem cells showing induction of Bmpr-1b and Msx1 within 2 to 4 h of treatment with 5 ng/mL BMP2, Bmpr-1a and GAPDH expression are unchanged. (D) Induction of Bmpr-1b and Msx1 by rncPCR in E12.5 VM stem cells after 4-h treatment with varying doses of BMP2. (E) Induction of Bmpr-1b and Msx1 at 4 h is inhibited by Shh cotreatment; inhibition of Shh by cyclopamine promotes induction of Bmpr-1b and Msx1. Bar, 500 µm.

Modulation and inhibition of the BMPR-IB termination response

We show that in early gestation embryos (before E12), caBmpr-1b caused mainly apoptosis (Fig. 5), as markers of patterning, proliferation, and differentiation remained unchanged (Fig. 6). However, as development proceeded, caBmpr-1b accelerated terminal differentiation of precursor cells (Fig. 7). This is consistent with our conclusion that BMPR-IB does not specify cell fate per se but instead mediates a generic terminal response by causing cell cycle arrest, leading to apoptosis or differentiation (Fig. 10B,D–G; Coffman and Studzinski 1999; Yoshikawa 2000).

These results suggest that additional competence signals are required to interpret the BMPR-IB termination response. BMP-treated precursors differentiate to a neuronal fate when neurogenin is expressed (Sun et al. 2001) but are directed to a glial fate when the Delta–Notch pathway is activated (Morrison et al. 2000). Changes in competence signal expression may explain why CNS stem cells in vitro show an age-dependent bias in their response to BMPs, ranging from apoptosis to neurogenesis and then gliogenesis (Gross et al. 1996; Mabie et al. 1999; Mehler et al. 2000; T.G. Hazel, D.M. Panchision, M.H. Sailer, and R.D. McKay, unpubl.).

We also show that as long as BMPR-IB expression was low or its activity blocked, BMP signaling maintained
cells in a dorsal precursor state (Figs. 9 and 10). There is evidence to support negative control of the BMPR-IB termination response. Apoptosis can be blocked by survival factors like insulin, which activates a PI3 kinase/Akt pathway to prevent TGFβ-mediated apoptosis (Chen et al. 1998). Differentiation may be blocked by mitogenic signals like EGF, which uses a p42/44ERK-mediated pathway to inhibit Smad activation (Kretzschmar et al. 1997). This would permit cell proliferation to continue despite the expression of BMPR-IB and would explain why the dorsal neural tube does not immediately differentiate once Bmpr-1b is induced (Fig. 1).

Generality of the mechanism

There is evidence that the feed-forward mechanism described here may also operate in invertebrates. In Caenorhabditis elegans, loss of function in the BMP receptor homolog sma-6 results in a small body phenotype (Krishna et al. 1999). The C. elegans BMP homolog db1-1 also controls body size by regulating cell growth and/or proliferation (Suzuki et al. 1999). Loss-of-function mutations in Drosophila Dpp and its receptor tkv result in patterning defects and decreased proliferation of wing imaginal disk precursors (Burke and Basler 1996; Edgar and Lehner 1996). These results suggest a BMPR-IA-like function for the invertebrate receptors. However, the tkv mutant also has a later defect in differentiation (Burke and Basler 1996), suggesting that it activates a BMPR-IB-like pathway.

We show how sequential activation of BMP receptors allows BMP ligands to control precursor cell production and fate. This may be one way to regulate tissue identity and size during development (Day and Lawrence 2000). Such a mechanism could act as a built-in safeguard against unrestricted growth during tissue renewal or repair (Sakou 1998) and may be one of the regulatory processes disrupted in cancer (Ishidou et al. 1995; Yamada et al. 1996; Kim et al. 2000).

Materials and methods

Construction of mutant type I BMPRs

cDNAs encoding mutant forms of the BMP receptors were generated by PCR-mediated site-directed mutagenesis and/or were the kind gift of L. Niswander (Memorial Sloan Kettering Cancer Center, New York). The ca forms contain a Glu to Asp substitution in the GS domain (Q233D caBmpr-1a, Q203D caBmpr-1b) to mimic phosphorylation by the type II receptor (Zou et al. 1997). The dn forms were generated by a Lys to Arg substitution in the ATP-binding site of the kinase domain (K261R dnBmpr-1a, K231R dnBmpr-1b), which abolishes kinase activity (Zou et al. 1997). Mutant Bmpr-1a cDNAs were tagged at the C terminus with the synthetic Flag epitope; mutant Bmpr-1b cDNAs were tagged at the C terminus with the hemagglutinin (HA) epitope derived from the human influenza virus.

Construction of expression vector

A neuroepithelial expression/reporter vector (pNERV) was constructed using a 5359-bp genomic fragment containing the nestin basal promoter and a 653-bp genomic fragment of the conserved portion of the nestin second intron (Josephson et al. 1998) to confer neural precursor-specific expression of the BMP receptor mutants. The nestin promoter was ligated upstream of a splice donor/acceptor site, lacZ reporter gene and SV40 polyadenylation sequence derived from a CMVβ expression vector (Clontech). The nestin second intron fragment was ligated downstream of the polyadenylation site in a reverse orientation. The lacZ reporter gene contained flanking NotI sites to allow
replacement of lacZ cDNA with Bmpr cDNAs. Staining for the β-galactosidase reporter in E11.5 to E15.5 mouse embryos indicated strong staining in the proliferative ventricular zone throughout the neural tube but not in the roof plate, floor plate, or prechordal plate (data not shown). No staining was seen in postmitotic mantle layers of the neural tube, consistent with previous studies (Josephson et al. 1998). Expression was detected in 94% of all embryos containing at least one transgene copy for any cDNA; ectopic expression was never observed.

**Generation of transient transgenic mice**

Transgenic mice were produced as described (Hogan et al. 1994). Mice used were C57Bl6 × C3H F1 embryo donors, C57Bl6 × C3H F1 stud males, and ICR foster mothers. Integration of the transgene was assayed by yolk sac PCR and/or Southern blot analysis (Hogan et al. 1994) using the 653-bp nestin second intron fragment as a probe. Copy number could be estimated by comparing intensity of pNERV transgene band versus endogenous nestin second intron band on Southern blot (Fig. 1D,E). Embryos were staged on dissection; the day of transfer roughly corresponded to 0.5 d postcoitum.

**Histological analysis**

Embryos were fixed in Bouin’s solution from 1 to 20 h, depending on the age of the embryo. Embryos were dehydrated in an ethanol series, clarified, and embedded in paraffin. Sections (10 µm) were mounted on Superfrost/Plus microscope slides (Fisher). Slides were de-paraffinized, rehydrated, and, for routine morphological analysis, stained with hematoxylin and eosin (H&E). For immunohistochemistry, slides were subjected to high-temperature antigen unmasking using 0.1M sodium citrate (pH 6.0) for 10 min, followed by a 15-min cooling period. For peroxidase staining, slides were incubated in 0.1% hydrogen peroxide in PBS for 30 min to quench endogenous peroxidase and then were rinsed in distilled water. Blocking was performed with 1x PBS containing 5% normal goat serum (NGS) and 0.1% Triton X-100. Primary antibodies were used against nestin (McKay lab, rabbit 130), activated Caspase3 [Iddn, CM11], Pax6 (gift of R. Reed, Johns Hopkins University, Baltimore, MD), Pax7 (DSHB), Islet1 (DSHB, clone 40.2D6), NF_{160} (Sigma N5264), smooth muscle α-actin (Sigma A2547), p75NGFR (Calbiochem GR10), and p21cip1 (Calbiochem PC55). Primary antibodies were diluted in blocking solution and incubated on sections for 16 to 72 h at 4°C. Sections were washed in 1x PBS.
BrdU+ cells were counted using Stereo Investigator (Micro-}

for 10 min, followed by 2N HCl in PBS for 30 min before block-

brightfield), and a Student

above. In situ hybridization using S 35-labeled riboprobes was

Embryos were fixed, processed, and sectioned as described

was performed as described [Hogan et al. 1994]. Template cDNAs

used for probe synthesis were generated by RT–PCR, and their

sequence was verified, including Wnt1 (mouse, accession no. M1943, bases 1053–1732), FoxJ1/Hand1 (mouse, accession no. L13204, 1633–2112), FoxJ1/Fli1 (mouse, accession no. U36760, 375–1167), Bmpr-1a (human, accession no. Z22535, 310–668),

and Bmpr-1b (mouse, accession no. Z23143, 1–742). Autoradi-

ographic emulsions were performed as described (Hogan et al.

and were exposed 4–14 d.

Figure 11. Induction–termination model for BMP receptor ac-

tions. In the induction phase, a precursor cell in the early post-

gastrulation embryo expresses only BMPR-IA and responds to

BMPs by expressing dorsal identity genes (including Bmpr-1b)

and by proliferating. This response is blocked in cells of the

ventral domain by the actions of Shh. The termination phase

occurs when cell surface levels of accumulating BMPR-IB ex-

ceed those of BMPR-IA, so that a cell responds to BMPs with

mitotic arrest. Changing competence signals (+) interpret the

response to BMPR-IB so that an early precursor cell terminations

by apoptosis while a later precursor cell terminates by differen-

tiation; inhibitory signals (−) can delay termination response.

for 3 × 20 min and incubated with secondary antibodies in 1×

PBS containing 1.5% NGS for 2 h. After washing, sections were

processed using Vectastain ABC Elite peroxidase kit (Vector)

and diaminobenzidine [DAB] with metal enhancer (Sigma). Sec-

tions were counterstained with eosin or cresyl violet, dehy-

drated, and mounted. Images were photographed in brightfield

using a Zeiss Axiosplan microscope and Spot digital camera [Di-

gnostic Instruments] and combined for figures using Photoshop

5.0 for Windows [Adobe].

BrdU staining and counting

Pregnant recipient mice were injected with 50 mg BrdU per

gram body weight 1 h before being killed. Embryos were col-

clected, processed, and sectioned as described above. Staining

was performed as described above using anti-BrdU (Becton-

Dickenson 347580), with the addition of an incubation in 0.1%

(w/v) trypsin (Sigma), 0.1% (w/v) CaCl2 in 0.1 M Tris (pH 7.5)

for 10 min, followed by 2N HCl in PBS for 30 min before block-

ing. BrdU+ cells were counted using Stereo Investigator [Micro-

brightfield], and a Student t-test was used to determine statis-

tical significance.

In situ hybridization

Embryos were fixed, processed, and sectioned as described

above. In situ hybridization using S55-labeled riboprobes was

performed as described [Hogan et al. 1994].
from the Developmental Studies Hybridoma Bank. We are grateful for the gift of Pax6 antibody from Dr. Randall Reed and cyclopamine from Dr. William Gaffield. We also thank Dr. James Nagle of the NINDS DNA-sequencing facility for sequence verification of constructs, Drs. Heather Cameron and Susan Wray for technical advice, and Dr. Martin Sailer, Helen Mitchell, Michael Nguyen, Emily Liu, and Martha Kimos for excellent technical assistance.

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