We conclude that BMP activity emanating from the lens is crucial for the maintenance of eye-field character, inhibition of dorsal neural tube/optic vesicle stages, BMP signals from the lens are not well defined. In the present study, we have analyzed these issues by manipulating signaling pathways in intact chick embryo and explant assays. Our results provide evidence that at blastula stages, BMP signals inhibit the acquisition of eye-field character, but from neural tube/optic vesicle stages, BMP signals from the lens are crucial for the maintenance of eye-field character, inhibition of dorsal telencephalic cell identity and specification of neural retina cells. Subsequently, our results provide evidence that a Rax2-positive eye-field state is not sufficient for the progress to a neural retina identity, but requires BMP signals. In addition, our results argue against any essential role of Wnt or FGF signals during the specification of neural retina cells, but provide evidence that Wnt signals together with BMP activity are sufficient to induce cells of retinal pigment epithelial character. We conclude that BMP activity emanating from the lens ectoderm maintains eye-field identity, inhibits telencephalic character and induces neural retina cells. Our findings link the requirement of the lens ectoderm for neural retina specification with the molecular mechanism by which cells in the forebrain become specified as neural retina by BMP activity.

KEY WORDS: BMP, Chick, Development, Eye, Lens, Neural retina

INTRODUCTION

During early development of the vertebrate central nervous system (CNS), the anterior neural domain becomes restricted into different regions, giving rise to the telencephalon, eye-field and hypothalamus (reviewed by Garcia-Lopez et al., 2009). The eye-field gives rise to most structures of the eye, such as the neural retina, the retinal pigment epithelium (RPE) and the optic stalk. By contrast, the lens of the eye derives from the lens ectodermal placode (reviewed by Gunhaga, 2011). Whether the specification of eye-field and tissue induction for over a century. Nevertheless, the molecular mechanisms that regulate the induction and maintenance of eye-field cells, and the specification of neural retina cells are poorly understood. Moreover, within the developing anterior forebrain, how prospective eye and telencephalic cells are differentially specified is not well defined. In the present study, we have analyzed these issues by manipulating signaling pathways in intact chick embryo and explant assays. Our results provide evidence that at blastula stages, BMP signals inhibit the acquisition of eye-field character, but from neural tube/optic vesicle stages, BMP signals from the lens are crucial for the maintenance of eye-field character, inhibition of dorsal telencephalic cell identity and specification of neural retina cells. Subsequently, our results provide evidence that a Rax2-positive eye-field state is not sufficient for the progress to a neural retina identity, but requires BMP signals. In addition, our results argue against any essential role of Wnt or FGF signals during the specification of neural retina cells, but provide evidence that Wnt signals together with BMP activity are sufficient to induce cells of retinal pigment epithelial character. We conclude that BMP activity emanating from the lens ectoderm maintains eye-field identity, inhibits telencephalic character and induces neural retina cells. Our findings link the requirement of the lens ectoderm for neural retina specification with the molecular mechanism by which cells in the forebrain become specified as neural retina by BMP activity. When describing development of any cell type, it is important to distinguish between cell fate and cell specification. Cell fate is the identity a cell adopts if left undisturbed in the embryo, whereas cell specification is defined as the step whereby cells have received sufficient signals to acquire a specific cell identity if cultured outside the embryo (Patthey and Gunhaga, 2011). Fate maps of chick embryos at neural plate stages have defined prospective eye-field cells in the anterior neural plate (Fernández-Garre et al., 2002; Sánchez-Arrones et al., 2009). Previous studies have revealed the importance of a set of overlapping transcription factors, such as Rax, Rax2, Six3, Six6, Pax6 and Lhx2, in promoting an eye-field identity (Grindley et al., 1995; Lagutin et al., 2003; Swindell et al., 2008; Zuber et al., 2003). The transcription factor genes Rax (retinal and anterior neural fold homeobox) and Rax2 are among the earliest markers of the eye-field, being initially expressed in the anterior neural region of head-fold stage embryos, but later becoming restricted to the neural retina and the ventral hypothalamus (Sánchez-Arrones et al., 2009). Inactivation of Rax in mouse or its ortholog rx3 in zebrafish within the anterior neural plate leads to complete absence of eyes as a result of failure to form the optic vesicles (Loosli et al., 2003; Mathers et al., 1997). However, how Rax2-positive eye-field cells progress to a neural retina character has not been determined. At neural tube closure, the optic vesicle comes in contact with the prospective lens ectoderm. In mammals and birds, the optic vesicle and the lens placode invaginate simultaneously. Following invagination, the optic vesicle transforms into a bilayered optic cup, in which the inner layer gives rise to the neural retina and the outer layer gives rise to the RPE (Fuhrmann, 2010). At optic vesicle stages, Vsx2 (visual system homeobox 2, previously known as Chx10) is upregulated in the prospective neural retina (Fuhrmann, 2010), whereas Mitf (microphthalmia associated transcription factor) is induced in the presumptive RPE (Mochii et al., 1998). Vsx2 mutants exhibit reduced proliferation of neural progenitors within the optic vesicle and, at later stages, bipolar cells are absent from within the retina (Burmeister et al., 1996). The signals that regulate the specification of neural retina cells and when this occurs have not yet been defined. Moreover, whether the specification of neural retina cells requires lens-derived signals and which one(s) remains controversial (Eiraku et al., 2011; Hyer et al., 1998).

Bone morphogenetic protein (BMP) signals have been shown to play important roles during eye formation. Several studies have shown that BMP activity is required for lens induction (Furuta and Hogan, 1998; Pandit et al., 2011; Rajagopal et al., 2009; Sjödal et al., 2007; Wawersik et al., 1999). In addition, both chick and mouse, BMP activity regulates the initial dorso-ventral patterning of the neural retina (Adler and Belecky-Adams, 2002; Kobayashi et al., 2010; Murali et al., 2005). In mouse, it has also been shown that BMP signaling is essential for retinal growth after embryonic day (E) 10.5 and for early retina neurogenesis (Murali et al., 2005). In vivo studies in chick have revealed that BMP activity is required for the development of the RPE (Muller et al., 2007), and that implanted BMP-soaked beads result in downregulation of neural...
retina markers and induction of RPE-like cells (Hyer et al., 2003; Muller et al., 2007). However, whether BMP signals are involved in the maintenance of eye-field identity and/or specification of neural retina cells has not been determined.

In the present study in chick, we show that eye-field cells become independent of adjacent tissues only at stage 13, coincident with the specification of neural retina cells. Prior to this stage, eye-field and optic vesicle cells cultured alone acquire dorsal telencephalic character. At the blastula stage, low levels of BMP signals prevent the generation of eye-field cells, whereas at neural tube/optic vesicle stages, BMP signals from the lens ectoderm are required and sufficient to maintain eye-field identity, block telencephalic character and specify neural retina cells. In addition, our results argue against any essential role for Wnt or FGF signals during the specification of neural retina cells.

RESULTS

Characterization of markers of the optic vesicle and other forebrain domains

To examine when cells of the eye-field acquire neural retina character, we analyzed the generation of neural retina cells in relation to other eye and forebrain cells. To achieve this, the expression of a panel of markers was monitored in chick (Gallus gallus) embryos from stage 9 to stage 21 of development on consecutive sections (Fig. 1; supplementary material Fig. S1).

In stage 9 chick embryos, Rax2 is expressed in the evaginating optic vesicle and in prospective hypothalamic cells (supplementary material Fig. S1A). At this stage FoxG1 (previously known as BF-1) is expressed in telencephalic cells (supplementary material Fig. S1A). By stage 11, Rax2 expression is restricted to the prospective optic vesicle (Fig. 1B). From this stage onwards, FoxG1 expression is detected in the periphery of the optic vesicle in addition to strong expression in the telencephalon (Fig. 1A; supplementary material Fig. S1B,C). At stage 13, Vsx2 is upregulated, and, in the forebrain, overlapping expression of Rax2 and Vsx2 is detected only in the neural domain of the optic vesicle, whereas Mitf expression is upregulated in the prospective RPE (supplementary material Fig. S1C). At stage 21, Rax2 and Vsx2 continue to be co-expressed only in the neural retina, whereas Mitf is expressed in the RPE (Fig. 1D). In addition, Rax2 is weakly expressed in the tuberal hypothalamus, but Vsx2 and Mitf are not expressed in the forebrain outside the neural retina and RPE, respectively (supplementary material Fig. S1D,E; data not shown). Strong expression of both FoxG1 and Emx2 marks the dorsal telencephalon, but no other regions of the forebrain (Fig. 1; supplementary material Fig. S2). Taken together, these results show that neural retina cells can be distinguished from

Fig. 1. Expression patterns of optic vesicle and anterior forebrain markers. (A,B) Expression patterns of various anterior forebrain markers were analyzed at stage 21 by in situ hybridization on consecutive sections. (A) At stage 21, Rax2 and Vsx2 are expressed in the neural domain, and Mitf expression is restricted to the RPE domain of the optic cup. FoxG1 expression is confined to the dorsal periphery of the optic cup. (B) At stage 21, FoxG1 and Emx2 are strongly expressed in the dorsal telencephalon. Rax2 and Vsx2 are not expressed in the dorsal telencephalon. (C,D) Optic vesicle (OV) explants cultured to approximately stage 21 and analyzed by in situ hybridization on consecutive sections. (C) Stage 9/10 OV explants generated FoxG1+ (25/25) and Emx2+ (25/25) dorsal telencephalic cells, but no Rax2+ (0/25), Vsx2+ (0/25) or Mitf+ (0/25) retinal cells. (D) Stage 13 OV explants generated Rax2+ (15/15) and Vsx2+ (15/15) neural retina cells, and a few FoxG1+ cells in a restricted region (0/15), but no Mitf+ (0/15) RPE cells or Emx2+ (0/15) cells were detected. Scale bars: 100 µm.
other eye and forebrain cell types by the co-expression of Rax2 and Vsx2 from early developmental stages.

The generation of eye-field cells requires inhibition of BMP signals at blastula stages

A recent study in zebrafish has suggested that at blastula to gastrula stages, BMP signals in the anterior neural ectoderm promote telencephalic identity at the expense of eye-field character (Bielen and Houart, 2012). To examine whether this molecular mechanism also acts in amniotes, we isolated late blastula stage (stage 2) medial (M) explants (Fig. 2A; Patthey et al., 2009) and cultured them alone or in the presence of the BMP inhibitor Noggin for 33 h, by which time intact embryos would correspond to approximately stage 10. Thereafter, the explants were processed and analyzed for marker expression on consecutive sections (Fig. 2A; see Materials and Methods for details).

Stage 2 M explants are specified as dorsal telencephalic cells (Patthey et al., 2009). Accordingly, stage 2 M explants cultured alone generated FoxG1+ cells, but no Rax2+ cells (Fig. 2B). By contrast, stage 2 M explants cultured together with Noggin generated Rax2+ and FoxG1+ cells (Fig. 2C). Taken together, these results indicate that: (1) eye-field cells are not specified at the late blastula stage; and (2) BMP activity represses eye-field character at blastula stages and needs to be inhibited in prospective forebrain cells for the generation of Rax2+ eye-field cells.

Isolated eye-field cells switch to dorsal telencephalic identity during culture

To determine whether the specification of eye-field cells and neural retina cells occur in a single or distinct inductive event, we tested whether eye-field cells differentiate into Rax2+ and Vsx2+ neural retina cells in culture. Guided by fate maps (Garcia-Lopez et al., 2009) and optic vesicle morphology (supplementary material Fig. S1A-C; Bell et al., 2001), at the onset (0 h) of culture the majority of stage 9, 10, 11 and 13 explants consisted of Rax2+ cells and a few FoxG1+ cells at one edge of the explants (supplementary material Fig. S4A-D) but no Emx2+ cells (data not shown). Thus, OV explants at stage 9 to 13 consist mainly of eye-field cells.

Unexpectedly, Rax2 expression was not maintained in stage 9, 10 or 11 OV explants, and no Vsx2+ cells or Mitf+ cells were induced when cultured for 46-54 h (~stage 21) (Fig. 1C; data not shown). Instead, stage 9-11 OV explants generated FoxG1+ and Emx2+ cells (Fig. 1C; data not shown), which is characteristic of a dorsal telencephalic identity (Gunhaga et al., 2003; Martynoga et al., 2005; von Frowein et al., 2006) (Fig. 1B). The absence of lens fiber cells in the cultured OV explants was verified by the lack of δ-crystallin+ cells (Pandit et al., 2011; Shinohara and Piatigorsky, 1976; data not shown). Thus, at stage 9-11, isolated and cultured optic vesicle cells acquire a dorsal telencephalic character.

By contrast, stage 13 OV explants generated Rax2+ and Vsx2+ neural retina cells and a few FoxG1+ cells, but no Mitf+ RPE cells or Emx2+ cells after 38-40 h culture (Fig. 1D), implying that neural retina cells are specified at stage 13. Taken together, these data provide evidence that, until stage 13, maintenance of Rax2-positive eye-field identity and induction of neural retina cells requires signals from adjacent tissues.

The generation of retinal cells is dependent on the lens ectoderm and on BMP signals

At stages 10-13, the prospective lens ectoderm lies in close apposition to the developing optic vesicle (supplementary material Fig. S1B,C). To determine whether the lens ectoderm is sufficient to specify retinal cells, we cultured stage 9/10 optic vesicle cells together with prospective lens ectoderm (OVL explants) for 52-54 h (~stage 21). In cultured OVL explants, Rax2+ and Vsx2+ cells, which are characteristic of the neural retina, and δ-crystallin+ cells, which are characteristic of lens fiber cells, were generated in distinct non-overlapping regions of the explants (Fig. 3A). No FoxG1+ and Emx2+ dorsal telencephalic cells (Fig. 3A) or Mitf+ RPE cells (data not shown) were generated. Thus, in the presence of prospective lens ectoderm, optic vesicle cells maintain the expression of Rax2 and acquire neural retina identity.

We and others have previously shown that around stage 11, Bmp4 is expressed in the prospective lens ectoderm (Pandit et al., 2011; Trousse et al., 2001) and that phosphorylated Smad1 (pSmad1) is enriched in the optic vesicle (Belecky-Adams et al., 2002). These data imply that BMP signals emanating from the lens are important for the activation of the BMP pathway in the optic vesicle and the subsequent development of the neural retina. To examine this hypothesis, we cultured stage 9/10 OVL explants in the presence of Noggin. After BMP inhibition, the generation of Rax2+ and Vsx2+ neural retina cells was suppressed and, instead, FoxG1+ and Emx2+ dorsal telencephalic cells were induced (Fig. 3B). Moreover, the spatial organization of the generated HuC/D+ post-mitotic neurons changed from a regularly aligned pattern that is similar to that shown by retina neurons in vivo (Fig. 3C,D), to a scattered telencephalon-like pattern throughout the CNS-derived part of the OVL explants (Fig. 3E,F). In addition, and in agreement with our previous publications (Pandit et al., 2011; Sjödahl et al., 2007), the generation of δ-crystallin+ lens cells was blocked by BMP inhibition (Fig. 3A,B). These results suggest that BMP signals emanating from the prospective lens ectoderm are required for proper
specification of neural retina cells, and in the absence of BMP activity, eye-field cells acquire dorsal telencephalic identity.

Lens-derived BMP signals are required for neural retina development in intact chick embryos

To examine the requirement for BMP signaling during the specification of neural retina cells in vivo, chick embryos were electroporated in ovo in the optic vesicle area (stage 9/10) to transfer a green fluorescent protein (GFP) vector alone or together with a Noggin-expressing vector (Timmer et al., 2002). The electroporated embryos were cultured to approximately stage 15-16, and embryos with GFP staining within the retina region were selected for further analyses. The optic vesicle region of Noggin-electroporated embryos was compared with the corresponding domain of the non-electroporated side as well as control GFP-electroporated embryos.

All control GFP-electroporated embryos and control sides of the Noggin-electroporated embryos exhibited normal morphology of the lens and retina, and a normal expression pattern of Rax2, Vsx2 and FoxG1 (Fig. 4A; supplementary material Fig. S5). By contrast, in embryos with Noggin-electroporated optic vesicles, the prospective retina failed to invaginate and did not form a bilayered structure (Fig. 4B). Moreover, the expression of Vsx2 was lost or severely inhibited, and Rax2 expression was downregulated in the malformed circular-shaped retina, whereas the expression of FoxG1 was expanded (Fig. 4B). In addition, the lens also failed to invaginate and did not form a proper vesicle (Fig. 4B). Thus, BMP activity is required for the early development of the neural retina in vivo.

To evaluate potential sources of BMP signals required for the specification of neural retina cells in vivo, we electroporated the Noggin construct in stage 9/10 chick embryos in different locations: (1) in the prospective lens ectoderm; (2) in the head ectoderm outside of the prospective lens ectodermal region; and (3) in the ventral midline of the forebrain. When the Noggin construct was electroporated in the prospective lens ectoderm, the optic vesicle failed to invaginate and did not form a bilayered structure (Fig. 4B). Moreover, the expression of Vsx2 was lost or severely inhibited, and Rax2 expression was downregulated in the malformed circular-shaped retina, whereas the expression of FoxG1 was expanded (Fig. 4B). In addition, the lens also failed to invaginate and did not form a proper vesicle (Fig. 4B). Thus, BMP activity is required for the early development of the neural retina in vivo.

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electroporated in the ventral midline of the forebrain or in the head ectoderm, no change in eye morphology or expression of Rax2 and Vsx2 in the neural retina was observed (Fig. 4D,E; supplementary material Fig. S5C,D). These data suggest that lens-derived BMP signals are required for the maintenance of eye-field identity and the induction of neural retina cells.

**Direct requirement of BMP activity for the specification of neural retina cells**

To test whether the specification of neural retina cells requires BMP signaling directly in the optic vesicle independently of the lens ectoderm, we used stage 13 OV explants that generate Rax2+ and Vsx2+ neural retina cells during culture in the absence of lens cells (Fig. 1D). Stage 13 OV explants cultured in the presence of Noggin all exhibited the same pattern: in half of the explant, the generation of Rax2+ and Vsx2+ neural retina cells was completely abolished and FoxG1 expression was upregulated (supplementary material Fig. S6); in the other half of the explant, the expression of Rax2 was strongly reduced and was combined with no or weak Vsx2 expression and no detectable FoxG1 expression (supplementary material Fig. S6). Emx2 expression was not detected in the Noggin-exposed stage 13 OV explants (supplementary material Fig. S6). This expression profile suggests that cells in half of the explants acquired the identity of FoxG1+ forebrain cells, although distinct from dorsal telencephalic identity, and cells in the other half acquired an expression profile reminiscent of the tuberal hypothalamus (supplementary material Fig. S1E). Regardless of the acquired cell identity, these results show that ongoing BMP activity is required directly for the specification of neural retina cells.

**BMP activity induces Fgf8 expression in neural retina cells, but FGF activity is not required or sufficient to specify neural retina identity**

FGF signals have been shown to play a role in the development and patterning of the neural retina (Hyer et al., 1998; Martinez-Morales et al., 2005; Pittack et al., 1997), and in chick embryos Fgf8 is upregulated in the medial part of the neural retina at stage 13-14 (Fig. 5A). Therefore, we addressed whether FGF signals act together with BMP activity during the specification of neural retina cells. During culture, stage 9/10 OVL explants generated Fgf8+ cells in the Vsx2+ and Rax2+ domain of the explants (Fig. 5B). By contrast, in the presence of Noggin, Fgf8 expression was inhibited in prospective retinal cells (Fig. 5C). Thus, inhibition of BMP activity in optic vesicle and lens co-cultures suppresses the upregulation of Fgf8 in neural retina cells, consistent with a loss of neural retina identity.

To test whether the loss of Rax2 and Vsx2 expression observed after BMP inhibition is a secondary effect due to downregulated FGF activity, we cultured stage 9/10 OVL explants together with Noggin and FGF8 (250 ng/ml). Addition of FGF8 could not rescue the generation of Rax2+ and Vsx2+ neural retina cells in
inhibited. No cells were cultured to approximately stage 21 and analyzed by stage 13-14 (stages demarcated by black line). (B-E) Stage 9/10 OVL explants dT explants cultured alone generated (Gunhaga et al., 2003) alone or in the presence of BMP4. Stage 9/10 culturing stage 9/10 prospective dorsal telencephalic (dT) explants dorsal telencephalic cell identity and induce retinal character by

Next, we analyzed whether BMP activity is sufficient to suppress retina character. BMP signals can replace the function of the lens ectoderm to induce neural retina cells. (Fig. 5E). Thus, BMP activity is not required or sufficient to induce neural retina cells. Next, we tested whether Wnt activity alone or in combination with BMP signals is sufficient to induce neural retina cells or RPE cells, by culturing stage 9/10 OV explants together with Wnt3A-conditioned medium alone or together with low levels of BMP4. Stage 9/10 OV explants cultured together with Wnt3A still generated FoxG1+ cells (supplementary material Fig. S7). Consistently, stage 13 OV explants cultured alone or in the presence of Frizzled also generated Rax2+ and Vsx2+ neural retina cells, and a few FoxG1+ cells in a restricted region (supplementary material Fig. S8C,D). Thus, Wnt signals are not required for the specification of neural retina cells.

BMP signals can replace the function of the lens ectoderm to suppress dorsal telencephalic cell identity and to promote neural retina character

Next, we analyzed whether BMP activity is sufficient to suppress dorsal telencephalic cell identity and induce retinal character by culturing stage 9/10 prospective dorsal telencephalic (dT) explants (Gunhaga et al., 2003) alone or in the presence of BMP4. Stage 9/10 dT explants cultured alone generated FoxG1+ and Emx2+ dorsal telencephalic cells, but no Rax2- or Vsx2- neural retina cells or Mitf- RPE cells (Fig. 6A). In the presence of BMP4, the generation of FoxG1+ and Emx2+ telencephalic cells was completely inhibited in stage 9/10 dT explants, and instead Rax2+ and Vsx2+ neural retina cells were induced, but no Mitf+ RPE cells were detected (Fig. 6B). Thus, BMP activity is sufficient to suppress dorsal telencephalic cell fate and induce Rax2+ and Vsx2+ neural retina cells.

To further assess whether BMP signals are sufficient to induce neural retina character, stage 9/10 OV explants, which, when cultured in neutral conditions, generate dorsal telencephalic cells (Fig. 2C), were exposed to low levels of BMP4 during culture. Exposure to BMP4 blocked the generation of FoxG1+ and Emx2+ dorsal telencephalic cells, and induced Rax2+ and Vsx2+ neural retina cells, but no Mitf+ RPE cells (Fig. 6C). We next examined whether higher levels of BMP signaling could induce Mitf+ RPE cells. However, even in the presence of a tenfold higher concentration of BMP4, Rax2+ and Vsx2+ neural retina cells, but no Mitf+ RPE cells, were induced (supplementary material Fig. S7). These data show that in isolated eye-field cells, BMP ligands can substitute for the missing lens ectoderm, and provide evidence that low levels of BMP activity are sufficient to induce neural retina identity.

Wnt activity is not required or sufficient to specify neural retina cells, but a combination of Wnt and BMP signals induces RPE cells

Wnt signals have been suggested to be important for the generation of RPE cells (Fujimura et al., 2009; Steinfeld et al., 2013), but whether Wnt activity plays any role in the specification of neural retina cells has not been determined. We therefore first tested whether Wnt activity is required for the specification of neural retina cells, by culturing stage 10 OV explants and stage 13 OV explants together with a soluble Frizzled receptor (Frizzled-conditioned medium) to inhibit Wnt activity (Gunhaga et al., 2003; Hsieh et al., 1999). Stage 10 OV explants cultured alone or in the presence of Frizzled generated Rax2+ and Vsx2+ neural retina cells, whereas no FoxG1+ cells were detected (supplementary material Fig. S8A,B). Consistently, stage 13 OV explants cultured alone or in the presence of Frizzled also generated Rax2+ and Vsx2+ neural retina cells, and a few FoxG1+ cells in a restricted region (supplementary material Fig. S8C,D). Thus, Wnt signals are not required for the specification of neural retina cells.

Next, we tested whether Wnt activity alone or in combination with BMP signals is sufficient to induce neural retina cells or RPE cells, by culturing stage 9/10 OV explants together with Wnt3A-conditioned medium alone or together with low levels of BMP4. Stage 9/10 OV explants cultured together with Wnt3A still generated FoxG1+ and Emx2+ dorsal telencephalic cells, but no Rax2- or Vsx2- neural retina cells or Mitf+ RPE cells (Fig. 7A). By contrast, the combination of Wnt and BMP activity inhibited the generation of FoxG1+ and Emx2+ dorsal telencephalic cells, and induced Mitf+ RPE cells, as well as Rax2+ and Vsx2+ neural retina cells (Fig. 7B). Thus, Wnt activity is not required or sufficient to specify neural retina cells, but a combination of Wnt and BMP signals can induce RPE cells.

DISCUSSION

In the present study, we have analyzed when and by what mechanisms anterior neural cells become specified as eye-field cells and neural retina cells. In summary, our results provide evidence that, around blastula stages, BMP signals repress the eye-field lineage, and that by early neural tube stages, lens-derived BMP activity maintains eye-field identity, inhibits telencephalic character and induces neural retina cells in the forebrain anlage.

Fate maps in chick have shown that prospective telencephalic cells and prospective retinal cells are situated in close proximity in the anterior neural plate (Cobos et al., 2001; Couly and Le Douarin, 1987; Sánchez-Arrones et al., 2009). Previous results in chick suggest that telencephalic cells of dorsal character are already specified by the late blastula stage (Patthey et al., 2009), and that ventral and definitive dorsal telencephalic identity are specified at the gastrula and early neural tube stages, respectively (Gunhaga et al., 2003).
et al., 2000, 2003). Using both retina and telencephalic markers, our results now provide evidence that Rax2-positive eye-field cells become independent of signals from adjacent tissues at stage 13, which coincides with the specification of neural retina cells. Our data show that, prior to this stage, eye-field cells acquire dorsal telencephalic character when cultured alone. Our study and others (Sánchez-Arrones et al., 2009) have shown that Rax2 is expressed in the prospective forebrain at early neural tube stages (stage 9/10 in chick) and appears to be a crucial factor in the cell choice between eye and telencephalic identity. Cell tracing experiments performed in the zebrafish rx3-null mutants have provided evidence that rx3-deficient retinal precursors acquire a telencephalic identity, and embryos exhibit an enlarged telencephalon and lack of eyes (Stigloher et al., 2006). Moreover, a recent study in Xenopus has also shown that Rax mutant embryos are eyeless and that tissue normally fated to form the retina acquires characteristics of the diencephalon and telencephalon instead (Fish et al., 2014). However, although a Rax2-expressing region of the forebrain is devoted to form the neural retina, our results provide evidence that a Rax2+ eye-field state is not sufficient for progression to a neural

![Fig. 6. BMP signals inhibit dorsal telencephalic identity and induce neural retina character.](image)

![Fig. 7. Combined Wnt and BMP signals induce cells of RPE identity.](image)
retina identity. Thus, prior to stage 13, additional signals from neighboring tissues are required for maintaining *Rax2* eye-field identity and inducing neural retina character.

During early neural tube stages, the evagination of the eye-field brings the optic vesicle in close contact with the prospective lens ectoderm. Our results show that optic vesicle cells cultured together with the prospective lens ectoderm maintain the *Rax2* state and upregulate neural retina identity. Moreover, our data suggest that after stage 13, neural retina cells develop independently of signals from the lens ectoderm. In agreement with this, surgical ablation of the lens placode prior to stage 13 resulted in failure of optic cup formation (Hyer et al., 1998), whereas ablation of the lens placode after stage 13 resulted in intact optic vesicles that initiated neural retina differentiation (Hyer et al., 2003). Moreover, in *Pax6* lens-specific mutants, in which lens induction occurs but further lens development is arrested, the differentiation of the neural retina proceeds as normal (Ashery-Padan et al., 2000). These results support our finding that signals from the prospective lens ectoderm are required for maintaining eye-field cells and for inducing neural retina cells.

Our *in vitro* gain and loss of BMP function results provide evidence that BMP signals are both required and sufficient to maintain *Rax2* identity, block dorsal telencephalic character and induce neural retina differentiation. Consistent with this, in retina-specific conditional BMP receptor knockout mice, *Vsx2* failed to be expressed in the mutant optic vesicle and retinal neurogenesis was suppressed (Murali et al., 2005). In addition, our *in ovo* BMP inhibition experiments in various forebrain tissues, suggest that BMP signals from the lens are required for induction of neural retina identity in the optic vesicle. Our finding links the requirement of the lens ectoderm for neural retina specification with the molecular mechanism by which cells in the forebrain become specified as neural retina by BMP activity. Although the optic vesicle and the lens ectoderm are in close contact with each other during early neural tube stages, the adjacent dorsal telencephalon is kept at a distance from the BMP-rich surface ectoderm by intervening neural crest-derived mesenchyme (Fig. 8). The cephalic neural crest cells express the endogenous BMP inhibitors *Noggin* and *Gremlin* (Creuzet, 2009), implying that the cephalic neural crest cells might act as a physical and molecular barrier to protect dorsal telencephalic identity from head-ectoderm-derived BMP signals (Fig. 8). In agreement with this, a recent study in chick has provided evidence that a reduction in *Noggin* and *Gremlin* expression in the cephalic neural crest cells results in abolished *FoxG1* expression in the telencephalon and subsequent microencephaly and partial holoprosencephaly (Aguiar et al., 2014). Our results, showing that BMP signals inhibit telencephalic identity while promoting neural retina character, highlight a new role for BMP signals during the development of the retina that is distinct from previously described roles in dorso-ventral patterning of the retina (Adler and Belecky-Adams, 2002; Kobayashi et al., 2010; Murali et al., 2005) and fate choice between neural versus RPE cells (Hyer et al., 2003; Muller et al., 2007).

It is noteworthy that the lens ectoderm and optic vesicle are so tightly attached that enzymatic treatments are required to dissect the two tissues apart (see Material and Methods). The tight connection can in part be explained by the existence of F-actin-rich filopodia protruding from the lens ectoderm to the prospective neural retina epithelium, which have been shown to be important for proper development of the eye (Chauhan et al., 2009). Specialized filopodia, termed cytonemes, have recently been suggested to function as conduits for morphogen dispersion between morphogen-producing cells and their target tissue (reviewed by Kornberg and Roy, 2014). Thus, existing filopodia might enable the transfer of BMP molecules between the prospective lens and the optic vesicle.

Our results show that, regardless of concentration, BMP activity is not sufficient to induce *Mitf* RPE cells in prospective optic vesicle explants. In addition, optic vesicle and lens co-cultures did not generate any *Mitf* RPE cells, indicating that additional signal(s) from other tissues than the lens ectoderm are required for the specification of RPE cells. A previous finding that BMP4-soaked beads implanted under the optic vesicle at stage 10-12 resulted in pigmentation in the entire optic vesicle (Hyer et al., 2003), does not exclude the possibility that BMP activity in combination with other signals from surrounding tissues induces ectopic pigmented cells. Consistent with this, other studies have suggested that the extra-ocular mesenchyme, which surrounds the prospective RPE domain (Fig. 8), is a source of RPE-inducing signals (Fuhrmann et al., 2000; Kagiyama et al., 2005). Our results suggest that a combination of BMP and Wnt activity is sufficient to induce *Mitf* RPE cells. This is in agreement with previous studies showing that both BMP and Wnt signals are required for the induction of RPE cells, and that neither BMP nor Wnt activity alone is sufficient to induce RPE identity (Steinfeld et al., 2013; Veien et al., 2008).

Our results provide evidence that the specification of neural retina cells is independent of FGF activity. Consistently, the early generation of retina cells appears to be unaffected in *Fgfl*, *Fgf2* and *Fgfl/Fgf2* double-knockout mice (Miller et al., 2000). Furthermore, our results show that inhibition of BMP activity in optic vesicle and lens co-cultures suppresses expression of *Fgfl* in prospective retinal cells, and that simultaneous ectopic addition of FGF8 cannot rescue retinal identity. In agreement with this, in *Bmp4* mouse mutants and in mice that lack *Bmpr1a* and *Bmpr1b* in the retina, *Fgf15* expression is drastically reduced or absent in the optic vesicle (Murali et al., 2005). Moreover, the *Fgf15* expression is restored by the application of BMP4-soaked beads in explanted *Bmp4*−/− optic vesicles (Murali et al., 2005). These results support
our conclusion that BMP signals, but not FGF activity, are required and sufficient to induce cells of neural retina identity.

Our results show that the role BMP signals play for the generation of eye-field cells changes between the blastula and neural tube stages. We find that endogenous BMP activity in neural tissue isolated at blastula stages inhibits Rax2 expression, without being required for FoxG1 expression. This is consistent with a recent study in zebrafish showing that, at late blastula to gastrula stages, BMP activity in the anterior neural ectoderm suppresses eye-field identity through the inhibition of Rx3, a homolog of Rax2 (Bielen and Houart, 2012). In addition, our previous studies at the blastula stage show that ectopic BMP signals induce epidermal character in prospective neural cells (Patthey et al., 2009). Taken together, this suggests that, at early stages, endogenous low levels of BMP signaling in the anterior neural plate are important for protecting prospective telencephalic cells from acquiring an eye-field identity, whereas, at neural stages, BMP signaling from the lens ectoderm promotes neural retina identity at the expense of a telencephalic character.

Studies from 3D stem cell differentiation assays have confusingly suggested that neural stem cells spontaneously generate cells of retinal identity (Eiraku et al., 2011) or cerebral cortical (dorsal telencephalic) character (Lancaster et al., 2013). In light of our model, it is possible that the presence or absence of low levels of endogenous BMP signals in these assays directs the generation of retinal versus cerebral cortical cells. Consistently, in human embryonic stem cell cultures eye-field and neural retina markers are rarely upregulated during conditions of constitutive BMP inhibition, whereas telencephalic markers are significantly enhanced (Lupo et al., 2013). Therefore, any protocols developed for the differentiation of stem cells into retina cells should take into account that in the embryonic forebrain anlage, neural retina cells are specified by BMP signals around optic vesicle stages.

MATERIALS AND METHODS

Embryos
Fertilized White Leghorn chicken eggs were obtained from Strömängs Ägg, Umeå, Sweden. Chick embryos were staged according to the protocol of Hamburger and Hamilton (1951).

Explant assay
Medial (M) explants were isolated from stage 2 chick embryos (Patthey et al., 2009). Prospective optic vesicle (OV) and optic vesicle/lens (OVL) explants were isolated from stage 9, 10, 11 and 13 chick embryos. Dorsal telencephalic (dT) explants were dissected from stage 10 chick embryos (Gunhaga et al., 2003). Explants were cultured in serum-free conditions with or without addition of factors. For details regarding the explant cultures, culture medium and factors, see the methods in the supplementary material.

In ovo electroporation
Stage 9/10 embryos were electroporated in tissues of interest. Noggin-electroporated tissues and the control non-electroporated side were compared with regions electroporated with GFP only. For vectors used and electroporation settings, see the methods in the supplementary material.

In situ hybridization and immunohistochemistry
In situ RNA hybridization and immunohistochemistry were performed essentially as described previously (Wilkinson and Nieto, 1993; Wittmann et al., 2014). Probes and antibodies are described in the methods in the supplementary material.

Statistics
Four to six explants in each of three to six separate experiments were evaluated with respect to the expression of various molecular markers on consecutive sections (supplementary material Fig. S9). Explants in which the majority of cells expressed the marker of interest were counted as positive and explants in which the marker was not expressed at all were counted as negative. The statistics are presented as number of explants with positive expression out of the total numbers of explants analyzed (e.g. 8/10).

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Competing interests
The authors declare no competing or financial interests.

Author contributions
T.P., C.P. and L.G. designed the experiments, performed the explant assays and related statistics. V.K.J. performed the in ovo electroporations and related statistics. All authors analyzed the data and contributed to figure preparation. L.G. wrote the paper with help from C.P. and T.P.

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Supplementary material
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References


Supplementary Figures

Fig. S1.

Expression patterns of optic vesicle and anterior forebrain markers

Expression patterns of various anterior forebrain markers were analysed at stages 9, 11, 13 and 21 by in situ hybridization on consecutive sections.
(A-C) White arrowheads in first column indicate eye-field cells (A) and optic vesicle cells (B, C).

(A,B) At stage 9 and 11, Rax2 is expressed within the evaginating eye-field (black arrowheads) and in prospective hypothalamic cells. FoxG1 expression is confined to prospective telencephalic cells. Neither Vsx2 nor Mitf expression is detected in the forebrain at these stages.

(C) At stage 13, Vsx2 expression is up-regulated in the neural domain of the optic vesicle together with Rax2 expression. Mitf expression is up-regulated in the prospective RPE of the optic vesicle, where weak FoxG1 expression also is detected. Neural and RPE domains indicated by black lines.

(D) At stage 21, FoxG1 is strongly expressed in the ventral telencephalon. Emx2 expression is not detected in the ventral midline of the telencephalon, and weak Emx2 expression is observed above the ventral most region of the telencephalon. Neither Rax2 nor Vsx2 expression are detected in the ventral telencephalon.

(E) At stage 21, Rax2 expression is weakly detected in the tuberal hypothalamus (indicated by brackets). Neither FoxG1, Emx2 nor Vsx2 expressions was detected in this region. Scale bars; 100µm.
Fig. S2.

**Strong FoxG1 and Emx2 expression are detected in the dorsal telencephalon**

Expression patterns of FoxG1 and Emx2 in the forebrain were analysed at stage 21 by in situ hybridization on consecutive sections.

(A) Strong expression of both FoxG1 and Emx2 are detected in the dorsal telencephalon.

(B) Strong expression of FoxG1 is observed in the ventral telencephalon. Emx2 expression is not detected in the ventral midline of the telencephalon, and weak Emx2 expression is observed above the ventral most region of the telencephalon.

(C) Weak expression of FoxG1, but no Emx2 expression, is detected in the dorsal periphery of the optic cup.

(D) Strong expression of FoxG1, but no Emx2 expression, is detected in the pre-optic area.

(E) In the walls of the diencephalon, weak expression of Emx2, but no FoxG1 expression, is observed. Scale bar; 100µm.
Fig. S3

Dissection of OV/OVL explants

(1) Whole head of a stage 10 embryo.

(2) Head divided into two halves and positioned with the optic vesicles up.

(3) Dissection around the optic vesicle with a tungsten needle.

(4) The optic vesicle/lens (OVL) explant is separated from the rest of the forebrain. Note the small pieces in the upper right corner from fine tuning the OVL dissection. OV explants require removal of the prospective lens ectoderm from OVL explants.

Red asterisk marks the optic vesicle region. A – anterior; P – posterior; V – ventral; D – dorsal. Scale bar; 100µm.
Fig. S4.

Expression of *Rax2* and *FoxG1* in OV explants at the onset of culture

Stage 9, 10, 11 and 13 OV explants fixed at 0 hours (=onset of culture) and analyzed by in situ hybridization on consecutive sections.

(A-D) At 0hr of culture, the majority of stage 9 (n=10), 10 (n=10), 11 (n=5) and 13 (n=5) OV explants consisted of *Rax2*⁺ cells, and a few *FoxG1*⁺ cells at one edge of the explants.

Scale bar; 100µm.
**Fig. S5.**

*In ovo GFP electroporation does not affect retinal development*

(A-D) *In ovo* electroporation of stage 9/10 embryos and cultured to stage 15/16, and thereafter analyzed by immunohistochemistry and in situ hybridization on consecutive sections. Electroporation of the prospective retina (B) or the ventral midline (D) and the control non-electroporated sides (A,C).

(A) Control non-electroporated retina with expression of *Rax2* and *Vsx2* in the neural retina, and weak expression of *FoxG1* in the dorsal part of the neural retina.
(B) Prospective retinal cells electroporated with a GFP vector (8/8) did not disturb retinal morphology, and did not result in altered expression of *Rax2, Vsx2* or *FoxG1*.

(C, D) No change in retina morphology or expression of *Rax2* and *Vsx2* was observed after Noggin was electroporated in the ventral midline of the forebrain (n=4). Scale bars; 100µm.
Fig. S6

**BMP requirement for the specification of neural retina cells independent of the lens**

(A, B) Stage 13 optic vesicle (OV) explants cultured to approximately stage 21 and analyzed by in situ hybridization.

(A) Same as Fig. 1D. Stage 13 OV explants generated Rax2⁺ (15/15) and Vsx2⁺ (15/15) neural retinal cells, and a few FoxG1⁺ cells in a restricted region (15/15), but no Mitf⁺ (0/15) RPE cells or Emx2⁺ (0/15) cells were detected.

(B) Stage 13 OV explants cultured together with Noggin generated two domains; cells in one region strongly expressed FoxG1 (15/15), but no Rax2 (0/15), Vsx2 (0/15) or Emx2 (0/15), whereas cells in the other domain expressed reduced levels of Rax2 (15/15), no or weak levels of Vsx2 (15/15), but no FoxG1 (0/15) or Emx2 (0/15). Scale bar; 100µm.
Fig. S7.

High levels of BMP4 do not induce RPE cells

Stage 9/10 R explants cultured to approximately stage 21 and analyzed by in situ hybridization.

In stage 9/10 R explants, BMP4 (35ng/mL) suppressed the generation of FoxG1$^+$ (0/10) telencephalic cells, and induced Rax2$^+$ (10/10) and Vsx2$^+$ (10/10) neural retinal cells, but no Mitf$^+$ (0/10) RPE cells. Scale bar; 100µm.
Fig. S8.

**Wnt activity is not required for the specification of neural retina cells**

(A-D) Stage 10 optic vesicle/prospective lens (OVL) explants, and stage 13 OV explants cultured to approximately stage 21 and analyzed by in situ hybridization on consecutive sections.

(A,B) Both stage 10 OVL explants cultured alone or in the presence of Frizzled generated $Rax2^+$ (25/25 OVL; 10/10 OVL+Fz) and $Vsx2^+$ (25/25 OVL; 10/10 OVL+Fz) neural retinal cells. No or a few weak $FoxG1^+$ (25/25 OVL; 10/10 OVL+Fz) cells were detected.
(C,D) Stage 13 OV explants cultured alone or in the presence of Frizzled generated \( Rax^2^+ \) (15/15 OV; 10/10 OV+Fz) and \( Vsx^2^+ \) (15/15 OV; 10/10 OV+Fz) neural retinal cells, and a few \( FoxG1^+ \) cells in a restricted region (15/15 OV; 10/10 OV+Fz). Scale bar; 100µm.
**Fig. S9.**

**Consecutive sections of a st10 OV explant cultured together with BMP4 for 50 hours**

An example showing all consecutive sections, labelled 1-18, of a st10 OV explant cultured together with BMP4 for 50 hr and analyzed by in situ hybridization for *Rax2, Vsx2, FoxG1* and *Emx2.*
Supplementary Materials and Methods

Explants

The explants were cultured in vitro in collagen in serum-free OPTI-MEM (GIBCO) containing N2 supplement (Invitrogen) and fibronectin (Sigma) to desired time points. To isolate optic vesicle (OV) and dorsal telencephalic (dT) explants, embryos were first incubated in Hanks balanced salt solution (without Ca$^{2+}$, Mg$^{2+}$) (GIBCO) for 3-5 minutes. Then the embryos were transferred to Hanks balanced salt solution (without Ca$^{2+}$, Mg$^{2+}$) containing collagenase (Sigma, 1000u/ml) for 2-4 minutes, followed by washing in Hanks balanced salt solution (with Ca$^{2+}$, Mg$^{2+}$) containing FCS after which they were transferred to L-15 medium (GIBCO). Noggin and control conditioned medium (CM) were obtained from stably transfected or un-transfected Chinese hamster ovary (CHO) cells (Lamb et al., 1993) and cultured in CHO-S-SFM II media (GIBCO). Soluble Wnt3A and control CM were obtained from stably transfected mouse L cells, and soluble Frizzled 8 and control CM were obtained from HEK-293 cells transfected with mFrz8CRD or LacZ reporter construct (Hsieh et al., 1999). The activity of Frizzled CM was tested on proven assays (Patthey et al., 2009). Noggin CM was used at an estimated concentration of 50 ng/ml, and Wnt3A at an estimated concentration of 30 ng/ml. BMP4 (R&D Systems) was used at 3.5-35ng/ml and FGF8 (R&D Systems) was used at 250ng/ml together with 0.5µg/ml heparin (Sigma). SU5402 (Calbiochem) was used at 5µM. Explants cultured in the presence of control CM generated the same combination of cells as explants cultured alone (data not shown).

In ovo electroporation

Vectors used for electroporation were: pCAβ-EGFP-m5 (1 µg/µl) and pMiwIII –Noggin (1 µg/µl) (Timmer et al., 2002). The DNA-constructs were transferred using an Electro Square
Porato ECM 830 (BTX.Inc) by applying 3 pulses (9-18 Volts, 25ms duration,) at 1-s intervals. After electroporation the embryos were cultured in ovo to stage 15-16. Electroporated domains in the Noggin-electroporated embryos were compared to the corresponding region of the non-electroporated side as well as control GFP-electroporated embryos.

**In situ hybridization and Immunohistochemistry**

For the use of in situ RNA hybridization and immunohistochemistry, embryos were fixed in 4% paraformaldehyde (PFA) in phosphate buffered saline (PBS) for 1.5 hours and explants for 25-30 minutes at 4°C. In situ hybridization was performed using the following Dig-labelled chick probes *Emx2, FoxG1* (McCarthy et al., 2001), *Fgf8, Mitf* (Mochii et al., 1998), *Rax2* (Sanchez-Arrones et al., 2009), *Vsx2* (Chen and Cepko, 2000). Antibodies used were: anti-sheep δ-crystallin (Beebe and Piatigorsky, 1981) and anti-mouse HuC/D (Molecular Probes). Nuclei were stained using DAPI (Sigma).
Supplementary References


