BMP4 acts as a dorsal telencephalic morphogen in a mouse embryonic stem cell culture system

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ABSTRACT

The ability of a morphogen – a molecule that specifies two or more cell fates in a concentration-dependent manner – is paradigmatic in developmental biology. Much remains unknown, however, about the existence of morphogens in the developing vertebrate central nervous system (CNS), including the mouse dorsal telencephalic midline (DTM). Bone morphogenetic proteins (BMPs) are candidate DTM morphogens, and our previous work demonstrated BMP4 sufficiency to induce one DTM cell fate – that of choroid plexus epithelial cells (CPECs) – in a mouse embryonic stem cell (mESC) culture system. Here we used BMP4 in a modified mESC culture system to derive a second DTM fate, the cortical hem (CH). CH and CPEC markers were induced by BMP4 in a concentration-dependent manner consistent with in vivo development. BMP4 concentrations that led to CH fate also promoted markers for Cajal–Retzius neurons, which are known CH derivatives. Interestingly, single BMP4 administrations also sufficed for appropriate temporal regulation of CH, CPEC, and cortical genes, with initially broad and overlapping dose-response profiles that sharpened over time. BMP4 concentrations that yielded CH- or CPEC-enriched populations also had different steady-state levels of phospho-SMAD1/5/8, suggesting that differences in BMP signaling intensity underlie DTM fate choice. Surprisingly, inactivation of the cortical selector gene Lhx2 did not affect DTM expression levels, dose-response profiles, or timing in response to BMP4, although neural progenitor genes were downregulated. These data indicate that BMP4 can act as a classic morphogen to orchestrate both spatial and temporal aspects of DTM fate acquisition, and can do so in the absence of Lhx2.

KEY WORDS: Cortical hem, Choroid plexus epithelial cell, ES cell

INTRODUCTION

By definition, a morphogen is an instructive molecule that can specify two or more cell fates in a concentration-dependent manner (Ashe and Briscoe, 2006). In metazoans, morphogens often share ability to act directly on cells at both short- and long-ranges (Grove and Monuki, 2013; Kicheva and Gonzalez-Gaitan, 2008; Tabata and Takei, 2004). Many such morphogens are now well known in invertebrate systems (Kicheva et al., 2007; Porcher and Dostatni, 2010). In vertebrate CNS model systems, classic morphogens are also thought to exist, including Sonic hedgehog (SHH) in the spinal cord, retinoic acid (RA) in the hindbrain, and fibroblast growth factors (FGFs) in the rostral-most telencephalon (Dessaud et al., 2007; Ericson et al., 1997; Schilling et al., 2012; Stamataki et al., 2005; Toyoda et al., 2010). These examples have relied largely on in vitro models, in which potentially overlapping, redundant, or interacting positional systems remain intact and are challenging to eliminate experimentally. In vivo systems, which allow for homogenization of positional information and enable formal testing for morphogen activity, have been more difficult to come by.

In this study we focus on the murine dorsal telencephalon, which consists of the dorsal telencephalic midline (DTM) and bilateral cerebral cortex. The DTM contains two distinctive bilateral structures that are derived from neuroepithelium: the choroid plexus (ChP) and the cortical hem (CH), which are separated at the immediate midline by the choroid plaque. The ChP is a distinctive papillary tissue with ChP epithelial cells (CPECs) that produce the cerebrospinal fluid (CSF) and form the blood-CSF barrier. The CH is a transient junctional tissue between the ChP and cerebral cortex, which functions as a hippocampal organizer (Mangale et al., 2008) and source of Cajal- Retzius (CR) neurons (Meyer, 2010; Molyneaux et al., 2007; Monuki et al., 2001).

Previous studies have implicated BMPs as potential morphogens for these DTM fates. BMPs are produced at high levels in the dorsal telencephalic roof plate of the early neural tube (Furuta et al., 1997), and an intact roof plate is required for the continuous BMP signaling gradient that characterizes the normal dorsal telencephalon (Cheng et al., 2006). This endogenous gradient is also evident from position- and orientation-dependent effects induced by BMP4-soaked beads in explants (Hu et al., 2008). Moreover, an intact roof plate and BMP receptors are required for CH and CPEC specification in mice (Cheng et al., 2006; Currie et al., 2005; Fernandes et al., 2007; Hébert et al., 2002). These studies demonstrate that the dorsal telencephalon possesses a BMP signaling gradient, and that BMP sources and receptors are required for DTM fates.

However, evidence for BMP sufficiency to specify DTM fates, the sine qua non for a morphogen, is more limited. When applied to dissociated cortical progenitors, exogenous BMP4 can modestly upregulate CPEC genes in a concentration-dependent fashion, but does not cause CPEC respecification (Hu et al., 2008). BMP4 also suffices to partially rescue CPEC fate in roof plate-ablated explants (Cheng et al., 2006) and to ectopically induce CPEC fate towards the rostral midline in wild-type explants (Srinivasan et al., 2014). In addition, BMP4 is sufficient to specify CPEC fate from mESC-
derived neuroepithelial cells (NECs) (Watanabe et al., 2012). These mESC-derived CPECs have properties that are indistinguishable from primary CPECs, and, consistent with in vivo and experimental studies (Thomas and Dziadek, 1993), CPEC competency is restricted to preneurogenic NECs rather than later-stage neurogenic radial glia. Likewise, the critical period for CH fate determination [embryonic day (E)8.5-10.5] maps to the preneurogenic NEC period (Mangale et al., 2008). The mESC culture system therefore provides an ideal in vitro model system to examine BMP4 sufficiency to induce CH in addition to CPEC fate. If consistent with in vivo development, CH cells should be specified at lower BMP4 concentrations than those required for CPEC fate.

In this paper, we utilize a modified mESC culture system to demonstrate BMP4 concentration dependence for CH and CPEC fates in vitro, consistent with in vivo development. In the modified system, aggregated mESC-derived NECs are dissociated into monolayers to allow for uniform exposure to exogenous BMP4. Importantly, single administrations of BMP4 also induce temporal patterns in CH and CPEC gene expression that mimic in vivo development, including sharpening of dose-response profiles over time. Thus, BMP4 alone can specify not only multiple DTM fates, but also appropriate temporal patterning of DTM gene expression.

RESULTS

BMP4 concentration-dependent induction of CH and CPEC markers in a modified mESC culture system

We first used our existing mESC aggregate cultures (Watanabe et al., 2012) to look for CH marker induction across an extensive range of BMP4 concentrations (0.1-512 ng/ml; Fig. S1B, M1 mESC line). In our previous study (Watanabe et al., 2012), we used five different CPEC markers that displayed similar BMP4 dose-response profiles (Tir, Msx1, Agp1, Cldn1, and Lmx1a) and confirmed CPEC identity by ultrastructural analyses and functional assays. Since Tir expression is particularly high and exclusive to CPECs, Ttr (or TTR::RFP) was used to follow CPEC differentiation here. In contrast to CPECs, CH cells should be negative for Ttr while being positive for Msx1, Lmx1a, and Wnt3a (Fig. 1A); however, we did not find a BMP4 concentration that induced a CH-like profile in the aggregate system, as all three genes displayed similar dose-response profiles (Fig. S1B). One potential explanation for this is that cells in aggregates are exposed to widely varying extracellular BMP4 concentrations depending on their radial positions, a natural limitation of aggregates in general.

We therefore developed a monolayer culture system to expose cells to more uniform BMP4 concentration. In this system, after 5 days in vitro (DIV) of neural induction, aggregates were dissociated into single cells and plated onto an adherent surface, then exposed to exogenous BMP4 at varying concentrations (0.15-150 ng/ml) for an additional 5 DIV (Fig. 1A). Using two different mESC lines, we found that, unlike the aggregate-only system, moderate BMP4 concentrations (0.5-5 ng/ml for M2 and 1.5-5 ng/ml for M1 cells) induced a CH-like profile, with relatively high Wnt3a, Lmx1a, and Msx1 compared to Ttr; this was particularly clear for the M2 line (Fig. 1B). In addition, at the highest BMP4 concentrations, CH-specific Wnt3a was strongly suppressed, while the Ttr was upregulated and high Msx1 was maintained, consistent with a CPEC profile (Fig. 1B). In the low BMP4 range (<0.15 ng/ml for M2 and <0.5 ng/ml for M1), DTM markers were not detectably induced; therefore, this range may correspond to the lower BMP signaling levels seen in the developing cortex (Fig. S1A) (Cheng et al., 2006). Thus, both CH and CPEC gene expression profiles could be induced in a BMP4 concentration-dependent manner in the same monolayer cultures.

To confirm CH identity at the single cell level, immunocytochemical analyses were performed. Based on the markers used, CH cells should express Lmx1a and Msx1, but not TTR::RFP. We found that, in the low BMP4 range (0 or 0.15 ng/ml), very few cells were positive for Msx1, Lmx1a, or TTR::RFP after 7 DIV (5 DIV aggregate plus 2 DIV monolayer) or 10 DIV (5 DIV aggregate plus 5 DIV monolayer; Fig. 1C-D,G-H), similar to the RT-qPCR studies (Fig. 1B). At moderate BMP4 concentration (1.5 ng/ml), Lmx1a/Msx1 double-positive cells were observed at 7 DIV (18.25±5.85%) and 10 DIV (13.83±1.23%), but few TTR::RFP-expressing cells were detected (Fig. 1E,I,K), which corresponds well to the CH gene expression profile (Fig. 1A). In cultures with the highest BMP4 concentration (15 ng/ml), Lmx1a/Msx1 double-positive cells were detected at 7 DIV (17.83±8.14%), but not at 10 DIV, while TTR::RFP expression displayed the converse pattern, low at 7 DIV (0.42±0.42%) but prominent at 10 DIV (12.00±6.46%) (Fig. 1F,J,K), with the TTR::RFP-expressing cells displaying a more flattened appearance typical of CPECs and other epithelial cells in culture (Fig. S2I-L). CPEC yield was approximately three to four times higher in this modified monolayer culture system compared to the aggregate system. Thus, the 10 DIV cultures most clearly distinguished the CH cells and CPECs, and further supported the BMP4 concentration dependence (1.5 ng/ml BMP4 for CH, 15 ng/ml BMP4 or higher for CPEC) seen by population RT-qPCR analysis (Fig. 1B). Interestingly, CPEC induction at 15 ng/ml BMP4 was preceded at 7 DIV by a more CH-like molecular phenotype; this raises the possibility that CPEC development involves an intermediate CH-like stage, a point that is further addressed below.

Co-enrichment for CH and Cajal-Retzius (CR) markers at moderate BMP4 concentration

In addition to being a hippocampal organizer (Mangale et al., 2008), the CH also generates CR neurons (Molyneaux et al., 2007). To further evaluate CH induction in our system we examined CR neuron markers. While Reelin-expressing CR cells have multiple origins in addition to the CH (Meyer, 2010; Molyneaux et al., 2007), p73 expression is restricted to CH-derived CR neurons and precedes Reelin expression (Meyer et al., 2002).

We therefore examined p73 and Reelin expression, and their temporal expression patterns over a longer time period (6-16 DIV). At 1.5 ng/ml BMP4 – the concentration yielding maximal CH marker expression (Fig. 1) – p73 and Reelin inductions were detected by 10 DIV (Fig. 1L). The p73 expression plateau was reached by 10 DIV, whereas the plateau for Reelin expression was not reached until 12 DIV (Fig. 1L). In contrast, at high BMP4 concentration (15 ng/ml), neither p73 nor Reelin expression increased above baseline levels during the 6-16 DIV period (Fig. S3A). Thus, the BMP4 dose-response profile for CR neuron markers matched that of the CH rather than CPECs. Moreover, the observed temporal profiles for p73 and Reelin are consistent with those seen in vivo for CH-derived CR cells (Meyer et al., 2002).

Temporal regulation of DTM markers in a BMP4 dose-dependent fashion

We then focused on other temporal aspects of DTM gene expression following a single application of BMP4, since in addition to determining spatial patterns, morphogens are thought to determine temporal aspects of patterning in vivo (Dessaud et al., 2007; Tozer et al., 2013). M2 5-day SFEBq aggregates were dissociated and plated with varying concentrations of BMP4 (0.15-15 ng/ml), with gene expression examined after 1-5 DIV (6-10 DIV total). RT-qPCR assays revealed that CH genes Wnt3a and Lmx1a were
induced quickly at both moderate and high BMP4 concentrations (0.15-15 ng/ml) within 1-3 DIV. Over time, these genes were maintained at the moderate BMP4 concentrations, but suppressed at the high ones (Fig. 2A,B). These temporal- and concentration-dependent patterns coincide with those seen in the immunocytochemical analyses (Fig. 1K).

Different temporal patterns were observed for Msx1 and CPEC-specific Tr. Msx1 was not only induced, but also maintained at moderate and high BMP4 (0.5-15 ng/ml) (Fig. 2C), consistent with Msx1 being a positional determinant of high BMP signaling (Cornell and Ohlen, 2000; Ramos and Robert, 2005) and its maintenance in adult CPECs in vivo (Ramos et al., 2004), unlike
intracellular signaling, we measured phospho-Smad1/5/8 (pSmad) and high (CPEC-inducing; 64 ng/ml) BMP4 concentrations on the levels and dynamics of Smad1/5/8 activation in our culture of Smad1/5/8 (Massague et al., 2005), leading us to wonder about the two cell fates.

Differential steady-state levels of BMP signaling correlate with the two cell fates

BMP signaling is directly transduced into the tail phosphorylation of Smad1/5/8 (Massague et al., 2005), leading us to wonder about the levels and dynamics of Smad1/5/8 activation in our culture system. To examine the effects of moderate (CH-inducing; 1.5 ng/ml) and high (CPEC-inducing; 64 ng/ml) BMP4 concentrations on intracellular signaling, we measured phospho-Smad1/5/8 (pSmad)

levels by western blot over time in 5-day SFEBq dissociated cells. At all time points examined, actin-normalized pSmad levels were higher in the cells treated with higher BMP4 (Fig. S3B). As in primary cortical progenitors (Hu et al., 2008), pSmad activation occurred rapidly in response to BMP4 (by 60 min, the earliest time point in these studies) and preceded the induction of DTM mRNAs (Fig. 2A, B). At later time points, pSmad levels remained elevated in cells treated with high BMP4 (Fig. S3B), although this could reflect excess BMP4 in the media rather a difference in cell biology per se (i.e. with high BMP4 concentrations in media intracellular BMP degradation could be limiting, whereas BMP4 availability could be the limiting factor at lower BMP4 concentrations). Regardless, the findings suggest that intracellular pSmad levels correlate positively with extracellular BMP4 concentration in this system.

BMP4 regulates neural progenitor markers in a dose-dependent fashion

In addition to DTM fates, BMP signaling has been implicated in regulating cortical progenitors, and specifically, their expression of transcription factors (TFs). Within the E10.5-E12.5 cortical anlagen, BMP/pSmad signaling exhibits a high dorsal-low ventral (DV) gradient, the same DV polarity as the gradients for TFs Lhx2 and Enmx2, while having the opposite polarity of the high ventral-low dorsal (VD)-graded TFs Pax6, Foxg1, and Ngn2 (Cheng et al., 2006). After reducing and flattening the BMP/pSmad gradient via roof plate ablation, the cortical TFs were differentially affected; the DV gradients (Lhx2 and Enmx2) were also reduced and flattened, while the VD gradients (Pax6, Foxg1, and Ngn2) were relatively unaffected (Cheng et al., 2006). This raised the possibility that among the cortical TFs examined, BMP/pSmad signaling selectively upregulates Lhx2 and Enmx2.

To address this potential selectivity, we examined the same DV- and VD-graded genes in our monolayer culture system using the M1 and M2 mESC lines. These analyses showed that the DV markers, Lhx2 and Enmx2, were upregulated at intermediate BMP concentrations, but interestingly, with different timing (Fig. 3A,B, best seen in Fig. 3F,G). Lhx2 activation occurred earlier than Enmx2, consistent with Lhx2 being upstream of Enmx2 and other cortical transcription factors (Mangale et al., 2008). At these same intermediate BMP4 concentrations there were mixed DV gene patterns. Pax6 was not consistently altered (Fig. 3C,H), Foxg1 was slightly downregulated (Fig. 3D,I), and Ngn2 was upregulated (Fig. 3E,J); generally high BMP4 concentrations downregulated all DV and VD markers. These results support the hypothesis that BMP signaling selectively upregulates DV markers, Lhx2 and Enmx2, at intermediate concentrations and can account for the Lhx2 and Enmx2 findings in roof plate-ablated mutants (Cheng et al., 2006) as well as the Lhx2 upregulation at a distance away from BMP4-soaked beads in cortical explants (Monuki et al., 2001). However, there were inconsistent effects on VD genes that cannot alone explain the DV gene expression phenotypes in these same mutants.

Lhx2 inactivation does not facilitate BMP4-mediated DTM induction in culture

We then studied the influence of the Lhx2 cortical selector gene on DTM induction by BMP4. Lhx2 is expressed by cortical progenitor cells (hippocampal and neocortical), but not by CPECs or CH cells, and constitutive Lhx2 absence results in excessive CH and CPEC fates (Monuki et al., 2001). This DTM fate suppression by Lhx2 is also seen in genetic mosaics, with Lhx2 null cells in the hippocampal primordium adopting CH, but not CPEC, fate in cell-autonomous fashion (Mangale et al., 2008). One possibility for the

Lhx2 (Zou et al., 2009). In contrast, Tr markers were relatively low at early time points and increased over time with the highest levels occurring at the highest BMP4 concentrations tested (Fig. 2D), consistent with Tr being a relatively late CPEC marker in vivo compared to Mxi1 and Lmx1a (Curtin et al., 2005; Hu et al., 2008). Similar temporal- and concentration-dependent expression patterns for these DTM markers were observed using a different mESC line (Fig. S4). Interestingly, the concentration-response profiles also sharpened (i.e. the slopes between points steepened) with increasing time. Since this sharpening occurs in monolayer cultures (rather than intact tissues with localized signaling sources and other positional cues), it raises the possibility of cell-intrinsic ultra-

sensitivity to BMP4 similar to that seen in primary cortical progenitors (Hu et al., 2008; Srinivasan et al., 2014). Taken together, these data indicate that a single BMP4 dose in the mESC-based system was sufficient for temporal DTM gene patterning and sharpening that recapitulates in vivo development.
ectopic CH fate is increased BMP signaling intensity in Lhx2 null cells, but pSmad and BRE-gal studies in Lhx2 mosaic embryos argue against this (Doan et al., 2012). Another possibility is that Lhx2 null cells are intrinsically biased to generate DTM fates even at lower BMP concentration and signaling intensity.

We explored this possibility here by performing BMP4 dose-response experiments after Lhx2 inactivation in cultured mESCs. Using M1 cells (R26CreER/+Lhx2cKO/cKO; Fig. S5A) (Watanabe et al., 2012), we developed a highly efficient and dose-dependent method for inactivating Lhx2 using 4-hydroxytamoxifen (4HT), with maximal inactivation achieved at and above 1 μM 4HT (Fig. S5B). 4HT addition before 4 DIV adversely affected neuroepithelial induction (data not shown). We therefore applied 4HT at 4 DIV (Fig. 4A). At 5 DIV SFEBq aggregates were dissociated, plated as monolayers with varying amounts of BMP4 (0.15-150 ng/ml), then analyzed after 2 or 5 DIV (7 or 10 DIV total). In these cultures Lhx2 was effectively inactivated at the genomic (Fig. S5C) and mRNA levels (96.47±2.92% reduction; Fig. S5D). Interestingly, the DTM gene profiles at both 7 and 10 DIV were quite similar between control and 4HT-treated cells (Fig. 4, solid and dashed lines, respectively) with no evidence of a left-shift or altered expression levels in the 4HT-treated cells (Fig. 4B,C). We performed similar studies with Lhx2 inactivation at 5 DIV, and again saw no obvious effects on the levels, timing, or concentration-dependence of the DTM genes to BMP4 despite highly efficient Lhx2 inactivation (data not shown). Thus, Lhx2

![Graphs](https://example.com/graphs.png)

Fig. 3. BMP4 mediated-neural progenitor marker regulation in a dose-dependent fashion. (A-J) 5-day SFEBq aggregates are dissociated and plated in monolayer with varying BMP4 (0.15-15 ng/ml for M2 and 0.15-150 ng/ml for M1) for another 1-5 DIV (6 to 10 DIV total). Usually, at higher BMP4 concentration, all neural progenitor expression is suppressed. Selectively, some neural progenitor markers are slightly upregulated in a temporal manner at lower concentration of BMP4. Data are presented as mean±s.e.m.; *P<0.05 compared to no BMP culture.
inactivation did not influence BMP4-mediated DTM induction in this mESC culture system.

Since BMP4 can also regulate DTM genes in primary cortical progenitors (Hu et al., 2008), we examined BMP4 concentration-response profiles in primary cortical progenitors lacking Lhx2 (Emx1Cre;Lhx2cKO/sKO; Mangale et al., 2008). In Lhx2 null E11.5 progenitors, Lhx2 mRNA levels were reduced by 81.76±9.05% compared to controls before plating (Fig. S5E). However, this Lhx2 reduction did not further promote DTM gene expression at any BMP4 concentration (Fig. S5F). Lhx2 inactivation therefore did not facilitate BMP4-mediated DTM induction.

Inactivation of Lhx2 downregulates other cortical progenitor markers

As a cortical selector gene, Lhx2 is at or near the top of the transcriptional hierarchy that specifies neuroepithelial cells with cortical identity (Mangale et al., 2008; Monuki et al., 2001). We
therefore asked whether Lhx2 inactivation (by 4HT addition at 4 DIV) influenced BMP4-mediated regulation of other cortical progenitor genes. These experiments revealed two consistent gene expression patterns by RT-qPCR analyses: first, at low to intermediate BMP4 concentrations (0-1.5 ng/ml), Lhx2 inactivation caused reductions for all of the other genes (Emx2 and VD genes) at both 7 and 10 DIV (Fig. 5A-J). This supports the view that Lhx2 is upstream of these cortical progenitor genes. Second, at both 7 and 10 DIV, Lhx2 inactivation had no effect on the suppression of Emx2, Foxg1, or Ngn2 at high BMP4 concentrations, but interestingly, abrogated the suppression of Pax6 (Fig. 5A-J). Thus, Lhx2 was not required for BMP4-mediated suppression of Emx2, Foxg1, or Ngn2, but was required for that of Pax6. Together, these data show that Lhx2 inactivation reduces cortical progenitor gene expression, but does not interfere with BMP4-mediated down-regulation of these genes, with the exception of Pax6.

**DISCUSSION**

In this study two DTM fates, CH and CPEC, were induced from mESC-derived neuroepithelial progenitors, with a single BMP4 dose sufficing to recapitulate in vivo spatial patterning. In addition, single BMP4 doses sufficed to recapitulate the normal temporal patterning of DTM gene expression, as well as some aspects of selective cortical TF regulation seen in vivo. Taken together, the results from a reduced in vitro system, which minimizes confounds inherent in in vivo studies, provide evidence for BMP4 activity as a classical morphogen in the dorsal telencephalon. Interestingly, inactivation of Lhx2, which leads to excessive DTM fates in vivo, did not affect the levels, timing, or dose-response profiles of...
BMP4-mediated DTM marker induction in vitro; however, Lhx2 inactivation downregulated cortical TF gene expression overall and abrogated BMP4-mediated suppression of Pax6. While consistent with Lhx2 being a cortical selector gene (Mangale et al., 2008), these studies suggest that Lhx2 neither mediates nor regulates the fundamental morphogenic activities of BMP4 during dorsal telencephalic development.

The mouse dorsal telencephalon as a classic BMP morphogen gradient system
CH inducibility by BMP4 complements previous findings on the induction (Srinivasan et al., 2014; Watanabe et al., 2012) or rescue of CPECs (Cheng et al., 2006). Previous unsuccessful attempts to induce CH fate with BMP4 (Furuta et al., 1997; Hu et al., 2008; Monuki et al., 2001) may relate to the early and short competency period for CH fate in vivo (E8.5-E10.5 in mice; Mangale et al., 2008). Using dissociated neuroepithelial aggregates BMP4 also recapitulated the concentration (CH fate at lower BMP4 concentration than CPEC, Fig. 1B) and temporal profiles (CH fate before CPEC fate, Figs 1 and 2) expected for a DTM morphogen. Unfortunately, CH grafts to test for hippocampal organizer activity have been challenging (Tole and Grove, 2001), due in part to the early critical period (Mangale et al., 2008), and further grafting or coculture advancements will be needed to further assess the derived CH cells. In addition, BMP4-mediated cell death and proliferation were not addressed here. CPECs and CH cells differentiate in domains of low proliferation and high death (Furuta et al., 1997; Currie et al., 2005), BMP4 can induce these fates (Furuta et al., 1997; Cheng et al., 2006; Watanabe et al., 2012), and BMP4 dose-response curves for these fates overlap (Mabie et al., 1999). Thus, proliferation and death effects are known and expected, and relevant BMP-mediated fates in our system, which are not accounted for in the normalized RT-qPCR data, reflect ‘per cell’ averages of viable populations.

Importantly, together with previous studies, our findings establish the mouse dorsal telencephalon as a classic morphogen gradient system (Grove and Monuki, 2013). In this system, BMP4 and other BMPs produced at the midline (Furuta et al., 1997; Monuki et al., 2001) lead to a continuous BMP/Smad signaling gradient (Cheng et al., 2006) with a length scale of 270-290 μm at E10.5 (Srinivasan et al., 2014). This gradient also gives rise to graded and oriented responses around BMP4-soaked beads (Hu et al., 2008). BMP-producing cells (Cheng et al., 2006; Currie et al., 2005) and BMP receptors (Hébert et al., 2002) are also required for DTM fates. The ability of BMP4 to induce two DTM fates (CPECs and CH cells) represents a final piece of evidence for the classic model.

An additional point worth noting is the apparent CH-like intermediate state during CPEC differentiation, i.e. CH markers Wnt3a and Lmx1a were initially induced then extinguished at CPEC-inducing BMP4 concentrations (Figs 1 and 2). This concept is consistent with Wnt3a and Lmx1a expression in vivo, as well as Wnt3a genetic lineage studies (Chizhikov et al., 2010; Louvi et al., 2007). While potentially consistent with temporal integration of BMP signaling, BMP4 time-response data were not clear in this regard (data not shown). A CH-like intermediate state would also complicate previous CH-CPEC lineage analyses and questions about CPECs being hem derivatives. Regardless, a possible CH-like intermediate state will be important to account for in future CPEC studies and models.

Selective BMP regulation of cortical patterning
Overall, our findings align well with previous BMP cortical gene studies. High BMP4 concentrations (i.e. the DTM regime) consistently suppressed the five cortical TF genes tested at both 7 and 10 DIV (Fig. 3), consistent with previous studies on primary cortical progenitors (Cheng et al., 2006; Hu et al., 2008), in vivo gene expression (e.g. Furuta et al., 1997; Currie et al., 2005), and roof plate or BMP receptor ablation phenotypes (Cheng et al., 2006; Hébert et al., 2002). Thus, high BMP consistently suppresses cortical gene expression and cortical fate.

Perhaps more interestingly, intermediate BMP4 concentrations (i.e. the dorsomedial cortex regime) upregulated DV-graded Emx2 and Lhx2, but not VD-graded genes Pax6 or Foxg1 (Fig. 3), which is concordant with the in vivo phenotypes of these four genes following roof plate ablation (Cheng et al., 2006). The concordant findings for Emx2 and Lhx2 are also consistent with the BMP-responsive enhancer in the Emx2 gene (Theil et al., 1999) and with Lhx2 upregulation at a distance from BMP4-soaked beads in explants (Monuki et al., 2001). However, Ngn2 findings are discordant; Ngn2 was upregulated by intermediate BMP4 here (Fig. 3), but unaffected by roof plate ablation (Cheng et al., 2006). Collectively, these findings support a model in which intermediate BMP signaling upregulates Emx2 and Lhx2 in dorsomedial cortex/hippocampal anlagen, thereby generating their DV gradients, while VD genes must rely on signals other than BMPs to sculpt their gradients (Cheng et al., 2006). (Note: the BMP4-mediated Ngn2 upregulation seen here would create an Ngn2 gradient of the wrong polarity, i.e. it would have DV rather than the VD polarity seen normally.)

Lhx2 independence of BMP morphogenetic activity
Lhx2 inactivation had no demonstrable effect on any aspect of BMP4-mediated CH and CPEC fate acquisition (levels, dose-response profiles, or kinetics). Thus, Lhx2 neither mediates nor regulates the DTM morphogenetic activities of BMP4. While the in vitro data are clear, they are perplexing given the DTM-related phenotypes of Lhx2 null mice. Constitutive Lhx2 null mice have excess CH and CPECs (Monuki et al., 2001), and mosaic Lhx2 inactivation in dorsomedial cortex leads to ectopic CH (Mangale et al., 2008), indicating that Lhx2 genetically suppresses DTM fates, but does not do so by regulating BMP signaling intensity (Doan et al., 2012).

As such, an explanation for the different Lhx2 null phenotypes will require further exploration. Spatial and temporal differences in Lhx2 inactivation could account for the differences, since early in vitro inactivations (more analogous to the constitutive Lhx2 state) were not possible here due to poor neuroepithelial induction, which is not apparent in vivo (Mangale et al., 2008; Monuki et al., 2001). This alone suggests important qualitative differences between the in vitro and in vivo systems. Positional determinants lost in culture could also be responsible, since Lhx2 null fates in vivo critically depend on position within the dorsal telencephalon (Mangale et al., 2008).

Lhx2 selector activity and BMP-mediated cortical patterning
In vivo, Lhx2 acts at or near the top of the genetic hierarchy for selecting cortical identity (Mangale et al., 2008), and our in vitro findings here are consistent with this concept. At intermediate BMP4 concentrations, Lhx2 upregulation (at 7 DIV) preceded the upregulations of other cortical genes (Emx2 and Ngn2 at 10 DIV) in both mESC lines (Fig. 3), and all cortical markers were reduced upon Lhx2 inactivation (Fig. 5). Abrogation of BMP4-mediated Pax6 suppression due to Lhx2 loss (Fig. 5) is also consistent with previous Lhx2-Pax6 epistasis studies (Hou et al., 2013; Mangale et al., 2008; Porter et al., 1997; Tetreault et al., 2009). Interestingly, published examples indicate that Lhx2 can regulate Pax6 either positively or negatively, highlighting the importance of Lhx2-Pax6 context. Other than Pax6, however, Lhx2 presence or absence had
no apparent effect on BMP4-mediated regulation of the other cortical genes. As with DTM fates, this implies that BMP morphogenetic activity in the dorsal telencephalon is largely Lhx2-independent; however, BMP consistently regulates Lhx2 expression (suppression at high concentrations, upregulation at intermediate ones), suggesting a BMP-Lhx2 pathway with little to no feedback.

**MATERIALS AND METHODS**

**Mice**

Mouse colonies and breeding were performed under Institutional Animal Care and Use Committee guidelines. Noon of the vaginal plug date was designated as day 0.5 for timed pregnancies. Crown-rump length was measured to verify embryonic ages. Wild-type mice (CD1) were obtained from Charles River Laboratories (Wilmington, MA) and *Emx1<sup>F<sup>Cre</sup>/E<sup>RFP</sup></sup>* were derived as described (Mangale et al., 2008).

**Mouse ESC line derivation and expansion**

The lines M1 (*R26<sup>CreER</sup>/E<sup>RFP</sup>*, C57BL/6J with minor CD1 background), M2 (*Tr:RFP* hemizygous, mostly CD1 background with C57BL/6J and ICR), and M4 (*R26<sup>CreER</sup>/E<sup>RFP</sup>*, C57BL/6J with minor CD1 background) were derived, cultured and verified for pluripotency, chromosomal numbers and mycoplasma negativity as described (Watanabe et al., 2012). The M2 line was derived by the UCI Transgenic Mouse Facility using the 2i method (Li et al., 2008). Prior to all experiments, frozen mESCs were cultured for at least two passages before thawing. Experiments were conducted on cells between passage numbers 11 and 40.

**Mouse ESC culture and differentiation**

Neural differentiations in 'SFEbq' aggregates were performed as described (Eiraku et al., 2008; Watanabe et al., 2012). For the 'aggregate' method, cells were maintained as aggregates for the entire culture period. After 5 DIV, differentiation media (Eiraku et al., 2008) was replaced with fresh media containing 0.5-150 ng/ml BMP4 (R&D Systems, Minneapolis, MN, USA) for another 5 days (10 days total). For the 'monolayer' method, 5-day aggregates were dissociated to single cells using TrypLE Express (Life Technologies, Carlsbad, CA, USA) and plated onto PDL/laminin-coated plates at 2.5-5×10<sup>5</sup> (M1) or 7-10×10<sup>5</sup> (M2) cells/cm<sup>2</sup> with BMP4 in fresh differentiation media for 5 days. Thus, for both the aggregate and monolayer methods, total culture time was 10 days, and BMP4 exposure duration was 5 days. BMP4 was applied only once in each experiment.

**Primary cortical progenitor culture**

For *Emx1<sup>F<sup>Cre</sup>/E<sup>RFP</sup></sup>* studies (Mangale et al., 2008), E11.5 cortical progenitor cells were dissected and cultured as described (Hu et al., 2008), except that BMP4 was added during cell resuspension prior to plating.

**RT-qPCR and immunostaining**

RT-qPCR was performed as described (Currie et al., 2005; Hu et al., 2008) on LightCycler® 480 System (Roche, Indianapolis, IN, USA) using 18S normalization, Microsoft Excel for statistical tests (two-tailed t-tests assuming equal variance, asterisks for *P* values <0.05), and KaleidaGraph (Synergy Software) for graphing. All primers and amplicons were validated (Synergy Software) for graphing. All primers and amplicons were validated (Supplementary information available online at http://bio.biologists.org/lookup/doi/10.1242/bio.012021.supplemental)

**Cell quantification**

Lmx1a, Msx1/2, Tr:RFP, and Hoechst were manually counted in Photoshop or ImageJ from two biological replicates and four different confocal planes in a blinded manner. Denominators included all Hoechst-stained cells. For blinded scoring of immunostains, matched unprocessed confocal images (Zeiss LSM 510) were scored. 1000-2000 cells were counted for each condition. Excel was used for statistical tests (two-tailed t-tests assuming equal variance, asterisks for *P* values <0.05).

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**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

M.W., E.S.F. and E.S.M. developed the concepts and approach. M.W., E.S.F., F.B.C., J.S.W., and M.C. performed experiments. M.W. and E.S.F. analyzed data. M.W., E.S.F., and E.S.M. developed the concepts and approach. M.W., E.S.F., and E.S.M. developed the concepts and approach. M.W., E.S.F., and E.S.M. developed the concepts and approach. M.W. and E.S.F. prepared and edited the manuscript prior to submission.

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**Supplementary information**

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