https://doi.org/10.1093/procel/pwad036 Advance access publication 10 June 2023 Research Article

Protein & Cell

Research Article

BMP7 expression in mammalian cortical radial glial cells increases the length of the neurogenic period

Zhenmeiyu Li^{1,‡}, Guoping Liu^{1,‡}, Lin Yang¹, Mengge Sun¹, Zhuangzhi Zhang¹, Dejun Xu¹, Yanjing Gao¹, Xin Jiang¹, Zihao Su¹, Xiaosu Li¹, Zhengang Yang¹, D

¹State Key Laboratory of Medical Neurobiology and MOE Frontiers Center for Brain Science, Institutes of Brain Science, and Department of Neurology, Zhongshan Hospital, Fudan University, Shanghai 200433, China

⁺These authors contributed equally.

*Correspondence: yangz@fudan.edu.cn (Z. Yang)

Abstract

The seat of human intelligence is the human cerebral cortex, which is responsible for our exceptional cognitive abilities. Identifying principles that lead to the development of the large-sized human cerebral cortex will shed light on what makes the human brain and species so special. The remarkable increase in the number of human cortical pyramidal neurons and the size of the human cerebral cortex is mainly because human cortical radial glial cells, primary neural stem cells in the cortex, generate cortical pyramidal neurons for more than 130 days, whereas the same process takes only about 7 days in mice. The molecular mechanisms underlying this difference are largely unknown. Here, we found that bone morphogenic protein 7 (BMP7) is expressed by increasing the number of cortical radial glial cells during mammalian evolution (mouse, ferret, monkey, and human). *BMP7* expression in cortical radial glial cells promotes neurogenesis, inhibits gliogenesis, and thereby increases the length of the neurogenic period, whereas Sonic Hedgehog (SHH) signaling promotes cortical gliogenesis. We demonstrate that BMP7 signaling and SHH signaling mutually inhibit each other through regulation of GLI3 repressor formation. We propose that *BMP7* drives the evolutionary expansion of the mammalian cortex by increasing the length of the neurogenic period.

Keywords radial glia, cortical neurogenesis, cortical gliogenesis, cortical evolution, BMP7, SHH

Introduction

Mammalian cortical radial glial (RG) cells are the source for all cortical pyramidal neurons (PyNs), most cortical glial cells, and a subpopulation of olfactory bulb interneurons (Rakic, 2006; Kriegstein and Alvarez-Buylla, 2009; Molnar et al., 2019; Lin et al., 2021). Three types of RG cells drive human cortical development; they are ventricular zone (VZ) full span radial glia (fRG), VZ truncated radial glia (tRG), and outer radial glia (oRG) which lie in the outer subventricular zone (OSVZ) (LaMonica et al., 2013; Nowakowski et al., 2016; Baburamani et al., 2020; Yang et al., 2022). Human cortical neuroepithelial cells start to convert into fRG cells around gestational week 7 (GW7), and then fRG cells undergo asymmetric cell division to self-renew and to produce PyN intermediate progenitor cells (IPCs) around GW8, which exclusively differentiate into cortical deep layer PyNs. After 9 weeks of neurogenesis, around GW16–GW17, human cortical fRG cells give rise to oRG and tRG cells. oRG cells inherit long basal fibers of fRG cells, while tRG cells inherit apical domains of fRG cells (LaMonica et al., 2013; Nowakowski et al., 2016; Yang et al., 2022). Both oRGs and tRGs can self-renew. The emergence of the cortical OSVZ in higher-order mammals is fundamentally different from mice (Smart et al., 2002). oRG cells in the OSVZ mainly generate upper layer cortical PyNs (Lui et al., 2011; Molnar et al., 2019; Cardenas and Borrell, 2020), and a recent study suggests that tRG cells in the VZ primarily

Received 16 April 2023; accepted 29 May 2023.

[©]The Author(s) 2023. Published by Oxford University Press on behalf of Higher Education Press.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

generate cortical glia (Yang et al., 2022). After tRG cells are generated from fRG cells, a subpopulation of putative primed tRG cells expresses ASCL1 and epidermal growth factor receptor (EGFR). Initially, this population of primed tRG cells generates PyNs, but then they undergo a neurogenesis-to-gliogenesis switch, and generate basal multipotent intermediate progenitor cells (bMIPCs) that express ASCL1, EGFR, and oligodendrocyte transcription factor 2 (OLIG2) (Yang et al., 2022). These bMIPCs, at the population level, undergo several rounds of mitosis and give rise to most of the cortical glial cells (oligodendrocytes and astrocytes) and a subpopulation of olfactory bulb interneurons (Yang et al., 2022). It is worth noting that this process in the human cortex is similar to that in the mouse cortex (Li et al., 2021), suggesting that mammalian cortical gliogenesis is evolutionarily conserved, and that the presence of EGFR-expressing progenitors in the cortical VZ/SVZ is a strong signal for the onset of cortical gliogenesis (Fu et al., 2021; Li et al., 2021; Yang et al., 2022).

On the other hand, humans have a longer neurogenic period compared to perhaps all other mammals (Lewitus et al., 2014). This allows human cortical RG cells to undergo more cell division and therefore produce more neurons (Lewitus et al., 2014; Picco et al., 2018; Stepien et al., 2021). However, how the length of the neurogenic period is regulated in the mammalian brain remains a fundamental unanswered question.

Results

Human cortical oRG cells are neurogenic while tRG cells are gliogenic

Recent studies reported that human cortical neurogenesis ceases around GW19 (Pebworth et al., 2021), and reported that cortical oRG cells start to generate IPCs of oligodendrocytes, astrocytes, and interneurons around GW20 (Huang et al., 2020; Allen et al., 2022; Liu et al., 2023). In contrast, we found that human cortical neurogenesis extends to GW26, based on analyzing published single-cell RNA sequencing (scRNA-Seq) data of human GW22 (12,567 cells), GW23 (12,557 cells), and GW26 (13,371 cells) cortical tissue (Figs. 1A–D, S1A; Table S1) (Trevino et al., 2021). PyNs are continuously produced by PyN-IPCs that expressed EOMES (TBR2), NEUROG2, PPP1R17, PAX6, NEUROG1, NEUROD4, INSM1, SSTR2, ELAVL2, ELAVL4, and PENK (Figs. 1A-D and S1A-C) (Pollen et al., 2015; Girskis et al., 2021). This result is consistence with our previous immunohistochemical analyses on GW23 human cortical samples (Yang et al., 2022).

To define the major transcriptional features that underlie neurogenesis versus gliogenesis in the developing human cortex, we performed differential gene expression analysis between oRG cells and tRG cells in GW22–GW26 cortex. Although both human cortical oRG and tRG cells are derived from fRG cells, they exhibit distinct transcriptional signatures; oRG cells had 374 genes while tRG cells had 325 genes that were expressed at relatively higher levels (Table S1). Selected differentially expressed genes are shown in the heat map (Fig. 1E). Gene Ontology (GO) analysis of differentially expressed genes revealed that, in general, cortical oRG cells are neurogenic while tRG cells are highly gliogenic (Fig. 1F). Briefly, compared to tRG cells, oRG cells expressed higher levels of BMP7, HOPX, LIFR, CXCL14, MFGE8, NOG (NOGGIN), TKTL1, integrin ITGB5, and WNT pathway inhibitors (SFRP1 and DKK3) (Fig. 1E). In addition, ~30% of oRG cells expressed FGF2 (Fig. S1D). Furthermore, FGF/ MAPK signaling is elevated in oRG cells based on their expression of FGFR1, ETV1, ETV5, SPRY1, and SPRY2 (Fig. 1E). We speculate that FGF2/MAPK activity in oRGs is a cell-autonomous mechanism that antagonizes autocrine quiescent bone morphogenic protein 7 (BMP7) signaling, through phosphorylation of the SMAD linker domain (Kretzschmar et al., 1997; Pera et al., 2003). Indeed, previous studies showed that while BMP signaling alone promotes terminal astrocytic differentiation, exposure to both BMP and FGF2 maintains the stem cell character of cultured progenitor cells (Sun et al., 2011; Martynoga et al., 2013).

Signaling pathways driving gliogenesis in tRG cells

PyN-IPCs derived from cortical neurogenic RG cells are slow-cycling cells (Girskis et al., 2021). In contrast, bMIPCs derived from cortical gliogenic tRG cells are rapid cycling (Li et al., 2021; Yang et al., 2022). We identified six molecular programs in tRG cells that drive gliogenesis: (i) tRG cells are in the cortical VZ, so their primary cilia contact cerebrospinal fluid (CSF). Sonic Hedgehog (SHH), a morphogen and mitogen, is in the CSF (Huang et al., 2009; Lun et al., 2015; Winkler et al., 2018). Furthermore, tRG cells express higher levels of BOC and CDON than oRG cells (Fig. 1E); they are coreceptors of PTCH1 that enhance SHH signaling (Allen et al., 2011). (ii) Transcription effects of SHH signaling are mediated by GLI transcription factors. In the absence of SHH signaling, GLI3 full-length (GLI3FL) protein is processed to a truncated repressor form of GLI3 (GLI3R) that represses SHH target genes. This process depends on adenylyl cyclase-mediated cAMP-dependent protein kinase A (PKA), which phosphorylates GLI3FL, stimulating the proteolytic processing of GLI3FL to GLI3R. Upon SHH pathway activation, the SHH signaling cascade decreases PKA activity, prevents GLI3R production, and promotes the formation of GLI activators that drive transcription of SHH target genes (Wang et al., 2000; Hui and Angers, 2011). We found that 50% of human cortical tRG cells expressed CXCL12 (Fig. 1E), whereas very few mouse cortical RG cells expressed Cxcl12. CXCR4, expressed by

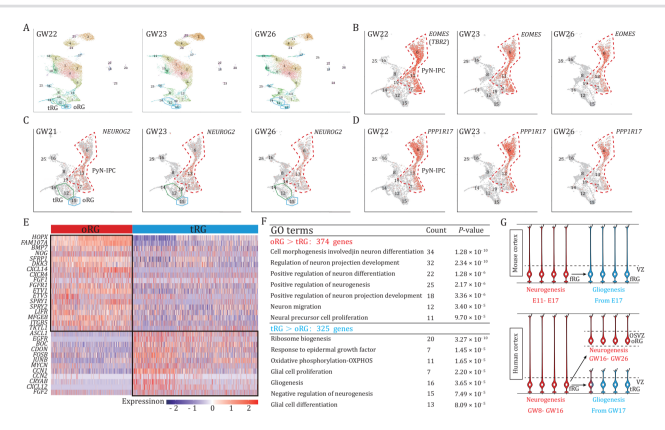


Figure 1. Human cortical oRG cells and tRG cells exhibit distinct transcriptional signatures. (A–D) scRNA-Seq analysis of molecular profiles of human cortical cells at GW22, GW23, and GW26. tSNE plot showed EOMES (TBR2)-, NEUROG2-, and PPP1R17-expressing PyN-IPCs, suggesting that cortical neurogenesis extends to GW26. (E) Heat map of selected differentially expressed genes for oRG cells versus tRG cells. Note that HOPX and FAM107A are enriched in oRG cells. (F) Selected GO terms implicating oRG cells are neurogenic while tRG cells are gliogenic. (G) Schematic summarizing mouse and human cortical RG cell lineage progression. Note that the human OSVZ is a duplicated neurogenic zone, with a greatly prolonged period of cortical neurogenesis. Thus, the length of the cortical neurogenic period is more than 130 days (GW8–GW26) in humans.

a subset of human cortical RG cells (Figs. 1E and S1E), is a G protein-coupled receptors (GPCR) that activates Gai protein and inhibits cAMP-PKA signaling. Thus, the autocrine activation of CXCR4 in human tRG cells further decreases GLI3R production, thereby increasing SHH signaling (Klein et al., 2001). CXCL12/CXCR4 also activates PI3K/AKT/mTOR and YAP-CCN1/2 signaling (Rao et al., 2016; Walenkamp et al., 2017; Shi et al., 2020). (iii) About 30% of tRG cells express FGF2, and most of them express FGFR1 and FGFR2 (Figs. 1E and S1D), thus FGF/MAPK pathway is also activated in tRG cells. (iv) Ribosomal biosynthesis genes are critical targets of cellular pro-growth signaling (Pelletier et al., 2018), and we found that more than 70 large and small ribosomal subunit genes were significantly upregulated in tRG cells (Fig. S2). (v) tRG cells also express higher levels of several key oxidative phosphorylation (OXPHOS) components (MT-ND1-5 and MT-CO1-3) (Fig. S2), suggesting their preferential use of OXPHOS instead of glycolysis (Fig. 1F). These data also indicate a metabolic shift from glycolysis to OXPHOS when cortical RG cells become gliogenic. (vi) Most importantly, ~40% of tRG cells are primed/poised for generating bMIPCs, as they express ASCL1 and EGFR (Fig. 1E), which then give rise to bMIPCs that express ASCL1, EGFR, and OLIG2 (Fig. S1F) (Yang et al., 2022). SHH signaling through *Smo* is required in cortical RG cells for gliogenesis (Winkler et al., 2018; Zhang et al., 2020b), as is MAPK signaling (Li et al., 2012, 2014). Both are also required for cortical RG cell proliferation (Komada et al., 2008; Lavoie et al., 2020). These data suggest that when proliferative and gliogenic signals (SHH + FGF/MAPK) overcome the threshold of quiescent and neurogenic signals (GLI3R + BMP7), tRG cells switch from neurogenesis to gliogenesis; this process in humans takes place over a much longer period than that in the mouse cortex (Yang et al., 2022).

Taken together, our analysis reveals that human cortical neurogenesis extends to at least GW26, due to the maintenance of the oRG cells' capacity to generate PyNs, and not due to tRG cells. The human neocortex has two germinal zones from GW16 to GW26: the gliogenic VZ and the neurogenic OSVZ (Fig. 1G) (Yang et al., 2022). This ensures that the onset of cortical gliogenesis normally occurs in the VZ around GW17, while oRGs continue to produce neurons in the OSVZ, thus increasing the length of the neurogenic period for another 10 weeks (GW16–GW26) (Fig. 1G).

An evolutionary increase in BMP7 expression in mammalian cortical RG cells

BMP7 is enriched in human but not in mouse cortical RG cells (Lui et al., 2014), and it is expressed relatively higher in oRG cells than that in tRG cells (Figs. 1E and S1D). We examined BMP7 expression in different mammalian cortical RG cells by integrating the analysis of scRNA-Seq datasets with mRNA in situ hybridization experiments. We found that between stages of cortical patterning and the onset of cortical neurogenesis, BMP7 expression pattern shows a startling similarity between mouse and human: BMP7 is expressed by the choroid plexus and cortical hem, and weak BMP7 expression extends to the medial cortex (Fig. 2A-D) (Abu-Khalil et al., 2004). Soon after, the growth of the human telencephalon accelerates, BMP7 expression begins in human cortical RG cells, whereas very few mouse dorsal cortical RG cells express Bmp7 (Fig. 2E-H). Comparative scRNA-Seq analyses reveal that there is an increase in BMP7 expression by cortical neurogenic fRG cells of mouse at E15, ferret at E39, rhesus monkey at E78, and human at GW14 (Figs. 2I, S3 and Table S2). As human development proceeds from GW12 to GW26, more cortical RG cells express BMP7 (from ~20% to ~90%) (Figs. 2J, S4 and Table S3), as well as BMP receptors BMPR1A, BMPR1B, and BMPR2 (Fig. S5 and Table S3) (Panchision et al., 2001; Hebert et al., 2002; Caronia et al., 2010). These results suggest the importance of BMP7 during mammalian cortical development and evolution.

BMP7 and SHH signaling inhibit each other to regulate mouse cortical gliogenesis

Because very few mouse dorsal cortical RG cells express Bmp7, we first explored its function using a gain-of-function method. Bmp7 was delivered (via pCAG-Bmp7-Gfp) into the mouse dorsal cortical VZ at E14.5 using in utero electroporation (IUE). Controls were electroporated with *pCAG-GFP*. At E17.5, while the control cortex expressed strong EGFR (Fig. 3A), the pCAG-Bmp7-Gfp-IUE cortex did not (Fig. 3B). BMP receptor signaling triggers the phosphorylation and activation of SMAD1/5/9 proteins, which form a complex with SMAD4 that together regulates gene expression (Wang et al., 2014b). Accordingly, cortical BMP7 overexpression resulted in strong expression of pSMAD1/5/9 and their downstream targets ID3, ID1, and HOPX (Figs. 3B and S6). However, in the absence of cortical Smad4 function (hGFAP-Cre;Smad4^{F/F} mice), Bmp7 overexpression did not inhibit EGFR expression (Fig. 3C and 3D). EGFR is autonomously required for cortical gliogenesis (Zhang et al., 2020a, 2023). Thus, BMP7-SMAD signaling inhibits cortical gliogenesis probably by downregulating EGFR expression.

We next examined BMP function in the medial cortex by overexpressing a repressor of BMP signaling. *hGFAP-Cre* is active in medial cortical RG cells from E12 (Fig. S7A and S7B); it was used to overexpress BMP inhibitor NOG (using Nog^{OE/+} mice). At E16.5, while EGFR expression was weak in the medial cortex of *hGFAP-Cre* control mice, its expression was upregulated in the medial and/or dorsal cortex of *hGFAP-Cre*;Nog^{OE/+} mice (Fig. S7C). In contrast, ID3 and pSMAD1/5/9 expression were downregulated in the medial cortex (Fig. S7C). Thus, BMP signaling inhibits gliogenesis mainly in the mouse medial cortex. Interestingly, the mouse medial cortex has human oRGlike cells (Vaid et al., 2018); perhaps, this is due to *Bmp7* expression.

BMPs and SHH are known to antagonize the downstream response to each other (Ulloa and Briscoe, 2007; Hebert and Fishell, 2008; Hoch et al., 2009). However, mechanisms for this remain elusive. We examined whether BMP7 signaling upregulates cortical GLI3R expression. *Bmp7* was electroporated into the mouse cortex at E14.5 or E15.5; the *Bmp7*-IUE cortex was collected 48 h later for GLI3 Western blot assay. *Bmp7* overexpression significantly increased the GLI3R/GLI3FL ratio in the cortex mainly due to excess production of GLI3R (Fig. 3E; see Western blot in Fig. S8), which antagonizes SHH signaling.

Next, we ectopically expressed non-cholesterol-modified SHH (SHHN) in the cortical VZ by IUE of pCAG-ShhN-GFP into wild-type (WT) mice at E15.5 and examined the cortex at E16.5. The increased ShhN expression upregulated EGFR, ASCL1, and OLIG2 expression compared to the pCAG-GFP-IUE (control) cortex (Fig. S9A-F). In contrast, ID3 and HOPX expression were downregulated upon ShhN overexpression (Fig. S9G-K). Expression of a constitutively active SMO protein that drives SHH signaling (hGFAP-Cre;SmoM2 mice) also increased in EGFR, ASCL1, and OLIG2 expression (Fig. 4A-C), and decreased in Bmp7, pSMAD1/5/9, ID3, HOPX, and EOMES expression in the E16.0 medial and/or dorsal cortex (Fig. 4D-H). Together, these results indicate that increased SHH signaling inhibits Bmp7 expression in the cortex and promotes cortical gliogenesis.

Then we tested whether EGFR expression in the dorsal cortex requires SHH signaling by deleting *Smo* (*hGFAP-Cre;Smo*^{F/F} mice). At E17.5, very few EGFR- and OLIG2-expressing cortical cells were detected in *hGFAP-Cre;Smo*^{F/F} mice (Fig. 5A); their expression partially recovered by P2 (Fig. S10). This result suggests that, without SHH signaling, cortical gliogenesis is greatly retarded.

To understand the molecular mechanism responsible for SHH-mediated gliogenesis, we examined *Bmp7* expression. mRNA in situ hybridization revealed that *Bmp7* expression was increased in the cortical VZ of *hGFAP*-*Cre;*Smo^{F/F} mice at E17.5 (Fig. 5F). Consistent with this observation, expression of BMP7 effector pSMAD1/5/9,

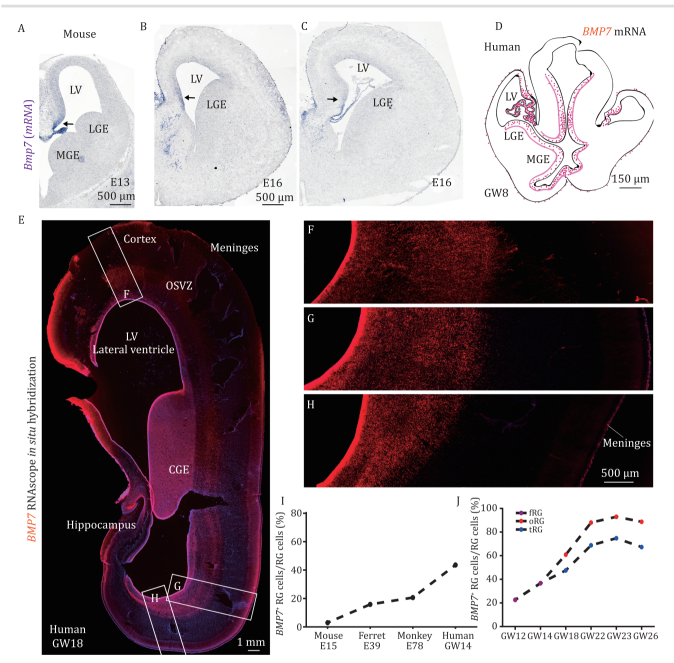


Figure 2. BMP7 is expressed by more and more cortical RG cells during mammalian development and evolution. (A–C) *In situ* hybridization shows that *Bmp7* mRNA is expressed in the mouse medial cortex, but not in the dorsal cortex at E13.0 and E16.0. (D) Maps of *BMP7*-expressing cells in the human cortex at GW8. The original picture of *BMP7* mRNA *in situ* hybridization on human GW8 cortical section is from previous studies (Abu-Khalil et al., 2004). (E–H) RNAscope *in situ* hybridization shows *BMP7* expression in germinal zones of the human cortex and caudal ganglionic eminence (CGE) at GW18. Note *BMP7* is also expressed in meninges. (I) Comparative scRNA-Seq analyses reveal an increase in the percentage of cortical RG cells that expressed *BMP7* in the E15 mouse cortex, E39 ferret dorsal cortex, E78 rhesus monkey visual cortex, and GW14 human prefrontal cortex. (J) Increasing the percentage of human cortical RG cells that expressing cell that expressed at least one UMI count. To compare UMI counts across samples, we took into account differences in sequencing depth between the samples according to the normalization procedure provided by 10× genomic. LGE, lateral ganglionic eminence; MGE, medial ganglionic eminence.

ID3 and HOPX was upregulated (Fig. S11). We also examined the E17.5 cortex of hGFAP- $Cre;Smo^{F/F}; Nog^{OE/+}$ mice and found that EGFR and OLIG2 expression were partially rescued (Fig. 5B), supporting the idea that upregulation BMP7 signaling contributes to downregulation of EGFR and OLIG2 expression in the cortex of hGFAP- $\mathit{Cre;Smo^{\text{F/F}}}$ mice.

Previous studies have demonstrated that the dorsal cortical identity and neurogenesis are protected by GLI3R-mediated maintaining *Bmp* and *Wnt* expression,

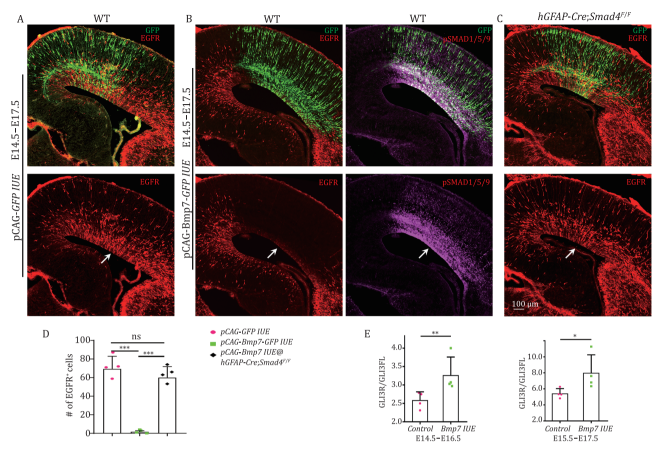


Figure 3. BMP7-SMAD signaling inhibits EGFR expression. (A and B) IUE of vectors that express either *GFP* (control) or *Bmp7-GFP* in E14.5 WT mouse cortex. At E17.5, EGFR expression was significantly downregulated and pSMAD1/5/9 expression was strongly upregulated in the *Bmp7-IUE* cortex (arrows). (*C*) In the *hGFAP-Cre;Smad4^{F/F}* cortex, *Bmp7-IUE* showed less EGFR inhibition. (D) Quantification of numbers of EGFR-expressing cells in the cortex. (E) *Bmp7* was overexpressed in the E14.5 or E15.5 cortex using IUE. The cortical GLI3R to GLI3FL ratio was significantly increased 48 h later (see Western blot in Fig. S8).

and suppression from the ventralizing effects of SHH and FGF signaling (Tole et al., 2000; Aoto et al., 2002; Kuschel et al., 2003; Rash and Grove, 2007). To investigate the function of GLI3R, we used hGFAP-Cre;Gli3^{F/F} mice. We examined E16.0 cortex and found that EGFR and OLIG2 expression were increased (Fig. S12). This result suggests that, in addition to BMP7, GLI3R also represses EGFR and OLIG2 expression. Furthermore, the GLI3R/GLI3FL ratio was significantly increased in the cortex of hGFAP-Cre;Smo^{F/F} mice (Fig. S8A). We therefore blocked the generation of GLI3R by ectopically expressing a dominant negative form of PKA (dnPKA). In the WT cortex that was electroporated with *dn*PKA, upregulation of EGFR, OLIG2, and ASCL1 was observed (Fig. S13). Similarly, in the cortex of hGFAP-Cre;Smo^{F/F} mice, we observed a rescue of EGFR, OLIG2, and ASCL1 expression in the *dnPKA* electroporated side compared to the contralateral side (Figs. 5C-E and S14). Moreover, Bmp7 upregulation disappeared upon dnPKA expression (Fig. 5F). This result suggests that an increase in GLI3R production could be key to inducing Bmp7 expression in the dorsal cortex of hGFAP-Cre;Smo^{F/F} mice. In other words, SHH signaling promotes cortical

progenitor proliferation and gliogenesis by reducing GLI3R production, which also contributes to preventing *Bmp7* expression to spread from the medial cortex to the dorsal cortex.

Increase in BMP7 signaling and loss of SHH signaling prolong mouse cortical neurogenic period

Finally, we performed scRNA-Seq transcriptome analysis of cortical cells in *hGFAP-Cre;Smo^{F/F}* mice and littermate *Smo^{F/F}* control mice at E18.0 (Fig. 6A–C and Table S4). First, we identified that expression of *Smo*, *Gli1*, and *Ptch1* was in the ground state in the cortex of *hGFAP-Cre;Smo^{F/F}* mice (Figs. 6D and S15A). Second, we confirmed that *Bmp7*, but not *Bmp2*, *Bmp4*, *Bmp5*, or *Bmp6* (Fig. 6D), was upregulated in cortical RG cells without SHH-SMO function. BMP7 response genes *Id3* and *Hopx* were also upregulated (Fig. 6D). Third, cell proliferation was greatly reduced because the expression of *Cdk6* and *Ccnd1* was significantly downregulated while *Cdkn1a* (P21) and *Trp53* were upregulated (Fig. 6D). Fourth, the generation of *Egfr-*, *Olig1-*, and *Olig2-*expressing bMI-PCs was greatly reduced (Fig. S15B), consistent with

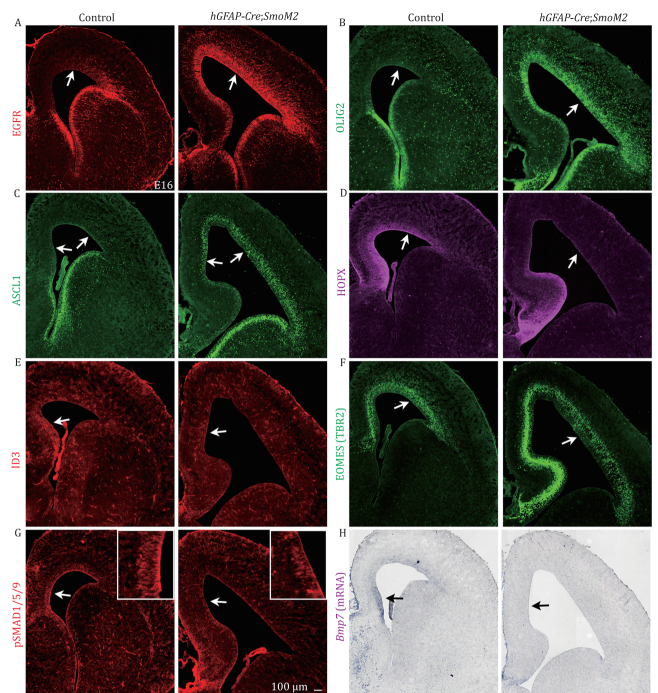


Figure 4. Constitutively active SmoM2 inhibits BMP7 signaling and promotes cortical gliogenesis. (A–H) The expression of EGFR, OLIG2, and ASCL1 was increased, whereas the expression of HOPX, ID3, EOMES (TBR2), pSMAD1/5/9, and Bmp7 was decreased in the medial and/or dorsal cortex of hGFAP-Cre;SmoM2 mice compared to control mice at E16 (arrows).

immunostaining results (Fig. 5A). Fifth, a set of WNT and BMP response genes in RG cells that promote cortical neurogenesis was upregulated, including *Wnt7b*, *Axin2*, *Lef1*, *Dmrta2*, *Lhx2*, and *Emx2* (Figs. 6D and S15A) (Monuki et al., 2001; Theil et al., 2002; Saulnier et al., 2013; Caronia-Brown et al., 2014), strongly indicating that neurogenesis persists in the cortex of *hGFAP*-*Cre;Smo^{F/F}* mice at E18.0, a time point that cortical RG cells do not undergo neurogenesis in the normal mouse brain. Sixth, consistently, while EOMES-expressing cells (they most likely are PyN-IPCs) in the cortical SVZ were reduced before birth due to reduced cell proliferation (Komada et al., 2008), EOMES-expressing cells increased in the *Smo*-mutant P4 and P6 cortex (Figs. 6E, 6F and S15C). Taken together, these results suggest that, without SHH signaling, cortical RG cell proliferation is greatly reduced and gliogenesis is retarded, but the length of the cortical neurogenic period is increased.

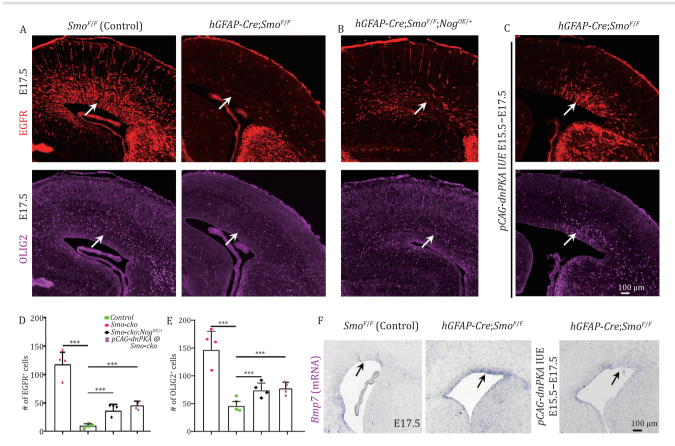


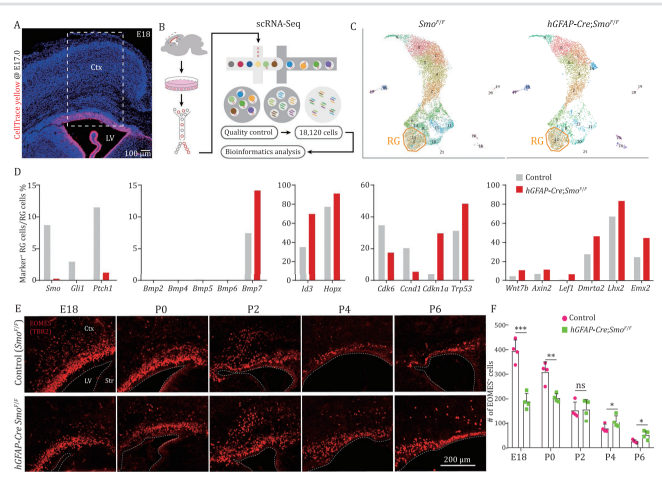
Figure 5. SHH-SMO signaling is required for cortical gliogenesis. (A) The expression of EGFR and OLIG2 was severely downregulated in the E17.5 cortex of hGFAP-Cre;Smo^{F/F} mice (Smo-cko). (B and C) Overexpression of Nog or dnPKA largely rescued the downregulation of EGFR and OLIG2 (arrows). (D and E) Quantification of numbers of cortical EGFR- and OLIG2-expressing cells. (F) Bmp7 mRNA expression was upregulated in the E17.5 cortex of hGFAP-Cre;Smo^{F/F} mice, but dnPKA-IUE blocked Bmp7 upregulation.

Discussion

BMP signaling primarily functions to maintain stem cells in a quiescent state (Li and Clevers, 2010). Here we suggest that BMP7, in combination with cell proliferation signals, sustains the self-renewal of cortical RG cells, maintaining cortical neurogenesis, and inhibiting gliogenesis. On the basis of these findings, we wish to suggest a principle of mammalian cortical development, expansion, and evolution (Fig. 7).

(1) The most recent ancestor to all mammals, i.e., 100 million years ago, is assumed to have a subset of BMP7-expressing fRG cells in the dorsal cortex, and is assumed to have already been in some degree gyrencephalic (Fig. S16) (Lewitus et al., 2014). Expression of BMP7 in cortical fRG cells increases GLI3R production, which further strengthens BMP7 expression in fRG cells. Therefore, BMP7 expression in cortical fRG cells participates in a positive feedback loop, which antagonizes SHH signaling, protects neurogenesis, inhibits gliogenesis, and thus increases the length of the neurogenic period (Fig. 7). Perhaps, SHH CSF concentration may be reduced in the larger lateral ventricle in bigger-brained mammals; that could further decrease SHH signaling within the dorsal cortex.

- (2) Around midterm gestation, the increase in SHH downstream signaling (including CXCL12/CXCR4 signaling), combined with FGF2/MAPK signaling in cortical fRG cells, is probably the major force that drives BMP7-expressing cortical fRG cells to give rise to neurogenic oRG and gliogenic tRG cells.
- (3) BMP7 expression in oRG cells also participates in a positive feedback loop. oRG cells in the OSVZ do not contact the lateral ventricle (largely insulated from SHH, similar to cortical RG cells in hGFAP-Cre;Smo^{F/F} mice), and express higher level of BMP7 and GLI3R, and maybe never express EGFR (Yang et al., 2022). The more fRG cells express BMP7, the more oRG cells are generated, lengthening the neurogenic period, the more PyNs are generated from the OSVZ, and expanding cortical size during development and perhaps evolution.
- (4) Although BMP7 is highly expressed in oRG cells, FGF2 and NOG expression in oRG cells suppress quiescency and differentiation of BMP7 signaling, thus sustaining self-renew in oRG cells. When



BMP7 increases the length of the neurogenic period | 29

Figure 6. Loss of SHH-SMO signaling in mouse cortical RG cells lengthens their neurogenic period. (A–C) Schematic of the workflow of scRNA-Seq analysis. CellTrace Yellow was injected into the lateral ventricle of $Smo^{F/F}$ or hGFAP-Cre; $Smo^{F/F}$ mice at E17.0. The cortex was collected at E18.0 for cell sorting and scRNA-Seq analysis. (D) Loss of Smo function in cortical RG cells resulted in significantly downregulation of Smo, Gli1, Ptch1, and upregulation of Bmp7, Id3, and Hopx, and inhibition of cell proliferation (downregulation of Cdk6 and Ccnd1, and upregulation of Cdkn1a and Trp53). A subset of WNT and BMP response neurogenic genes (Wnt7b, Axin2, Lef1, Dmrta2, Lhx2, and Emx2) was upregulated in cortical RG cells. (E and F) Numbers of EOMES-expressing cells were decreased in the E18 and P0 cortex, whereas increased in the P4 and P6 cortex of hGFAP-Cre; $Smo^{F/F}$ mice, compared to $Smo^{F/F}$ (control) mice, indicating that the length of the cortical neurogenic period is increased. Ctx, cortex; LV, lateral ventricle; Str, striatum.

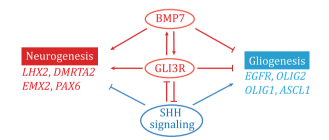


Figure 7. BMP7 and GLI3R coordinately protect cortical neurogenesis in human brain. Diagram showing that BMP7/ GLI3R and SHH signaling mutually inhibit each other. BMP7 and GLI3R coordinately protect the genetic program of cortical neurogenesis and inhibit cortical gliogenesis. In contrast, SHH signaling promotes cortical gliogenesis and opposes BMP7/ GLI3R functions during cortical development.

BMP7 expression gets stronger, oRG cells may stop generating PyN-IPCs via asymmetric division, directly transform into astrocytes that also express BMP7 (Sun et al., 2011), and do not produce oligodendrocyte IPCs (Yang et al., 2022).

- (5) On the other hand, for unknown reasons, during development and evolution, there was a subpopulation of mammals (i.e., ancestors of hamsters, mice, and rats) (Fig. S16), whose cortical fRG cells perhaps received higher SHH signaling, which represses *Bmp7* expression in dorsal RG cells, and restricts *Bmp7* expression to the medial cortex. Smaller brains with the smaller lateral ventricle may further increase SHH signaling within the dorsal cortex.
- (6) Mouse cortical neurogenesis ceases and gliogenesis occurs when SHH signaling is further increased at ~E17.0. The increase in SHH and MAPK signaling in the mouse dorsal cortical VZ induces a subset of fRG cells to directly transform into cortical astrocyte-IPCs (transforming RG cells) that express EGFR, ASCL1, and OLIG2 (Li et al., 2021),

but not oRG cells, as lack of *Bmp7* expression, and induces most cortical RG cells to generate bMIPCs that also express EGFR, ASCL1, and OLIG2 (Li et al., 2021). This population of bMIPCs gives rise to most of the cortical glial cells (oligodendrocytes and astrocytes) and a subpopulation of olfactory bulb interneurons (Yang et al., 2022). Therefore, mouse cortical neurogenesis is mainly protected by Gli3R, whereas human cortical neurogenesis is protected by GLI3R and BMP7; both of them directly or indirectly repress SHH target genes and EGFR expression (Fig. 7).

- (7) It is worth noting that BMP7 expression also extends from the cortex to subcortical ganglionic eminences in higher-order mammalian brains (Fig. 2D-H), thus it may prolong the neurogenic period in most regions of the telencephalon. The study reports that heterozygous mutations in the BMP7 gene in patients cause small head (Wyatt et al., 2010), providing additional genetic evidence for BMP7 function in brain development.
- (8) BMPs, WNTs, FGFs, and SHH, are used over and over again across ontogeny and phylogeny. In the nervous system, these molecules regulate neural induction, pattern dorsal, ventral, rostral and caudal structures, and regulate neurogenesis versus gliogenesis (Hebert and Fishell, 2008; Hoch et al., 2009). Therefore, it is not surprising that they also play a leading role in driving cortical evolution.
- (9) We believe that this basic principle (Fig. 7) is still driving human brain to evolve and to generate a cortex with more neurons.

Materials and methods Animals

All procedures and animal care, including ferrets and mice, were approved and performed in accordance with the Fudan University, Shanghai Medical College Laboratory Animal Research Center guidelines. WT CD-1, hGFAP-Cre (Zhuo et al., 2001) (JAX no. 004600), Smo flox (Long et al., 2001) (JAX no. 004526), R26SmoM2 (JAX no. 005130), Gli3 flox (Blaess et al., 2008) (JAX no. 008873), Smad4 flox (Yang et al., 2002) (Jax no. 017462), and pMes-Noq (Nog overexpression, Noq^{OE/+}) transgenic (Xiong et al., 2009) mice were described previously. The Nog^{OE/+} mice were generously provided by Professor Yanding Zhang from Fujian Normal University. The day of the vaginal plug detection was designated as Embryonic day 0.5 (E0.5). The day of birth was designated as PO. The genders of the embryonic mice were not determined, and both male and female postnatal mice were used. One pregnant ferret (Mustela putorius furo) was purchased from Wuxi Sangosho Biotechnology Co., Ltd, Wuxi, China, and we used one embryo at E39

in this study. The bilateral cortex from one E39 embryo was used for scRNA-Seq.

Plasmid construction

pCAG-ires-GFP plasmid was from Addgene (Addgene #11150), and pCAG-ShhN-ires-GFP was used in the previous study (Zhang et al., 2020b). Mouse Bmp7 cDNA and dnPKA cDNA (Clegg et al., 1987) were cloned and inserted into pCAG-GFP vector to construct pCAG-Bmp7-ires-GFP and pCAG-dnPKA-ires-GFP overexpression plasmids.

In utero electroporation (IUE)

Overexpression of plasmids was performed in the cortex using IUE at E14.5 or E15.5. About 0.5 μ L of 1–2 μ g/ μ L plasmid solution with 0.05% Fast Green (Sigma) was injected into the lateral ventricle of the embryos using a cable-drawn glass micropipette. Five electrical pulses (duration: 50 ms) were applied across the uterus to E14.5 embryos at 33 V and E15.5 embryos at 35 V. The interval between pulses is 950 ms. Electroporation was performed using a pair of 7 mm platinum electrodes (BTX, Tweezerrode 45-0488, Harvard Apparatus) connected to an electrocleaner (BTX, ECM830).

Fixation and sectioning of the brain tissue

Embryos were harvested from deeply anesthetized pregnant mice. Each embryo was separated from the placenta, the brain was dissected out, and then fixed in 4% diethylpyrocarbonate and paraformaldehyde (DEPC-PFA) overnight. Postnatal mice were deeply anesthetized and perfused intracardially with 4% paraformaldehyde (PFA). All brains were fixed overnight in 4% PFA at 4°C and cryoprotected in 30% sucrose for at least 24 h, embedded with O.C.T. (Sakura Finetek) in ethanol slush with dry ice. All mouse brains in this study were sectioned in a coronal plane at 20 µm. The GW18 human cortical sections were 60 µm as we used in our previous studies (Ma et al., 2013; Wang et al., 2014a; Yang et al., 2022).

Immunohistochemistry

All immunohistochemical stains in this study were performed on 20 μ m coronal cryostat sections of mouse brains as previously described (Li et al., 2021; Yang et al., 2022). Sections were rinsed with TBS (0.01 mol/L Tris–HCl + 0.9% NaCl, pH = 7.4) for 10 min, incubated in 0.5% Triton-X-100 in TBS for 30 min at room temperature (RT), and then incubated with block solution (5% donkey serum + 0.5% Triton-X-100, pH = 7.2) in TBS for 30 min at RT. For double immunostaining, primary antibodies from different species were incubated simultaneously followed by secondary antibodies. Primary antibodies were diluted in donkey serum block solution and incubated overnight at 4°C, incubated for an additional 30 min at RT, and rinsed three times with TBS. Sections were then incubated with secondary antibodies (1:600, all from Jackson ImmunoResearch) for 2 h at RT, rinsed three times with TBS for 10 min, incubated with 4',6-diamidino-2-phenylindole (DAPI, 1:5,000, Sigma) diluted in TBS for 5 min, and then finally rinsed three more times with TBS. Primary antibodies used in this study include: goat anti-EGFR (1:1,000, R&D System, BAF1280), rabbit anti-OLIG2 (1:500, Millipore, AB9610), rabbit anti-pSMAD1/5/9 (1:100, Cell signaling Technology, 13820S), guinea pig anti-EOMES (1:500, Asis Biofarm, OB-PGP022); rat anti-EOMES (1:500, Thermo Fisher, 12-4875-82), rabbit anti-ASCL1 (1:1,000, Cosmo Bio, SKT01-003), rabbit anti-ID3 (1:5,000, Biocheck Inc, BCH-4/17-3), rabbit anti-HOPX (1:500, Proteintech, 11419-1-AP), rabbit anti-ID1 (1:2,000, Biocheck Inc, BCH-1/37-2), and chicken anti-GFP (1:3,000, Aves labs, GFP-1020).

Mouse Bmp7 mRNA in situ hybridization

Mouse *Bmp7* mRNA *in situ* RNA hybridization experiment was performed using digoxigenin riboprobes on 20 µm cryostat sections as described previously (Zhang et al., 2020b). Riboprobes were made from cDNAs amplified by polymerase chain reaction using the following primers: *Bmp7*-F: GGGCCAGAACTGAGTAAAGGAC; *Bmp7*-R: GAAGCTCATGACCATGTCGG.

Human BMP7 mRNAscope in situ hybridization

Human BMP7 mRNA in situ hybridization on GW18 human brain sections was performed on 60 µm cryostat sections using the RNAscope[™] assay as described previously (Wang et al., 2012). The human BMP7 probe (424401) was purchased from Advanced Cell Diagnostics, Newark, CA. The RNAscope Multiplex Fluorescent Reagent Kit v2 (Advanced Cell Diagnostics, Newark, CA, 323110) was used according to the instruction manual. The slides were dehydrated in a series of ethanol and loaded onto the Leica Bond RX automated stainer, followed by the pretreatments (protease), probe incubation, amplification steps, fluorophores, and DAPI counterstain. Quality control for RNA integrity was completed using probes specific to the housekeeping genes PPIB. Negative control background staining was evaluated using a probe specific to the bacterial DapB gene. The original picture of BMP7 mRNA in situ hybridization on human GW8 cortical section is from Abu-Khalil et al. (2004).

Western blot

Bmp7 was delivered (via *pCAG-Bmp7-Gfp*) into the mouse dorsal cortical VZ at E14.5 or E15.5 using IUE. Controls were electroporated with *pCAG-GFP*. The IUE-mouse cortex was collected 48 h later under a microscope. The cortex was lysed with RIPA lysis buffer (Beyotime, P0013K); main components of which are 50 mmol/L Tris (pH 7.4), 150 mmol/L NaCl, 1% Triton-X-100, 1% sodium deoxycholate, 0.1% SDS, and 1% PMSF (Beyond, ST506-2). Protein concentration was determined using a bicinchoninic assay kit (EpiZyme, ZJ101). Equal amounts of protein were loaded onto 8% SDS-PAGE gels, and Western blot was performed with GLI3 antibody (Goat anti-GLI3, 1:1,000, R&D System, AF3690) overnight at 4°C. Then membranes were washed three times with TBST, and secondary antibodies (Horseradish Hydrogen Peroxide Labeled Donkey Anti-Goat IgG (H+L), Beyotime, A0181, 1:1,000) was incubated at RT for 1 h. Next, membranes were washed by TBST and placed in BeyoECL Sta detection solution. Blots were imaged using e-blot biomolecular imager and bands were quantified using ImageJ software. β -Actin rabbit mAb (monoclonal antibody) was from ABClonal, Ac026, and Horseradish Hydrogen Peroxide Labeled Goat Anti-Rabbit IgG (H+L) was from Beyotime, A0208.

Cortical single-cell dissociation

CellTrace Yellow (Life Technologies, #C34567, 0.5 µL of 10 mmol/L) (Telley et al., 2016; Govindan et al., 2018) was injected into the lateral ventricle of Smo^{F/F} or hGFAP-Cre;Smo^{F/F} mice at E17.0. The cortex was collected 24 h later (E18.0) for cell sorting and scRNA-Seq analysis. Briefly, mouse embryos were dissected out and immediately submerged in fresh ice-cold Hanks balanced salt solution (Gibco 12175-095). The cortex was then cut into pieces and dissociated into a single-cell suspension using a Papain Cell Dissociation Kit (Miltenyi Biotec, catalog no. 130-092-628) according to the manufacturer's instructions. CellTrace Yellow labeled single-cells were sorted using a BD FACSAriaII (BD Biosciences). One pregnant ferret at E39 was anesthetized and embryos were dissected. The bilateral cortex from one embryo were then cut into pieces and dissociated into a single-cell suspension using a Papain Cell Dissociation Kit.

Construction of 10× Genomic scRNA-Seq libraries and sequencing

The Chromium droplet-based sequencing platform (10× Genomics) was employed to generate scRNA-Seq libraries, following the manufacturer's instructions (manual document part number: CG00052 Rev C). The cDNA libraries were purified, quantified using an Agilent 2100 Bioanalyzer, and sequenced on an Illumina Hiseq4000. High-quality sequences (clean reads) of samples were processed using Cell Ranger to obtain quantitative information on gene expression. Cellular quality control thresholds were set at 750-5,000 genes and <10% mitochondrial transcripts per cell. After filtering, the number of cells in our dataset was as follows: E18 Smo^{F/F} (control) cortex, 10,524 cells, 1,954 genes/cell; hGFAP-Cre;Smo^{F/F} cortex, 7,596 cells, 2,104 genes/cell; E39 ferret cortex, 23,482 cells, 1,759 genes/cell. scRNA-Seq data from E18 Smo^{F/F} (control) and hGFAP-Cre;Smo^{F/F} mice, and E39 ferret cortical scRNA-Seq data have been deposited in the Gene Expression Omnibus (GEO) under the accession number GSE221389. On the other hand, E15.5 mouse cortex

scRNA-Seq data is from (Di Bella et al., 2021), E78 rhesus monkey visual cortical scRNA-Seq data is from (Micali et al., 2020), and GW14 human cortex scRNA-Seq data is from (Ma et al., 2021), and GW18, GW22, GW23, and GW26 human cortex scRNA-Seq dataset is from (Trevino et al., 2021). Raw read counts were processed using the global scale normalization method Log Normalize. The normalized datasets were then combined using the Find Integration anchor and Integration Data functions. Statistically significant principal components identified by resampling tests were retained for unified manifold approximation and projection (UMAP) analysis. The enriched genes in different clusters were identified by Wilcoxon rank sum test (ad p value < 0.05, $|\log_2 FC|$ > 0.25). For cluster annotation, we searched for the most comprehensive and reliable cell type markers through an extensive literature review. All these analyzes were performed in the Seurat package v4.0.6.

Gene Ontology (GO) enrichment analysis

The GO analysis was performed on statistically significant upregulation genes in cortical oRG cells and tRG cells, using the ClusterProfiler, GO.db, DOSE, org Mm.eg. db, org.hs.eg.db packages in R, and the BH algorithm was used to control the P-value adjustment. The P value of GO item <0.05 is a significant item. Our analysis was performed separately on upregulated genes in human cortical oRG cells only, or upregulated genes in human cortical tRG cells only at GW22–GW26. The results of these analyses are reported in Table S1.

Microscopy and imaging

All images of stained sections in this study were collected by Olympus VS120 Automated Slide Scanner with X20. Adobe Photoshop software was used to combine multiple visual fields, if needed. Both Adobe Photoshop and Adobe Illustrator were employed to process or adjust images without destroying the original details.

Cell counts

In this study, all cell counts were performed on the imaged sections collected by Olympus VS120 Automated Slide Scanner with X20. For the quantification of numbers cells in the control cortex, vector-IUE-cortex, or transgenic mouse cortex, cells were always counted within a certain area of known size in the control or experimental cortex. Normally, at least three sections per cortex and three cortices were analyzed.

Statistical analysis of cell counts

Individual data points in the figure panels were plotted along with the mean and standard error of the mean (SEM). Unpaired Student's t-test was used to calculate the significance of differences between the two condition analyses. One-way analysis of variance followed by the Tukey–Kramer post-hoc test was used for assessing the different significance of the analyses containing more than three conditions. Significance is stated as follows: P > 0.05 (ns), P < 0.05 (*), P < 0.01 (***), P < 0.001 (***).

Supplementary information

The online version contains supplementary material available at https://doi.org/10.1093/procel/pwad036

Acknowledgements

We thank Professor J. L. Rubenstein for comments, suggestions, and edits. We are grateful to Professor Y. Zhang at Fujian Normal University for sharing the pMes-Noggin (Nog^{OE/+}) mouse lines.

Abbreviations

bMIPCs, basal multipotent intermediate progenitor cells; CSF, cerebrospinal fluid; DEPC, diethylpyrocarbonate; dnPKA, dominant negative form of protein kinase A; fRG, full span radial glia; GLI3FL, GLI3 fulllength; GLI3R, repressor form of GLI3; GO, gene ontology; GPCR, G protein-coupled receptor; GW, gestational week; IPCs, intermediate progenitor cells; IUE, in utero electroporation; oRG, outer radial glia; OSVZ, outer subventricular zone; OXPHOS, oxidative phosphorylation; PFA, paraformaldehyde; PKA, protein kinase A; PyNs, pyramidal neurons; RG, radial glia; scRNA-Seq, single-cell RNA sequence; SHHN, non-cholesterol-modified SHH; tRG, truncated radial glia; VZ, ventricular zone.

Data availability

The E18 Smo^{F/F} (control) and hGFAP-Cre;Smo^{F/F} mouse cortex (mainly including RG cells and progenitors), and E39 ferret cortex scRNA-seq data used in this study have been deposited in the Gene Expression Omnibus (GEO) under the accession number GSE221389. All datasets generated in this study are available upon request.

Conflict of interest

None declared.

Funding

This work was supported by the Ministry of Science and Technology of China (STI2030-2021ZD0202300), National Natural Science Foundation of China (NSFC 31820103006, 32070971, 32100768, 32200776, and 32200792), Shanghai Municipal Science and Technology Major Project (No. 2018SHZDZX01), ZJ Lab, and Shanghai Center for Brain Science and Brain-Inspired Technology.

Ethical approval

All procedures and animal care, including ferrets and mice, were approved and performed in accordance with the Fudan University, Shanghai Medical College Laboratory Animal Research Center guidelines.

Consent to participate

All authors give their consent to participate.

Consent for publication

All authors give their consent to participate.

Author contributions

Z.Y. and Z.L. designed the experiments. Z.L., G.L., and M.S. conducted majority of experiments and analyzed the data. L.Y. performed bioinformatics analysis. Z.Z., Z.X., X.J., and X.L. contributed to cells counting, partial immunostaining experiments. Y.G. performed Western blot analysis and genotyping. Z.S. performed in situ hybridization. Z.Y. initiated the project, provided the resources, supervised the whole project and wrote the manuscript.

References

- Abu-Khalil A, Fu L, Grove EA *et al*. Wnt genes define distinct boundaries in the developing human brain: implications for human forebrain patterning. *J Comp Neurol* 2004;**474**:276–288.
- Allen BL, Song JY, Izzi L *et al.* Overlapping roles and collective requirement for the coreceptors GAS1, CDO, and BOC in SHH pathway function. *Dev Cell* 2011;**20**:775–787.
- Allen DE, Donohue KC, Cadwell CR *et al.* Fate mapping of neural stem cell niches reveals distinct origins of human cortical astrocytes. *Science* 2022;**376**:1441–1446.
- Aoto K, Nishimura T, Eto K *et al*. Mouse GLI3 regulates Fgf8 expression and apoptosis in the developing neural tube, face, and limb bud. *Dev Biol* 2002;**251**:320–332.
- Baburamani AA, Vontell RT, Uus A *et al*. Assessment of radial glia in the frontal lobe of fetuses with Down syndrome. Acta Neuropathol Commun 2020;**8**:141.
- Blaess S, Stephen D, Joyner AL. Gli3 coordinates three-dimensional patterning and growth of the tectum and cerebellum by integrating Shh and Fgf8 signaling. *Development* 2008;**135**:2093–2103.
- Cardenas A, Borrell V. Molecular and cellular evolution of corticogenesis in amniotes. *Cell Mol Life Sci* 2020;**77**:1435–1460.
- Caronia G, Wilcoxon J, Feldman P *et al*. Bone morphogenetic protein signaling in the developing telencephalon controls formation of the hippocampal dentate gyrus and modifies fear-related behavior. *J Neurosci* 2010;**30**:6291–6301.
- Caronia-Brown G, Yoshida M, Gulden F et al. The cortical hem regulates the size and patterning of neocortex. Development 2014;**141**:2855–2865.

- Clegg CH, Correll LA, Cadd GG et al. Inhibition of intracellular cAMP-dependent protein kinase using mutant genes of the regulatory type I subunit. J Biol Chem 1987;**262**:13111–13119.
- Di Bella DJ, Habibi E, Stickels RR *et al.* Molecular logic of cellular diversification in the mouse cerebral cortex. *Nature* 2021;**595**:554–559.
- Fu Y, Yang M, Yu H et al. Heterogeneity of glial progenitor cells during the neurogenesis-to-gliogenesis switch in the developing human cerebral cortex. Cell Rep 2021;34:108788.
- Girskis KM, Stergachis AB, DeGennaro EM et al. Rewiring of human neurodevelopmental gene regulatory programs by human accelerated regions. *Neuron* 2021;**109**:3239– 3251.e7.
- Govindan S, OberstJabaudon D. In vivo pulse labeling of isochronic cohorts of cells in the central nervous system using FlashTag. Nat Protoc 2018;**13**:2297–2311.
- Hebert JM, Fishell G. The genetics of early telencephalon patterning: some assembly required. *Nat Rev Neurosci* 2008;**9**:678–685.
- Hebert JM, Mishina Y, McConnell SK. BMP signaling is required locally to pattern the dorsal telencephalic midline. *Neuron* 2002;**35**:1029–1041.
- Hoch RV, Rubenstein JL, Pleasure S. Genes and signaling events that establish regional patterning of the mammalian forebrain. *Semin Cell Dev Biol* 2009;**20**:378–386.
- Huang X, Ketova T, Fleming JT et al. Sonic hedgehog signaling regulates a novel epithelial progenitor domain of the hindbrain choroid plexus. *Development* 2009;**136**:2535–2543.
- Huang W, Bhaduri A, Velmeshev D *et al*. Origins and proliferative states of human oligodendrocyte precursor cells. *Cell* 2020;**182**:594–608.e11.
- Hui CC, Angers S. Gli proteins in development and disease. Annu Rev Cell Dev Biol 2011;**27**:513–537.
- Klein RS, Rubin JB, Gibson HD et al. SDF-1 alpha induces chemotaxis and enhances Sonic hedgehog-induced proliferation of cerebellar granule cells. *Development* 2001;**128**:1971–1981.
- Komada M, Saitsu H, Kinboshi M et al. Hedgehog signaling is involved in development of the neocortex. *Development* 2008;**135**:2717–2727.
- Kretzschmar M, Doody J, Massague J. Opposing BMP and EGF signalling pathways converge on the TGF-beta family mediator Smad1. *Nature* 1997;**389**:618–622.
- Kriegstein A, Alvarez-Buylla A. The glial nature of embryonic and adult neural stem cells. *Annu Rev Neurosci* 2009;**32**:149–184.
- Kuschel S, Ruther U, Theil T. A disrupted balance between Bmp/ Wnt and Fgf signaling underlies the ventralization of the Gli3 mutant telencephalon. *Dev Biol* 2003;**260**:484–495.
- LaMonica BE, Lui JH, Hansen DV *et al.* Mitotic spindle orientation predicts outer radial glial cell generation in human neocortex. Nat Commun 2013;**4**:1665.
- Lavoie H, Gagnon J, Therrien M. ERK signalling: a master regulator of cell behaviour, life and fate. *Nat Rev Mol Cell Biol* 2020;**21**:607–632.

- Lewitus E, Kelava I, Kalinka AT *et al*. An adaptive threshold in mammalian neocortical evolution. *PLoS Biol* 2014;**12**:e1002000.
- Li L, Clevers H. Coexistence of quiescent and active adult stem cells in mammals. *Science* 2010;**327**:542–545.
- Li X, Newbern JM, Wu Y et al. MEK is a key regulator of gliogenesis in the developing brain. *Neuron* 2012;**75**:1035-1050.
- Li S, Mattar P, Dixit R et al. RAS/ERK signaling controls proneural genetic programs in cortical development and gliomagenesis. J Neurosci 2014;**34**:2169–2190.
- Li X, Liu G, Yang L et al. Decoding cortical glial cell development. *Neurosci Bull* 2021;**37**:440–460.
- Lin Y, Yang J, Shen Z et al. Behavior and lineage progression of neural progenitors in the mammalian cortex. *Curr Opin Neurobiol* 2021;**66**:144–157.
- Liu DD, He JQ, Sinha R et al. Purification and characterization of human neural stem and progenitor cells. *Cell* 2023;**186**:1179–1194.e15.
- Long F, Zhang XM, Karp S *et al.* Genetic manipulation of hedgehog signaling in the endochondral skeleton reveals a direct role in the regulation of chondrocyte proliferation. *Development* 2001;**128**:5099–5108.
- Lui JH, Hansen DV, Kriegstein AR. Development and evolution of the human neocortex. *Cell* 2011;**146**:18–36.
- Lui JH, Nowakowski TJ, Pollen AA et al. Radial glia require PDGFD-PDGFRbeta signalling in human but not mouse neocortex. Nature 2014;**515**:264–268.
- Lun MP, Johnson MB, Broadbelt KG et al. Spatially heterogeneous choroid plexus transcriptomes encode positional identity and contribute to regional CSF production. J Neurosci 2015;**35**:4903–4916.
- Ma T, Wang C, Wang L et al. Subcortical origins of human and monkey neocortical interneurons. Nat Neurosci 2013;**16**:1588–1597.
- Ma L, Du Y, Hui Y *et al.* Developmental programming and lineage branching of early human telencephalon. *EMBO J* 2021;**40**:e107277.
- Martynoga B, Mateo JL, Zhou B *et al*. Epigenomic enhancer annotation reveals a key role for NFIX in neural stem cell quiescence. *Genes Dev* 2013;**27**:1769–1786.
- Micali N, Kim SK, Diaz-Bustamante M *et al.* Variation of human neural stem cells generating organizer states *in vitro* before committing to cortical excitatory or inhibitory neuronal fates. *Cell Rep* 2020;**31**:107599.
- Molnar Z, Clowry GJ, Sestan N *et al*. New insights into the development of the human cerebral cortex. *J Anat* 2019;**235**:432–451.
- Monuki ES, Porter FD, Walsh CA. Patterning of the dorsal telencephalon and cerebral cortex by a roof plate-Lhx2 pathway. *Neuron* 2001;**32**:591–604.
- Nowakowski TJ, Pollen AA, Sandoval-Espinosa C *et al.* Transformation of the radial glia scaffold demarcates two stages of human cerebral cortex development. *Neuron* 2016;**91**:1219–1227.
- Panchision DM, Pickel JM, Studer L *et al.* Sequential actions of BMP receptors control neural precursor cell production and fate. *Genes Dev* 2001;**15**:2094–2110.

- Pebworth MP, Ross J, Andrews M *et al.* Human intermediate progenitor diversity during cortical development. *Proc Natl Acad Sci U S A* 2021;**118**.
- Pelletier J, Thomas G, Volarevic S. Ribosome biogenesis in cancer: new players and therapeutic avenues. *Nat Rev Cancer* 2018;**18**:51–63.
- Pera EM, Ikeda A, Eivers E *et al.* Integration of IGF, FGF, and anti-BMP signals via Smad1 phosphorylation in neural induction. *Genes Dev* 2003;**17**:3023–3028.
- Picco N, Garcia-Moreno F, Maini PK *et al.* Mathematical modeling of cortical neurogenesis reveals that the founder population does not necessarily scale with neurogenic output. *Cereb Cortex* 2018;**28**:2540–2550.
- Pollen AA, Nowakowski TJ, Chen J et al. Molecular identity of human outer radial glia during cortical development. Cell 2015;**163**:55–67.
- Rakic P. A century of progress in corticoneurogenesis: from silver impregnation to genetic engineering. *Cereb Cortex* 2006;**16**:i3–17.
- Rao R, Salloum R, Xin M *et al*. The G protein Galphas acts as a tumor suppressor in sonic hedgehog signaling-driven tumorigenesis. *Cell Cycle* 2016;**15**:1325–1330.
- Rash BG, Grove EA. Patterning the dorsal telencephalon: a role for sonic hedgehog? J Neurosci 2007;**27**:11595–11603.
- Saulnier A, Keruzore M, De Clercq S et al. The doublesex homolog Dmrt5 is required for the development of the caudomedial cerebral cortex in mammals. *Cereb Cortex* 2013;**23**:2552–2567.
- Shi Y, Riese DJ, 2nd, Shen J. The role of the CXCL12/CXCR4/ CXCR7 chemokine axis in cancer. Front Pharmacol 2020;**11**:574667.
- Smart IH, Dehay C, Giroud Petal. Unique morphological features of the proliferative zones and postmitotic compartments of the neural epithelium giving rise to striate and extrastriate cortex in the monkey. *Cereb Cortex* 2002;**12**:37–53.
- Stepien BK, Vaid S, Huttner WB. Length of the neurogenic period-a key determinant for the generation of upperlayer neurons during neocortex development and evolution. Front Cell Dev Biol 2021;**9**:676911.
- Sun Y, Hu J, Zhou L *et al.* Interplay between FGF2 and BMP controls the self-renewal, dormancy and differentiation of rat neural stem cells. *J Cell Sci* 2011;**124**:1867–1877.
- Telley L, Govindan S, Prados J *et al.* Sequential transcriptional waves direct the differentiation of newborn neurons in the mouse neocortex. *Science* 2016;**351**:1443–1446.
- Theil T, Aydin S, Koch S *et al*. Wnt and Bmp signalling cooperatively regulate graded Emx2 expression in the dorsal telencephalon. *Development* 2002;**129**:3045–3054.
- Tole S, Ragsdale CW, Grove EA. Dorsoventral patterning of the telencephalon is disrupted in the mouse mutant extra-toes(J). *Dev* Biol 2000;**217**:254–265.
- Trevino AE, Muller F, Andersen J et al. Chromatin and gene-regulatory dynamics of the developing human cerebral cortex at single-cell resolution. *Cell* 2021;**184**:5053– 5069.e23.
- Ulloa F, Briscoe J. Morphogens and the control of cell proliferation and patterning in the spinal cord. *Cell Cycle* 2007;**6**:2640–2649.

- Vaid S, Camp JG, Hersemann L et al. A novel population of Hopx-dependent basal radial glial cells in the developing mouse neocortex. *Development* 2018;**145**:dev169276.
- Walenkamp AME, Lapa C, Herrmann K et al. CXCR4 ligands: the next big hit? J Nucl Med 2017;**58**:77S–82S.
- Wang B, Fallon JF, Beachy PA. Hedgehog-regulated processing of Gli3 produces an anterior/posterior repressor gradient in the developing vertebrate limb. *Cell* 2000;**100**:423–434.
- Wang F, Flanagan J, Su N et al. RNAscope: a novel in situ RNA analysis platform for formalin-fixed, paraffin-embedded tissues. J Mol Diagn 2012;**14**:22–29.
- Wang C, You Y, Qi D *et al.* Human and monkey striatal interneurons are derived from the medial ganglionic eminence but not from the adult subventricular zone. *J* Neurosci 2014a;**34**:10906–10923.
- Wang RN, Green J, Wang Z *et al.* Bone Morphogenetic Protein (BMP) signaling in development and human diseases. *Genes Dis* 2014b;**1**:87–105.
- Winkler CC, Yabut OR, Fregoso SP et al. The dorsal wave of neocortical oligodendrogenesis begins embryonically and requires multiple sources of Sonic hedgehog. J Neurosci 2018;**38**:5237–5250.
- Wyatt AW, Osborne RJ, Stewart H et al. Bone morphogenetic protein 7 (BMP7) mutations are associated with variable

ocular, brain, ear, palate, and skeletal anomalies. Hum Mutat 2010;**31**:781–787.

- Xiong W, He F, Morikawa Y et al. Hand2 is required in the epithelium for palatogenesis in mice. *Dev Biol* 2009;**330**:131-141.
- Yang X, Li C, Herrera PL et al. Generation of Smad4/Dpc4 conditional knockout mice. *Genesis* 2002;**32**:80–81.
- Yang L, Li Z, Liu G et al. Developmental origins of human cortical oligodendrocytes and astrocytes. *Neurosci Bull* 2022;**38**:47–68.
- Zhang X, Mennicke CV, Xiao G et al. Clonal analysis of gliogenesis in the cerebral cortex reveals stochastic expansion of glia and cell autonomous responses to Egfr dosage. Cells 2020a;9:2662.
- Zhang X, Xiao G, Johnson C *et al*. Bulk and mosaic deletions of Egfr reveal regionally defined gliogenesis in the developing mouse forebrain. iScience 2023;**26**:106242.
- Zhang Y, Liu G, Guo T et al. Cortical neural stem cell lineage progression is regulated by extrinsic signaling molecule Sonic hedgehog. *Cell Rep* 2020b;**30**:4490– 4504.e4494.
- Zhuo L, Theis M, Alvarez-Maya I et al. hGFAP-cre transgenic mice for manipulation of glial and neuronal function in vivo. Genesis 2001;**31**:85–94.