Untangling Cortical Complexity During Development

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ABSTRACT: The cerebral cortex is composed of billions of morphologically and functionally distinct neurons. These neurons are produced and organized in a regimental fashion during development. The ability of neurons to encode and elicit complex cognitive and motor functions depends on their precise molecular processes, identity, and connectivity established during development. Elucidating the cellular and molecular mechanisms that regulate development of the neocortex has been a challenge for many years. The cerebral cortical neuronal subtypes are classified based on morphology, function, intrinsic synaptic properties, location, connectivity, and marker gene expression. Development of the neocortex requires an orchestration of a series of processes including the appropriate determination, migration and positioning of the neurons, acquisition of layer-specific transcriptional hallmarks, and formation of precise axonal projections and networks. Historically, fate mapping, genome-wide analysis, and transcriptome profiling have provided many opportunities for the characterization of neuronal subtypes. During the course of this review, we will address the regimental organization of the cerebral cortex, dissect the cellular subtypes that contribute to cortical complexity, and outline their molecular hallmarks to understand cellular diversity in the cerebral cortex with a focus on the excitatory neurons.

KEYWORDS: Brain development, neural stem cells, neurogenesis, cerebral cortex

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Neuroepithelium to Neural Stem Cell Transition and Beyond

Early during vertebrate embryonic development, neural fate is induced in the ectoderm. The consequent patterning of the neural plate results in the formation of the central nervous system. The process of neurulation induces formation of the neural tube, a pseudostratified epithelial sheet of neuroepithelial cells (NEPs). It is the NEPs that are the precursors of the central nervous system including cerebral cortex, which is formed over an extended period of development. Important biological questions remain about how the complex structure of the cerebral cortex, which is composed of diverse neuron subtypes, is generated from a simple epithelial sheet of cells to form the most complex tissue of the body. At embryonic day 9 (E9), the neuroepithelium gives rise to neural stem cells (NSCs) that line the luminal surface of the vesicles of the neural tube. In mice, NSCs are located in the ventricular zone (VZ) and the ends of their basal processes remain in contact with the outer (pial) surface of the neural tube. This apical-basal polarity, which spans the thickness of the neural tube, requires the integrity of adherens junctions to segregate the apical and basolateral cell membrane and adhere neighboring NSCs to each other. The importance of adherence in NSC polarity is exemplified by the knockdown of the adherens junction-associated protein Afad1 (Af6). Af6 depletion leads to a loss of adherens junctions and disturbed cell polarity.

At the onset of neurogenesis, the NEPs generate radial glial cells (RGCs) and short neural precursors. The somata of these cells remain within the VZ but migrate radially along the apical-basal process through the zone during cell division in a process referred to as interkinetic nuclear migration (INM). The location of the soma within the VZ is cell cycle dependent. During M-phase, the cell body is positioned apically at the luminal surface of the neural tube (Figure 1). As the cell progresses through G1-phase of the cell cycle, the cell body moves radially to the VZ boundary with the overlying subventricular zone (SVZ) and forming cerebral cortex. S-phase and DNA replication occur at the basal boundary of the VZ followed by migration of the cell body back to the luminal surface of the neural tube during G2 to initiate mitosis. Primary cilia in the apical membrane project into the vesicles and detect factors and signals in the fluid filling the neural tube and these support apical-basal polarity. The oriented cell polarity is important for determining the structure of the cerebral cortex. Disruption of the small GTPase, ADP ribosylation factor–like GTPase 13B (Ar13b), results in loss of cell polarity and the cortical wall is generated in an inverted fashion. M-phase of Ar13b-deficient RGCs is no longer restricted to the luminal surface but also occurs at the basal, pial surface and neurons migrate centripetally to the VZ.

During early phases of neurogenesis, embryonic days 10.5 to 11.5 (E10.5-11.5) in mice, NSCs undergo symmetric stem cell divisions, expanding the pool (Figure 1). This is referred to as the “neural expansion” phase of cortical development. Later, NSCs progressively undergo asymmetric cell divisions, allowing for both self-renewal and the generation of committed daughter cells (Figure 1). The transition from symmetric stem cell to asymmetric neurogenic divisions during neurogenesis is...
associated with a lengthening of primarily G1-phase of the cell cycle. However, the S-phase of the NSCs in the symmetric dividing, expansion phase is longer than those in the asymmetric dividing neurogenic phases. Hence, although the precise function of INM and the changing in cell cycle phase length are not understood, it seems that they play an important role on the control of the sequential switching of NSCs from a symmetric self-renewing mode to the asymmetric division mode that drives the production of neurons.

During the neurogenic phase of cortical development, the self-renewal and generation of committed daughter cells have to be tightly controlled. Loss of self-renewing NSC daughter cells would purge the stem cell pool. Conversely, a failure to generate sufficient neuronal-determined precursors would severely affect neuronal composition and cortical layering. During early stages of cortical development, some asymmetric stem cell divisions generate one NSC daughter and a neuron directly. This is referred to as direct neurogenesis. However, as neurogenesis progresses, the daughter cell that is committed to differentiate and leaves the stem cell pool becomes a basal progenitor (BP) and migrates to the forming SVZ (Figure 1).

Distinct Stem and Progenitor Populations Contribute to Cortical Development
Throughout neurogenesis, another VZ population of dividing cells called the short neural precursors contributes to the progenitor pool. Short neural progenitors have either a short or no basal process at all but retain the apical process and contact to the lumen of the neural tube. These cells are morphologically, ultrastructurally, and molecularly different from the NSCs and have been observed to undergo direct neurogenesis, generating neurons without passing through a BP state.

In higher mammals including ferrets, primates, and humans, additional intermediate progenitor populations have evolved, and although they also reside in the SVZ, they have different morphologies and larger cell fate potentials compared with the classic BPs in mice. In fact, in primates, some of these intermediate progenitors even display NSC potential and are even referred to as outer RGCs (oRGCs). The oRGCs are morphologically distinct, unipolar, and retain only the basal process with no connection to the VZ and neural tube lumen (Figure 2). They also do not express the apical membrane constituents associated with VZ NSCs and RGC including prominin1 (CD133), Par3 family cell polarity regulator (Par3), or atypical protein kinase C (aPKC). They have a long basal phospho-Vimentin (pVim)-positive process that extends toward the pia and retain the basal fiber throughout the duration of cell cycle. Similar to VZ RGCs, the soma of oRGCs also moves during cell divisions but this movement is distinct to the INM of VZ NSCs. The soma of oRGCs moves basally and once translocation is complete, they divide mostly by self-renewing, asymmetric divisions, and push the boundary of the outer SVZ (OSVZ) outward expanding the SVZ (Figure 2). Self-renewing oRGCs continue to proliferate, whereas the daughter cells differentiate into neurons.

These SVZ progenitors in primates are the major source of expansion and neurogenesis in the cerebral cortex and are responsible for the massive evolutionary expansion of the cortical gray matter, neuron number, and cortical surface. Indirectly, these SVZ progenitors are responsible for the increase in functional capacity of the cerebral cortex in primates. The coexistence of oRGC cells and VZ RGCs demonstrates the distinct germinal zones in higher mammals, highlighting the mechanisms of increased neuron production, relevant for the formation of bigger brains (Figure 2). Here, we will focus on cortical development in the mouse and refer to excellent review focusing on primate and human cortical development.

Basal progenitors are intermediate, transient amplifying cells that undergo 1 or 2 divisions before giving rise to neurons (Figure 2). The BPs are in one of the main zones of amplification.
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and neurogenesis in the developing mouse cortex. As neurogenesis reaches completion, the NSCs start to generate other cell lineages, oligodendrocytes, astrocytes, and ependymal cells.\textsuperscript{21,22} This is referred to as the “gliogenesis phase” of cortical development. The transition from neurogenesis to gliogenesis is associated with a downregulation of the Golgi-derived apical trafficking and VZ NSCs lose tight junctions while keeping intact the adherens junctions.\textsuperscript{23,24} This is followed by the gradual expression of the astroglial hallmarks including glial fibrillary acidic protein (GFAP) in the mouse.\textsuperscript{22,24–26} Although the mechanisms of the neurogenic to gliogenic phase transition are not clearly understood, Notch signaling and its downstream targets, the bHLH (basic helix-loop-helix) transcription factors including the Hes proteins, and the growth factor Fgf10 are necessary for this transition.\textsuperscript{27–29}

Because the generation of neurons from BPs results in the expansion of the neuronal progenitor pool enabling the production of many neurons from a restricted population of NSCs, their role is crucial in the expansion of the cortex.\textsuperscript{28,30,31} In the mouse, BPs can undergo symmetric divisions and generate 2 neuronal daughter cells.\textsuperscript{31} However, evidence suggests that some, if not all, may also undergo 1 or 2 rounds of self-renewing

\textbf{Figure 2.} Scheme illustrating the composition and laminar organization of the developing human cortex, in comparison with mouse cortex. The human cerebral cortex develops in a similar fashion to that of the mouse. One exception is the expansion of the subventricular zone (SVZ) to form the outer SVZ (oSVZ). The oSVZ in humans is the main zone of amplification. In addition to the neural stem cells (NSCs) and basal progenitors (BPs) of the developing mouse cerebral cortex, the human has addition progenitors, outer radial glial cells. CP indicates cortical plate; IZ, intermediate zone; MZ, marginal zone; SP, subplate; SVZ, subventricular zone; VZ, ventricular zone.
cell divisions. Basal progenitors are defined based on their position in the SVZ, their lack of polarized morphology and expression of the transcription factors, Eomesodermin (Eomes or Tbr2), Btg antiproliferation factor 2 (also called Tis21), Cut-like homeobox 1/2 (Cux1/Cux2), and special AT-rich sequence binding protein (Satb2) and the non-coding RNA Svet1. Because the different progenitor cell types are localized to different niches and thus likely exposed to different combinations of cues from their microenvironment, it is imperative to study the role of their niche in controlling their proliferation and fate commitment. This cellular heterogeneity requires a deeper understanding of the cell fate identities and commitments.

Symmetric and Asymmetric Cell Divisions

Neural stem cells of the developing cerebral cortex display multiple modes of cell division. Initially, the major form of divisions is symmetric stem cell divisions, generating 2 daughter cells that retain stem cell potential and reenter cell cycle. As development progresses, the stem cell divisions are slowly superseded by asymmetric neurogenic divisions where 1 daughter remains a stem cell and reenters the cell cycle within the VZ, whereas the other is committed to differentiate and will leave the VZ (Figure 1). The third mode is the symmetric neurogenic division where both daughter cells will differentiate thereby depleting the stem cell pool. The balance between these different forms and outcomes of cell division are temporally and spatially regulated which is necessary to control correct cortical development.

The molecular basis of symmetric and asymmetric divisions and the transition from self-renewing to differentiating modes of cell division are not understood. It has become clear that the orientation of the mitotic spindle plays an important role in the type of division and the fate of the respective daughter cells generated (Figure 1). A cleavage plane bisecting the apical membrane of the NSCs, including inheritance of junctional complexes by both daughters, contributes heavily to maintenance of stem cell potential (Figure 1). During symmetric divisions of NSCs, the cleavage plane is oriented perpendicular to the ventricular surface (Figure 1). This spatial organization of the mitotic spindle requires a deeper understanding of the cell fate identities and commitments.

This results in an unequal partition of Par3 into the 2 daughter cells, and the sibling cell receiving less Par3 protein exits cell cycle and differentiates. In addition, asymmetric cell division is accompanied by an unequal distribution of fate determinants between the daughter cells. These components include mediators of Notch signaling, the Notch ligand delta-like 1 (Dll1), Mind bomb, and Numb. Segregation of Notch components including inhibitors of the pathway leads to differential Notch signaling between daughter cells. Notch signaling plays a critical role in NSC maintenance and differentiation by regulating cell proliferation and fate determination. Notch activates the expression of Hes genes which encode bHLH transcriptional regulators. Hes-related proteins repress expression of the proneurogenic transcription factors including neurogenins (Ngns) and Ascl1. Thus, activation of Notch signaling inhibits differentiation of NSCs by suppressing transcription factors required for neurogenesis. In addition, Notch signaling regulates cell cycle progression via regulation of Ascl1 expression. Ascl1 not only controls neurogenic differentiation but is also involved in entry of NSCs into cell cycle.

In addition to Notch, some cytoplasmic proteins show differential distribution on asymmetric division. Staufen is a double-stranded RNA-binding protein which is pivotal in asymmetric cell fates in Drosophila neural development. Staufen is selectively segregated into the differentiating daughter cells on asymmetric self-renewing cell division. Staufen binds messenger RNAs that encode proteins crucial in cell cycle exit and differentiation. Furthermore, the transcription factor Pax6 promotes asymmetric neurogenic cell division. Pax6-mutant NSCs show a defective cell cycle exit and an increase in self-renewing capacity.

In addition to molecular segregation, the orientation of the mitotic spindle plays an important role in fate determination. In NSCs, the mitotic spindle poles oscillate around their final positions before anaphase is initiated. This dynamic movement of the spindle seems to be important in determining the cleavage plane and then the segregation of intracellular components. Only subtle changes in spindle orientation can cause major shifts in the plane of cytokinesis and thereby the inheritance of membrane compartments and cell fate determinants. Mutations in the abnormal spindle-like microcephaly-associate (Aspm) gene severely affect cerebral cortical size and reduce the volume of the cerebral cortex in primates. Aspm is important for spindle orientation and control in the division modes of symmetric versus asymmetric cells.

Inheritance of the apical plasma membrane of NSCs has an influence on cell fate. During symmetric cell divisions, both daughters acquire apical membrane and junctional components. When only 1 daughter cell inherits the apical plasma membrane, for example, when the cleavage plane is parallel to the neural tube luminal surface, that daughter remains as an NSC, whereas the other sibling that does not receive apical
membrane and adherens junctions from the mother cell will exit the VZ and commit to differentiation. The SNARE-mediated membrane fusion machinery controls NSC fate specification. A hypomorphic missense mutation in α-SNAP (α-soluble N-ethylmaleimide-sensitive fusion protein [NSF] attachment protein) causes NSCs to prematurely switch from symmetric proliferative to asymmetric neurogenic divisions. This is primarily due to an impaired apical protein localization affecting the Golgi-derived membrane traffic necessary for NSC proliferation. In addition, NSCs in these mice show distribution of apical β-catenin along the adherens junctions and phenocopying of conditional β-catenin null mutant mice. Hence, β-catenin plays a role not only in the control of cell cycle but also in the choice between symmetric and asymmetric divisions.

Regulation and Cell Fate Commitment
The stem and progenitor cells in the dorsal VZ of the anterior neural tube generate the multiple classes of projection neurons that make up the future cerebral cortex in sequential waves. In the dorsal cerebral cortex, neurogenesis commences around E10.5 in mice. The earliest born neurons segregate from the NSCs in the VZ and migrate radially to the pial surface forming the preplate. Later born neurons migrate into the preplate, splitting it into the marginal zone and the subplate (Figure 2). Throughout neurogenesis, newborn neurons migrate into the cortical plate (CP), through the preformed layers of earlier born neurons, and as such the early born neurons form the deep layers and the later born neurons form upper layers. The detailed molecular cascade that determines neuronal cell fate commitment, an excitatory neuron subtype specification, is largely unknown. Different models have been proposed to explain the temporal dynamics of neuronal specification in the dorsal cortex. The “common progenitor model” implies that a single type of NSC sequentially gives rise to the different neuron subtypes overtime during neurogenesis and that neuron fate is determined by time (Figure 3A). According to the “multiple progenitor model,” multiple stem cell types coexist and are predetermined to generate specific neuron subtypes. In the multiple progenitor model, the fate of the stem cell and the type of neuron generated are determined by the NSC type (Figure 3B). There is experimental evidence supporting both models.

The Common Progenitor Model
McConnell demonstrated that NSC fate becomes restricted over time during development. By performing elegant heterochronic transplantation experiments initially in ferrets, they demonstrated that early developmental stage progenitors have a greater potential and can generate early and late neuronal subtypes when grafted into hosts. Conversely, late–stage progenitors have a more restricted potential and a reduced capacity to form early neuronal types in host embryos. This implies that NSCs lose their potential to generate deep layer neurons with time. In support of this model, clonal in vitro experiments showed the sequential generation of deep and upper layer neurons from NSCs supporting initial multipotency and subsequent fate restriction over time. Finally, retroviral labeling and lineage tracing of individual NSCs supported progressive fate restriction in vivo. More recently, genetic labeling...
in the developing mouse cerebral cortex following expression of the transcription factor Fezf2 (enriched in deep cortical layer V neurons) revealed that Fezf2-expressing NSCs generate deep, upper layer neurons and glial cells. Instructive roles of factors such as Fezf2 in NSCs can have major implications in cell fate commitment during neurogenesis. Ectopic expression of Fezf2 can direct NSCs to differentiate into deep layer neurons and reverse late fate commitment.

The Multiple Progenitor Model

An alternative model for fate specification proposes independent, fate-restricted lineages of NSC that generate specific neuronal subtypes and have limited potentials. Evidence suggests that many transcription factors expressed during cortical development instruct fate commitment during neurogenesis. Ectopic expression of Fezf2 can direct NSCs to differentiate into deep layer neurons and reverse late fate commitment.

Neuronal Diversity and Transcriptional Dynamics in Cortical Layering

The cerebral cortex is an isocortex composed of 6 clearly defined layers. The newborn excitatory neurons migrate out of the VZ along the radial processes of the NSCs (Figures 4 and 5). The immature neurons reach the CP by migrating through their earlier born siblings. On reaching the pial surface, the immature neurons leave the RGC process and differentiate and form neurons of their specific cortical layer. Hence, the isocortex of the cerebrum is formed in an inside-out fashion, with early born neurons forming the deep layers while the later born neurons generating the upper layers (Figure 5). The neuronal type and their location within the isocortex are critical for function. The interneurons of the cerebral cortex originate from the ventral telencephalon and migrate to their final destination in the cerebral cortex (we refer the reader to excellent recent reviews on cortical interneuron development and will not cover the topic here).

The major types of cortical projection neurons can be defined by their connectivity and projection patterns depending on whether they project through associative, commissural, and corticofugal projection fibers. Associative projection neurons project their axons within a single cerebral hemisphere connecting local areas or proximal gyri. Commisural, callosal projection neurons are localized primarily in layers II/III, V, and VI of the 6-layered isocortex. They extend their axons from 1 hemisphere to neurons in the contralateral hemisphere. The axons project through the corpus callosum, the major commissural connection between the hemispheres, or through the anterior or posterior commissures. The commisural neurons are further subdivided based on the projection destinations. Corticofugal neurons include the subcerebral neurons,
which are the largest pyramidal neurons and extend projections to subcortical structures including the brainstem and spinal cord. The corticofugal projections include corticopontine, corticospinal, and corticotectal neurons. Another subtype of corticofugal neurons is the corticothalamic neurons, which populate the layer VI of the isocortex with a small population in layer V and extend their projections to different nuclei of the thalamus. Hence, the regulation of neuron subtype formation and the temporospatial control of neurogenesis are critical for brain function. Numerous neocortical determinants are expressed along the dorsolateral wall of the cortex, on the induction of neurogenesis. Key factors including forkhead box G1 (FoxG1), LIM homeobox 2 (Lhx2), Emx2, and Pax6 define and control the neocortical progenitor domains along the dorsal and ventral axis. Ablation of the dorsal progenitor domain determinants Pax6 and Emx2 results in expansion of the ventral domains. Pax6 and T-cell leukemia homeobox (Tlx) regulates the cell fate decisions in the VZ and the loss of function of these factors leads to a thicker superficial cortex. Hence, the NSCs and progenitors of the cerebral cortex are determined by their expression of axial-specifying factors but these fates are not fixed or restricted as loss of these determinants results in alternate fate acquisition.

The transcription factors Tbr1, Ctip2, Sox5, Fez2, Satb2, Cux1, Cux2, Brn1, Brn2, and others have been studied extensively and determined to be key determinants of neuronal specification. These transcription factors are often used as markers of specific cortical neuron populations and layers and are expressed in waves during cortical development. Some of these markers are expressed in specific neuronal subtypes within a layer or are expressed in more than one neuronal subtype and layer. For example, the Ets-related protein B1 (Er81/Etv1) is expressed in cortico-cortical and subcerebral projection neurons of layer V. Conversely, LIM domain only 4 (Lmo4) is selectively expressed in callosal neurons of layers II/III.

Some of these fate-determining and defining factors seem to be coexpressed initially and their expression becomes restricted and refined later in neuronal differentiation. For example, in mice, postmitotic deep layer neurons coexpress Ctip2 and Satb2 at E13.5. As development progresses, these deep layer neurons express either Ctip2 or Satb2 and become fate restricted to form subcerebral projection neurons.
or corticothalamic projection neurons, respectively.\textsuperscript{35,72,82,83} Ngn1 and Ngn2 are 2 proneural transcription factors and induce neurogenesis; however, Tbr1- and Er81-expressing deep layer neurons are still generated in their absence.\textsuperscript{84,85} It is likely that other proneural transcription factors are able to compensate for the loss of Ngn1 but the exact mechanism remains to be defined.

**Epigenetic and Transcriptional Interplay During Neurogenesis**

The gene expression in NSCs regulated not only by transcription factors but also by epigenetic mechanisms. DNA methylation and histone modifications are involved in spatial and temporal gene expression during neurogenesis and the switch from neuronal to glial fate.\textsuperscript{86-88} New methods for genome-wide methylation mapping facilitates investigation of epigenetic landscapes that control lineage commitments and fate decisions during neuronal specification. Epigenetic regulation of critical transcription factors in NSCs play important roles in the regulation of cell fate and neurogenesis. The expression of epigenetic regulators including the high-mobility group (HMG) proteins during early phases of cortical development regulate chromatin state and methyltransferase activity including enhancer of zeste 2 polycomb repressive complex 2 subunit (Ezh2).\textsuperscript{86-88} This suggests that the chromatin in early NSCs is in a more "open" state than that of later NSCs.\textsuperscript{87} The transcription factor Tbr2, which is critical for BP generation and differentiation, associates with the histone demethylase Jmjd3 (also called Kdm6b). Tbr2 directs Jmjd3-dependent chromatin remodeling to specific gene loci promoting the removal of H3K27me3 chromatin marks.\textsuperscript{89} This emphasizes the additional level of control that Tbr2 has on neuronal specification by regulating the epigenetic marks at specific promoter and enhancer sites.\textsuperscript{89}

Hes5 is a pivotal mediator of Notch signaling and inducer of maintenance of NSCs by blocking proneural transcription factor expression. However, the expression of Hes5 also depends on glial cell missing homolog (Gcm) and active DNA demethylation during neurogenesis.\textsuperscript{90} Loss of Gcm prevents upregulation of Hes5 and the formation of definitive NSCs between E7.5 and E8.5.\textsuperscript{90} Pax6 interacts with BAF155 and BAF170, components of the ATP (adenosine triphosphate)—dependent multi-subunit mSWI/SNF nucleosome remodeling complexes in NSCs. At the onset of neurogenesis, BAF155 and BAF170 compete and modify euchromatin structure.\textsuperscript{91,92} This leads to the recruitment of the Pax6/RE1 silencing transcription factor (REST)—corepressor complex to the Pax6 targets, transducin-like enhancer of split 1 (Tle), Eomes, and Cux2, and repressing their expression. This prevents the formation of BPs and late NSCs.

During peak of neurogenesis, the chromatin remodeler sucrase nonfermenting—like protein 1 (Snf2l) represses expression of FoxG1, which leads to the derepression of the cell cycle regulator p21 and promotes neuronal differentiation by inducing cell cycle exit.\textsuperscript{91,92} During later stages of neurogenesis, the polycomb proteins repress Ngn1 expression to trigger the NSC fate switch from neurogenesis to astrogliogenesis.\textsuperscript{86} The NSC switch to gliogenesis is associated with the expression of the astrocytic protein GFAP. DNA methylation of the \textit{Gfap} promoter prevents its premature activation. Notch signaling induces demethylation of the \textit{Gfap} promoter through the transcription factor nuclear factor I (NFI), by dissociating associated DNA methyltransferases and thereby supports generation of astrocytes.\textsuperscript{28,93,94} Further analysis of the interplay between epigenetic and transcriptional dynamics during cortical development may contribute to a greater understanding of novel mechanisms and dysregulation during brain disorders.

**Signaling Dynamics During Neurogenesis**

Various signaling pathways impinge on downstream effectors and regulate NSC fate decisions during neurogenesis. Among these pathways are Notch, Wnt, Shh, fibroblast growth factors (FGFs), transforming growth factor \(\beta\) (TGF-\(\beta\)), retinoic acid (RA), and Hippo. How the cross talk between these signaling pathways and the integration of their signals on target genes governs complex cell fate choices is unclear.

Notch was first discovered in \textit{Drosophila}, and the signaling pathway is evolutionarily highly conserved.\textsuperscript{95} It plays critical roles in NSC maintenance and differentiation. Conditional loss of Notch receptors results in the precocious differentiation, impaired survival, and aberrant migration of NSCs.\textsuperscript{45,48} Notch signaling regulates neurogenesis through lateral signaling between neighboring cells. In the presence of Notch ligands, delta-1 (Dll1) and Jagged1 (Jag1), the receptor is cleaved at S2 and S3 sites by ADAM metallopeptidase domain 10 (ADAM10) and \(\gamma\)-secretase, respectively.\textsuperscript{96,97} This results in the release of the Notch intracellular domain (NICD). The NICD has nuclear localization signals and mediates the transcription of \textit{Hes} genes via DNA-binding factor recombination signal-binding protein for immunoglobulin kappa J region (RBPJ) complex.\textsuperscript{96,98} On interaction of NICD with RBPJ, the preformed repressor complex on target genes is disrupted and replaced by a coactivator complex including mastermind-like protein-1 (Maml) and histone acetyltransferase (HAT, p300).

These changes facilitate the recruitment of RNA polymerase II and initiate transcription of target genes including \textit{Hes} and \textit{Hes5}.\textsuperscript{99} The Hes factors in turn downregulate expression of the genes encoding the proneural bHLH factors Ngn and Achaete-Scute family bHLH transcription factor 1b (Ascl).\textsuperscript{100,101} Hence, active Notch signaling maintains NSCs in a proliferative state and represses differentiation in the mammalian brain.\textsuperscript{49,101,102} The expression of Hes1 and Hes5 oscillates in the VZ due to an auto-inhibitory feedback loop.\textsuperscript{28,103} The oscillations in the expression of the Hes factors in turn induce oscillatory expression of proneural factors.\textsuperscript{50,97,104} This dynamic expression seems to be critical in modulating the functions of
the proneural transcription factors in controlling proliferation and neurogenic differentiation. Although the source of Notch ligands is not clear, NSCs, BPs, and newborn neurons express Delta and Jaggeds as well as the E3 ligase Mind bomb that potentially promote Notch signaling in the NSCs.

Wnt signaling is involved in the patterning and development of many tissues including the nervous system. Wnt1 and Wnt3a are expressed by cells at the dorsal midline of the developing neural tube. In the absence of Wnt1, midbrain structures fail to form and Wnt3a-mutant mice do not form a hippocampus likely due to the reduced proliferation of hippocampal precursors. Wnt ligands bind the receptor complex Frizzled/LRP5/6 leading to stabilization of cytoplasmic β-catenin. β-catenin translocates to the nucleus and binds to target genes via LEF/TCF factors and recruits histone acetyltransferases. Transgenic overexpression of β-catenin induces enhanced proliferation of cortical neural progenitors leading to an increase in cortical neurons and surface area. During early neurogenesis, Wnts play an active role in symmetric divisions, whereas later, during neurogenesis, Wnts are implicated in neuronal differentiation through expression of N-myc, which in turn represses the Notch signaling. Wnt signaling can also induce the expression of Ngn1 and NeuroD1 thereby counteracting Notch signaling and promoting neuronal differentiation. Thus, Wnt and Notch compete to regulate proneural gene expression and the maintenance and differentiation of NSCs. The differential dynamics of signaling pathways impinging on same or different downstream effectors could be cell type or phase specific.

Fibroblast growth factor signaling has been long known to be involved in area specification in the brain. Many Fgfs are expressed in the developing cerebral cortex. Fgfs are expressed along the rostral midline of the neocortex in the commissural plate between E9.5 and E12.5, suggesting the presence of a rostral, Fgf-secreting signaling center. The Fgf signals play important roles in anterior-posterior patterning of NSCs and promote proliferation. In addition, Fgf signaling can regulate Hes1 transcription thereby synergizing with and promoting Notch signaling. Fgf18 is expressed in the CP between E13.5 and E16.5, although its role remains unclear.

Three of the classical receptor tyrosine kinase Fgf receptors (Fgfr1-3) are expressed by NSCs. Fgfr1 is expressed higher by rostral NSCs than caudal NSCs, whereas Fgfr2 and Fgfr3 are expressed higher caudally than rostrally. In mice, loss of Fgfr1 function results in the loss of rostral identity, indicating that Fgfr1 acts as a secreted rostral morphogen. Conversely, Fgfr2 is expressed higher in the dorsal forebrain than in the ventral, thus contributing to the dorsoventral patterning of the developing brain. Loss of function of Fgfr2 changes dorsal cortex specification. Pea3-ETS transcription factors are downstream of the Fgf signaling pathway and ectopic expression of Fgf18 induces their expression with phenotypic changes in neuronal migration. Pea3-ETS transcription factors are expressed in gradients high rostral to low caudally implying a role in axial patterning in the cerebral cortex.

Transforming growth factor β/bone morphogenetic protein (BMP) signaling pathways play important roles in neurogenesis. Both TGF-β and BMP are expressed in the dorsal cerebral cortex during embryonic neurogenesis and regulate proliferation, survival, differentiation, and migration in the cerebral cortex. The BMP binds to the BMP receptor (BMPRI) on the cell surface and induces phosphorylation of Smad family transcription factors. The BMP signaling inhibits neuronal differentiation and promotes glial differentiation during corticogenesis. Both BMP and Notch may converge on some cellular processes, for example, they could impinge on some similar targets such as Hes3 and inhibitor of DNA-binding factor genes (Ids), as observed during adult neurogenesis.

Retinoic acid, a derivative of vitamin A, is involved in neuronal differentiation. Retinoic acid binds nuclear receptors of the RA receptor family (RARs α, β, and γ) that regulate the expression of target genes that contain an RA response element. The interaction of RA with the RAR bound as a repressor complex to target genes releases corepressor proteins and recruits histone acetyltransferases. However, RA can also induce rapid and transient activation of a cascade of kinases including the MAPK and ERK pathways which contribute to coregulation of the RAR target genes by phosphorylation of cofactors and histones. Dietary depletion of vitamin A in pregnant mothers results in embryonic defects, including delayed development and reduced cortical hemispheres, with a reduction in neuron-specific class III β tubulin (β-tubulinIII) expression and lower levels of Harvey rat sarcoma viral oncogene (HRas) protein in the intermediate zone and CP regions. The reduction in HRas levels is rescued by supplementing the embryos with RA indicating a stabilization of HRas by RA. Retinoic acid deficiency affects neuronal migration to cortical layers V to III during development. This impaired migration also results in neuronal fate switching to layer II neuron subtypes. The RA pathway also cross-talks with Wnt signaling at the level of β-catenin. The Wnt-RA axis is most prominent at the rostral end of the developing cerebral cortex, implying a potential role of RA in arealization of the forebrain.

Hippo signaling regulates size and homeostasis in many organs and tissues. The Hippo signaling pathway is a cascade of kinases that converge onto the control of the transcriptional coregulators Yap and Taz. The Hippo kinases and Yes-associated protein/transcriptional coactivator with PDZ-binding motif (Yap1/Taz) are regulated at different levels by different stimuli including G protein–coupled receptor signaling, cell adhesion, mechanical stress, and changes in cellular energy status. The Hippo kinase cascade can be activated by activation of the macrophage-stimulating 1/2 (Mst1/2) and
large tumor suppressor 1/2 (Lats1/2) kinases. These serine/threonine kinases phosphorylate Yap1 and Taz. Phospho-Yap/Taz is targeted to degradation. If Lats1/2 is inactive, then Yap/Taz is dephosphorylated and translocated to the nucleus where they interact with multiple transcriptional regulators.130 Yap/Taz interacts with β-catenin and Smads and thus coregulates both the Wnt and TGF-β pathways to regulate gene expression.130 The TEA domain transcription factors (Tead) are key targets and mediators of the Hippo pathway and critical effectors of Hippo-regulated target gene expression.125,129 In NSCs, the Hippo pathway plays a niche role and regulates the communication between neighboring cells. The expression of fat tumor suppressor homologue (Fat4) and Dachsous (Dchs), the upstream receptor, and ligand of the pathway increases NSC proliferation and reduces differentiation.132 However, the targets and mediators of the Hippo pathway in NSCs and cortical development remain unclear. Hence, future analysis of the Hippo pathway and its control of NSC maintenance, commitment, and differentiation could uncover novel interactions and functions.

Although the origins of many of the factors described above are not clear, the cerebral spinal fluid is an obvious source. Growth factors and morphogens released into the cerebral spinal fluid can influence NSC proliferation and fate. Some of these factors are produced and released by the choroid plexus and their expression is dynamic during cortical development (reviewed by Lehtinen and Walsh133 and Johansson134). In addition, neuronal inputs from subcortical regions of the brain have also been shown to influence neural progenitor proliferation and maintenance.135,136 This suggests that not only the local environment of the developing cerebral cortex is affecting the production of neurons in the dorsal cortex but more distant brain regions may also play a critical role in the control of cortical NSC fate.

In summary, rather unsurprisingly, development of the brain and particularly the cerebral cortex incorporates many different signaling pathways. Here, we just cover a few but due to the complexity of the cerebral cortex and the need for precise NSC proliferation, fate commitment and differentiation, the balance, and interaction of these pathways will be critical. Hence, a deeper understanding of the signaling pathways and their underlying downstream mechanisms is required to develop a model of how NSCs integrate different signals to regulate development of the brain.

Conclusions
In this review, we have tried to highlight some of the main developmental processes and signaling mechanisms controlling cerebral cortical development. From decades of work, it is clear that transcription factors and signaling are key regulators of NSC generation of the cerebral cortex. However, there remains much to be learnt about how these pathways interact and converge to impose the precise regulation needed to form the complex structure of the cortical isocortex from a simple pseudostratified sheet of NEPs. State-of-the-art technologies employing high-throughput single-cell RNA sequencing platforms have allowed a considerable increase in resolution to the single-cell level. These techniques provide a comprehensive understanding of single cells isolated from the brain, facilitating the extrapolation of intrinsic molecular architecture to function. With the advent of high-throughput single-cell omics and lineage tracing in vivo, the future looks demanding but bright and exciting for elucidating the mechanism of development of the cerebral cortex.

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