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Temporal and sequential transcriptional dynamics define lineage shifts in corticogenesis

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Abstract

The cerebral cortex contains billions of neurons, and their disorganization or misspecification leads to neurodevelopmental disorders. Understanding how the plethora of projection neuron subtypes are generated by cortical neural stem cells (NSCs) is a major challenge. Here, we focused on elucidating the transcriptional landscape of murine embryonic NSCs, basal progenitors (BPs), and newborn neurons (NBNs) throughout cortical development. We uncover dynamic shifts in transcriptional space over time and heterogeneity within each progenitor population. We identified signature hallmarks of NSC, BP, and NBN clusters and predict active transcriptional nodes and networks that contribute to neural fate specification. We find that the expression of receptors, ligands, and downstream pathway components is highly dynamic over time and throughout the lineage implying differential responsiveness to signals. Thus, we provide an expansive compendium of gene expression during cortical development that will be an invaluable resource for studying neural developmental processes and neurodevelopmental disorders.

Keywords cortical development; lineage specification; networks; signaling pathways; transcriptional landscape

Subject Categories Chromatin, Transcription & Genomics; Development; Neuroscience

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Introduction

The cerebral cortex of vertebrates is an isocortex, composed of six layers of morphologically and functionally distinct neurons. During development, cortical NSCs pass through consecutive stages of mitotic expansion, deep- to upper-layer neurogenesis and then gliogenesis. Most neurons are generated from NSCs through a transient progenitor population, the BPs. Maintenance of progenitor potential and control of cortical fate commitment are regulated through the integration of dynamic signaling pathways organized in space and time, which induces an elaborate interplay between downstream transcriptional networks. Although the molecular nature of mature neurons within the six cortical layers has been described, their corresponding progenitors have not been clearly characterized.

Different hypotheses have been proposed to explain the heterogeneity in the cortical precursor cells in terms of temporal expansion and differentiation potential (Hevner *et al*, 2003; Molyneaux *et al*, 2007; Woodworth *et al*, 2012; Lodato & Arlotta, 2015). One hypothesis states that NSCs switch their fate temporally in

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coherence with the time points of neurogenesis and thus generate neurons of successive layers of the cortex followed by glial cells (Guo *et al*, 2013). An alternate hypothesis proposes that NSCs are a multipotent cell pool, wherein each cell would be guided by intrinsic and extrinsic signals to generate a specific selection of neuronal subtypes or glial cells, and these different progenitors are recruited in a sequential manner (Franco *et al*, 2012). Whether one or both hypotheses are correct remains a major debate.

As RNA sequencing (RNA-Seq) technology increased over recent years, so has our acceptance of an increasing repertoire of cell types present during cortical development. Particularly single cell sequencing techniques have allowed an ever more detailed transcriptomic analysis of cortical precursor cells (Desai & McConnell, 2000; Fode *et al*, 2000; Hevner *et al*, 2003; Haubensak *et al*, 2004; Arlotta *et al*, 2005; Gotz & Huttner, 2005; Molyneaux *et al*, 2007; Stancik *et al*, 2010; Lui *et al*, 2011; Greig *et al*, 2013; Han & Sestan, 2013; Paridaen & Huttner, 2014; Arber *et al*, 2015; Chuang *et al*, 2015; Lodato & Arlotta, 2015; Pollen *et al*, 2015; Liu *et al*, 2016; Telley *et al*, 2016, 2019; Ecker *et al*, 2017; Johnson & Walsh, 2017; Nowakowski *et al*, 2017; Mukhtar & Taylor, 2018; Rosenberg *et al*, 2018; Zeisel *et al*, 2018; Di Bella *et al*, 2021; La Manno *et al*, 2021). Frequently, cells are isolated based on positional information or temporal labeling, and this is used to delineate cell type and predict potential (Telley *et al*, 2019; Di Bella *et al*, 2021). Although these approaches have been very successful in providing a framework, our understanding of transcriptional programs during brain development and cortical patterning is not complete and some critical points remain. One major challenge is the extreme complexity of the system and the differences in technical approaches undertaken. As RNA-Seq takes a snapshot in time of gene expression in a population or of single cells, it is challenging to predict the past and future gene expression profile of a cell population. Elegant labeling procedures have provided some insight into cell diversity in the NSC pool and allowed analysis of specific gene function (Telley *et al*, 2016, 2019). However, it remains unclear how gene expression within the defined populations of NSCs and progenitors in the developing mammalian cortex *in vivo* change over time and through the lineage as the fate decisions are being made.

In order to compare like-with-like and circumvent some of the challenges of random cell selection, we took advantage of the knowledge about murine cortical development and transgenic mice that allow isolation of defined progenitor populations at each day between embryonic day 10.5 (E10.5) and birth (Hebert & Fishell, 2008). We performed bulk and single cell RNA-Seq to generate gene expression profiles of NSCs, BPs, and NBNs from the dorsal cortex, spanning the critical periods of NSC expansion (E10.5-11.5), neurogenesis (E12.5-16.5), and gliogenesis (E17.5-PN1). From these data catalogs, we elucidated the transcriptional landscapes of NSCs, BPs, and neuronal subtypes and systematically followed robust temporal dynamics in their gene expression through cortical development. We determined an amazing dynamic heterogeneity within these progenitor populations at the single-cell level, identifying individual clusters of NSC, BP, and NBN subtypes and providing gene signatures for each of these clusters. We evaluated the changes in signaling pathway component expression during cortical development and identified receptors, ligands, and downstream signaling pathways that potentially play critical roles in cortical development. Finally, we found that the transcriptional programs that define specific cortical

neuron types are active in NSCs prior to the birth of the neurons. Our work provides a versatile and comprehensive resource that will be useful to address gene expression but also novel aspects of NSCs fate choice and neuronal cell subtype generation.

Results

Transcriptional analyses validate the selection and sorting procedure

Canonical Notch signaling in the developing cortex suppresses NSC differentiation by repressing expression of proneurogenic transcription factors while promoting proliferation and survival (Gaiano & Fishell, 2002; Dang *et al*, 2006; Mason *et al*, 2006; Kageyama *et al*, 2009; Imayoshi *et al*, 2010). *Hes5* is a transcriptional target of Notch signaling and labels NSCs at all stages of development and in the adult (Lugert *et al*, 2010, 2012; Basak *et al*, 2012; Bansod *et al*, 2017). Conversely, *Eomes* (*Tbr2*) is expressed by BPs and committed neural progenitors (Arnold *et al*, 2008; Sessa *et al*, 2017). *Hes5::GFP* labels NSCs in the ventricular zone and *Tbr2::GFP* BPs and NBNs in the subventricular zone and developing cortical plate (Fig EV1A and B) (Basak & Taylor, 2007; Arnold *et al*, 2009).

To address changes in gene expression within the NSC, BPs, and early neurons of the cortical neural lineages, cells were sorted from individual *Hes5::GFP* and *Tbr2::GFP* embryos at each day of development between embryonic day 10.5 (E10.5) and birth (PN), and RNA-Seq performed on the samples from each embryo separately (Figs 1A and EV1A and B). E10.5-PN1 covered the embryonic stages of cortical development from NSC expansion (E10.5-E11.5), through neurogenesis (E12.5-E16.5) to gliogenesis (E17.5-PN). We showed previously that *Hes5::GFP*⁺ cells in the VZ of the developing dorsal telencephalon express Pax6 and radial glial proteins including GLAST and Nestin (Fig EV1B; Basak & Taylor, 2007). A few newly generated *Tbr2*⁺ BPs in the apical VZ express residual *Hes5::GFP*, which is lost before the cells reach the SVZ (Basak & Taylor, 2007). All Pax6 cells in VZ express *Hes5::GFP* but those in the SVZ do not (Fig EV1B; Basak & Taylor, 2007; Lugert *et al*, 2012). Conversely, *Tbr2::GFP* is not expressed by radial glia in the VZ but localizes with *Tbr2* in the SVZ.

We gated on *Hes5::GFP*^{high} cells at all time points excluding low and negative cells (Fig EV1C). Immunostaining of acutely sorted *Hes5::GFP*⁺ cells showed expression of the NSC-associated proteins Sox2 and Pax6 but not *Tbr2* (Fig EV1D). *Tbr2::GFP*⁺ cells were first detectable by FACS at E12.5 corresponding to the prominent appearance of BPs in the developing dorsal cortex (Arnold *et al*, 2009). From E15.5 on, the *Tbr2::GFP*⁺ population was divided into GFP^{high}, GFP^{low}, and GFP⁻ populations (Fig EV1C). We separated *Tbr2::GFP*^{low} and *Tbr2::GFP*^{high} cells and analyzed these populations separately. Sorted *Tbr2::GFP*^{high} cells expressed low levels of Sox2 and Pax6 and high levels of *Tbr2* denoting them as BPs (Fig EV1E). Sorted *Tbr2::GFP*^{low} cells did not express Sox2 and Pax6 and had lower levels of *Tbr2* than the BPs (Fig EV1F). We reasoned that the *Tbr2::GFP*^{low} cells were immature NBNs labeled by low levels of *Tbr2* and perduring GFP.

As proof of concept, *Hes5* RNA levels were high in the NSCs populations at all developmental time points and low in the *Tbr2::GFP*⁺ samples (Fig 1B). As expected, the transcripts of the BP markers *Tbr2* and *Btg2* were highly expressed by *Tbr2::GFP*^{high} cells at all

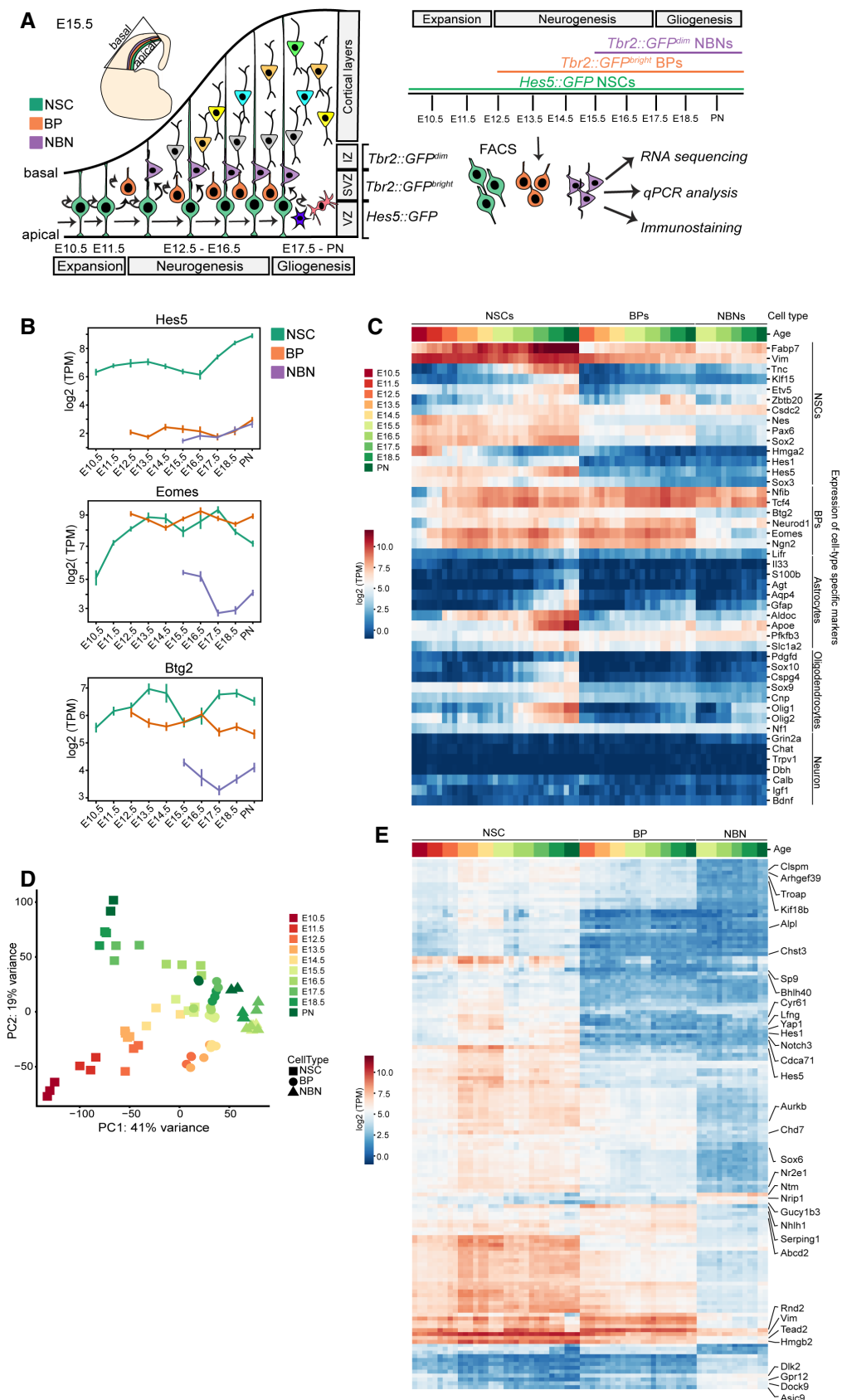


Figure 1.

Figure 1. Overview and validation of the transcriptional analyses.

- A Overview of the biological system with experimental paradigm, illustrating NSCs, BPs, and NBNs were isolated at each day during development from E10.5 to PN.
- B Notch signaling effector *Hes5* is expressed high in NSCs while *Eomes* (*Tbr2*) and *Btg2* are expressed high in both NSCs and BPs at the mRNA level. Each dot defines the mean and lines define the Standard Deviation (SD). Three to four biological replicates were collected for each time point.
- C Heatmap validating the known cell-type specific marker gene expression from RNA sequencing data.
- D Principal Component Analysis (PCA) projection of all samples of NSCs, BPs, and NBNs throughout development onto the first two principal components (PCs) covering maximum variance reveals that PC1 sorts samples by cell type and PC2 sorts samples by developmental stage.
- E Heatmap illustrating the novel marker genes identified from the RNA sequencing data, as signature genes for NSCs, BPs, and NBNs. BPs, basal progenitors; E, embryonic day; IZ, intermediate zone; NBNs, newborn neurons; NSCs, neural stem cells; PN, post natal; SVZ, subventricular zone; VZ, ventricular zone. Expression values on the heatmaps are log₂ (transcripts per million).

stages from E12.5–PN1 consistent with being dorsal cortical BPs and at lower levels by *Tbr2::GFP^{low}* NBNs (Fig 1B). *Tbr2* and *Btg2* transcripts were detected in the NSC samples without detectable protein, confirming previous observations (Figs 1B and EV1B and D–F; Pollen et al, 2015; Mukhtar et al, 2020). These points demonstrate the challenges of allocation of cell type based on transcriptional activity of a few “marker” genes.

Interrogation of the RNA-Seq data revealed that NSCs, BPs, and NBN transcriptomes were remarkably similar, and few genes were differentially expressed between NSCs and BPs (Fig 1C). We analyzed the dynamics in expression of known NSC, BP, and NBN markers between E10.5 and PN1 (Figs 1C and EV1G and H). NSC markers were highly expressed throughout cortical development with characteristic temporal dynamics in the NSC populations (Gotz & Huttner, 2005; Molyneaux et al, 2007; Ohtsuka et al, 2011; Pollen et al, 2015; Mukhtar & Taylor, 2018).

Known BP markers, including *Nfib*, *Ngn2*, *Tcf4*, and *Neurod1*, were highly expressed throughout cortical development by BPs. Astrocytic markers including *S100b*, *ApoE*, *Gfap*, and *Aldoc* were expressed highly by NSCs isolated late in development corresponding to the onset of gliogenesis indicating that the glial transcriptional program had already been initiated (Zhang & Barres, 2010; Molofsky et al, 2012; Liddelw & Barres, 2015). Similarly, key markers for oligodendrocytes including *Pdgfd*, *Sox10*, *Cspg4*, and *Sox9* were expressed higher by late-stage NSCs corresponding to the last wave of oligodendrogenesis originating in the ventricular zone of the dorsal cortex (Ono et al, 2008; Zhang & Barres, 2010; Takebayashi & Ikenaka, 2015). The mature neuronal markers *Grin2a*, *Chat*, *Bdnf*, and *Igf1* were expressed at very low levels by the *Hes5::GFP* and *Tbr2::GFP* sorted cells indicating that the selection process isolated progenitors and excluded mature neurons (Figs 1C and EV1G; Sarnat, 2013).

Unbiased computational analyses revealed extensive transcriptional dynamics within the different cell types. Principal component analysis (PCA) showed that the first two principal components (PC) capture 60% of the total variance in gene expression (Fig EV1I). Remarkably, the first PC largely separated the samples based on cell type (NSC, BP and NBN), whereas the second PC orders samples by differentiation state for all three cell types (Fig 1D). This latter observation indicates that NSCs, BPs, and NBNs change their transcription in a similar direction with developmental stage. The projection onto the first two PCs indicated that BPs and NBNs are transcriptionally closer to NSCs in the neurogenic phase of cortical development than those in the expansion and gliogenic phases. While the NBNs cleanly separate from the other cell types along the first PC, the BPs were positioned between NBNs and the NSCs in

the neurogenic phase, which they partially overlapped, consistent with the BPs being a transient neuronal precursor population (Fig 1D). Compared with BPs and NBNs, the NSCs displayed much larger variation in gene expression across time on the first two PCs, with a fluid separation from the expansion to neurogenesis and gliogenesis phases. Subsequently, we performed pairwise differentially expressed gene analyses (DEG) between the different cell types (Figs 1E and EV1J–O). To exclude substantial contamination of the *Hes5::GFP^{high}* sorted cells with BPs and NBNs, we identified transcripts that were highly expressed by BPs and NBNs but not by NSCs (Fig EV1J–O). Thus, although the transcriptomes of NSCs and BPs are remarkably similar, the sorting procedures were effective in enriching stem from progenitor populations throughout cortical development. We identified novel markers for NSCs, BPs, and NBNs using two independent methods—DEGs and Z-score log₂ (TPM) expression values (Dataset EV1). Genes including *Sp9*, *Cyr61*, *Yap1*, *Hes1*, *Lfng*, and *Notch3* are highly expressed by NSCs. Identification of these signature genes using an unbiased approach is an independent validation of the approach as the functions of some have been studied in NSCs. The Hippo co-activator *Yap1*, Notch signaling components *Hes1*, *Lfng* are involved in NSC proliferation and maintenance (Pourquie, 2003; Bray, 2006; Takebayashi & Ikenaka, 2015). *Gucy1b3*, *Nhlh1*, and *Serping1* were highly expressed by BPs and novel markers of the cell type in the lineage but not much is known about their function (Lipkowitz et al, 1992). *Ntm*, *Nrip1* are expressed higher in NBNs than in BPs or NSCs also providing novel markers (Gil et al, 1998). Interestingly, DEG analyses revealed that the majority of the highly expressed genes in NSCs are downregulated by BPs and reduced further by NBNs.

Temporal dynamics in transcriptional landscapes of NSCs, BPs, and NBNs based on gene expression

Our analyses showed that NSCs displayed maximum variance over time and therefore contribute heavily to the first two PCs. To understand the transcriptional dynamics in the NSCs, we performed PCA focusing only on the NSCs. The first two PCs covered almost 70% of the total variance and exposed a dynamic transcriptional path among the phases of expansion, neurogenesis, and gliogenesis (Figs 2A and EV2A). Although the NSCs were isolated using the same characteristic, *Hes5::GFP^{high}* expression, we observed striking, stage-related dynamic movement through transcriptional space. PCA indicated that NSCs could follow a continuous path from expansion through neurogenesis to gliogenesis, consistent with the common origin model of sequential cell specification over time. Surprisingly, *Hbb-bh1*, *Hba-x*, and *Hbb-y* expression distinguished NSCs

in the expansion phase (PC1 negative axis) from those in the neurogenic and gliogenic phases (Fig 2B). Although hemoglobin subunits are predominantly associated with erythrocytes and oxygen transport from the lungs, hemoglobin subunits are also expressed in the brain and by neurons (Brown et al, 2016). Hemoglobin subunits are found in the mitochondria in neurons and may assist oxygen

transport across mitochondria membranes. As NSCs are mitotically highly active and require extensive energy for cell division, it is possible that hemoglobin supports the energy requirements during the expansion phase.

By contrast, *Neurod6*, *Cntn2*, *Slc17a7*, and *Nfix* separated NSCs in the neurogenic phase along the PC2 negative axis from those in

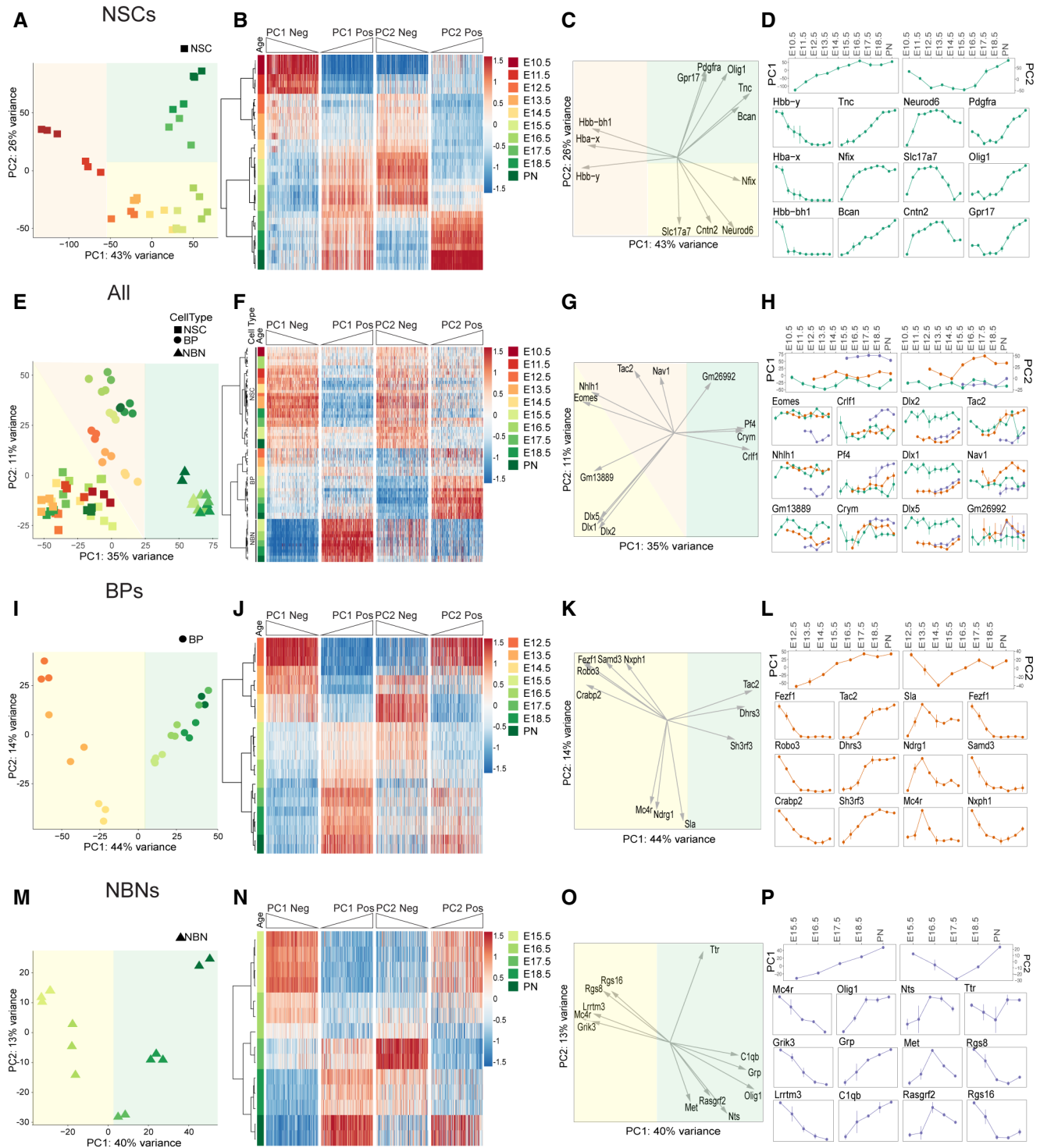


Figure 2.

Figure 2. Dynamics of transcriptional profile changes in different populations over time.

- A PCA of NSCs from E10.5 to PN showing their transcriptional dynamics.
- B Heatmap of genes that have the highest contribution to the PC1 and PC2 for NSCs, sorted by their projection on the PC axis (250 most positive and 250 most negative, Dataset EV2).
- C Projections of the expression profiles of example genes onto the first two PCs, showing the developmental stage at which, they are most highly expressed.
- D Expression profiles of PC1 and PC2 along developmental time (top) and expression profiles of the top three genes with highest negative and positive projection on PC1 (first and second column) and highest negative and positive projection on PC2 (third and fourth column) for NSCs.
- E PCA of all samples after removal of the first two principal components of the expression profiles of NSCs separates samples by cell type.
- F Heatmap of genes that have the highest contribution to the PC1 and PC2, sorted by their projection on the PC axis (250 most positive and 250 most negative).
- G Projections of the expression profiles of example genes onto the first two PCs, showing their differential expression across cell types.
- H Expression profiles of PC1 and PC2 across all samples with time from left to right and NSCs in green, BPs in red, and NBNs in purple (top), and the gene expression profiles of top three genes with highest negative and positive projection on PC1 (first and second column) and highest negative and positive projection on PC2 (third and fourth column).
- I PCA of BPs from E12.5 to PN showing their transcriptional dynamics.
- J As in panel (B) but now for BPs.
- K As in panel (C) but now for BPs.
- L As in panel (D) but now for BPs.
- M PCA of NBNs from E15.5 to PN showing their transcriptional dynamics.
- N As in panels (B) and (J) but now for NBNs.
- O As in panels (C) and (K) but now for NBNs.
- P As in panels (D) and (L) but now for NBNs.

Data Information: In (D), (H), (L), and (P) (bottom), the x-axis is embryonic days, and the y-axis is log₂(TPM).

the expansion and gliogenic phases. *Neurod6* is a Helix-Loop-Helix (HLH) transcription factor (TF) that plays a prominent role in neuronal differentiation (Sommer *et al*, 1996). *Neurod6* is transcriptionally activated by *Neurog1* and *Neurog2*, two proneural HLH TFs that are downstream of Notch signaling (Ross *et al*, 2003). *Neurod6* is associated with familial temporal lobe epilepsy and attention deficit hyperactivity disorder in humans (Tutukova *et al*, 2021). *Cntn2* is a member of the Contactin family of immunoglobulin cell adhesion molecules and functions in neuronal differentiation, determination, and migration as well as axon guidance (Mohebiany *et al*, 2014). *Cntn2* is located at 1q32.1, a region associated with microcephaly, and mutations in *Cntn2* cause familial adult myoclonic epilepsy 5 (FAME5) (Rickman *et al*, 2001; Stogmann *et al*, 2013; Mohebiany *et al*, 2014). *Slc17a7* is a transmembrane channel and urea transporter. It is selectively expressed in NSCs compared with BPs and NBNs, and its expression increases with developmental stages. The function of *Slc17a7* in NSCs remains to be shown. *Nfix* is a member of the nuclear I family of TFs. *Nfix* regulates NSC proliferation and differentiation both during embryonic development and in the adult and has been proposed to be a tumor suppressor in gliomas (Heng *et al*, 2015; Stringer *et al*, 2016). Loss of *Nfix* is associated with increased proliferation in the SVZ of the embryonic brain and delayed gliogenesis (Heng *et al*, 2015; Stringer *et al*, 2016). In summary, the unbiased analysis of gene expression revealed novel markers of stage specific NSCs and potential regulators of differentiation in the dramatic switch from the expansion to neurogenic phases of cortical development.

The unbiased computational approach identified genes important in NSC maintenance and differentiation as well as a plethora of novel and dynamically expressed NSC genes (Pollen *et al*, 2015; Telley *et al*, 2016, 2019). We validated the expression of the novel signature genes predicted by the PC separations by RT-qPCR on independent biological replicates (Figs 2B and EV2B and C; Dataset EV2). We randomly selected genes differentially expressed by NSCs during the expansion, neurogenic and gliogenic phases. *Ccnd1*, *Crabp2*, *Hbb-bh1* are highly expressed during the expansion

phase; *Bcl11b*, *Cntn2*, *Id2*, *Satb2* during neurogenic phase; and *ApoE*, *Aqp4*, *Sparcl1*, *Tril* during the gliogenic phase (Fig EV2C; Dataset EV2). By clustering the gene expression profiles, we identified pools of genes that follow the same transcriptional trajectory in NSCs over time. This implied either co-regulation at the transcriptional level or gene expression associated with distinct cell states (Fig EV2D–G). Some genes within these profiles are typical markers of the different phases of NSC development. For example, *Neurog2* and *Cspg4* mark the neurogenic and gliogenic phases of NSCs, respectively. Others, including *Shh*, mark specifically the early expansion phase and their expression is: low upon the onset of fate determination (Fig EV2D–G).

To address the transcriptional changes among the BPs and NBNs, we excluded the predominant variance resulting from the NSCs shifts in gene expression by computing PCs of all the samples orthogonal to the first two PCs of the NSCs. PCA of the remaining variables clustered all NSCs together indicating their underlying identity. BPs and NBNs separated in transcriptional space. Thus, the data orthogonal to the first PCs of NSCs enhanced the differences between NSCs, BPs, and NBNs (Fig 2E) and increased the separation of BPs and NBNs, revealing a clear separation of BPs at early (E12.5–E14.5) and late (E15.5–PN) time points of cortical development (Figs 2E and F and EV2H). In these analyses, NBNs showed less transcriptional dynamics over time. We performed pairwise comparisons to reveal DEGs genes contributing to PCs in the orthogonal analyses. Genes including *Dlx1*, *Dlx5*, and *Dlx2* separate NSCs while *Tbr2*, *Nhlh1* represent the highest loading along the orthogonal PC1 negative axis. *Crym*, *Pf4*, and *Crlf4* separated BPs and NBNs along the orthogonal PC1 positive axis (Fig 2F–H; Dataset EV2).

To investigate the stage-correlated changes in gene expression by BPs and NBNs, we performed PCAs on BPs and NBNs separately. Despite being selected based on differences in level of *Tbr2::GFP* expression, PCA displayed continuous dynamics in these populations over time. However, the first PC was sufficient to separate BPs based on developmental stage (Figs 2I–L and EV3A). From these analyses, we identified *Fezf1*, *Samd3*, *Robo3* to be highest in early

BPs while *Tac2*, *Dhrs3*, *Sh3rf3* were expressed higher by late BPs. Therefore, we could define distinct gene profiles for BPs that reflected their developmental stage implying that BPs are also a heterogeneous population of intermediate cells. Similarly, the first PC was also sufficient to separate NBNs over the course of development (Figs 2M–P and EV3B). In order to validate the novel signature genes separating BPs and NBNs, we performed RT-qPCR on independent biological replicates (Fig EV3C and D; Dataset EV2), which confirmed the differential expression of the signature genes between early BPs, late BPs, and their corresponding NBNs. *Cckar*, *Kif2c*, *Uncx*, *Robo3* were highly expressed by early BPs while *Loxl1*, *Unc5d*, *Ezr* were highly expressed by late BPs. On the contrary, NBNs displayed high expression of *Mef2c*, *Usp43*, *Lrfr5*, *Ntsr1*, and *Gucy1a3* (Fig EV2D). A more comprehensive list of these DEGs is available in Dataset EV2. Thus, by our preliminary analyses of gene expression, we demonstrate dynamics in NSCs, BPs, and NBNs and have identified novel signature genes, which are binary and unique for these populations.

Temporal dynamics in transcriptional landscapes of NSCs, BPs, and NBNs is based on TF nodes and networks

To characterize the transcriptional states of NSCs, BPs, and NBNs and map the activities of TFs throughout cortical development, we used the Integrated System for Motif Activity Response Analysis (ISMARA) (Balwierz et al, 2014; Artimo et al, 2016). ISMARA infers the regulatory states of samples by computationally predicting transcription factor binding sites (TFBSs) genome-wide and modeling the observed gene expression state of each sample in terms of the predicted TFBS and “activities” of the TF binding motifs (Fig 3A, <https://ismara.unibas.ch/NeuroStemX/>). PCA of the motif activities revealed that, in agreement with the observations made based on mRNA expression (Fig 2A), the majority of the variance across samples was dominated by differences within the NSC samples. Therefore, we first analyzed the NSCs from all time points in isolation and then analyzed NSCs from the neurogenic phase of development together with the BPs and NBNs.

PCA of the motif activities in NSCs showed that 80% of the variance was captured by the first two PCs, with the plane spanned by these PCs clearly separating the NSC samples into segments corresponding to the expansion, neurogenic and gliogenic phases (Fig 3B; Dataset EV3). We identified the top 20 TF binding motifs that contributed most to the variance in these PCs and projected their motif activity vectors onto the same plane in order to show which TFs contribute most to gene expression at each of the stages (Fig 3C; Appendix Fig S1A). For example, *Scrt1* and *Hdx* contribute strongly to the expansion phase, *REST* to the neurogenesis phase, and *Nfib/c* contribute to the gliogenesis phase. Using the gene expression profiles and inferred motif activities, ISMARA also infers which genes are targeted by each TF motif (Fig 3A, Balwierz et al, 2014). For example, ISMARA predicts that *Neurod1* targets the *Neurod6* gene. This implies that the upregulation of the *Neurod6* gene during cortical development is due, at least partially, to increasing *Neurod1* TF activity over time (Fig 3D). In-depth analyses of the top TF motifs and their predicted target genes show a strong coherence with those genes identified by PCA of the RNA-Seq data (Fig 2B–D).

Of particular interest are so called motif–motif interactions in which ISMARA predicts a TF motif to target a gene, which encodes a TF, since these interactions correspond to edges of the ISMARA predicted regulatory network. To extract a core regulatory network for NSCs in the three phases of cortical development, we selected the motif–motif interactions with the strongest statistical strength (Fig 3E, see Materials and Methods). A large proportion of TF motifs cross-regulate each other and include TFs that are known to be involved in cortical development. For example, *Scrt1* and *Scrt2* are predicted to be active in NSCs during the expansion, while *Hoxb7* and *Sox5* are active during the neurogenic phase, and *Nfi* family members are involved in glial cell specification (Bel-Vialar et al, 2002; Lai et al, 2008; Paul et al, 2014; Zhou et al, 2015). Intriguingly, the TF motifs that are most active in the neurogenesis phase (depicted in the center of Fig 3E) are predicted to repress many of the TF motifs involved in NSC expansion (left side of Fig 3E) and activate many of the TF motifs that induce gliogenesis (right side of

Figure 3. Dynamics of transcriptional network changes with ISMARA in different populations over time.

- Conceptual illustration of ISMARA, which models gene expression profiles in terms of the activities of TF binding motifs. ISMARA models the matrix of gene expression E as a linear function of computationally predicted TF binding sites in the promoters of each gene (site count matrix N) and activities of TF binding motifs across samples (matrix A).
- PCA projection of the motif activity profiles of the NSCs for all time points onto the first two PC components, which capture 80% of the total variance in motif activities. The background color represents segments of the plane spanned by the first two PCs corresponding to the expansion (red, E10.5–E11.5), neurogenesis (green, E12.5–E15.5), and gliogenesis (purple, E16.5–PN) phases.
- Top 20 motifs contributing the most to the first two PCs, projected on the first two PCs, illustrating which TFs are active in each developmental stage.
- Example of a predicted regulatory interaction between two TFs. ISMARA predicts that the *Neurod1* motif upregulates expression of the *Neurod6* gene through a binding site in its promoter. Note that the *Neurod1* motif activity (bottom) is upregulated in parallel with its gene expression (top right), indicating it acts as an activator, and that *Neurod6* gene expression is: similarly upregulated across developmental time (bottom right).
- Core regulatory network of most significant motif–motif interactions in the NSC dynamics, together with overrepresented gene ontology categories among the target genes of different core regulatory motifs. The motif activity profile of each core regulatory motif is shown framed with a color representing the developmental stage at which its most active (red for expansion, green for neurogenesis, and blue for gliogenesis). Edges from motifs A to B correspond to predicted regulatory interactions with motif A either activating or repressing activity of motif B. Ellipses indicate top Gene Ontology categories overrepresented among the target genes of different motifs.
- PCA projection of the motif activity profiles of NSCs from the neurogenesis phase (day 12.5–16.5), BPs and NBNs, which capture 69% of the total variance.
- Top 20 motifs contributing the most to the first two principal components, projected on the first two PCs of panel, illustrating which TFs distinguish cell types and developmental stages.
- Core regulatory network and overrepresented gene ontology categories as in panel E but now for analysis of the samples of the BPs, NBNs and NSCs from the neurogenesis stage. Motif activity profiles are shown as a function of time for NSCs (green), BPs (red), and NBNs (purple) and are framed with a color indicating whether the motif activity varies mainly across time (blue) or cell type (pink).

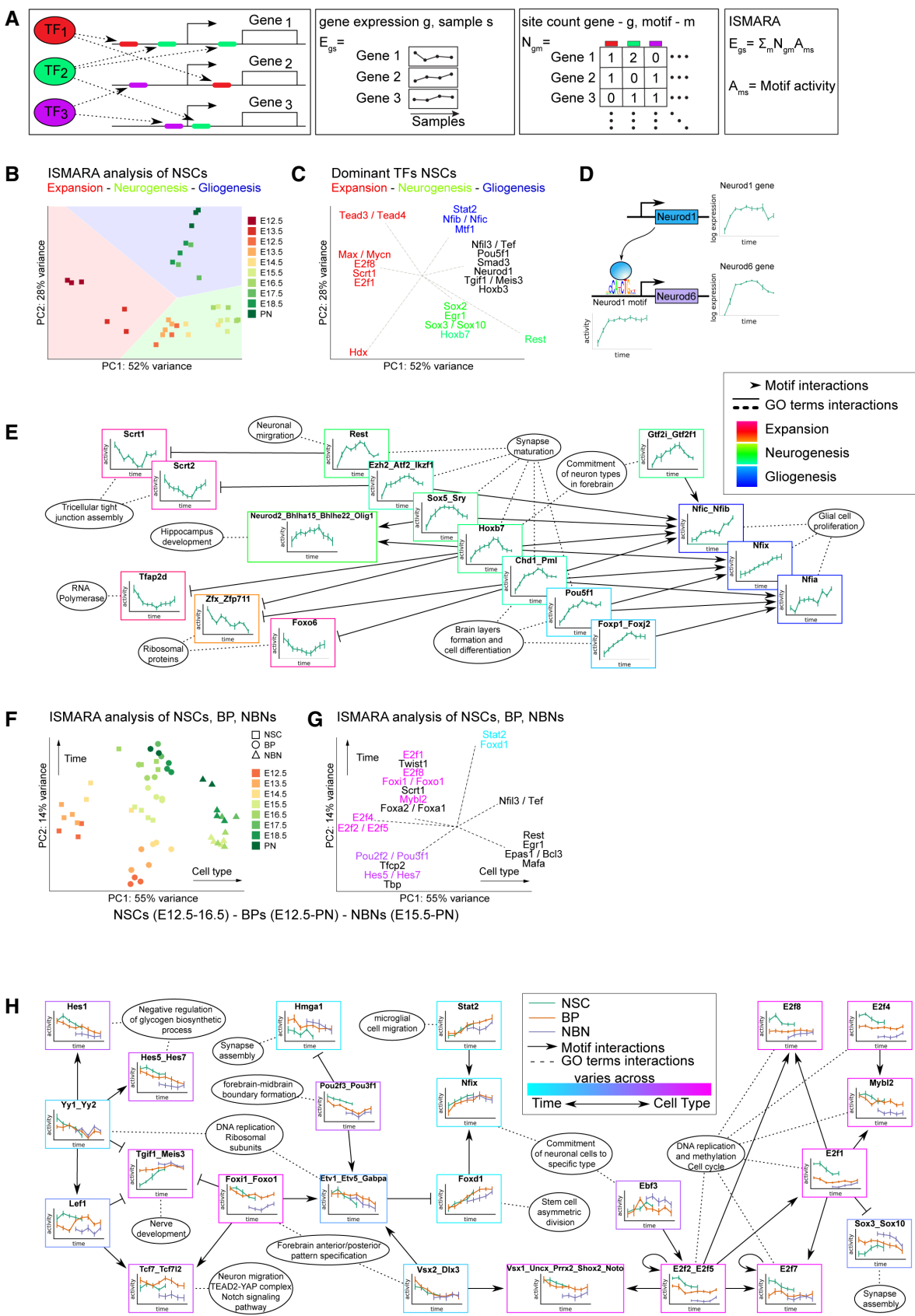


Figure 3.

Fig 3E). These analyses predict complex interconnected gene regulatory networks in the dynamics of the NSC lineage with TFs that can act independently but show a large degree of synergy. Perhaps most striking, virtually all of the top regulatory interactions involve motifs that are most active during neurogenesis, either repressing or activating motifs associated with the expansion and gliogenesis phases. This suggests a key role for the TFs associated with neurogenesis in orchestration of the developmental dynamics of NSC in the dorsal cortex (Fig 3E).

Next, we analyzed the TF motif activities in NSCs, BPs, and NBNs together. Since the variance in the first two PCs is dominated by the time dynamics of the NSCs, we removed the contribution of these two components from the motif activities for subsequent analyses, in order to reveal activities that may be masked. We then identified the top TF motifs determining the separation of NSCs, BPs, and NBNs by PCA (Appendix Fig S1B and C). Similar to our gene expression analyses, we also analyzed TF motif activities along the developmental trajectory separately for BPs and NBNs and identified which TFs and motif activities contribute most to these developmental dynamics (Appendix Fig S1D–I). To better understand the relationship between the different progenitor cell types in the dorsal cortex, we jointly analyzed the motif activity dynamics of neurogenic NSCs, BPs, and NBNs. Remarkably, and similar to what we observed based on the gene expression, the PC1 of the motif activities separated the three cell types (NSCs on the left, BPs in the middle, and NBNs on the right; Fig 3F). The PC2 ordered the samples of each cell type by developmental stage (Fig 3F). It is particularly noteworthy that the temporal dynamics of the motif activities along PC2 is shared by NSCs, BPs, and NBNs (earliest stages at the bottom, oldest stages at the top; Fig 3F). We defined the key TFs and motifs that separate the cell types and define the developmental lineage of NSCs, BPs, and NBNs over time (Fig 3G).

We identified the motif–motif interactions with strongest statistical power within the neurogenic NSCs, BPs, and NBNs and represented these as a predicted core regulatory network (Fig 3H). A subnetwork of E2f family motifs (E2f1, E2f2, E2f4, E2f5, and E2f8), which all target genes involved in DNA replication, methylation, and cell cycle, positively regulate Mybl2 (a cell cycle regulator) and the paired-like homeodomain Vsx1-like TF activity but repress Sox3/Sox10 activity. Note that the activities of these motifs remain relatively constant over time and mainly characterize differences between the cell types, with highest activity in NSCs and lowest

activity in NBNs. We also observed TF motifs that show similar activities across the progenitor cell types but change their activity over time, such as Foxd1 and Stat2 whose motifs regulate neural and glial development through Nfix (Fig 3H). Conversely, some TF motifs show cell-type specificity, for example, those for Hes1, Hes5, Meis3, Tcf7, and Foxo1. Interestingly, Hes1, Hes5, and Tcf7 are all primary regulators of Notch signaling in NSCs (Fig 3H).

Single-cell RNA sequencing reveals underlying heterogeneity in NSCs, BPs, and NBNs

PCA at the population level revealed extensive changes in the transcriptome of NSCs, BPs, and NBNs over time. We addressed heterogeneity within each cell population by analyzing the transcriptional landscapes at the single-cell level by single-cell RNA-Seq (scRNA-Seq) (Appendix Fig S2A). The single-cell transcriptomes of highly variable genes (HVGs) revealed a low heterogeneity within the NSCs during expansion and gliogenesis (Fig 4A). By contrast, NSCs during the neurogenesis phase (E13.5 and E15.5) were heterogeneous (Fig 4A). To validate that the scRNA-Seq data were representative of the population data, we averaged the single-cell transcriptomes of a specific time point and projected them on the PC matrices of the population samples. The averaged single-cell data superimposed on the population samples and followed the same transcriptional trajectory over time and confirmed that the single-cell transcriptomes reflected the heterogeneity of the population at the respective time point (Fig 4B). Therefore, the single-cell heterogeneity in NSCs during cortical development is representative of the biological changes in single NSC gene expression over time.

k-Means clustering divided the NSCs into five cell clusters and revealed DEGs across these clusters (Fig 4C; Appendix Fig S2C–F and Dataset EV4). The five NSC types were unequally represented over time. NSC type 1 (cluster 1) was present almost exclusively at E10.5 and E11.5 and represents the major NSC transcriptional status in the expansion phase. NSC cluster 5 was the predominant NSC type during the later, gliogenic phases of corticogenesis. NSC clusters 2–4 were found during multiple phases of development from E11.5 and expansion through neurogenesis to gliogenesis (Fig 4C).

Visualization of the single cell data by t-SNE also showed separation of the five NSC cell types (clusters 1–5; Fig 4D). Projection of gene expression onto the t-SNE identified cluster-specific expression. *Crabp2* and *Tnc* marked clusters 1 and 5, respectively, while

Figure 4. Heterogeneity of NSCs, BPs and NBNs at single cell level.

- A PCA of NSC single cells, using the top 2,000 highly variable genes (HVGs) obtained from bulk NSCs.
- B Projection of average single cells of NSCs at each time point on the first two PCs of bulk NSCs using the top 2,000 HVGs obtained from bulk NSCs.
- C Clustering of assignment matrix of NSC single cells using k-means and hierarchical clustering.
- D Marker genes that are up/downregulated in each cluster of NSCs.
- E PCA of BP single cells, using the top 2,000 highly variable genes obtained from bulk BPs.
- F Projection of average BP single cells on the first two PCs of bulk BPs using the top 2,000 HVGs obtained from bulk BPs.
- G Clustering of assignment matrix of NBN single cells using k-means and hierarchical clustering.
- H Marker genes that are up/down regulated in each cluster of NBNs.
- I PCA of NBN single cells, using the top 2,000 HVGs obtained from bulk NBNs.
- J Projection of average single cells of NBNs on the first two PCs of bulk NBNs using the top 2,000 HVGs obtained from bulk NBNs.
- K Clustering of assignment matrix of NBN single cells using k-means and hierarchical clustering.

Data Information: In (C), (G), and (K), heatmaps represent the hierarchical clustering of assignment matrix of single cells after 500 times applying k-means clustering. The optimal number of clusters is selected based on the Silhouette coefficient. It is “1” (red) when two cells are always clustered together, “0” (blue) when two cells never fall in the same cluster. Pie charts represent the percentage of single cells at each time point in each cluster.

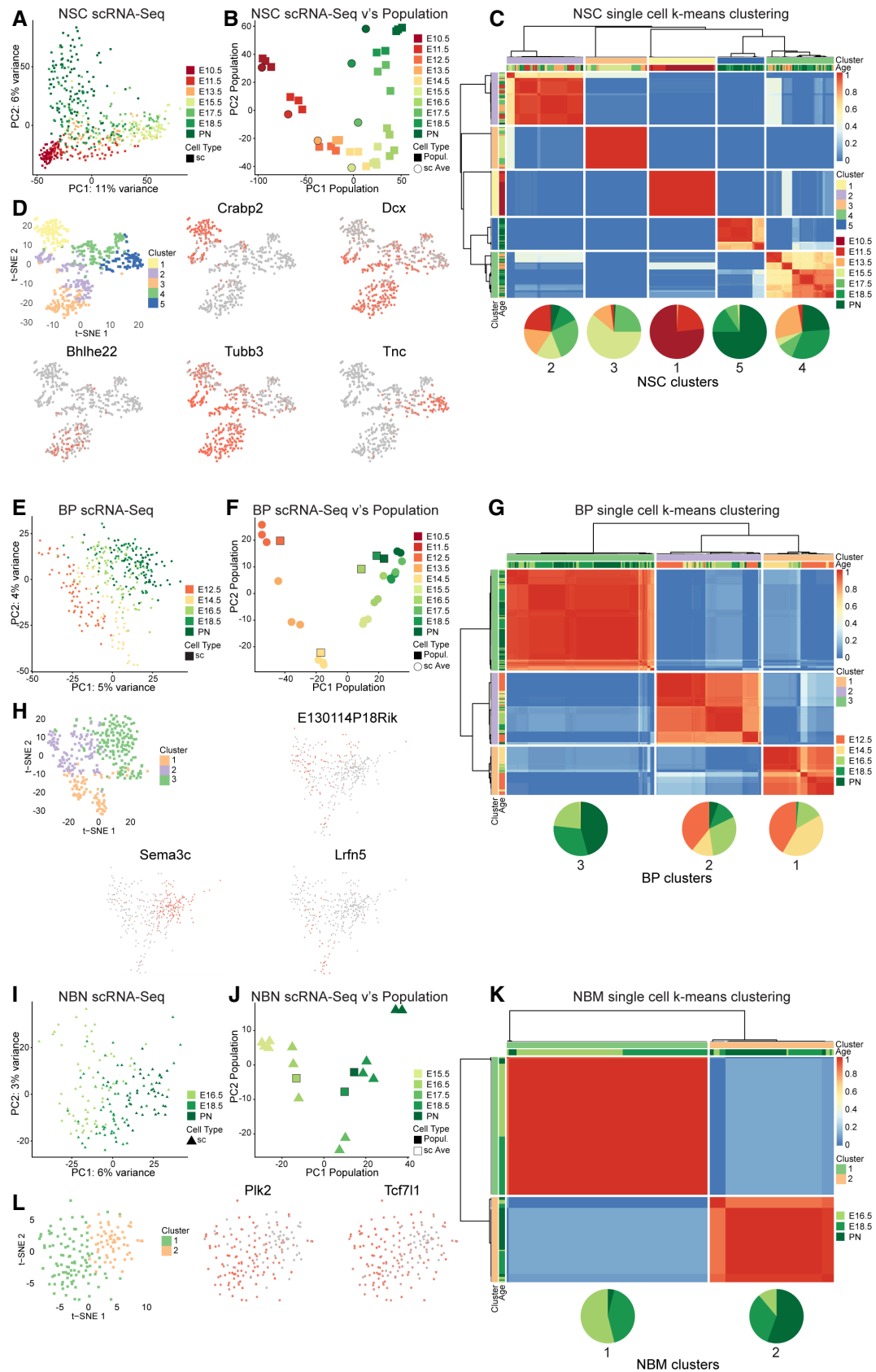


Figure 4.

Tubb3 and *Dcx* were expressed in a more expanded domain across multiple NSC clusters (Fig 4D). The heatmap shows a more comprehensive list of distinct signature genes for the five clusters (Appendix Fig S2F; Dataset EV4). The GO analyses and process networks for gene expression by the NSC clusters are shown in Dataset EV5.

We analyzed heterogeneity within the BPs from E12.5-PN1 by scRNA-Seq. BPs showed that an age-related difference is gene expression with heterogeneity distributed along the PC2 (Fig 4E). We pooled the single-cell sequences at each time point and plotted these averaged values on a PCA defined by the HVGs identified from the BP analysis at the population level (Fig 4F). Strikingly, the averaged single-cell data superimposed on the biological replicates at population level supporting that the individual BP scRNA-Seq data are representative of the populations (Fig 4F). The analysis indicated a distinct, time-dependent dynamic in gene expression from E12.5-PN1. Clustering the single BPs based on k-means revealed three distinct clusters (Fig 4G; Appendix Fig S2G–I and Dataset EV4). Cluster cell-type 3 cells are present mostly at later developmental stages (E16.5-PN1). Conversely, cluster 1 and 2 cells were more prominent at earlier times (Fig 4G). We visualized the BP clusters by t-SNE and plotted some signature genes for each cluster (Fig 4H; Appendix Fig S2J and Dataset EV4). We performed similar analyses for NBNs expecting a large heterogeneity over time due to the plethora of neuron types generated in the dorsal cortex. scRNA-Seq confirmed a broad spread in the NBNs with a component of time over the PC1 (Fig 4I).

We pooled the single-cell NBN data at each time point and plotted these averaged values on a PCA defined by the HVGs from the NBN analysis at the population level (Fig 4J). Both the population and averaged single-cell samples followed a time-dependent trajectory in gene expression consistent with the sequential generation of neurons forming the deep and superficial layers of the isocortex (Fig 4J). k-Means clustering revealed the reduced heterogeneity in the NBNs compared with NSCs and BPs and identified two major clusters (Fig 4K). Cells belonging to cluster type 1 NBNs were present at earlier time points (E15.5 and E18.5), while cells belonging to cluster type 2 were underrepresented at E15.5 and were mostly present at PN1 (Fig 4K; Dataset EV4).

t-SNE representation of the NBNs belonging to the two cell clusters showed the sparse distribution of type 1 cells reflecting the single-cell heterogeneity within this cluster. Type 2 NBNs clustered more tightly than cluster 1 cells. Due to this heterogeneity, it was difficult to pinpoint single-gene signatures for the two NBN clusters (Fig 4L; Appendix Fig S2N and Dataset EV4). These findings demonstrate an unprecedented heterogeneity in NSCs and BPs over time and a dynamic shift in gene expression of these cells at the

population and single-cell levels. The scRNA-Seq data enabled a high-resolution definition of gene signatures for each cluster (cell subtype) of NSCs, BPs, and NBNs.

We repeated the analyses of the scRNA-Seq data using KNN graph-based clustering and UMAP visualizations. UMAP visualization of NSCs showed their temporal distribution within five distinct clusters (Appendix Fig S3A). Feature plots of *Hbb-bh1*, *Hba-x*, and *Hbb-y* replicate the bulk expression patterns in NSCs, where expression is: higher during the early developmental stages (E10.5 and E11.5; Appendix Fig S3B). Additional feature plots of known markers of expansion (*Sox2*, *Pax6*, *Crabp2*), excitatory (*Dcx*, *Tubb3*, *Bcl11b*, *Pou3f2*), inhibitory (*Gad1*, *Dlx1*, *Dlx5*), astrocytic (*Aldoc*, *Aqp4*, *Sox10*), and oligodendrocytic (*Olig1*, *Olig2*, *Cspg4*) lineages show unique and consistent patterns of expression (Appendix Fig S3C–G). We performed similar analyses for the BPs and NBNs and visualized the cells on UMAPs. We identified three BP and two NBN clusters over developmental time and present example feature plots for BP genes such as *Tbr2*, *Ascl1*, *Nes*, *Lrln5* (Appendix Fig S4A–C). Reanalysis of the NBNs using KNN-graph-based methods confirmed only two discernable NBN clusters with the expression of single neuronal subtype genes such as *Bcl11b*, *Cux1*, *Cux2*, *Foxp2*, *Pfn2*, and *Pou3f2* poorly defining the heterogeneity (Appendix Fig S4D–F).

Pseudo-time analysis reveals potential trajectories among NSCs, BPs and NBNs

KNN-graph-based clustering and UMAP visualizations of pooled single NSCs, BPs, and NBNs segregated eight cell clusters (Fig 5A and B; Appendix Fig S3A). NSCs segregated in a temporal pattern as was seen with the PCA analysis. NSCs segregate into four major clusters correlated with their prospective fate – NSC1 (expansion), NSC2 (late, astrocytic), NSC3 (late, oligodendrocytic), and NSC4 (intermediate, neurogenic). BPs segregated into two clusters, BP1 enriched in *Bcl11b*⁺ cells (*Ctip2*⁺, deep layer neuron marker), and BP2 enriched in *Pou3f2*⁺ cells (*Brn2*⁺, upper layer neuron marker). NBNs divided into two clusters (NBN1 and NBN2), and similar to the individual NBN analyses, the clustering was not driven by differential expression of cortical layering markers (Fig 5C and D). The markers for each cluster are listed in Dataset EV6. We also performed Sling-shot for pseudotime analysis of the scRNA-Seq data to elucidate the neurogenic trajectories from NSCs to BPs to NBNs (Fig 5E–G). We identified three neurogenic lineage pathways starting from our early expansion NSCs to NBNs through BPs (Fig 5G).

We compared our single-cell C1 data with the extensive 10X genomics Linnarsson developing mouse brain dataset (La Manno et al, 2021; Fig EV4). First, we compared our NSCs, BPs, and NBNs

Figure 5. Pseudo-time analysis reveals potential trajectories among NSCs, BPs and NBNs.

- UMAP clustering of all NSCs, BPs, and NBNs, visualized based on cell type and time point.
- UMAP clustering showing all cell types, four clusters of NSCs, two of BPs, and two of NBNs.
- Violin plots of highly enriched genes in each cluster, labeled with gene and cluster names.
- Feature plots showing examples of highly enriched genes in the neuronal lineage, identifying excitatory neuron, inhibitory neuron, astrocytic and oligodendrocytic clusters. Y axis is the log normalized expression.
- Visualization of inferred trajectory based on UMAP embeddings showing a neurogenic lineage from NSCs, to BPs to NBNs, with a time component. Colors represent the cell types shown in (B).
- Visualization of inferred trajectory overlaid on cell type, capturing neurogenic lineage.
- Smooth representations of lineage trajectory using principal curves overlaid on cell types showing a neurogenic lineage from NSCs to BPs to NBNs.

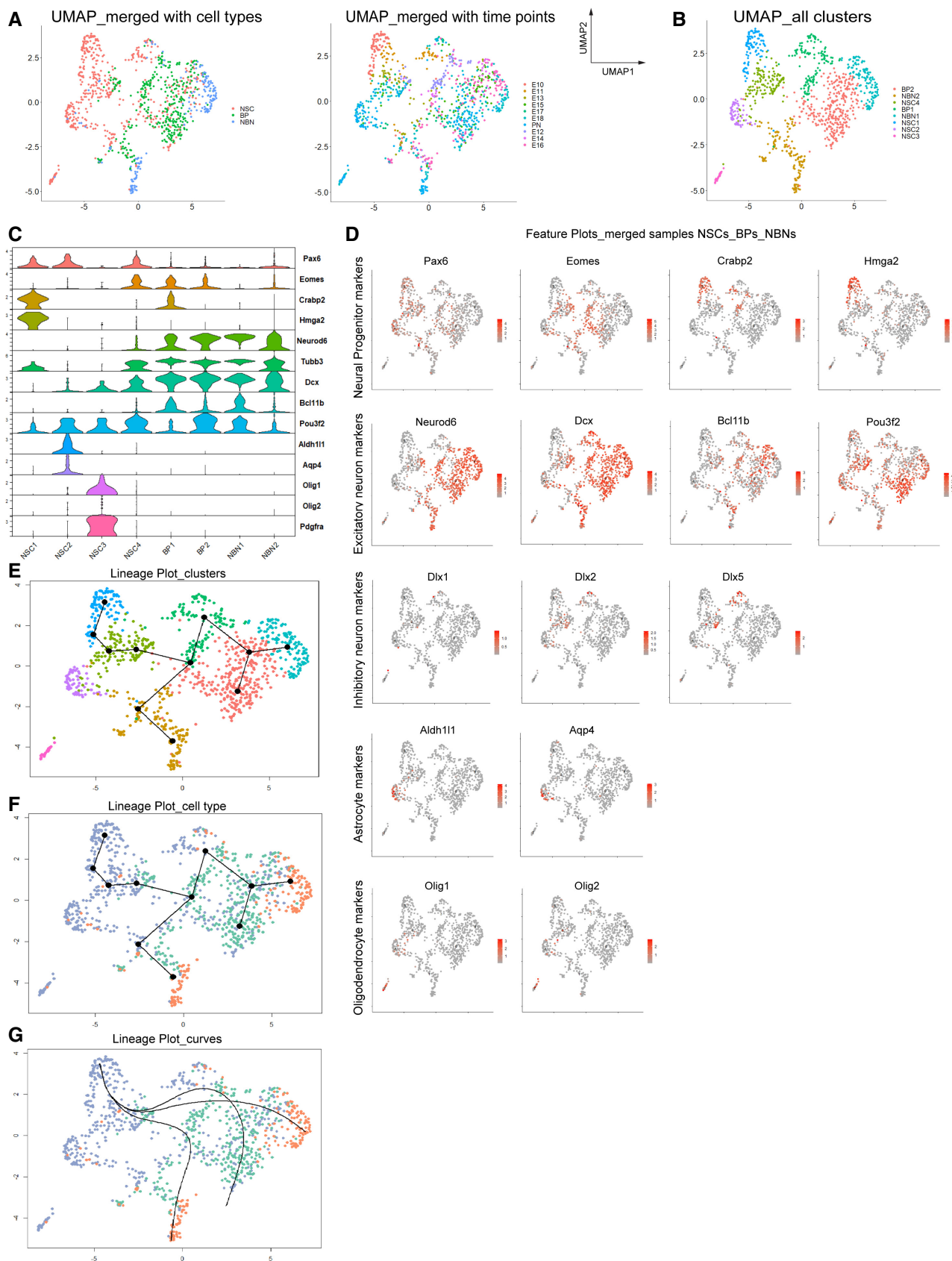


Figure 5.

to their forebrain and dorsal forebrain cells (E9-E18); second, we compared our NSCs with the cells defined as radial glia in the Linnarsson dataset from the same brain regions. We performed CCA integration analyses with our C1 scRNA-Seq data and the Linnarsson cells. Our cells integrated as expected into the Linnarsson dataset with the expression of distinct marker genes. Our cells maintained their distinct groupings in these analyses and segregated into 10 distinct clusters (Fig EV4B). Visualization revealed segregation of our cells within the Linnarsson dataset post-CCA integration. These analyses also revealed that the cells we defined as NSCs and NBNs fall within the clusters defined as radial glia and neurons by Linnarsson, respectively (Fig EV4C–E). Hence, both C1 and 10X genomics sequencing approaches correctly identify the same cell populations. However, the increased sequence depth of our C1 data allowed identification of cell subtypes that were not discernable by 10X genomics. These findings also indicate that the *Hes5::GFP* and *Tbr2::GFP* selection procedure we used does not bias the analysis of the progenitors.

Interestingly, the Linnarsson dataset does not identify BPs as a separate population. Our BPs fall into the neuron clusters defined by the Linnarsson analysis and overlap with *Tbr2* expressing cells in these clusters (Fig EV4F). These findings support the added value and power of the deeper SmartSeq2 C1 sequencing approach. We selected the cells classified as radial glial cells in the Linnarsson dataset and integrated our NSCs (Fig EV4G–I). This revealed distinct subpopulations of mainly dividing early and late progenitors defined by the expression of marker genes. We also found that the Linnarsson 10X dataset had low expression of the “mature” genes of radial glia cells compared with our C1 data, which we interpret as being due to the increased depth and reproducibility of the C1 sequencing approach (Fig EV4J). However, *Hbb* subunits that defined subtypes of NSCs in our analyses were also detectable in the radial glia in the Linnarsson dataset (Fig EV4K).

Neuronal specification markers show sequential waves in gene expression and massive heterogeneity at the single cell level

During cortical development, morphologically and physiologically unique classes of neurons are formed sequentially in waves throughout neurogenesis (Molyneaux et al, 2007; Greig et al, 2013; Telley et al, 2016; Fig 6A). Several gene combinations have been identified that classify the distinct subtypes of projection neurons in the cortex (Molyneaux et al, 2007; Greig et al, 2013). We selected and curated an extensive list of known patterning and neuronal

subtype marker genes from the literature and analyzed their expression dynamics in the NSCs, BPs, and NBNs at the population and single cell levels.

Surprisingly, transcription of the neuronal subtype genes showed sequential and developmental wave-like patterns of expression even in NSCs, at the population level (Figs 6B and EV5A). At the single cell level, these developmental waves were recapitulated in NSCs, albeit with a pronounced heterogeneity at each time point (Fig 6C). Particularly those genes commonly used to define neuronal subtypes and cortical layers later in development (*Tbr1* and *Ctip2*—Layers V and VI; *Satb2* and *Cux2*—Layers IV and II/III) showed characteristic and transient dynamics in expression 1–2 days prior to the established birth date of the neurons (Molyneaux et al, 2007). We plated the freshly FACsorted *Hes5::GFP* positive NSCs but could not detect expression of these neuronal specification factors at the protein level by immunocytochemistry (Fig 6D). This suggested that the transcriptional program that defines cortical neuron subtypes is initiated in NSCs long before their exit from cell cycle.

We performed similar computational analyses on the BPs and NBNs and identified similar sequential waves of cortical neuron gene expression correlating with the birth date of the respective neuron subtype (Figs 6C, E, and F and EV5B). Similarly, we could not detect protein expression in the BPs acutely isolated from the developing cortex (Figs 6C, E, and F and EV5B). However, and as expected, NBNs expressed proteins associated with neuron subtypes of definitive cortical layers (Figs 6C, G, and H and EV5C).

These striking findings indicate that neuronal specification programs start early in the lineage, in NSCs and BPs, and continue into the NBNs. At the single-cell level, some E10.5 NSCs expressed high levels of deep layer neuronal markers including *Cux2* while later NSCs expressed both deep and upper-layer neuronal markers (Figs 6C and EV5D). This explains the seemingly controversial *Cux2* lineage tracing experiments described previously (Franco et al, 2012). We observe similar expression in BPs, from a deep to both deep and upper layer marker expression. While the NBNs match upper layers, corresponding to their later time points of collection (Fig EV5E and F).

Signaling pathway effectors show dynamic expression in the neurogenic lineage

Signaling pathways impinge on downstream effectors to regulate NSC fate decisions during corticogenesis. The cross talk between signaling pathways and the integration of their target effectors

Figure 6. Dynamic expression of neuronal specification factors in NSCs, BPs, and NBNs.

- A Illustration of distinct projection neurons born sequentially during neurogenesis.
- B Heatmap illustrating the dynamics of expression of cortical layering markers in NSCs at population level.
- C Examples of expression dynamics of deep layer markers *Tbr1*, *Ctip2* and upper layer markers *Satb2*, *Cux2* in NSCs, BPs, and NBNs, profiles at population level (left) and single cell level (right). Each dot defines the mean and lines define the SD. Three to four biological replicates were collected for each time point.
- D Experimental validation of NSCs isolated at E13.5 using *Hes5::GFP* transgenic embryos, showing no detectable protein for *Tbr1*, *Ctip2* and *Satb2*. NSCs do express *Brn2* (*Pou3f2*) *in vitro* and *in vivo* at protein level. Scale bar = 20 μ m.
- E Heatmap illustrating the dynamics of expression of cortical layering markers in BPs at population level.
- F Experimental validation of BPs isolated at E16.5 using *Tbr2::GFP* transgenic embryos, showing no detectable protein for *Tbr1*, *Ctip2* and *Satb2*. Scale bar = 20 μ m.
- G Heatmap illustrating the dynamics of expression of cortical layering markers in NBNs at population level.
- H Experimental validation of NBNs isolated at E16.5 using *Tbr2::GFP* transgenic embryos, showing protein expression for *Tbr1*, *Ctip2*, *Satb2* and *Brn2* (*Pou3f2*). Scale bar = 20 μ m.

Data information: In (B), (E) and (F), heatmaps are based on z-score of \log_2 (TPM) expression values.

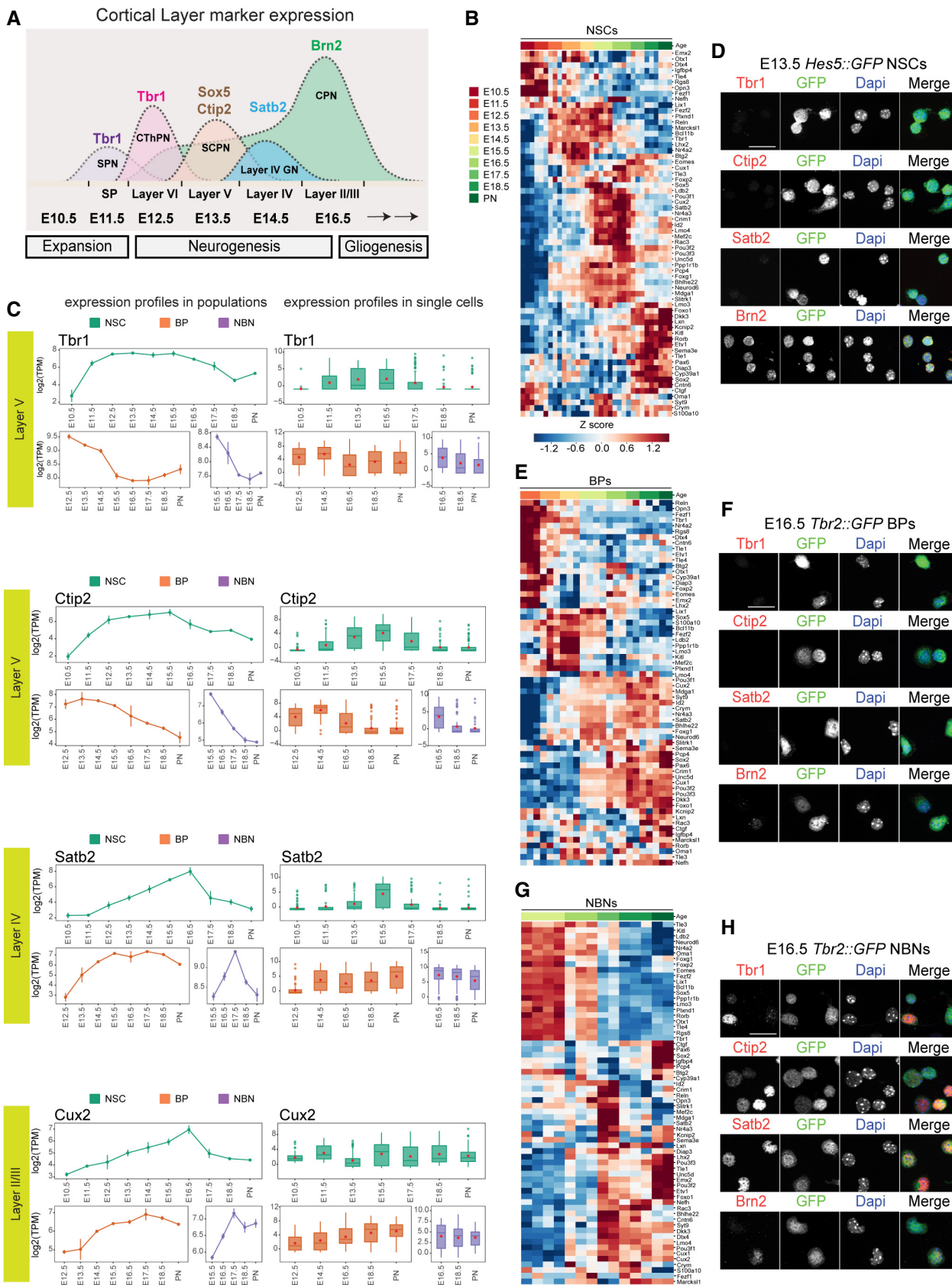


Figure 6.

governs stem cell maintenance and fate. However, it remains unclear to which signals NSCs, BPs, and NBNs are competent to respond *in vivo*. In order to evaluate susceptibility to paracrine signaling molecules and the dynamics in this responsiveness, we selected those genes designated to be receptors in the databases and analyzed the expression of the 440 receptors that showed variable gene expression throughout the neurogenic lineage. The resulting extensive gene expression profiles could be divided into three groups (Fig 7A and B):

- 1 Receptors that are highly expressed by NSCs through the most of cortical development (Fig 7A and B). These receptors, including those for Wnt (*Fzd5*, 7, 9), Notch (*Notch1*, 2, 3), Fgf (*Fgfr2*, *Fgfr3*), and Shh signaling (*Smo*, *Ptch1*), are part of pathways involved in stem cell maintenance (Blaschuk & French-Constant, 1998; Fukuchi-Shimogori & Grove, 2001; Gaiano & Fishell, 2002; Itoh & Ornitz, 2004; Bray, 2006; Louvi & Artavanis-Tsakonas, 2006; Shimojo *et al*, 2008, 2011; Iwata & Hevner, 2009; Sahara & O'Leary, 2009; Imayoshi *et al*, 2010; Rash *et al*, 2011; Wang *et al*, 2011).
- 2 Receptors that are highly expressed by NSCs during neurogenesis and later stages and by BPs and NBNs, which we refer to as neurogenic (Fig 7A).
- 3 Receptors that are highly expressed predominantly at later stages of development in the NSCs during the gliogenic phase. These we refer to as gliogenic pathways and include the receptors for known ligands involved in gliogenesis, including Tgf-beta/BMP signaling (*Tgfr2*, *Bmpr1a*, *Bmpr1b*) and Il6/Lif signaling (*Lifr*, *Il6st*) (Ebendal *et al*, 1998; Gomes *et al*, 2005; Rodriguez-Martinez & Velasco, 2012; Pollen *et al*, 2015).

The neurogenic niche during corticogenesis provides local autocrine and paracrine signals but also responds to blood-born ligands and factors in the fluid of the telencephalic vesicles. We assessed the potential local signals in the NSCs, BPs, and NBNs by examining the expression of ligands for the top, regulated receptors (Fig 7C). Similar to the expression profile of their cognate receptors, the expression of some ligands could be divided into three clusters. Ligands expressed predominantly by NSCs during the expansion phase of corticogenesis, ligands expressed predominantly by NSCs in the gliogenic phase, and ligands expressed mostly by neurons that act as paracrine signals back to the progenitors. Many Wnt ligands and their receptors are expressed by NSCs suggesting autocrine signaling. One notable exception being *Wnt7b*, which is prominently expressed by NBNs and its canonical receptor *Fzd7* also by NSCs at the expansion and gliogenesis phases. By contrast, although their cognate receptors were mostly expressed by NSCs, the Fgf ligands were

divided into two major groups: those expressed mainly by NBNs and those expressed mainly by NSCs (Fig 7A and C).

As a proof of concept, we also evaluated selected modulators and effector targets of some of the key signaling pathways including Bmp, Wnt, Notch, and Shh signaling (Fig 7D). The expression of many target genes of these pathways reflected the expression of their respective receptors suggesting that not only are the ligands available and the receptors expressed but the pathways may be active throughout the neurogenic lineage (Fig 7D).

bHLH TFs are dynamically and heterogeneously expressed by NSCs

bHLH TFs are notably involved in the control of neurogenesis and brain development downstream of many pathways including Notch, BMP, TGF β , and Wnts. We analyzed the expression profile of the bHLH family genes. The bHLH TFs could be grouped into three classes based on their expression profiles:

- 1 bHLH factors related to NSCs maintenance, including *Hes1*, *Hes5*, *Hey1*, and *Id4*, are highly expressed by NSCs (Fig 8A and B).
- 2 bHLH factors related to neuronal commitment and differentiation. For example, the proneural differentiation bHLH genes including *Neurog2*, *Neurod2*, and *Neurod6*, which are expressed by NSCs during the neurogenic phase and by BPs, but their expression is lower in NBNs (Fig 8A and C).
- 3 bHLH genes with expression associated prominently with gliogenesis, which are expressed at low levels by BPs and NBNs including *Olig1*, *Olig2*, and *Id1* (Fig 8A and D).

We also identified a group of bHLH TFs expressed moderately by NSCs during the neurogenic phase of corticogenesis, but which are expressed by BPs and NBNs suggesting a role in neural commitment and differentiation (Fig 8C and D). At the single cell level, expression of the bHLH factors by the NSCs was highly heterogeneous, even at the same embryonic time point (Fig 8E). As expected at E10.5, most NSCs expressed high levels stemness markers (*Hes1*, *Hey1*, and *Id4*) and low or no neurogenic-associated (*Neurog2*, *Neurod2*, and *Neurod6*) and gliogenic-associated bHLH TFs (*Olig1*, *Olig2*, and *Id1*). As neurogenesis initiated at E13.5, more cells started to express neurogenic markers and the number of cells expressing *Hes1*, *Hey1*, and *Id4* reduced, but very few cells were expressing the gliogenic bHLHs. At E13.5 two major NSC populations were evident based on bHLH expression, one expressing high stemness markers and low neurogenic markers, and another expressing low stemness markers and high neurogenic markers. However, there was also a subpopulation of cells that expressed

Figure 7. Dynamic expression profile of signaling receptors during corticogenesis.

- A Heatmaps representing the expression profile of signaling receptors that can be divided into three main groups based on k-means clustering of z-scored $\log_2(\text{TPM})$ expression values: stem cell maintenance (121 receptors), neurogenic (180 receptors), and gliogenic (139 receptors). Names of selected receptors are displayed. For the complete list please see Dataset E7. Expression profiles are represented by their z-score.
- B Average expression profile of each cluster for NSCs (green), BP (orange), and NBN (purple). Solid line represents the average z-score, while the area represents the SD estimated from different biological samples. Three to four biological replicates were collected for each time point.
- C Heatmap representing the expression profile of ligands from selected signaling pathways, based on the z-scored $\log_2(\text{TPM})$ expression values.
- D Expression profile of selected target or modulator of key signaling pathways: BMP, Wnt, Shh and Notch signaling. Each dot defines the mean and lines define the SD. Three to four biological replicates were collected for each time point.

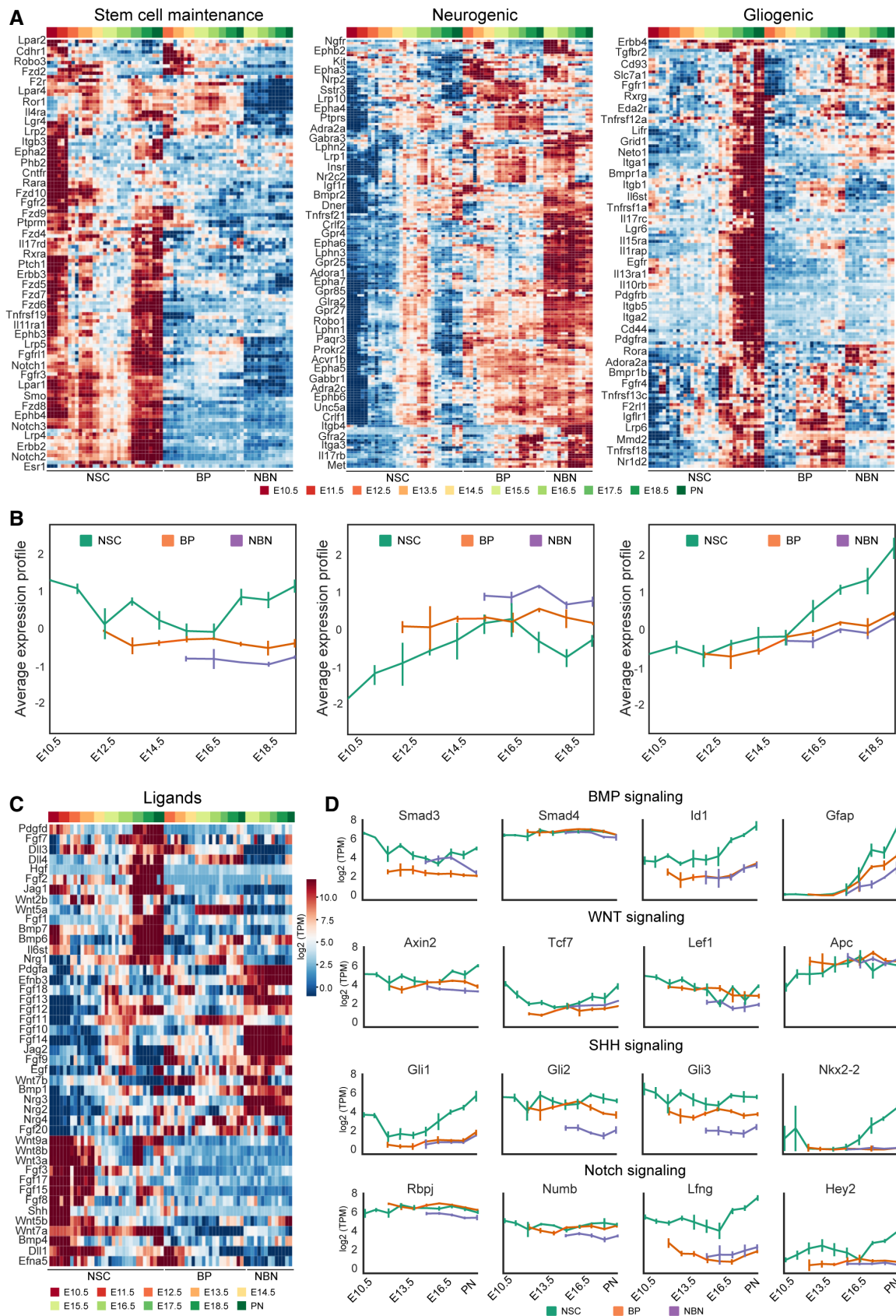


Figure 7.

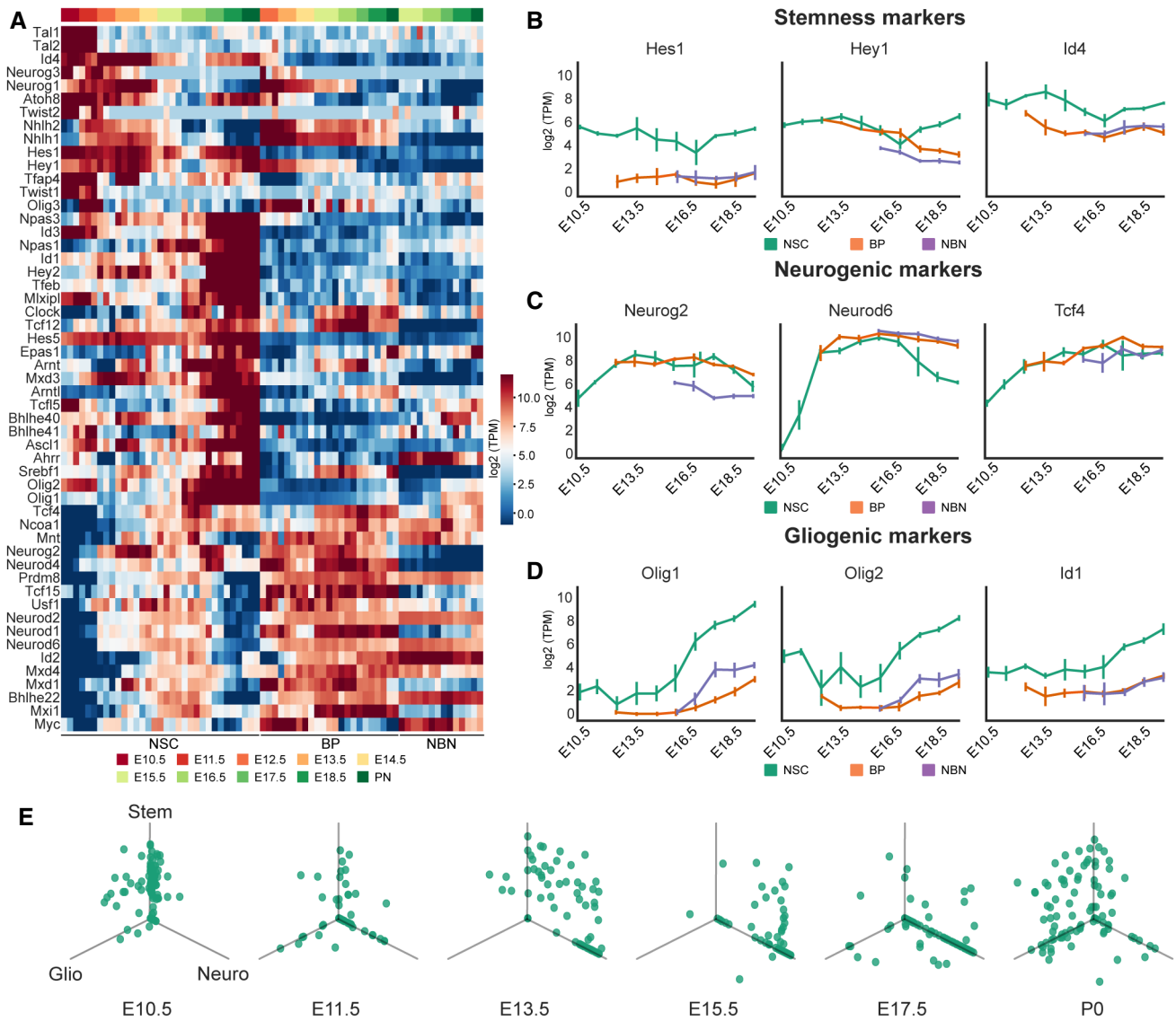


Figure 8. Dynamic and heterogenic expression profile of bHLH factors during forebrain development.

- A Heatmaps representing the expression profile of bHLH factors. Three main groups are observed based on k-means clustering of z-scored $\log_2(\text{TPM})$ of expression value: stem cell maintenance (high expression in the NSCs at early embryonic times and low in BPs and NBNs), neurogenic (high expression in the NSCs during neurogenesis and high expression in BPs and NBNs), and gliogenic (high expression in the NSCs at late embryonic times and low in BPs and NBNs). Expression profiles are represented by their z-score.
- B Expression profile of selected stem cell maintenance markers Hes1, Hey1 and Id4. Each dot defines the mean and lines define the SD. Three to four biological replicates were collected for each time point.
- C Expression profile of selected neurogenic markers Neurog2, Neurod2, and Neurod6. Each dot defines the mean and lines define the SD. Three to four biological replicates were collected for each time point.
- D Expression profile of selected gliogenic markers Olig1, Olig2 and Id1. Each dot defines the mean and lines define the SD. Three to four biological replicates were collected for each time point.
- E Expression of stem cell markers (Hes1, Hey1 and Id4), neurogenic markers (Neurog2, Neurod2, and Neurod6) and gliogenic markers (Olig1,2 and Id1) in NSCs during different embryonic time points in the single-cell levels. Each point represents the expression value of one single cell in $\log_2(\text{TPM})$.

both maintenance and neurogenesis-associated bHLH TFs. One explanation for this populations is the oscillatory expression of stemness factors *Hes1* and *Hes5* and their repression of the neurogenic targets including *Neurog2*. At later stages, when NSCs exit the neurogenic phase and enter gliogenesis (E15.5-E17.5), the proportion of NSCs expressing the neurogenic bHLHs rather than the stem

cell maintenance-associated TFs increased. As the NSCs transitioned into gliogenesis and toward PN1, the proportion of NSCs expressing *Neurog2*, *Neurod2*, and *Neurod6* diminished with a concomitant increase in gliogenic factor (*Olig1*, *Olig2*, and *Id1*) expressing cells (Fig 8E). Strikingly, some NSCs in the gliogenic state coexpressed the maintenance bHLH TFs. This confirms previous observations

that Notch signaling and the expression of Hes-related TFs are linked to glial commitment of NSCs.

Discussion

The temporal dynamics in gene expression during lineage commitment throughout ontogeny of the cerebral cortex remains unclear. Advances in scRNA-Seq and gene cluster analysis have given unprecedented insights into cellular heterogeneity in the mammalian cortex, as well as in primary human samples. Particularly when trying to understand the genetic regulation of cell diversification from stem and progenitor cells, these snapshots of cellular transcriptomics are used to define cellular state and therefore predict fate potential. One challenge for transcriptome analysis is to predict not only the future of a particular cell and its offspring but also its history. This is particularly confounded by highly dynamic gene expression over time windows ranging from days to minutes.

Here we posed the questions of how gene expression changes in stem cells, progenitors, and newly formed neurons over time, and whether we can reveal distinct gene expression patterns within specific cell types that may allude to their differentiation potential. We characterized gene expression of the dorsal cortical neural lineages over time, focusing on the phases of expansion, neurogenesis, and gliogenesis using NSC, BP, and NBN populations with definitive characteristics. We have created an extensive resource of dorsal cortical ontogeny, which can be mined through an interactive web-based browser (Appendix Fig S5; <http://neurostemx.ethz.ch/>).

Development of the cerebral cortex is a dynamic process, however, and remarkably, our understanding of the lineage heterogeneity and changes in gene expression that accompany the formation of the different neuron subtypes and subsequent cortical layers is limited. The most widely accepted model of cortical development utilizes a common multipotent progenitor, which becomes progressively restricted in its fate over time. Unbiased computational analysis of our data revealed distinctive, stage-specific changes in gene expression not only in NSC, but also BPs and NBNs. These shifts in transcriptional space at the population and single cell levels reveal a heterogeneity in each of these cell populations and establish novel gene signatures defining five NSC, three BP and two NBN types. Remarkably, we show the presence of different NSC, BP, and NBN types at the same developmental stage, and that these constitute different proportions of the particular populations at each stage. Although these findings do not disprove a common progenitor mechanism, they imply that the populations of NSC, BPs, and NBNs at any point in developmental time are composed of different proportions of cells with distinct transcriptomes, which can be predicted by a panel of signature genes.

We analyzed our dataset using two separate clustering methods and identified similar distinct clusters of NSCs, BPs, and NBNs, highlighting the robustness of our data. We also integrated our C1 data with 10X genomics data from the developing mouse forebrain (La Manno *et al*, 2021). Our cells integrated into the 10X genomics data as expected, while maintaining their distinct groupings, again validating their broader utility. We consider the use of transgenic *Hes5::GFP* and *Tbr2::GFP* lines as a major strength of the paper because we were able to enrich clean populations of NSCs, BPs, and NBNs using these lines. To understand the neurogenic trajectories,

we performed the pseudo-time analyses, which revealed three lineages, from NSCs to BPs to NBNs following a strict time component. These observations again emphasize the tight regulation of neural fate dynamics.

Our analyses of cells types with definitive characteristics exemplify potential dangers in *a priori* allocation of cell type based on comparative gene expression and limited gene sets. For example, the transcriptomes of NSC in the neurogenic phase of cortical development are closer to BPs at this stage than they are to NSCs at E10.5 (expansion phase) or PN (gliogenic phase). Only by separating the NSCs, BPs, and NBNs and analyzing their transcriptomes in isolation, was it possible to uncover specific signatures and determine cellular heterogeneity. We also found that NSCs express many neuronal RNAs but with no detectable protein expression. This suggests that essential posttranscriptional regulation programs are active in the neurogenic lineage during differentiation. Several mechanisms of posttranscriptional regulation have previously been reported during embryonic neurogenesis including by Drosha (Knuckles *et al*, 2012) and m⁶ mRNA methylation (Yoon *et al*, 2017).

The conjunction of gene expression dynamics and predicted transcriptional networks by ISMARA identified active TFs and nodes in NSCs, BPs, and NBNs over time. These TF motifs and activities also revealed the same directional trajectory and pathway of each cell type through transcriptional space as predicted by the mRNA expression. As ISMARA predicts the activity of more than 800 TFs and their targets in all cell types, this dataset and resource will be valuable to explore and extrapolate the known regulatory networks to the missing novel nodes. Recently, we validated the Tead TFs in cortical development and elucidated different functions of Tead factors in NSCs (Mukhtar *et al*, 2020). The dynamic changes in the TFs in NSCs, for example, reflect the sequential changes these cells undergo during corticogenesis. Our analyses determining the relationship among the neuronal lineage demonstrate a naturally occurring directionality, indicating strong intrinsic control. Moreover, among all the cell types, NSCs seem to be most dynamic, be it at gene expression level or the level of transcriptional networks. The NSCs follow a continuous path through the three phases of corticogenesis, supporting the neuronal origin from “common progenitors,” sequentially changing in transcriptional space. To this end, we identified signature genes of NSCs in expansion, neurogenic and gliogenic phases. As all the known genes depicted the dynamics of expression as expected, we believe that we provide more extensive lists of novel signature genes, which could be used to identify NSCs in different phases. These signatures are like a “scorecard” for the NSC population undergoing corticogenesis, some of which we have also validated experimentally (Figs EV2B and C, and EV3C and D). Similar analyses for BPs and NBNs have yielded key signature markers, which hold promise for further biological exploration. The up- and downregulations of genes could be presumed to be the result of active or inactive downstream programs in these cells and their progeny. It is crucial to differentiate between early and late BPs, or different NBN populations across time in order to consolidate our knowledge about their downstream fate and function.

The microenvironment of the cells plays critical roles in regulating cell fate choices. We used these resources and explored some of the signaling pathways defining logic in NSC differentiation using a high-throughput microfluidic approach (Zhang *et al*, 2019). This validation was the tip of the iceberg and together with the recent

developments in the field, we provide a consolidated resource, a comprehensive and systematic characterization of major progenitor pools in cortical development. Further biological validations of our

predicted signaling and transcriptional nodes will provide more promise toward the deeper exploration of mechanisms controlling corticogenesis.

Materials and Methods

Reagents and Tools table

Antibodies	Source	Identifier
Chick anti-GFP (1:300)	Millipore	Cat# 06-896, RRID:AB_11214044
Rat anti-Ctip2 (1:500)	Abcam	Cat# ab18465, RRID:AB_2064130
Goat anti-Brn2 (1:250)	Santa Cruz	Cat# sc-6029, RRID:AB_2167385
Sheep anti-GFP (1:250)	AbD Serotec/Biorad	Cat# 4745-1051, RRID:AB_619712
Rabbit anti-Tbr2 (1:500)	Abcam	Cat# ab23345, RRID:AB_778267
Rabbit anti-Pax6 (1:500)	Covance	Cat# PRB-278P, RRID:AB_291612
Mouse anti-Satb2 (1:200)	Abcam	Cat# ab51502, RRID:AB_882455
Rabbit anti-Tbr1 (1:500)	Abcam	Cat# ab31940, RRID:AB_2200219
Donkey anti-Sheep, Alexa 488 (1:500)	Jackson ImmunoResearch Labs	Cat# 713-545-147, RRID:AB_2340745
Donkey anti-Rabbit, Cy3 (1:500)	Jackson ImmunoResearch Labs	Cat# 711-165-152, RRID:AB_2307443
Donkey anti-Mouse, Cy3 (1:500)	Jackson ImmunoResearch Labs	Cat# 715-165-151, RRID:AB_2315777
Donkey anti-Rat, Cy3 (1:500)	Jackson ImmunoResearch Labs	Cat# 712-166-153, RRID:AB_2340669
Donkey anti-Goat, Cy3 (1:500)	Jackson ImmunoResearch Labs	Cat# 705-165-147, RRID:AB_2307351
Donkey anti-Chicken, Alex488 (1:500)	Jackson ImmunoResearch Labs	Cat# 703-545-155, RRID:AB_2340375
Chemicals	Source	Identifier
Formaldehyde Solution (w/v)	Sigma	47608(47673/33220)
DNase I, RNase-free	Sigma	04716728001
DNase I Grade II	Roche	10104159001
L_Cysteine	Sigma	168149
Papain	Sigma	P3125-100MG
Trypsin inhibitor from Glycine max (soybean)	Sigma	T6522-5x100MG
L15 Medium	Invitrogen	31415029 (31415086)
PBS cell culture	Dulbecco	14080089 (14080048)
Triton X-100	Fisher	BPE151-500
TRizol	Invitrogen	VX15596018
Glycoblue Co-precipitate	Life Technologies	D1417005
Poly L- Lysine hydrobromide	Sigma	P9155-5MG
B27 supplement+A26	Gibco	17504-044
DMEM/F12	Gibco	31966-047
Chloroform	Sigma	288306
Normal Donkey Serum	Jackson ImmunoResearch Labs	017-000-121
Agarose	Fisher Scientific	BPE1356-100
RNase free water	Ambion	AM9906
TE buffer	Invitrogen by Thermo Fisher Scientific	AM9849
NaOH	Roth	6785.1
Chamber slides	Lab-Tek	177402
Critical Commercial Assays	Source	Identifier
2x Assay Loading Reagent	Fluidigm	85000736

Reagents and Tools table (continued)

Critical Commercial Assays	Source	Identifier
20x DNA Binding Dye Sample Loading Reagent	Fluidigm	100-3738
20x GE Sample Loading Reagent	Fluidigm	85000746
Sso Fast EvaGreen SuperMix with low Lox	BioRad	172-5211
Dynamic Array 48.48	Fluidigm	BMK-M-48.48
Dynamic Array 96.96	Fluidigm	BMK-M-96.96
PreAmp and Reverse Transcription Master Mix	Fluidigm	100-6300
Human Brain Reference RNA	Life Technologies	AM6050
Exonuclease I	New England Biolabs	M0293L
C1 Single Cell Auto Prep Array for mRNA Seq Multipack —contains:	Fluidigm	100-6041 100-5518 100-6209 100-5757
• Module 1 Single Cell Auto Prep Kit		
• Module 2 mRNA Seq		
• C1 Single Cell Auto Prep Array for mRNA Seq (5–10 µm)		
SMARTer Ultra Low RNA Kit for Illumina Sequencing	Clontech/Takara	634936
Advantage 2 PCR Kit	Clontech/Takara	639206
Nextera XT DNA Library Preparation Kit	Illumina	FC-131-1096
Nextera XT Index Kit v2 Set A	Illumina	FC-131-2001
Nextera XT Index Kit v2 Set B	Illumina	FC-131-2002
Nextera XT Index Kit v2 Set C	Illumina	FC-131-2003
Nextera XT Index Kit v2 Set D	Illumina	FC-131-2004
Agencourt AMPure XP	Beckman Coulter	A63882
DNA Suspension Buffer, pH 8.0	Teknova	T0221
HS NGS Fragment 35-6000bp.mthds	Labgene Scientific SA	DNF-486-0500
HS NGS Fragment 1-6000bp.mthds	Labgene Scientific SA	DNF-474-0500
SS NGS Fragment 35-6000bp.mthds	Labgene Scientific SA	DNF-479-0500
SS NGS Fragment 1-6000bp.mth	Labgene Scientific SA	DNF-473-0500
RNA 6000 Pico Complete Kit	Agilent Technologies	5067-1513
Agilent RNA 6000 Nano Kit	Agilent Technologies	5067-1511
Quant-IT RiboGreen® RNA Assay Kit	Life Technologies	R11490
Quant-IT PicoGreen® dsDNA Assay Kit	Life Technologies	P11496
GREINER-384-Well plate, black	Greiner	784076
TruSeq RNA Library Preparation Kit v2, Set A	Illumina	RS-122-2001
TruSeq RNA Library Preparation Kit v2, Set B	Illumina	RS-122-2002
Superscript II Reverse Transcriptase	Life Technologies	18064-014
twin.tec PCR Plate 96, semi-skirted	Vaudaux-Eppendorf AG	0030 128.575
Ethanol absolute Honeywell 1L	Honeywell	02860-1L
10mM TRIS-HCl with 0.1% TWEEN-20, pH 8.5	TEKNOVA	T7724
Experimental models	Source	Identifier
Mouse: <i>Hes5::GFP</i>	Verdon Taylor (Basak & Taylor, 2007)	N/A
Mouse: <i>Tbr2::GFP</i>	Arnold et al (2009)	N/A
Oligonucleotides	Source	Identifier
Ccnd1_Forward_5'- TGCCGAGAAGTTGTCATCTA-3'	This paper	N/A
Ccnd1_Reverse_5'- TGTTACCAGAAGCAGTTCCA-3'	This paper	N/A
Crabp2_Forward_5'-ATGCCTAACTTTTCTGGCAACT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Crabp2_Reverse_5'-GCACAGTGGTGGAGTTTGA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Hbb-bh1_Forward_5'-GAAACCCCGATTAGAGCC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A

Reagents and Tools table (continued)

Oligonucleotides	Source	Identifier
Hbb-bh1_Reverse_5'-GAGCAAAGTCTCCTTGAGGT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Bcl11b_Forward_5'-CCCAGCCTGATCTACTCAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Bcl11b_Reverse_5'-CTCCTGCTGGACAGATGCC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Bhlhe22_Forward_5'-AAGCGCATCAAGTGGAGAA-3'	This paper	N/A
Bhlhe22_Reverse_5'-CTTGGTTGAGGTAGGCGACTAA-3'	This paper	N/A
Cabp1_Forward_5'-GAGCTGTCTCAGCAGATCAAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cabp1_Reverse_5'-TTTAGGGCCCATCAGTTCCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cntn2_Forward_5'-GCTGATGCCATGACCATGAA-3'	This paper	N/A
Cntn2_Reverse_5'-ACTTAAGGCTGAGCTGGAA-3'	This paper	N/A
Id2_Forward_5'-ACCCTGAACACGGACATCA-3'	This paper	N/A
Id2_Reverse_5'-TCGACATAAGCTCAGAAGGAA-3'	This paper	N/A
Satb2_Forward_5'-GCCGTGGGAGGTTTGATGATT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Satb2_Reverse_5'-ACCAAGACGAACCTCAGCGTG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tubb3_Forward_5'-GCGCATCAGGTATACTACA-3'	This paper	N/A
Tubb3_Reverse_5'-AGGTTCCAAGTCCACCAGAA-3'	This paper	N/A
Fezf2_Forward_5'-GTCACCGGCCACTCTAAAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Fezf2_Reverse_5'-GTCTGCCTTAACGCAGCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
ApoE_Forward_5'-CTGACAGATGCCTAGCCG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
ApoE_Reverse_5'-CGCAGTAATCCAGAAGC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Aqp4_Forward_5'-CTTTCTGGAAGGCAGTCTCAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Aqp4_Reverse_5'-CCACACCGAGCAAAACAAGAT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cspg4_Forward_5'-GGGCTGTGCTGTCTGTGA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cspg_Reverse_5'-TGATTCCTCAGTAAGGCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Hmgb2_Forward_5'-GTGGCAGGTACATGCAATCC-3'	This paper	N/A
Hmgb2_Reverse_5'-GTACTTTGGTGGTGTCTCA-3'	This paper	N/A
Olig1_Forward_5'-CTGTATGAGCTGGTGGTTACA-3'	This paper	N/A
Olig1_Reverse_5'-GAGAAGGATCGCGTGGAA-3'	This paper	N/A
Pdgfra_Forward_5'-AGAGTTACAGTGTGAGCTGTC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Pdgfra_Reverse_5'-GTCCCTCCACGGTACTCCT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Sparcl1_Forward_5'-GGCAATCCGACAAGTACAAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Sparcl1_Reverse_5'-TGGTTTTCTATGTCTGTGAGC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tril1_Forward_5'-CTATGTATGCCGTTGGGTAGG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tril1_Reverse_5'-AGCTTTTCACTATTTCCGCCAT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cckar1_Forward_5'-CTTTCTGCCTGGATCAACCT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Cckar1_Reverse_5'-ACCGTGATAACCAGCGTGTTC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ccnb1_Forward_5'-AAGGTGCCTGTGTGAACC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ccnb1_Reverse_5'-GTCAGCCCATCATCTGCG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Dhrs4_Forward_5'-CCTGTGCTCCTCCATCCTA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Dhrs4_Reverse_5'-GCAAGGTGTCTTTTGTGGGA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tbr2_Forward_5'-GCGCATGTTCTTTCTTGAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tbr2_Reverse_5'-GGTCGGCCGAACCACTTC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Kif2c_Forward_5'-ATGGAGTCGCTTACGCAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Kif2c_Reverse_5'-CCACCGAAACACAGGATTTCTC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Mcm2_Forward_5'-ATCCACCACCGCTTCAAGAAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Mcm2_Reverse_5'-TACCACCAACTCTCAGGTT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Uncx1_Forward_5'-ACCCGACCAACTTTACCG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A

Reagents and Tools table (continued)

Oligonucleotides	Source	Identifier
Uncx_Reverse_5'- TGAAGCTCGGACTCGACCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Robo3_Forward_5'- AGATGAACTGTTCGCGGACT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Robo3_Reverse_5'- GGAAGCAGACTAGGGTTGAGC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Nde1_Forward_5'-ATGGAGGACTCGGAAAGACC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Nde1_Reverse_5'-TCAGTCTCGTATTCTCGGCTT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tpx2_Forward_5'- GATCCCCACCGACTTTATC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Tpx2_Reverse_5'- CTTGTCTCCAAGTTGGCCTT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Lox11_Forward_5'- GAGTGCTATTGCGCTTCCC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Lox11_Reverse_5'- GGTGCCGAAGTACACAGT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Unc5d_Forward_5'- TGGCTAGGACTCTTTTTCTGGG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Unc5d_Reverse_5'- GCTCCTCGATGAAATGAGCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ezr_Forward_5'- CAATCAACGTCCGGTGAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ezr_Reverse_5'-GCCAATCGTCTTTACCACCTGA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Mef2c_Forward_5'- GTCAGTGGGAGCTTGCACTA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Mef2c_Reverse_5'- CGGTCTCTAGGAGGAAACA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Usp43_Forward_5'- AGCTCACGGCTGGTATCT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Usp43_Reverse_5'- AAGACCTGTACTGTCTTGAAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Lrfr5_Forward_5'- TGTTTCTATTGGCATAGCTGT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Lrfr5_Reverse_5'- TGGTGGAACAATAGAAGCCCT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ntsr1_Forward_5'-CAGTTCGACTGGAGACGATG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ntsr1_Reverse_5'- ACCAGCACCTTGAATAAATGTC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Gucy1a3_Forward_5'- CCCCTGGTCAGGTTCTAAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Gucy1a3_Reverse_5'- GGAGACTCCCTTCTGATTCT-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
β-actin_Forward_5'- AGGTGACAGCATTGCTTCTG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
β-actin_Reverse_5'- GGGAGACCAAGCCTTCATA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ubb_Forward_5'- TCTGAGGGTGGCTATTAA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Ubb_Reverse_5'-TGCTTACCATGCAACAAAAC-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Topp_Forward_5'-GGCTGTACAGACTAGAAGAGCA-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Topp_Reverse_5'-CCTCTCGATCTGTGGCTTG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Gapdh_Forward_5'- CTCCACTCTCCACCTTCG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Gapdh_Reverse_5'- CCACCACCTGTGTGTAG-3'	https://pga.mgh.harvard.edu/primerbank/	N/A
Resource	Source	Identifier
Fiji	Hosted by University of Wisconsin	https://imagej.net/Fiji/Downloads
Photoshop	Adobe	N/A
Illustrator	Adobe	N/A
Prism 7	GraphPad Software, Inc	https://www.graphpad.com/scientific-software/prism/
R	R Core Team	https://www.r-project.org
MATLAB R2016a 9.0.0.341360	MathWorks	mathworks.com/products/matlab
Python 2.7.11	Python Software Foundation	www.python.org
Python 3.6	Python Software Foundation	www.python.org
goatools		https://github.com/tanghaibao/goatools
InChIlib		https://openscreen.cz/software/inchlib/home/
MGI_Gene_Model_Coord.rpt		http://www.informatics.jax.org/downloads/reports/index.html
fastcluster		http://www.danifold.net/fastcluster.html
Inkscape	The Inkscape Project	inkscape.org

Methods and Protocols

Experiment model and subject details

Hes5::GFP (Basak & Taylor, 2007) and *Tbr2::GFP* (Arnold et al, 2009) transgenic lines have been described previously. Mice were maintained on a 12-h day–night cycle with free access to food and water under specific pathogen-free conditions and according to the Swiss federal regulations. All procedures were approved by the Basel Cantonal Veterinary Office (license number ZH_Tay).

Method details

Tissue preparation and fluorescence-assisted cell sorting (FACS)

Dorsal cortices from embryonic day (E10.5) to postnatal day 1 (PN) were micro-dissected and dissociated into single-cell suspensions using Papain and Ovo-mucoid mix (as described previously Giachino et al, 2009). Cells were washed with L15 medium and FAC-sorted for GFP-positive NSCs using FACSariaIII (BD Biosciences) derived from *Hes5::GFP* transgenic embryos for NSCs and *Tbr2::GFP* transgenic embryos for BPs and NBNs. For each time point, 3–4 biological replicates were generated.

RNA isolation and RNA sequencing

Total RNA was isolated from FAC-sorted GFP-positive cells from *Hes5::GFP* and *Tbr2::GFP* transgenic lines using TRIzol reagent. A time course was performed with NSCs, BPs, and NBNs isolated at each time point during development from E10.5 to postnatal day 1 (PN) or as specified in the Fig 1A. Samples were analyzed for their integrity and concentration using Agilent 2100 Bioanalyzer and Quant-IT RiboGreen RNA Assay Kit. Sequencing libraries were prepared with the Illumina TruSeq RNA Library Prep Kit v2 according to Illumina's instructions. After quality control (Fragment Analyzer, AATI), libraries were pooled and loaded on an Illumina flow cell for cluster generation (HiSeq SR Cluster Kit v4 cBot). Libraries were sequenced SR50 on the HiSeq 2500 system (HiSeq SBS Kit V4) following the manufacturer's protocols.

Single-cell RNA sequencing

Single-cell capture, lysis, and cDNA preparation were performed with the Fluidigm C1 system. Cells were loaded on a microfluidic C1 Single Cell Auto Prep Array for mRNA Seq (5–10 µm), and capture efficiency evaluated using microscopy. Lysis, reverse transcription, and cDNA amplification were performed with the SMARTer Ultra Low RNA Kit for Illumina Sequencing (Clontech/Takara) according to Fluidigm's guidelines for single-cell RNA-seq on the C1 system. cDNA was harvested, profiles checked on the Fragment Analyzer (AATI), and their concentration determined using Quant-iT PicoGreen dsDNA Assay Kit. For subsequent library preparation using Nextera XT DNA library preparation kit (Illumina) following the Fluidigm manual, cDNAs were normalized to 0.3 ng/µl. Libraries were pooled and sequenced SR75 on an Illumina NextSeq 500 system (75 cycles High Output v2 kit).

qPCR validation

Total RNA was isolated from FAC-sorted GFP positive cells from *Hes5::GFP* (E11.5, E15.5 and E18.5) and *Tbr2::GP* (E13.5 BPs, E15.5 BPs and E15.5 NBNs) transgenic embryos using TRIzol reagent. Independent biological replicates were generated for RT-qPCR validation. Samples were analyzed for their integrity and concentration

using Agilent 2100 Bioanalyzer and Quant-IT RiboGreen RNA Assay Kit. DNase treatment was done using Roche DNase kit and cDNA prepared using the PreAmp and Reverse Transcription Master Mix from Fluidigm. Deltagene Assay primers (Fluidigm) and EvaGreen (BioRad) were used for real-time qPCR. Gene expression was assayed using Dynamic Array IFC chips and the BioMark system (Fluidigm). Fluidigm real-time PCR analysis software was used to calculate cycle threshold (C_t) values for each qPCR.

Tissue preparation and immunohistochemistry

Hes5::GFP and *Tbr2::GFP*-positive brains at E17.5 were isolated and fixed with 4% PFA in 0.1 M phosphate buffer (PBS). Brains were embedded in 3% agarose, sectioned 40 µm thick using a Vibrotome. Sections were mounted in mounting media containing diazabicyclooctane (DABCO; Sigma) as an anti-fading agent on SuperFrost glass slides and visualized using Zeiss Apotome 2 microscope. For immunostainings, fixed brains were equilibrated serially with 15% and then 30% sucrose solution in PBS overnight at 4°C. Brains were embedded in 100% OCT and 20 µm sections collected on SuperFrost glass slides.

Adherent NSC culture in vitro and immunocytochemistry

Primary NSCs were isolated from E13.5 dorsal cortices from *Hes5::GFP* transgenic embryos and BPs, NBNs were isolated at E16.5 from *Tbr2::GFP* transgenic embryos. Following FAC-sorting, the cells were seeded in 100 µg/ml Poly L-Lysine pre-coated 8-well Lab-Tek chamber slides and cultured in DMEM/F12 + Glutamax medium (with 2% B27). The cells were incubated for 1 h at 37°C, 5% CO₂. The cells were fixed with 4% PFA, at RT for 15 min and blocked with 5% Normal donkey serum and 0.1% Triton X-100. Primary antibody incubations were performed overnight at 4°C. Secondary antibody incubations were performed at RT, for 1 h. The cells were incubated with 1:1000 Dapi for 30 min at RT and rinsed with PBS. Slides were mounted with DABCO and imaged using Zeiss Apotome 2 microscope.

Quantification and statistical analysis

Images taken by Zeiss Apotome 2 were processed with FIJI software. Contrast and image size of IF images were adjusted with Adobe photoshop. Expression profiles of genes of interest were produced in R. Bar graphs were generated by GraphPad Prism 7. All figures were made in Adobe Illustrator CS6.

Sample size is mentioned in the excel sheets for the quantifications. For FACS analysis, for *Hes5::GFP* transgenic embryos, only the bright GFP-positive cells were collected. For *Tbr2::GFP* transgenic embryos, both bright and dim GFP-positive cells were collected and analyzed. For IF images, three fields of views were analyzed and quantified per sample. Unpaired *t*-tests were used for qPCR validation experiments. The cutoff value for statistical significance was indicated in corresponding tables.

Read mapping and data preprocessing

Reads from single cell and cell population mRNA-Seq were mapped to the transcriptome (GENCODE Release M2 GRCm38.p2) with kallisto 0.43.0[*]. The option --pseudobam was used to save the pseudoalignments to transcriptome in BAM file. The reads mapping to multiple transcripts were uniformly distributed. To obtain the expression per transcript, we first divided the number

of reads mapping to each transcript by the length of the transcript in nucleotides and then transformed the length-normalized read counts in transcript per million (TPM). Gene expression was obtained by summing for each gene the TPM of the transcripts corresponding to the gene. Promoter expression was obtained by summing for each promoter the length-normalized count of the transcripts associated with the promoter and then transformed in TPM. We added a pseudo-count of 0.5 to express transcript, gene, and promoter expression in logarithmic space ($\log_2(\text{TPM} + 0.5)$). For the population mRNA-Seq, we computed replicate averages in $\log_2(\text{TPM} + 0.5)$. The method used is adapted from Bray *et al* (2016a, 2016b).

Differentially expressed genes in different cell types

A pairwise comparison between each two cell types is applied using tximport and Deseq2 packages in R. Next, the first 50 top DEGs (differentially expressed genes) for each cell type have been selected considering fold change of more than 2 and adjusted *P*-value < 1e-3. Finally, the common DEGs of each cell type are used for visualization. The complete list of DEGs of each comparison is given in (Dataset EV1).

The goal of our analysis was to find the most optimal marker genes. That is, if we were to only make gene expression measurements of a few genes (using qPCR, for example), those that give us the most information about the sample. When we are only interested in knowing whether the sample belongs to one of two classes (e.g., NSC vs. non-NSC), this information content is given by the conditional entropy described below. Hence, we use it as a score to find good marker genes. In the derivation, we account for the fact that the empirical expression variance from a few data does not necessarily reflect its true variance by using a prior that makes very small and very large variances unlikely.

Assuming that the probability $P(x|w, \mu)$ to measure log-expression x of a gene follows a Gaussian distribution with mean μ and inverse variance w , and using a uniform prior for μ and a gamma-distribution prior $P(w|\alpha, \beta) = \beta^\alpha w^{\alpha-1} \exp(-\beta w) / \Gamma(\alpha)$ for w , we find the likelihood of getting a set of measurements $D^c = (x_1, x_2, \dots, x_{n^c})$ for samples of class c to be

$$P(D^c|\alpha, \beta) = \frac{\beta^\alpha}{(\beta + n^c v^c / 2)^{\alpha + (n^c - 1)/2}} \frac{\Gamma(\alpha + (n^c - 1)/2)}{\Gamma(\alpha)}$$

where v^c is the empirical variance of D^c . Hence, we numerically find the maximum-likelihood estimates α^*, β^* from maximizing the sum of log-likelihoods across all genes. Finally, the inferred probability distribution of x in class c is:

$$P(x|c) = Z^c \left(1 + \frac{(x - x^c)^2}{V^c} \right)^{-\gamma^c}$$

where $x^c = \langle x \rangle_c$, $V^c = (n^c + 1)(v^c + 2\beta^*/n^c)$, $\gamma^c = \alpha^* + (n^c - 1)/2$ and $Z^c = \frac{\Gamma(\gamma^c)}{\sqrt{\pi V^c} \Gamma(\gamma^c - 1/2)}$. This distribution is approximately Gaussian with variance $\sigma^2 = V^c / (2\gamma^c)$, which provides us with a more accurate estimate of the true variance of a gene rather than simply taking v^c .

Furthermore, we can take the expression of $P(x|c)$ and $P(c) = 1/|c|$ to calculate the conditional entropy $H(c|x) = H(x, c) - H(x)$. While

$$H(x, c) = -\sum_c \int dx P(x|c)P(c) \log [P(x|c)P(c)] \\ = -\log(Z^c) + \gamma^c(\psi(\gamma^c) - \psi(\gamma^c - 1/2)),$$

with ψ being the digamma function, has an analytical solution, $H(x) = \int P(x) \log(P(x))$ with $P(x) = \sum_c P(x|c)P(c)$ can be calculated through numerical integration.

In an experiment that only measures the expression of a single gene, $H(c|x)$ serves as a measure for how much information the result provides about the class of the sample. With only two classes, we can write $H(x|c) = -p_e \log p_e - (1 - p_e) \log(1 - p_e)$, which we can numerically invert to find p_e , the probability to falsely classify a sample based on gene expression. The table below summarizes the classes for which we looked for such marker genes:

	Class 2	Number of marker genes $p_e < 0.01$
By cell type		
NSC	Non-NSC	37
BP	Non-BP	0
NBN	Non-NBN	136
NSC	BP	49
NSC	NBN	469
BP	NBN	249
NSC by phase		
Expansion NSC	Non-expansion NSC	222
Neurogenesis NSC	Non-neurogenesis NSC	4
Gliogenesis NSC	Non-gliogenesis NSC	102
Expansion NSC	Neurogenesis NSC	207
Expansion NSC	Gliogenesis NSC	759
Neurogenesis NSC	Gliogenesis NSC	117
Neurogenic (E12.5–16.5) NSC by day		
E12.5	Other days	0
E13.5	Other days	3
E14.5	Other days	0
E15.5	Other days	1
E16.5	Other days	50
E12.5-E13.5	E14.5-E16.5	9
E12.5-E14.5	E15.5-E16.5	16
BP by day		
E12.5	Other days	248
E13.5	Other days	0
E14.5	Other days	29
E15.5	Other days	0
E16.5	Other days	1
E17.5	Other days	3
E18.5	Other days	1
PN	Other days	38
E12.5-E13.5	E14.5-PN	54

	Class 2	Number of marker genes $p_e < 0.01$
E12.5-E14.5	E15.5-PN	83
E12.5-E15.5	E16.5-PN	0
E12.5-E16.5	E17.5-PN	15
E12.5-E17.5	E18.5-PN	4
Other		
BP E12.5-E14.5	NBN	354
BP E15.5-PN	NBN	650

Gene regulatory networks

Using a set of curated binding motifs for 503 families of transcription factors, ISMARA predicts TF binding sites in the promoters (TSS \pm 500 bp) of all genes' genome-wide and summarizes these predictions by total count N_{pm} of the number of binding sites for motif m on the promoter region of promoter p . Using a matrix of measured gene expression of promoter p in sample s E_{ps} (measured in logarithms of transcripts per million transcripts), ISMARA infers the matrix A_{ms} of activities of each motif m in each sample s , using the likelihood

$$P(\mathbf{E} | \mathbf{A}, \mathbf{N}, \sigma) \propto \prod_{ps} 1/\sigma \exp \left[- (E_{ps} - \sum_m N_{pm} A_{ms})^2 / 2\sigma^2 \right]$$

together with a Gaussian prior on the motif activities. To quantify the significance of the regulatory interaction from each motif m to each target promoter p (where all promoters that contain at least one binding site for motif m are potential targets), ISMARA calculates the log-likelihood ratio of the model without and with the specific interaction N_{pm} . The core regulatory networks of Fig 3E and H were obtained by taking all interactions with a log-likelihood of at least 28 for Fig 3E and at least 15 for Fig 3H.

Furthermore, motif activities at each time point and cell type are given as a mean and a standard deviation of the posterior distribution over motif activities. The significance of each motif is characterized by a Z-score, which is calculated as follows: for each sample, the ratio of the estimated motif activity and its standard deviation is squared, averaged across the samples, and finally the square root of the average is calculated. That is, a motif's Z-score corresponds to the average number of standard deviations the motif activity is away from zero across the samples. For Fig 3H, we further restricted the core regulatory network to motifs with a Z-score of at least 1.

When analyzing the NSCs, we noticed that motifs can be roughly distinguished as being active in either the expansion, neurogenic or gliogenic phases. To make this more visible, we have color-coded each motif of the graph (Fig 3E) in red for motifs active in expansion phase, in green for motif active in neurogenesis phase, and in blue for motifs active in gliogenesis phase. Some motifs are active in multiple phases, and we colored motifs using an HSV wheel to interpolate the motif activity profile between those three states. For example, Pou5f1 in Fig 3E is active in both neurogenesis and gliogenesis phases and is therefore colored in cyan, i.e., between green and blue on the HSV wheel.

When analyzing the Neurogenic NSCs, BPs, and NBNs, we noticed that motifs would either mostly vary along the first PC axis,

i.e., distinguishing samples of cell types, or along the second PC axis, i.e., distinguishing samples from different time points. We therefore color-coded the motifs of the graph (Fig 3H) from cyan to magenta, cyan representing genes that vary across time but that are constant across cell types and magenta representing motifs that vary across cell types but that are constant across time. We use a continuous gradient to report motifs varying across both time and cell types, as for instance, motifs Pou2f2_Pou3f1 on Fig 3H.

Each node on the networks of Fig 3E and H represents a TF motif. ISMARA also reports interactions between motifs and genes that are not transcription factors. To summarize the predicted target genes of each motif of the network, we ran a GO analysis (The Gene Ontology resource: enriching a Gold mine; Gene Ontology, 2021), and we use dotted edges to associate each motif with the main gene ontology biological process categories that are overrepresented among its predicted target genes.

Selection of highly variable genes

To select the most highly variable genes (HVGs), we have defined a score for each gene based on the contribution of each gene on each principal component and the variance that each component explains considering the first two components, as following:

$$Score(g_i) = \sqrt{(w_{g_i}(pc_1) * var(pc_1))^2 + (w_{g_i}(pc_2) * var(pc_2))^2}$$

Where $w_{g_i}(pc_k)$ refers to the weight (contribution) of gene i in pc_k , and $var(pc_k)$ denotes the percentage of variance that is covered by pc_k . Next, the first 2000 genes with the highest scores are selected as the highly variable genes, HVGs.

Clustering of single cells

First, single cells at each time point are clustered by applying 500 times k-means clustering to avoid the dependency of k-means clustering on the random initialization number (seed value). To implement k-means clustering, *clustering* package considering Euclidean distance as metric in R is used. Next, the assignment matrix is estimated based on the frequency of observing each two single cells in the same clustering at each iteration. Next, the hierarchical clustering is used to sort the assignment matrix using Euclidean distance as metric and *ward.D2* as method in *hclust* function in R.

Selection of differentially expressed gene in single cells

Kruskal–Wallis non-parametric test is used to select differentially expressed genes, and genes with adjusted P -value of less than $1e-3$ are considered as significantly differentially expressed genes.

Visualization

PCA is applied using *prcomp* function in R after centering the log transformed data. Heatmap are illustrated using *pheatmap* package in R on log transformed data.

Secondary KNN clustering and UMAP visualization

Raw counts were globally normalized; the expression of each gene was normalized against the total expression for each cell and multiplied by a scale factor of 10,000 using *LogNormalize* in *Seurat* v4.0. The most variable genes were then identified using *FindVariableFeatures*; this used local polynomial regression to fit log-variance to

log-mean value with gene variance being calculated from the standardized values. The top 2,000 most variable genes were scaled and used for downstream analyses. Significant principal components were carried forward, and a K-nearest neighbor graph was constructed on the Euclidean distance in PCA space. Edges were weighted based on shared overlap in neighborhoods using Jaccard index and graph-based Louvain clustering was conducted using FindClusters. For UMAP visualization, UMAP coordinates were calculated in PCA space for significant PCs and cells were assigned by previously derived cluster identities.

Trajectory inference analysis

Cell embeddings from established clustering were then used to infer lineage structures in low-dimensional space using Slingshot. First, the global lineage structure was identified constructing a cluster-based minimal spanning tree in a semi-supervised manner using *a priori* knowledge using GetLineages. This was then converted into smooth lineages to infer pseudotime by fitting simultaneous principal curves for each lineage with the getCurves function.

Linnarsson comparisons

Our C1 dataset was compared against (La Manno *et al*, 2021) single-cell developmental time course dataset, all cells from tissue labeled as “Forebrain” and “Dorsal Forebrain” by (La Manno *et al*, 2021) were clustered to capture cells spanning E9-E18 developmental timepoints. For radial glial exclusive comparisons, the cells were further subset to include only cells classed as “Radial Glia” in the original analysis. Canonical correlation analysis (CCA) was used to integrate this dataset with our C1 data. Log-normalized and scaled variable features that were repeatedly variable in both datasets were called and used to define cell anchors between the datasets, these anchors were then used to integrate the two datasets together. The integrated datasets were then clustered and visualized using UMAP as described above.

NeuroStemX data exploration web app

The NeuroStemX data exploration web app makes it possible to navigate data produced in the NeuroStemX project. The site supports viewing data by focusing on one of several parameters: gene list, biological sample, or measurement type (single cell vs. population). The website allows entry of a list of mouse genes (either as gene symbol or Ensembl ID) to focus on the data acquired for those genes. It alternatively allows viewing data on individual samples. When looking at a sample, a list of genes that have been determined to be outliers are shown. A gene is considered an outlier for a sample if the expression value of the gene either exceeds the 75th percentile +1.5*iqr or is less than the 25th percentile -1.5*iqr, where percentiles and inter-quartile range are computed based on the expression values for the given gene over all samples within the measurement type (single cell or population).

When viewing all data for a measurement type, data are displayed using hierarchical clustering. The InChLib widget displays the clustered data. Clustering is performed using the fastcluster package in python with distance (both row and column) calculated using the Euclidean metric and linkage (both row and column) performed using Ward’s method (Mullner, 2013; Skuta *et al*, 2014). The site supports performing gene ontology enrichment analysis either locally, using goatools, or with PANTHER. For local

enrichment analysis, we use the MGI_Gene_Model_Coord annotations based on the GRCh38 assembly (Mi *et al*, 2017).

Goatools: <https://github.com/tanghaibao/goatools>

Haibao Tang *et al* GOATOOLS: Tools for Gene Ontology. Zenodo. [10.5281/zenodo.31628](https://doi.org/10.5281/zenodo.31628).

MGI_Gene_Model_Coord.rpt

<http://www.informatics.jax.org/downloads/reports/index.html>

Data availability

The RNA sequencing datasets have been deposited in Gene Expression Omnibus (GEO) with accession number GEO:

GSE134688

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE134688>

and

GSE134738

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE134738>

For ISMARA, data can be found here:

<https://ismara.unibas.ch/NeuroStemX>

Expanded View for this article is available [online](#).

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Disclosure and competing interests statement

The authors declare that they have no conflict of interest.

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