Designing signaling environments to steer transcriptional diversity in neural progenitor cell populations

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Abstract

Stem cell populations within developing embryos are diverse, composed of many different sub-populations of cells with varying developmental potential. The structure of stem cell populations in cell culture remains poorly understood and presents a barrier to differentiating stem cells for therapeutic applications. In this paper we develop a framework for controlling the architecture of stem cell populations in cell culture using high-throughput single cell mRNA-seq and computational analysis. We find that the transcriptional diversity of neural stem cell populations collapses in cell culture. Cell populations are depleted of committed neuron progenitor cells and become dominated by a single pre-astrocytic cell population. By analyzing the response of neural stem cell populations to forty distinct signaling conditions, we demonstrate that signaling environments can restructure cell populations by modulating the relative abundance of pre-astrocyte and pre-neuron subpopulations according to a simple linear code. One specific combination of BMP4, EGF, and FGF2 ligands switches the default population balance such that 70\% of cells correspond to the committed neurons. Our work demonstrates that single-cell RNA-seq can be applied to modulate the diversity of in vitro stem cell populations providing a new strategy for population-level stem cell control.

Highlights

- Natural progenitor diversity in the brain collapses during in vitro culture to a single progenitor type
- Loss of progenitor diversity alters fate potential of cells during differentiation
- Large scale single-cell signaling screen identifies signals that reshape population structure towards neuronal cell types
- Signals regulate population structure according to a simple log-linear model
Introduction

Stem cells remain an exciting resource for regenerative medicine and tissue replacement. Work over many decades has shown that a wide range of stem cell populations, including embryonic stem cells and neural stem cells, can be extracted from an embryo and then expanded and differentiated in cell culture using applied signaling molecules [1]. Expansion of cells in culture provides a strategy for generating a wide range of functional cell types from a small initial pool of precursors that could be used to replace or repair damaged tissue in the brain, heart, and other organs that do not have intrinsic regenerative capacity.

However, despite the promise, the control of stem cell populations in cell culture, outside of the embryonic environment, has proved to be difficult and limited in many cases. Practically, it has been challenging to generate homogenous populations of functionally differentiated cells from stem cell precursors [2]. Although, for example, we can isolate and grow cells from the developing brain that express stem cell marker genes, the extracted cells often fail to differentiate into functional neurons both in vitro [3] or upon transplantation [4] limiting the use of these cells for treating neurodegenerative diseases. In other cases, differentiation occurs in cell culture but is highly variable, so that the resulting cell populations contain many different types of cells [2]. While the nature and source of variability is poorly understood, variability has led to practical bottlenecks in research and costly failures in clinical trials [5, 4, 6].

The practical challenges that complicate stem cell therapies raise fundamental questions about the generation and maintenance of stem cell populations in the embryo and the subsequent dynamics of cell populations within cell culture signaling environments. The combinatorial signaling environment within mammalian embryos generates a broad spectrum of different stem cell populations [7, 8, 9] that each generate specific differentiated cell-types. For instance, the mammalian brain develops through the coordinated proliferation and differentiation of multiple subpopulations of stem cells, which range in commitment and propensity for distinct differentiated neural cell fates [7]. Clonal tracking of neural stem cells has revealed a clonal bias in astrocytic versus neuronal fate[10], and suggests that propensity for distinct layers of the cortex is based on combinatorial patterns of transcription factor expression[11, 12]. Despite the importance of stem cell population diversity in the developing embryo, we do not know how natural progenitor diversity changes within cell culture signaling environments or how to shape population diversity to amplify stem cell types of practical interest.

In this paper, we apply single cell mRNA-seq to demonstrate a new approach to programming stem cell fate by controlling the balance of stem cell subpopulations primed for different lineages. By comparing stem cell populations from the developing mouse brain to neural stem cells growing in culture, we discover that the natural population diversity of neural stem cells collapses in culture. In culture, the critical pre-neuron progenitor cell subpopulations that generate neurons in the brain are lost. By exposing neural stem cells to forty different signaling environments, we show
that signaling environments can be designed using a simple log-linear model to alter the balance of pre-neuron and pre-astrocyte cell populations enabling amplification of the committed neurons that are lost under canonical conditions. Our work points to population structure and dynamics as playing an important and previously hidden role in impacting the outcome of stem cell differentiation. Broadly, we demonstrate an experimental and computational strategy for designing signaling environments that can restructure cell populations to overcome challenges implicit in stem cell therapies.

Results

Neural stem cell populations experience collapse in transcriptional diversity in cell culture

We wanted to understand how the diversity of neural stem cell populations changes when cells are extracted from the embryonic brain and placed in cell culture signaling environments. Therefore, we applied single cell mRNA-seq to create a map of the stem cell populations contained in the developing mouse brain, and then compared the underlying transcriptional diversity to neural stem cells grown in cell culture. Specifically, we profiled 23,438 cells at three time points of embryonic development (embryonic day 18 subventricular zone, postnatal day 4, and Adult Cortex) (Fig 1a) and compared these cells to populations of subventricular zone (SVZ) stem cells dissociated from the ventricular zone of the E18 mouse embryonic cortex and cultured with EGF and FGF2 proteins. To analyze the data, we used a probabilistic modeling platform, PopAlign, [13] to build, align, and compare Gaussian mixture models (GMMs) of the embryonic and cell culture populations (Fig 1b). Cell type markers for mature differentiated cell types (Fig 1a - inset) are used to type subpopulations in the adult cortex (Fig 1b), which provide a reference for assessing differentiation potential. For each sample, the resulting PopAlign models were low error both qualitatively (SI Figure 1) and quantitatively (SI Figure 2) and so provided a computational substrate for performing quantitative comparison between the embryonic and in vitro cells.

Our computational analysis revealed that embryonic day 18 (E18) brain contains a broad range of neuronal and astrocyte progenitor cells but that population diversity collapses in cell culture, so that cultured cell populations become dominated by a single pre-astrocyte cell state. Specifically, in the E18 SVZ, we identified three major categories of cells as the centroids of the GMMs identified by PopAlign (Fig 1b, 1c), committed neurons, pre-neuronal stem cells, and pre-astrocytic stem cells, as 71.4%, 13.8%, and 12.4% of the population respectively. Committed neurons (CNs), the dominant cell populations, expressed the canonical neuron markers Tubb3 and Map2 (Fig. 1d). The pre-neuronal progenitors (NPs) expressed stem cell markers Sox2, Pax6, and the neuronal transcription factor Neurog2, while the pre-astrocyte progenitors (APs) expressed glial transcription factor Sox9[14] in addition to Sox2 and Pax6 (Fig. 1d).
Each identified cell population also expressed blocks of lineage specific genes, supporting the idea that the identified cell populations could exhibit distinct differentiation potential. Both the pre-neuronal and pre-astrocyte progenitor cells expressed a base gene expression program of stem cell-associated genes including Glast(SLC1A3), Vim, Fabp7 (Blbp), and Dbi. These genes are known to be glial and reflect the radial-glial origin of neural stem cells in the brain [15]. However, in addition to this base stem cell program, pre-neuronal progenitors (NPs) specifically expressed a group of 13 neurogenic genes including neuron-specific TFs such as Tbrs (Eomes) Neurod6, and Neurog2 (Fig. 1e). Pre-astrocytic progenitors (APs) expressed a large block of 45 genes associated with mature astrocytes, including Aldh1l1, a mature astrocytic marker [16], and Glul (GS), an astrocyte-specific glutamine synthetase. While all progenitor cells expressed the core radial glial program, the astrocyte and neurogenic programs are mutually exclusive within the progenitor populations suggesting that they might repress each other and foreshadow downstream fate. Committed neurons (CN) expressed neuron specific gene expression programs, but not the stem cell program or the mitotic gene expression program (Fig. 1e).

When placed into the cell culture signaling environment with EGF and FGF2, the diversity of the progenitor population dramatically shifted undergoing a qualitative loss of the neural progenitor (NP) and committed neuron (CN) subpopulations within five days of culture (Fig. 1f). By day five of culture, the cell population had become dominated by the pre-astrocytic progenitor (AP) cells (96% of cells at day 5 but 18% at day 0) (Fig. 1g) and the CN cells have been lost (4% day 5 vs 66% day 0) (Fig 1h). Further, pre-neuronal progenitors (NPs) diminish from 16% to 0% at day 5 (Fig 1i). In this way, although the developing brain at E18 contains a diverse population of both pre-neuronal and pre-astrocytic progenitors, only the pre-astrocytic progenitors are expanded in EGF/FGF2 cell culture.
Decreased population diversity is associated with limited differentiation potential

The collapsed pre-astrocytic populations derived from the SVZ and grown in EGF and FGF2 for five days (Fig 2a) had limited differentiation potential. Upon differentiation in serum-containing media, the cells generated uniform cultures of astrocytes with no neurons generating no density in the neuronal branch of a PCA fate map (Fig 2b). The fate map was constructed using populations at postnatal day 4 (P4), which temporally corresponds to 5 days of in vitro differentiation. Differentiated populations quantitatively align most closely to the P4 astrocytic population, and produce matching gene expression signatures (Fig 2c). However, under the same conditions, cells extracted from SVZ tissue (Fig 2d) and immediately differentiated could generate mixed cultures of neurons and astrocytes based on the fate map (Fig 2e), and gene expression profiles (Fig. 2f). Although we cannot definitively conclude that the AP cells lack neuronal potential under all possible contexts, our results show that they do not generate neurons in conditions that support neuronal differentiation from a broader range of progenitor cell types from the brain. Thus these findings suggest that the collapse in progenitor cell diversity during cell culture limits the capacity of the population to generate neurons, and imply that increasing the progenitor diversity could expand the neurogenic potential of stem cells in vitro.

Combinatorial signaling environments modulate balance between pre-astrocytic progenitors and committed neurons

While EGF and FGF2 at fixed concentration represent just a single signaling condition, the embryonic brain contains combinatorial gradients of signaling molecules. We hypothesized that EGF and FGF2 likely select for the pre-astrocytic progenitor (AP) cells, thus restricting the differentiation potential of the cell population. Therefore, we asked whether we could stabilize a more diverse progenitor population by employing additional signaling factors. We performed a signaling screen where we exposed the SVZ stem cell populations across a range of different signaling conditions and analyzed the impact of signaling combinations on population structure using single-cell transcriptional profiling.

We selected signaling cues from the EGF, FGF, Wnt, PDGF, and BMP families whose receptors are all present in the developing brain (SI Figure 3) and are known to be associated with diverse functions such as proliferation (EGF, FGF2 [17], Wnt pathway activation via small molecule agonist CHIR, PDGF-AA [18]), neuronal differentiation, (WNT, [19] PDGF), astrocytic differentiation (BMP4 [20]), and the maintenance of quiescence [BMP4 [21]]. We also included two immune signaling cytokines, GM-CSF and IFN-gamma as outgroups that are not normally active during neural development, but whose receptors are present in our cells (SI Figure 3).

We combined all signals with EGF/FGF2 in varying doses ranging from 20ng/mL, a high concen-
Figure 2: Decreased progenitor diversity is associated with limited differentiation potential. (a) PCA rendering of cell populations extracted from the SVZ and cultured for 5 days in EGF/FGF and (b) after differentiating in serum containing media for 5 days. Above: cartoon maps showing relative positions of populations from corresponding stages of natural brain development (E18 and P4). (c) Gene expression heatmaps of population centroids from differentiated cells, compared to aligned populations from the in vivo brain at P4. Astrocytic, mitotic, and neuronal gene expression programs are indicated in the key. (d) Model renderings of cells dissociated directly from the SVZ and (e) their differentiated progeny after 5 days of in vitro differentiation. (i) Gene expression heat-map comparing the centroids of populations from (e) to their aligned in vivo populations.
tration that is the standard culture condition, down to 0.8 ng/mL, a low concentration that we found was necessary to maintain cell survival. Signals were added immediately to cells after SVZ dissociation. Cells were profiled after 5 days of exposure, and the Clicktags multiplexing platform [22] applied for sample labeling. We profiled an average of 500 cells per signaling condition leading to a data set with 18,700 cells, and applied PopAlign to dissect and analyze population diversity and to compare identified cell populations to those in the E18 brain.

We found that signaling conditions could shift the balance between pre-neuronal and pre-astrocyte cell populations. Across all signaling conditions, we identified three major subpopulations of progenitor cells that aligned to the three cell populations we previously identified (Figure 1) in the E18 mouse brain: committed neuroblasts (CN), pre-astrocytic progenitors (AP) and pre-neuronal progenitors (NP) (Fig 3a, SI Fig 4). Individual signals altered the relative abundances of the pre-neuronal and pre-astrocyte subpopulations, with the subset of conditions including BMP showing particularly strong impact. Specifically, BMP4 shifted the population structure towards more pre-neuronal cells (CN - pink), which could be observed qualitatively in terms of increasing density in the CN population (Fig 3b) and quantitatively in a phase-diagram (Fig 3c). Increasing the concentration of EGF/FGF2 had the opposite effect, shifting the population away from CN, towards pre-astrocytic progenitors (AP) (Fig 3b, 3d). CHIR boosted the proportions of pre-astrocytic progenitors (AP) (SI Fig 5a), while having no effect on pre-neuronal CNs (SI Figure 5b). Other signals such as FGF9, PDGF-AA, GM-CSF, and IFNG predominantly reduced the proportions of pre-astrocytic progenitors (AP) (SI Fig 5c) and boosted pre-neuronal CNs (SI Figure 5c).

Simple log-linear model enables design of signaling environments to achieve target population structure

The combinatorial impact of the signaling environment on the relative abundance of the three underlying progenitor cell populations could be captured by a simple log-linear model. To capture the relationship between signaling inputs and population size, we consider a simple model where the relative growth rates of specific cell populations respond independently to signaling inputs (Fig 4a). In this model, the log of the relative proportion of population (e.g. CN/NP) depends on a linear combination of signaling molecule concentrations (Fig 4b). The coefficients A, can be interpreted as the difference in relative growth rate between population i versus reference population k for each signal j. Critically, Aij can be encode both positive and negative values and can reveal antagonistic relationships between signals (Fig 4c). For instance, BMP4 and EGF compete to modulate the relative population sizes of both CN and AP cells (relative to the NP population) (Fig 4c). Model parameters reveal that each additional unit of BMP4 increases the relative proportion of CNs by 4.4x, while EGF/FGF reverses this trend, scaling down the relative CN proportion by 0.4x (Fig 4c).

The simple log-linear model was sufficient to recreate observed population abundances across all experiments. The accuracy of the model is both qualitative and quantitative; Model-generated pre-
Figure 3: Signals reshape global population diversity. (a) 2D PCA rendering of common model built using single-cell data sampled across 40-condition signaling screen. Identified subpopulations align to the main subtypes from the in vivo SVZ-D0 population: AP, CN, NP. (b) Renderings show population shifts in response to varying EGF/FGF and BMP4. The density of the CN population, highlighted in dashed white box, increases with increasing BMP4, and decreases with increasing EGF/FGF. (c) Heatmap of CN proportions with varying concentrations of BMP4 and EGF/FGF. Increasing BMP4 increases the CN proportion, while increasing EGF diminishes it. (d) Heatmap of AP proportions, showing AP proportions increasing with EGF/FGF, and decreasing with BMP4.
Figure 4: Population structure responds to signaling input according to a simple log-linear model. (a) Model equation relating relative population size \( n_i/n_k \) to a linear function of signaling inputs, \( x \). Coefficients \( A_i \) can be considered the relative growth rate of population \( i \) to population \( k \). The bias encodes the natural difference in growth rates in the absence of signal (\( x=0 \)). (b) The relative proportion of CN to AP populations (logged) responds linearly to signaling inputs. Grey dots: model output, red dots: observed proportions, blue line: error between observed data and prediction. (c) Coefficient matrix \( A \) linking relative population ratio (columns) to signals (rows). The exponent of the coefficient \( (e^{A_{i,j}}) \) is denoted in brackets and reports how much the ratio \( n_i/n_k \) goes up for each unit increase in signal. (d) Observed and predicted (colored) subpopulation proportions reported for each experimental sample (column). The standard deviation is calculated from bootstrapping samples from experimental data (\( n=100 \) cells each, iterations=20). Heatmap represents signaling concentration used within the column. (e-f) Model (colored lines) predicts combinations of signals that push CN and AP proportions towards target levels, such as that seen in original tissue (dotted line - SVZ-D0). Note: Signal concentrations across all panels in this figure is normalized so that 1 unit equals the maximum concentration (all 20 ng/mL except CHIR, whose maximum concentration is 3 uM).
dictions match patterns seen in the measured subpopulation abundances (Fig 4d). Additionally, the RMS error between the model-predicted proportions and the observed proportions is small, averaging only 5.58% of the observed value. The accuracy of the simple log-linear model suggests that additive impact of signaling molecules on cell-type specific proliferation rates might be sufficient to explain population size changes observed across the complex combinatorial signaling conditions considered in our screen (Methods).

Using the idea of a linear signaling code, we were able to design combinatorial signaling conditions to achieve desired population structures. For instance, although none of the signaling combination we tested could completely recreate the transcriptional diversity of the SVZ tissue, some conditions could approximate it by suppressing unwanted pre-astrocytic populations and boosting desired pre-neuronal populations. The original SVZ tissue was composed of \( \approx 70\% \) CNs (Fig 4e-dotted line). By reducing EGF/FGF2 alone, we boosted the population of CNs to 26% of the population (Fig 4e). Adding BMP4 boosts the population percentage to 56%. Finally, adding another CN-boosting and AP-reducing signal such as FGF9 or IFN-G can boost the CNs again up to \( \approx 80\% \), while also reducing the number of pre-astrocytic progenitors to 4-6% (Fig 4g). Globally, the conditions satisfying these constraints (low EGF/FGF2, high BMP4, and an additional AP-reducing signal) produced populations of cells that most closely matched the original SVZ tissue, based on a statistical metric which calculates the divergence of the two models (SI Fig 5 - Ranking).

**Tracking transcriptional changes shows that BMP4 also induces neuronal genes in the pre-astrocytic progenitor cell type**

Broadly, our results demonstrate that the structure of neural progenitor cell populations can be steered through the modulation of a relatively simple combinatorial signaling environment. In addition to altering the broad population structure, we also note that the signaling environment had an impact on the detailed transcriptional state of specific cell populations. In addition to altering population structure by boosting the CN population, both CHIR and BMP4 had a large impact on the transcriptional state of both CN (Fig 5a) and AP (Fig 5b) cells, resulting in the most differentially expressed genes.

Interestingly, BMP4 altered the gene expression program of the AP cells inducing a neuronal gene expression program in the pre-astrocyte progenitor cells. The gene expression program contained genes specific to neuron populations in different layers of the cortex [23] (Fig 5c), and are not induced by other signals. This result suggests that the pre-astrocytic progenitor cell type may be potentially capable of generating neurons, despite not readily doing so under differentiating conditions within our experiments. Overall, our results show that BMP4 induces greater neuronal potential in the entire population in two different ways: by restructuring the population to increase the proportion of committed neurons, as well as by inducing the neuronal target gene program in the background of the pre-astrocytic progenitor cell-type.
Figure 5: BMP and CHIR impact AP populations but not CN. Scatter plot of gene means for both (a) CN and (b) AP populations in response to BMP4, CHIR, and PDGF. Gene means for control population (20ng/mL EGF/FGF, 5 days) are plotted on the x-axis. Red points denote genes which have log2(fold change) ≥ 1, and statistically significant p-values (T-test, Benjamini-Hochberg corrected, alpha=0.01). Gene means are normalized by total transcript count per cell and rescaled. Number of statistically significant genes reported on upper left. (c) Gene expression plot of layer-specific genes [23] in AP cells across samples. Gene levels (y-axis) is individually normalized (rescaled by max), and 500 random cells are displayed for each sample. All populations maintain expression of progenitor markers Pax6, Sox2, and SLC1A3(Glast), but only BMP4 reinvigorates expression of layer-specific genes broadly.
Discussion

The embryo naturally regulates distinct pools of multipotent stem and progenitor cells to generate a diversity of mature cell types in the correct proportions. Changes in the population diversity of lineage-primed progenitors can determine the relative abundance of neurons versus astrocytes during brain development, or, in the immune system, the relative abundance of lymphoid versus myeloid cells in disease or aging [9]. The ability to manipulate the relative proportions of distinct stem cell and progenitor subpopulations provides an important route to controlling the differentiation potential of any stem cell population and also to modulating the state of tissues in the body during development and disease. In this work, we show that signaling ligands can directly steer the population structure of stem and progenitor cells isolated from the embryonic mouse brain. We identify conditions that amplify specific pre-neuronal and pre-astrocytic populations. In future work, it will be interesting to explore whether additional factors allow the expansion of other cell types including proliferative pre-neuronal stem cells.

Our approach has implications for understanding the behavior of stem cell populations broadly. Cells in the embryo are exposed to combinations of signaling molecules generated by morphogen gradients. An important question is to determine how stem cells can decode combinatorial signaling environments to select their fate and proliferation rate. Our findings point towards a potentially simple signaling code in which different signals linearly regulate the proliferation rate of distinct transcriptionally primed stem cell populations. Interestingly, few signals in our screen created cells with entirely new transcriptional signatures (only BMP4 and CHIR), suggesting that transcriptional patterns within these subtypes could be hardwired by epigenetics and difficult to alter. Future work using dynamical measurements that can accurately reflect subpopulation state will be important for determining the possibility of direct transition between subpopulations with different transcriptional lineage priming.

Our work emphasizes the role of local signaling environments in generating and maintaining population diversity both in the embryo and in artificial cell culture contexts. The embryo maintains population diversity naturally, by controlling signaling environments on cellular length scales to allow divergent cell-types to be generated in close spatial proximity [24]. The population diversity of the embryo is thus directly enabled by the self-organization of local combinatorial signaling environments. An important challenge for cell engineering is to control combinatorial signaling environments outside of the body with similar precision. In engineering applications, signals have traditionally been applied globally, to an entire cell culture, and therefore, might be fundamentally limited in their ability to control the generation of different cell types in close spatial proximity. Organoid based approaches solve the signaling control problem by generating signals locally through cell-cell interactions [25] It might also be possible to engineer synthetic signaling environments where signals are modulated on cellular length scales to generate cell populations where different cell-types can emerge through distinct, micron-scale, local signaling environments.

Broadly, our approach can be applied to study the combinatorial impact of a wide range of sig-
naling molecules on heterogeneous populations of stem and progenitor cells. The key technical innovations we apply are single-cell mRNA-seq, population level gene expression analysis and computational modeling. We anticipate that this strategy can be applied to uncover the underlying signaling code for engineering the population structure of heterogeneous multicellular constructs across both synthetic biology and tissue engineering applications.

References

[1] A. L. Perrier, V. Tabar, T. Barberi, M. E. Rubio, J. Bruses, N. Topf, N. L. Harrison, and L. Studer, “Derivation of midbrain dopamine neurons from human embryonic stem cells,” *Proceedings of the National Academy of Sciences*, vol. 101, no. 34, pp. 12543–12548, 2004.

[2] R. Tsunemoto, S. Lee, A. Szűcs, P. Chubukov, I. Sokolova, J. W. Blanchard, K. T. Eade, J. Bruggemann, C. Wu, A. Torkamani, *et al.*, “Diverse reprogramming codes for neuronal identity,” *Nature*, vol. 557, no. 7705, p. 375, 2018.

[3] V. Balasubramaniyan, A. De Haas, R. Bakels, A. Koper, H. Boddeke, and J. Copray, “Functionally deficient neuronal differentiation of mouse embryonic neural stem cells in vitro,” *Neuroscience research*, vol. 49, no. 2, pp. 261–265, 2004.

[4] S. E. Marsh, S. T. Yeung, M. Torres, L. Lau, J. L. Davis, E. S. Monuki, W. W. Poon, and M. Blurton-Jones, “Hucns-sc human nscs fail to differentiate, form ectopic clusters, and provide no cognitive benefits in a transgenic model of alzheimer’s disease,” *Stem cell reports*, vol. 8, no. 2, pp. 235–248, 2017.

[5] A. J. Anderson, K. M. Piltti, M. J. Hooshmand, R. A. Nishi, and B. J. Cummings, “Preclinical efficacy failure of human cns-derived stem cells for use in the pathway study of cervical spinal cord injury,” *Stem Cell Reports*, vol. 8, no. 2, pp. 249–263, 2017.

[6] S. Temple and L. Studer, “Lessons learned from pioneering neural stem cell studies,” *Stem cell reports*, vol. 8, no. 2, pp. 191–193, 2017.

[7] S. J. Franco and U. Müller, “Shaping our minds: stem and progenitor cell diversity in the mammalian neocortex,” *Neuron*, vol. 77, no. 1, pp. 19–34, 2013.

[8] R. A. Nimmo, G. E. May, and T. Enver, “Primed and ready: understanding lineage commitment through single cell analysis,” *Trends in cell biology*, vol. 25, no. 8, pp. 459–467, 2015.

[9] C. E. Muller-Sieburg, H. B. Sieburg, J. M. Bernitz, and G. Cattarossi, “Stem cell heterogeneity: implications for aging and regenerative medicine,” *Blood, The Journal of the American Society of Hematology*, vol. 119, no. 17, pp. 3900–3907, 2012.
[10] M. A. Bonaguidi, M. A. Wheeler, J. S. Shapiro, R. P. Stadel, G. J. Sun, G.-I. Ming, and H. Song, “In vivo clonal analysis reveals self-renewing and multipotent adult neural stem cell characteristics,” *Cell*, vol. 145, no. 7, pp. 1142–1155, 2011.

[11] S. Lodato and P. Arlotta, “Generating neuronal diversity in the mammalian cerebral cortex,” *Annual review of cell and developmental biology*, vol. 31, pp. 699–720, 2015.

[12] L. Telley, G. Agirman, J. Prados, S. Fievre, P. Oberst, I. Vitali, L. Nguyen, A. Dayer, and D. Jabaudon, “Single-cell transcriptional dynamics and origins of neuronal diversity in the developing mouse neocortex,” *bioRxiv*, p. 409458, 2018.

[13] S. Chen, J. H. Park, P. Rivaud, E. Charles, J. Haliburton, F. Pichiorri, and M. Thomson, “Dissecting heterogeneous cell-populations across signaling and disease conditions with popalign,” *bioRxiv*, p. 421354, 2018.

[14] S. Martini, K. Bernoth, H. Main, G. D. C. Ortega, U. Lendahl, U. Just, and R. Schwanbeck, “A critical role for sox9 in notch-induced astrogliogenesis and stem cell maintenance,” *Stem Cells*, vol. 31, no. 4, pp. 741–751, 2013.

[15] S. C. Noctor, A. C. Flint, T. A. Weissman, W. S. Wong, B. K. Clinton, and A. R. Kriegstein, “Dividing precursor cells of the embryonic cortical ventricular zone have morphological and molecular characteristics of radial glia,” *Journal of Neuroscience*, vol. 22, no. 8, pp. 3161–3173, 2002.

[16] J. D. Cahoy, B. Emery, A. Kaushal, L. C. Foo, J. L. Zamanian, K. S. Christopherson, Y. Xing, J. L. Lubischer, P. A. Krieg, S. A. Krupenko, *et al.*, “A transcriptome database for astrocytes, neurons, and oligodendrocytes: a new resource for understanding brain development and function,” *Journal of Neuroscience*, vol. 28, no. 1, pp. 264–278, 2008.

[17] H. G. Kuhn, J. Winkler, G. Kempermann, L. J. Thal, and F. H. Gage, “Epidermal growth factor and fibroblast growth factor-2 have different effects on neural progenitors in the adult rat brain,” *Journal of Neuroscience*, vol. 17, no. 15, pp. 5820–5829, 1997.

[18] K. Funa and M. Sasahara, “The roles of pdgf in development and during neurogenesis in the normal and diseased nervous system,” *Journal of Neuroimmune Pharmacology*, vol. 9, no. 2, pp. 168–181, 2014.

[19] M. Y. S. Kalani, S. H. Cheshier, B. J. Cord, S. R. Bababeygy, H. Vogel, I. L. Weissman, T. D. Palmer, and R. Nusse, “Wnt-mediated self-renewal of neural stem/progenitor cells,” *Proceedings of the National Academy of Sciences*, vol. 105, no. 44, pp. 16970–16975, 2008.

[20] M. A. Bonaguidi, T. McGuire, M. Hu, L. Kan, J. Samanta, and J. A. Kessler, “Lif and bmp signaling generate separate and discrete types of gfap-expressing cells,” *Development*, vol. 132, no. 24, pp. 5503–5514, 2005.
[21] D. A. Lim, A. D. Tramontin, J. M. Trevejo, D. G. Herrera, J. M. García-Verdugo, and A. Álvarez-Buylla, “Noggin antagonizes bmp signaling to create a niche for adult neurogenesis,” Neuron, vol. 28, no. 3, pp. 713–726, 2000.

[22] J. Gehring, J. H. Park, S. Chen, M. Thomson, and L. Pachter, “Highly multiplexed single-cell rna-seq for defining cell population and transcriptional spaces,” bioRxiv, p. 315333, 2018.

[23] B. J. Molyneaux, P. Arlotta, J. R. Menezes, and J. D. Macklis, “Neuronal subtype specification in the cerebral cortex.” Nature reviews neuroscience, vol. 8, no. 6, p. 427, 2007.

[24] C.-H. L. Eng, M. Lawson, Q. Zhu, R. Dries, N. Koulena, Y. Takei, J. Yun, C. Cronin, C. Karp, G.-C. Yuan, et al., “Transcriptome-scale super-resolved imaging in tissues by rna seqfish+,” Nature, vol. 568, no. 7751, p. 235, 2019.

[25] A. Warmflash, B. Sorre, F. Etoc, E. D. Siggia, and A. H. Brivanlou, “A method to recapitulate early embryonic spatial patterning in human embryonic stem cells,” Nature methods, vol. 11, no. 8, p. 847, 2014.

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