## Article

# **Cell Stem Cell**

## Stage-Specific Transcription Factors Drive Astrogliogenesis by Remodeling Gene Regulatory Landscapes

## Graphical Abstract



## **Highlights**

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- Distinct transcriptional programs define consecutive stages of astrogliogenesis
- Stage- and lineage-specific enhancers promote the distinct gene expression programs
- ATF3 and NFIA drive astrocyte differentiation, while RUNX2 promotes astrocyte maturation
- $\bullet$  These TFs switch the enhancers from a primed to an active chromatin state

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[benedikt.berninger@kcl.ac.uk](mailto:benedikt.berninger@kcl.ac.�uk) (B.B.)

Authors

In Brief

Sophie Péron, ...,

**Correspondence** 

Laura López-Mascaraque,

distinct stages. The authors found that the transcription factors NFIA, ATF3, and RUNX2 play pivotal roles in establishing these stages.



## Stage-Specific Transcription Factors Drive Astrogliogenesis by Remodeling Gene Regulatory Landscapes

Neha Tiwari,<sup>[1,](#page-1-0)[9](#page-1-1)</sup> Abhijeet Pataskar,<sup>[2,](#page-1-2)9</sup> Sophie Péron,<sup>1,[10](#page-1-3)</sup> Sudhir Thakurela,<sup>2,[3,](#page-1-4)[4,](#page-1-5)10</sup> Sanjeeb Kumar Sahu,<sup>[2](#page-1-2)</sup> María Figueres-Oñate, $^5$  $^5$  Nicolás Marichal, $^1$  $^1$  Laura López-Mascaraque, $^5$  Vijay K. Tiwari, $^{2,6,11,12,\ast}$  $^{2,6,11,12,\ast}$  $^{2,6,11,12,\ast}$  $^{2,6,11,12,\ast}$  $^{2,6,11,12,\ast}$  $^{2,6,11,12,\ast}$ and Benedikt Berninger<sup>[1,](#page-1-0)[6,](#page-1-7)[7,](#page-1-11)[8,](#page-1-12)[11,](#page-1-8)[12,](#page-1-9)[13,](#page-1-13)\*</sup>

<span id="page-1-2"></span><span id="page-1-0"></span>1Institute of Physiological Chemistry, University Medical Center Johannes Gutenberg University Mainz, 55128 Mainz, Germany 2Institute of Molecular Biology (IMB), 55128 Mainz, Germany

<span id="page-1-4"></span>3Broad Institute of MIT and Harvard, Cambridge, MA, USA

<span id="page-1-5"></span>4Department of Stem Cell and Regenerative Biology, Harvard University, Cambridge, MA, USA

<span id="page-1-6"></span>5Instituto Cajal, CSIC, Madrid, Spain

<span id="page-1-7"></span>6Focus Program Translational Neuroscience, Johannes Gutenberg University Mainz, 55131 Mainz, Germany

<span id="page-1-12"></span><span id="page-1-11"></span>7Centre for Developmental Neurobiology, Institute of Psychiatry, Psychology & Neuroscience, King's College London, London SE1 1UL, UK 8MRC Centre for Neurodevelopmental Disorders, Institute of Psychiatry, Psychology & Neuroscience, King's College London,

London SE1 1UL, UK

<span id="page-1-1"></span>9These authors contributed equally

<span id="page-1-3"></span>10These authors contributed equally

<span id="page-1-8"></span>11These authors contributed equally

<span id="page-1-9"></span>12Senior author

<span id="page-1-13"></span>13Lead Contact

<span id="page-1-10"></span>\*Correspondence: [vijay.tiwari@yahoo.com](mailto:vijay.tiwari@yahoo.com) (V.K.T.), [benedikt.berninger@kcl.ac.uk](mailto:benedikt.berninger@kcl.ac.uk) (B.B.) <https://doi.org/10.1016/j.stem.2018.09.008>

#### **SUMMARY**

A broad molecular framework of how neural stem cells are specified toward astrocyte fate during brain development has proven elusive. Here we perform comprehensive and integrated transcriptomic and epigenomic analyses to delineate gene regulatory programs that drive the developmental trajectory from mouse embryonic stem cells to astrocytes. We reportmolecularly distinct phases of astrogliogenesis that exhibit stage- and lineage-specific transcriptomic and epigenetic signatures with unique primed and active chromatin regions, thereby revealing regulatory elements and transcriptional programs underlying astrocyte generation and maturation. By searching for transcription factors that function at these elements, we identified NFIA and ATF3 as drivers of astrocyte differentiation from neural precursor cells while RUNX2 promotes astrocyte maturation. These transcription factors facilitate stage-specific gene expression programs by switching the chromatin state of their target regulatory elements from primed to active. Altogether, these findings provide integrated insights into the genetic and epigenetic mechanisms steering the trajectory of astrogliogenesis.

#### **INTRODUCTION**

A comprehensive understanding of the molecular mechanisms underlying cellular diversity and cell-fate specification during CNS development remains elusive. In the developing CNS, neural precursor cells (NPCs) are known to give rise first to neurons and then to glia such as astrocytes and oligodendrocytes [\(Krieg](#page-14-0)[stein and Alvarez-Buylla, 2009\)](#page-14-0). The molecular circuitry underlying the differentiation of NPCs into astrocytes is only beginning to be untangled. Astrocytes are involved in various functions that are important for the establishment, maintenance, and plasticity of the brain [\(Tiwari and Berninger, 2017\)](#page-15-0). Because of these important roles, malfunctions in astrocytes have been implicated in many neurological diseases ([Molofsky et al., 2012](#page-15-1)).

Despite these advances, the gene regulatory mechanisms that control the differentiation and maturation of astrocytes from NPCs, hereafter referred to as astrogliogenesis, remain to be uncovered. A plethora of signaling pathways are known to play a critical role in the differentiation of NPCs into astrocytes ([Kanski](#page-14-1) [et al., 2014\)](#page-14-1). For instance, BMP2 promotes STAT3-mediated astrogliogenesis by forming a STAT3-SMAD1-p300 co-activating complex that initiates expression of astrocyte-specific genes by binding to their promoters [\(Fukuda et al., 2007; Nakashima](#page-14-2) [et al., 1999\)](#page-14-2). In addition to STAT3 ([Bonni et al., 1997; Moon](#page-14-3) [et al., 2002\)](#page-14-3), several other transcription factors (TFs), including SOX9, NFIA ([Kang et al., 2012](#page-14-4)), ETV5/ERM ([Li et al., 2012\)](#page-15-2), and ZBTB20 [\(Nagao et al., 2016](#page-15-3)), have been implicated as important in astrogliogenesis. Emerging evidence implies that these TFs interplay with epigenetic mechanisms to regulate astrocytespecific genes. At the onset of astrogliogenesis, STAT3 binds to the Gfap promoter and promotes DNA demethylation, resulting in *gfap* expression ([Fan et al., 2005\)](#page-14-5). Similarly, NOTCH signaling induces the expression of Nfia in neural progenitor cells, which then targets the promoters of astrocyte-specific genes and causes DNA demethylation at these promoters by displacing DNMT1 [\(Namihira et al., 2009\)](#page-15-4). Furthermore, at the onset of gliogenesis, the Polycomb group (PcG) proteins repress



pro-neuronal genes, such as *Neurog1*, thereby restricting the neurogenic competence of NPCs ([Hirabayashi et al., 2009](#page-14-6)). Finally, proteins of the high-mobility group nucleosome-binding family, HMGN1, HMGN2, and HMGN3, also play a critical role in promoting differentiation of astrocytes [\(Nagao et al., 2014](#page-15-5)). Despite these exciting advances, a comprehensive understanding of the transcriptional and epigenetic mechanisms acting along the differentiation trajectory from NPCs to astrocytes remains elusive. In addition, although the enhancers have a known role in cell-fate specification [\(Buecker et al., 2014](#page-14-7)), no studies have performed an in-depth investigation to identify the regulatory elements that are involved in defining the distinct phases of astrogliogenesis and the epigenetic mechanisms and TFs that operate at these sites during this process.

Here we used a model of astrogliogenesis from mouse embryonic stem cells in combination with next-generation sequencing and computational approaches to pinpoint distinct stages along the differentiation trajectory by identifying stage-specific transcriptional programs and epigenetic states. This quest allowed us to discern stage- and lineage-specific regulatory elements that represent putative targets of the TFs Nfia, Atf3, and Runx2.

#### RESULTS

#### Distinct Gene Expression and Epigenetic Landscape Define Stages of Astrogliogenesis

To investigate the gene regulatory mechanism underlying astrogliogenesis, we first adapted an experimental model that uses mouse embryonic stem cells (ESCs) to generate highly enriched astrogliogenic neural precursor cells (aNPCs) that subsequently differentiate into astrocytes (early astrocyte [eA] and mature or late astrocyte [lA]) [\(Figure 1A](#page-3-0); Figure S1A) ([Conti et al., 2005;](#page-14-8) [Pollard et al., 2006](#page-14-8)). qRT-PCR and immunoblot analyses showed that these astrocytes express established markers (e.g., Gfap, S100b, Nfia, Glast, Aqp4, and Cx43) (Figures S1B and S1C; [Fig](#page-3-0)[ure 1A](#page-3-0)). Using markers for neuronal cells (TUBB3 and DCX) and progenitor cells (NES, OLIG2, and ASCL1), we found that these cultures exhibit a minimal presence of these cells (Figure S1D). However, we observed differences in the expression levels of AQP4, GLAST, and CX43, indicative of astroglial heterogeneity (Figure S1D). The differentiation of aNPCs into astrocytes was accompanied by a cessation of cell proliferation (Figure S1E). Furthermore, we found that similar to astrocytes *in vivo*, mature astrocytes generated calcium signals upon mechanical stimulation that propagated to neighboring astrocytes, indicative of their functional maturation (Figures S1F and S1G). In line with previous studies, aNPCs were able to generate neurons when leukemia inhibitory factor (LIF) and fetal bovine serum (FBS) were withdrawn from the differentiation medium (Figure S1H) ([Conti et al., 2005; Pollard et al., 2006\)](#page-14-8).

To measure the global gene expression changes that accompany astrogliogenesis, we performed high-coverage transcriptome profiling (RNA sequencing [RNA-seq]) of ESCs, aNPCs, and astrocytes at various stages of differentiation, spanning the early and later phases (1, 5, and 21 days, hereafter referred as eA, lA\_1, and lA\_2, respectively). A principal-component analysis (PCA) of the transcriptome datasets uncovered a progressive separation of the transcription profiles during astro-

gliogenesis, suggesting that the acquisition of distinct gene expression programs gave rise to stage-specific cellular identity [\(Figure 1B](#page-3-0)). The differential expression analysis of the contiguous stages of astrogliogenesis revealed major differences among the early stages, while the stages lA\_1 and lA\_2 were transcriptionally similar, indicating that terminal astroglial gene expression was largely established at lA\_1 [\(Figure 1](#page-3-0)B). Thus, in the subsequent analysis, we focused on the mature or late astrocytes, hereafter referred to as lA. We then identified genes that were differentially expressed using pairwise comparisons of subsequent stages during astrogliogenesis (Figure S1I; Table S1) and grouped them into five clusters ([Figure 1C](#page-3-0); Table S2), in temporal order, according to their expression kinetics during astroglial differentiation. Moreover, genes that were highly expressed during astrogliogenesis from mouse ESCs were highly expressed in astrocytes *in vivo* [\(Cahoy et al., 2008; Zhang](#page-14-9) [et al., 2016](#page-14-9)) and at higher levels than those found in neurons [\(Fig](#page-3-0)[ure 1C](#page-3-0)). This analysis showed progressive changes in the expression profiles and reflected the developmental trajectory as supported by gene ontology (GO) term analysis of these gene clusters, showing enrichment of stage-relevant biological pathways (Table S2). Cluster 1 (ESC genes) and cluster 2 (ESC/aNPC genes) were enriched with genes involved in cellcycle and metabolic pathways, reflecting the proliferative activity of these cell types and the associated high metabolic rate (Table S2; [Figure 1](#page-3-0)C; Figures S1J and S1K). Cluster 3 (aNPC/eA) and cluster 4 (eA) showed enrichment with genes involved in nervous system development, reflecting the ongoing process of astrogliogenesis (Table S2; [Figure 1C](#page-3-0); Figures S1L and S1M). Cluster 5 (lA\_1 and lA\_2) was enriched with GO terms such as signaling and cytokine response, which are features that have been linked to mature astrocytes [\(Michelucci et al., 2016](#page-15-6)) (Table S2; [Figure 1](#page-3-0)C; Figure S1N). Consistent with the progressive differentiation and maturation of astrocytes, we found that Gfap was induced at the onset of astroglial differentiation (eA), while the maturation marker Aqp4 only appeared during later stages [\(Figure 1](#page-3-0)D). In contrast, the mature neuronal marker Rbfox3 was never expressed during astrogliogenesis ([Figure 1](#page-3-0)D). In addition, a comparison of our datasets with the astrocyte gene expression data retrieved from Network Glia [\(http://www.](http://www.networkglia.eu) [networkglia.eu](http://www.networkglia.eu)), which includes published datasets [\(Cahoy](#page-14-9) [et al., 2008](#page-14-9)), revealed similar expression kinetics in signature genes of *in vivo*- and *in vitro*-differentiated astrocytes (Figure S1O). Altogether, these features confirm that our *in vitro* astrogliogenesis model is highly suitable for studying gene regulatory mechanisms underlying astrocyte differentiation. Furthermore, our observations clearly highlight that the consecutive stages of astrogliogenesis can be defined by distinct gene expression profiles.

Acetylation of lysine 27 at histone H3 (H3K27ac) marks active proximal and distal regulatory elements [\(Creyghton et al., 2010;](#page-14-10) [Shlyueva et al., 2014\)](#page-14-10). To identify these regulatory regions, we generated H3K27ac chromatin immunoprecipitation sequencing (ChIP-seq) profiles during the stages of astrogliogenesis (aNPC, eA, and lA) (Figure S1P). Single-gene visualizations showed an expected enrichment in H3K27ac levels at astroglial genes (e.g., Gfap and Aqp4), while neuronal genes (e.g., Rbfox3) failed to show enrichment of this mark during astrogliogenesis [\(Fig](#page-3-0)[ure 1E](#page-3-0)). Although the total number of H3K27ac-enriched sites

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#### Figure 1. Distinct Gene Expression Programs and Distal Epigenetic Footprints Define Distinct Stages of Astrogliogenesis

(A) Schematic representation of astroglial differentiation from ESCs. Immunofluorescence staining of DAPI and GFAP in the ESC-derived astrocytes during phases of astrogliogenesis, including aNPC, eA,  $IA_1$ , and  $IA_2$ . Scale bar is 100  $\mu$ m.

(B) Principal-component analysis plot depicting the distribution of the transcriptomes of ESC (pink), aNPC (light blue), eA (dark blue), and two stages of lA (shades of green) within the scope of the first two components after RNA-seq in these stages. The x and y axes show the percentage of variance explained by the first 2 principal components.

(C) Left side: clustered heatmap depicting the expression of differentially expressed genes during any stage of the *in vitro* astroglial differentiation, along with the names of known key genes in each of these clusters. Expression in the heatmap is scaled from blue (lowest) to red (highest). Right side: locally weighted scatterplot smoothing (lowess) regression line for these clusters in the same order as the published astrocyte transcriptomes by [Cahoy et al. \(2008\)](#page-14-9) and [Zhang](#page-15-7) [et al. \(2016\)](#page-15-7).

(D and E) University of California Santa Cruz (UCSC) browser track example of expression (D) and H3K27ac enrichment (E) at the astroglial genes, i.e., Gfap and Aqp4, and the neuronal gene Rbfox3 during the stages of the *in vitro* astroglial differentiation (ESC, aNPC, eA, and lA).

(F) Stacked-bar plot depicting the number of common and unique (across astroglial differentiation) H3K27ac peaks under each condition of astroglial differentiation sorted by their genomic location, i.e., intergenic, promoter, exon, and intron.

(G) Venn diagram representing the overlap of the H3K27ac peaks within a distance of 50 kb from the nearest genes called in aNPC, eA, and lA.

(H) Boxplot representing the expression of the genes, as determined as log2-normalized read counts, that are associated with the aNPC, eA, and lA unique H3K27ac peaks depicted in (G).

remained in the same range for all stages, they exhibited a substantial number of uniquely acquired H3K27ac-enriched regions. These uniquely enriched sites were particularly conspicuous at the aNPC and lA stages, whereas astrocytes at the eA stage showed far fewer unique H3K27ac-enriched sites, possibly reflecting an intermediate epigenetic profile at this transitory stage (Figures S1Q and S1R).

The genomic distribution of H3K27ac-enriched sites revealed its very similar occurrence at promoters and non-promoter (or 'distal', including intergenic, exons, and introns) regions in aNPC, eA, and IA stages ([Figure 1](#page-3-0)F). However, the stage-specific H3K27ac sites were predominantly located in non-promoter regions, except for eA [\(Figure 1F](#page-3-0)). We also validated that the H3K27ac enrichment at promoter and distal regions of several known astrocyte genes increased during astroglial differentiation using ChIP-qPCR experiments (Figures S1S and S1T). To identify genes putatively regulated by the distal H3K27ac sites, we assigned genes according to proximity  $(-50$  kb) to the stagespecific distal H3K27ac sites. Although many of these sites shared the same nearest gene, many genes were uniquely asso-ciated with stage-specific H3K27ac-enriched sites ([Figure 1G](#page-3-0)). These non-overlapping genes were expressed at significantly higher levels at their corresponding stages, except for the eA stage [\(Figure 1H](#page-3-0)).

#### Epigenetic Priming Precedes Stage-Specific Acquisition of Active Chromatin at Regulatory Regions during Astrogliogenesis

Mono-methylation of lysine 4 at histone H3 (H3K4me1) marks primed or active enhancers in the absence or presence of H3K27ac, respectively [\(Creyghton et al., 2010](#page-14-10)). Given the critical role of the crosstalk among various chromatin marks at regulatory elements in defining cell-type-specific gene expression programs, we investigated whether H3K27ac enrichment is related to the occurrence of H3K4me1 at these sites during astrogliogenesis. Toward this end, we generated H3K4me1 ChIP-seq profiles during the stages of astrogliogenesis (aNPC, eA, and lA) (Figure S2A). Further analysis revealed a substantial number of stage-specific H3K4me1-enriched regions, which were nearly three-fold higher in lA (Figure S2B). The genes with H3K4me1-enriched promoters included Gfap and Aqp4, which already exhibited high levels of H3K4me1 in aNPC before acqui-sition of H3K27ac ([Figure 1E](#page-3-0)) and transcription ([Figures 2A](#page-5-0) and 2B) at the eA and lA stages, respectively, indicative of priming. High levels of H3K4me1 were also detected at the promoters of neurogenic genes, such as Rbfox3 in aNPC ([Figure 2C](#page-5-0)). These H3K4me1-enriched regions were not only at the promoters of astrocyte-specific genes but also at the distal regions, as validated by ChIP-qPCR assays (Figures S2C and S2D). Astrocyte-specific genes were either not primed or minimally primed in ESCs, and they acquired priming only upon commitment to the astrocyte lineage, i.e., in astrocyte progenitor cells (aNPCs) (Figures S2C and S2D).

We next investigated the H3K4me1 enrichment patterns at the stage-specific H3K27ac sites during astrogliogenesis. A profile plot depiction of these H3K27ac sites revealed their stage-specific high enrichment during astrogliogenesis ([Figures 2D](#page-5-0)–2F). The aNPC-specific H3K27ac sites showed the highest enrichment of H3K4me1 at the aNPC state, and upon astrocyte differ-

entiation, H3K27ac enrichment was lost in these regions and H3K4me1 levels were reduced [\(Figures 2D](#page-5-0) and 2G). eA-specific H3K27ac sites were already marked by H3K4me1 in aNPC, and H3K4me1 enrichment patterns were maintained through the eA stage while gaining H3K27ac ([Figures 2](#page-5-0)E and 2H). Furthermore, these regions lost H3K27ac at the lA stage, and the levels of H3K4me1 were reduced. Similarly, lA-specific H3K27ac sites gained H3K4me1 at the eA stage and increased this mark at the lA stage ([Figures 2F](#page-5-0) and 2I). These data suggest that epigenetic priming of astroglial genes occurs before their acquisition of an active chromatin state and transcriptional activation.

We then attempted to unravel the distinct set of genes that show different dynamics of H3K27ac at the H3K4me1 primed sites during astrogliogenesis versus neurogenesis and extended these analyses to reveal the gene regulatory landscapes that contribute to the divergence of these two lineages. Therefore, we adapted an established system in which ESCs differentiate into neurogenic neural precursor cells (nNPCs) and subsequently into terminally differentiated neurons (TNs) via different stages (early neuron [eN] and late neuron [IN]) ([Bibel et al.,](#page-14-11) [2004\)](#page-14-11) (Figure S2E). Previous studies have shown distinct epigenomic and transcriptomic changes in this model of neurogenesis that closely mimicked *in vivo* [\(Mohn et al., 2008; Stadler et al.,](#page-15-8) [2011; Thakurela et al., 2015\)](#page-15-8). A clustering analysis of the regions primed in aNPC produced seven sets of patterns [\(Figure 2](#page-5-0)J; Figure S2F; Table S3). Clusters 1, 3, and 5 showed minimal dynamics of H3K4me1 and H3K27ac during both astrogliogenesis and neurogenesis. Cluster 2 represented regions primed in aNPC that gain H3K27ac only during neurogenesis. Further analysis of cluster 2 showed that this cluster contains genes that are specific to neuronal differentiation, such as NeuroD1, Neurog1, and Neurog2 [\(Figure 2K](#page-5-0)), supporting multipotency at the aNPC stage. Cluster 4 contained genes that are marked by H3K4me1 without subsequent enrichment of H3K27ac in either lineage. Finally, clusters 6 and 7 contained sites that are primed in aNPC and gain H3K27ac specifically during astrogliogenesis. These clusters comprise astrocyte-specific genes, such as Gfap, Aqp4, Aldh1l1, Aldoc, Cd44, and S100b ([Figure 2L](#page-5-0)). The switch from a primed to an active state is accompanied by a transcriptional induction of genes in the nearest to these sites (Figure S2G). GO analysis of cluster 2 showed enrichment of genes related to neuronal differentiation (Figure S2H), whereas cluster 6- and cluster 7-specific genes were related to cell signaling and cell morphogenesis (Figure S2I). Furthermore, cluster 6and cluster 7-contained sites were primed in nNPC, eN, and lN, whereas they never gained H3K27ac during neuronal differentiation ([Figures 2](#page-5-0)M and 2N). Altogether, these data reveal epigenetic priming in regulatory elements, which precedes the stage-specific acquisition of active chromatin and transcriptional activation during astrogliogenesis.

#### Distinct TFs Shape the Epigenetic Landscape to Demarcate Astrocyte versus Neuronal Fate

Prompted by our finding that these regulatory elements are dynamically used during astrogliogenesis, we were interested in identifying TFs that function at these elements and how they differ from the factors that orchestrate neurogenesis. Using datasets from the neuronal differentiation model described earlier [\(Bibel et al., 2004; Thakurela et al., 2015](#page-14-11)), our analysis revealed

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Figure 2. Epigenetic Priming of Lineage- and Stage-Specific Regulatory Elements before Their Activation during Astrogliogenesis (A–C) UCSC browser track showing H3K4me1 enrichment at the regulatory regions of astroglial genes (A: Gfap, B: Aqp4) and neuronal gene (C: Rbfox3). (D–I) Profile plots depicting the enrichment of H3K27ac (D–F) and H3K4me1 (G–I) at the aNPC (D and G), eA (E and H), and lA (F and I) unique H3K27ac peaks. (J) Unsupervised clustering-based heatmap depicting the ChIP-seq enrichment of H3K4me1 during all stages of astrogliogenesis and H3K27ac during all stages of astrogliogenesis and neurogenesis at the H3K4me1 peaks in aNPC.

(K and L) Line plot depicting the enrichment of H3K4me1 during all stages of astrogliogenesis and H3K27ac during all stages of astrogliogenesis and neurogenesis in the regulatory regions of neuronal marker genes (K) and astroglial marker genes (L) that belong to clusters 2 and 6 or 7 from (J), respectively. (M and N) H3K4me1 enrichment during neuronal differentiation at cluster 6 (M) and cluster 7 (N) sites.

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Figure 3. Distinct TFs Set Up the Active Distal Regulatory Landscape Underlying Astroglial Differentiation

(A) Workflow that includes the outline of the computational pipeline for how and which TFs were selected, as well as the follow-up experimental part, including knockdown of these TFs and accompanied phenotypic and/or marker analysis.

(C–D) Venn diagram depicting the overlap of motifs that are enriched at unique H3K27ac peaks, including promoters, in aNPC and nNPC (A), eA and eN (B), and lA and lN (C). Shown inset are two representative motifs from each set.

(E–I) Profile plots representing H3K27ac enrichment at the genomic location of the motifs of the respective TFs within the scope of the unique H3K27ac peaks in eA (Nfia and Atf3 in E and F) and lA (Stat3, Prdm9, and Runx2 in G–I) upon depletion of TFs. The enrichment is depicted as a log two-fold change above the genomic input.

(J–L) UCSC browser track example from (E), (F), and (I).

strikingly different gene expression programs during astrocytic differentiation compared to neuronal differentiation (Figure S3A) and a large number of genes differentially expressed for aNPC versus nNPC, eA versus eN, and lA versus lN (Figures S3B– S3D). GO analysis revealed that highly induced genes in aNPC were enriched for nervous system development, including gliogenesis (aNPC versus nNPC) (Figure S3E). Furthermore, the genes highly upregulated in eAs compared to eNs were enriched for proliferation-, migration-, and cell adhesion-related genes (Figure S3F). Finally, the genes that were more higher expressed in lAs versus lNs were enriched with functions such as signaling and cytokines (Figure S3G). In addition to the TFs and epigenetic regulators (ERs) known to be relevant for astrogliogenesis, such as Nfia [\(Deneen et al., 2006; Piper et al., 2010](#page-14-12)), this list contained several TFs and ERs whose function in astrogliogenesis is unknown (Figures S3H–S3M; Table S1).

To reveal the differential chromatin landscape during astroglial versus neuronal differentiation, we compared the uniquely enriched H3K27ac sites during various stages of astrogliogenesis to those of the corresponding stages of neurogenesis [\(Figure](#page-6-0) [3](#page-6-0)A; Figure S3N). This comparison revealed that stage-specific H3K27ac-enriched regions were largely unique to this lineage

and did not occur during neurogenesis (Figure S3N). Similar findings were observed in comparisons with tissues from other lineages (Figures S3O and S3P). We then investigated whether H3K27ac-enriched genomic regions during astrogliogenesis recruit TFs that are similar to or different from those that are recruited during neurogenesis. Thus, we performed a motif enrichment analysis of the unique H3K27ac peaks during each stage of astrogliogenesis, compared these peaks with those that occurred during the corresponding stages of neurogenesis, and extended similar analysis to common peaks ([Figure 3A](#page-6-0)). Although several enriched TF motifs were common between the astrogliogenesis and the neurogenesis stages, a remarkable number of motifs were unique to the astrocyte or the neuronal differentiation stages [\(Figures 3B](#page-6-0)–3D; Figure S3Q; Table S4). These motifs included binding sites for many TFs that have previously been shown to be critical for astrogliogenesis (e.g., Nfia and Runx2) and neurogenesis (e.g., Lhx2 and Brn1) (Table S4) [\(Deneen et al., 2006; Dominguez et al., 2013; Okawa](#page-14-12) [et al., 2016; Subramanian et al., 2011](#page-14-12)). This analysis revealed several additional TFs that have not been previously implicated in astrogliogenesis and neurogenesis, thus warranting further investigation.

TFs that function at distal regulatory elements often contribute to the activation of these sites [\(Shlyueva et al., 2014; Thakurela](#page-15-9) [et al., 2015](#page-15-9)). Therefore, we investigated whether the TFs that were predicted to be enriched at the active regulatory elements during astrogliogenesis contribute to the active chromatin state in these regions [\(Figure 3A](#page-6-0); Figure S3Q). Thus, we performed small interfering RNA (siRNA)-mediated depletion of selected stage-specific TFs during the eA and lA stages and shortlisted the TFs whose knockdown impaired the acquisition of astrocyte markers [\(Figure 3A](#page-6-0); Figures S3R–S3T). This approach led to the identification of Nfia and Atf3 at the eA stage and Stat3, Runx2, and Prdm9 at the lA stage of astrogliogenesis ([Figure 3A](#page-6-0); Figures S3R–S3U). While Nfia was expressed during each stage of astroglial differentiation, Atf3 and Stat3 are specifically upregulated in eA and lA, respectively (Figures S3V and S3W). In contrast, Runx2 (Figures S3V and S3X) and Prdm9 showed highest expression in lA and aNPC, respectively (Figure S3V). However, while Nfia, Atf3, and Stat3 transcription was low during neurogenesis, Prdm9 and Runx2 showed higher expression (Figure S3V). We then investigated whether knockdown-induced defects in the astrocytic gene expression were due to an aberrant epigenetic state at the target distal regulatory elements. Therefore, we assessed the levels of H3K27ac at putative target sites using ChIP-seq analysis after their depletion during astrogliogenesis. The loss of Nfia and Atf3 in eA and the loss of Runx2 in lA led to a reduction in the H3K27ac levels at putative distal target sites, while the depletion of the other TFs (Stat3 and Prdm9) did not have such an effect [\(Figures 3E](#page-6-0)–3I). These global observations were validated by single-locus visualizations [\(Figures 3](#page-6-0)J–3L). In summary, these findings reveal lineage-specific TFs that shape the epigenetic landscape of distal regulatory elements during astrocyte differentiation and maturation.

#### Stage- and Lineage-Specific Activation of Regulatory Elements during Astrogliogenesis and Neurogenesis

Prompted by our findings that distinct TFs are required for conferring an active chromatin state to distal regulatory elements, we next investigated the activation dynamics of the TFtarget elements during astrogliogenesis and neurogenesis by measuring H3K27ac levels specifically at these sites. Putative Nfia and Atf3 sites were highly enriched with H3K27ac only during the eA stage of astrogliogenesis; these sites remained devoid of H3K27ac during neurogenesis ([Figures 4A](#page-8-0) and 4B; Figures S4A–S4F). Conversely, a similar analysis of the eN-specific TF Lhx2 showed that its putative target sites gain H3K27ac in eN, but not in eA ([Figure 4](#page-8-0)C; Figures S4G–S4I). The expression of the genes nearest to these sites also tended to be higher in eA compared to that in eN for both Nfia and Atf3, while in the case of Lhx2, the nearest genes were induced in eN, but not in eA [\(Fig](#page-8-0)[ures 4D](#page-8-0)–4I). Similarly, putative Runx2 and Brn1 sites showed elevated H3K27ac only in lA and lN, respectively [\(Figures 4J](#page-8-0) and 4K; Figures S4J–S4O). In keeping with this, the genes nearest to these Runx2 and Brn1 sites were significantly induced in IA and lN, respectively ([Figures 4L](#page-8-0)–4O). In addition, by comparing the sites putatively occupied by Nfia, Atf3, and Runx2 at all H3K27ac peaks throughout astrogliogenesis, we observed eA stage-specific H3K27ac enrichment at Nfia and Atf3 motifs and lA stage-specific H3K27ac enrichment at the Runx2 motif (Figure S4P). Consistent with our previous observations, Nfia, Atf3, and Runx2 sites were already primed during the preceding stage, before they acquired H3K27ac in eA and lA, respectively [\(Figures 4](#page-8-0)A, 4B, and 4J). Altogether, our data suggest that Nfia, Atf3, and Runx2 are crucial for switching the chromatin state of their putative target elements from primed to active, thereby driving the gene expression program underlying astrocyte differentiation and maturation.

#### Nfia, Atf3, and Runx2 Mediate Gene Expression Programs Underlying Astrogliogenesis

To further investigate the target genes that are under the direct transcriptional control of Nfia, Atf3, and Runx2, we performed transcriptome profiling (RNA-seq) after performing an siRNAmediated depletion of these TFs at the eA stage (Nfia and Atf3) or the lA stage (Runx2). Our analysis revealed a large number of differentially expressed genes after their depletion [\(Figures](#page-9-0) [5A](#page-9-0)–5C; Table S1). Depletion of Nfia and Atf3 led to changes in astrocyte morphology, as well as downregulation in astroglial markers, such as Gfap, Aldh1l1, Aldoc, and Aqp4, while Runx2 knockdown resulted in the downregulation of genes such as Nfia, Cx43, and Aqp4, but not of other markers, such as Gfap, which was even upregulated, suggesting a potential role for Runx2 in preventing a reactive state [\(Figure 5](#page-9-0)D; Figures S5A and S5B). While Nfia depletion had no effect on proliferation, apoptosis, or cell cycle (Figures S5C–S5E), Atf3 and Runx2 knockdown reduced apoptosis (Figures S5C–S5E) and Runx2 depleted cells exhibited G0/G1 arrest (Figures S5C–S5E).

The genes that were downregulated or upregulated after the Nfia depletion exhibited enrichment in extracellular matrix organization- and neuronal differentiation-related GO, respectively, suggesting that Nfia may repress the neurogenesis-related program ([Figure 5E](#page-9-0)). However, genes that were downregulated or upregulated after the Atf3 knockdown were enriched with terms related to cell adhesion or mitotic cell-cycle processes, respectively, implying that these genes are relevant for exiting the cell cycle and entering astrocyte differentiation [\(Figure 5](#page-9-0)F). In contrast, after Runx2 depletion, the downregulated genes were enriched with GO terms related to cell morphogenesis and differentiation, while the upregulated genes were enriched with genes related to cytokine response, suggesting that Runx2 promotes differentiation and may counteract acquisition of a reactive phenotype [\(Figure 5G](#page-9-0)). Single-gene examples substantiated our observations that Nfia and Atf3 promote astrogliogenesis by suppressing neurogenesis and promoting cell-cycle exit of progenitors, respectively, while Runx2 counteracts activation of a reactive phenotype to promote astrocyte maturation [\(Fig](#page-9-0)[ures 5](#page-9-0)H–5J). Genes that were downregulated upon Nfia, Atf3, and Runx2 depletion were upregulated during astrogliogenesis, corroborating the specific role of these TFs in astrogliogenesis (Figures S5F–S5H). In addition, the genes downregulated upon Nfia and Atf3 knockdown are expressed at a higher level in astrocytes compared to neurons, and an opposite pattern was observed for the upregulated genes (Figures S5I and S5J). We did not observe differential expression in astrocytes versus neurons in the case of Runx2-deregulated genes (Figure S5K).

To determine whether the regulatory landscape of the deregulated genes is also altered following TF depletion, we assessed H3K27ac enrichment at stage-specific regulatory sites nearest to these genes (Atf3 and Nfia for eA and Runx2 for lA). In

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#### Figure 4. Distinct TF-Target Regulatory Elements Are Stage- and Lineage-Specifically Activated during Astrogliogenesis versus Neurogenesis

(A–C) Heatmaps depicting the enrichment of H3K27ac and H3K4me1 during stages of either astroglial or neuronal differentiation at the genomic locations of the Nfia (A), Atf3 (B), and Lhx2 (C) motifs within the scope of unique eA, in the cases of Nfia and Atf3, and eN, in the case of the Lhx2, H3K27ac peaks, including the promoters.

(D–I) Expression of the genes nearest to and within 50 kb of the sites depicted in (A)–(C) in astroglial (D, F, and H) and neuronal (E, G, and I) differentiation (log2 normalized read counts).

(J and K) Same as (A)–(C) except for the Runx2 and Brn1 motif sites within the scope of the lA and lN H3K27ac unique peaks, respectively, including the promoter. (L–O) Same as (D)–(I) except for the Runx2 and Brn1 motif sites within the scope of the lA and lN unique H3K27ac peaks, respectively, including the promoter.

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#### Figure 5. Astroglial TFs Regulate Distinct Gene Expression Programs during Astrogliogenesis

(A–C) Volcano plots depicting the differential expression of genes between the control conditions and the conditions with depletion of Nfia in eA (A), Atf3 in eA (B), and Runx2 in lA (C) (green, up- and downregulated; black, no change).

(D) Bar plots representing the expression of astroglial markers (Gfap, Nfia, Aldh1l1, Aldoc, Aqp4, and Cx43) under the control and depleted conditions. Values are plotted as normalized read counts.

(E–G) Heatmap depicting the top gene ontologies enriched with downregulated (marked in red) and upregulated (marked in green) genes upon Nfia knockdown (E) and Atf3 knockdown (F) in eA and Runx2 knockdown (G) in lA.

(H–J) Examples of genes upregulated upon depletion of Nfia (H), Atf3 (I), and Runx2 (J).

(K–M) Profile plot of H3K27ac enrichment under the control conditions and the conditions with depleted Nfia (K), Atf3 (L), and Runx2 (M) in the regulatory regions (H3K27ac peaks near deregulated genes) of the upregulated genes and downregulated genes upon Nfia and Atf3 reduction in eA and Runx2 reduction in lA.

each case, downregulated genes lost H3K27ac enrichment upon knockdown of these factors, indicating that these TFs have an activating function at these sites [\(Figures 5K](#page-9-0)–5M). Strikingly, in the case of upregulated genes, H3K27ac enrichment at their regulatory sites did not change for Atf3 and Runx2 and only slightly increased for Nfia, suggesting that the activation of these genes was likely a result of secondary cascades. Overall, our data suggest that Nfia, Atf3, and Runx2 are critical for mediating the gene expression programs underlying astrogliogenesis.

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Figure 6. Nfia, Atf3, and Runx2 Directly Bind Predicted Target Elements and Induce Chromatin Accessibility

(A) Motif enrichment analysis of the set of commonly deregulated and H3K27ac-enriched genes (47 genes) upon Nfia and Atf3 depletion.

(B–E) ChIP-qPCR following Nfia, Atf3, and Runx2 ChIP to detect the Nfia and Atf3 co-occupied (B), Nfia (C), Atf3 (D), and Runx2 (E) binding at predicted Nfia, Atf3, and Runx2 motif sites in the candidate regulatory regions. Enrichments are plotted as a ratio of precipitated DNA (bound) to total input DNA and then further normalized to an intergenic region.

(F–I) qPCR following ATAC in aNPC, eA, and lA and after Atf3 and Nfia (F; eA), Nfia (G; eA), Atf3 (H; eA), and Runx2 (I; lA) depletion at regulatory regions listed in (B)–(E), depicted as a heatmap. Enrichments are plotted as in (B)-(E).

#### Nfia, Atf3, and Runx2 Directly Bind Predicted Target Elements and Induce Chromatin Accessibility

Given the known cooperativity among TFs that control cell-fate decisions, we assessed whether Nfia and Atf3 collaborate to regulate early astrogliogenesis. A comparison of the transcriptome data after Nfia and Atf3 knockdown revealed a significant overlap among the genes that were downregulated after their depletion ( $n = 198$  genes), suggesting similar aberrations in the astrogliogenesis program after their knockdown (Figure S6A; Table S5). Based on H3K27ac enrichment, a significant set of genes that were commonly downregulated after Nfia and Atf3 depletion exhibited binding sites for both TFs ( $n = 47/198$ ), suggesting their potential direct cooperativity during astrocyte differentiation ([Figure 6](#page-10-0)A; Figure S6A).

Given our findings that Nfia, Atf3, and Runx2 are crucial for switching the chromatin state of distal regulatory elements from primed to active, we next investigated whether these factors bind the predicted target elements at the respective stages of astrogliogenesis. Toward this end, we reanalyzed our datasets to validate the reduction in expression and H3K27ac enrichment at selected target genomic loci following depletion of these TFs (Figures S6B–S6I) Next, ChIP assays for Nfia and Atf3 in eA and Runx2 in lA to demonstrated binding at the selected target sites ([Figures 6B](#page-10-0)–6E).

Finally, we investigated whether the target sites of these TFs gain in accessibility while they are bound by these TFs and whether this increase in accessibility depends on the presence of these factors. Assay for transposase-accessible chromatin (ATAC) showed that these sites become more open in the corresponding stages concomitant to TF binding and acquisition of H3K27ac ([Figures 6F](#page-10-0)–6I; Figures S6J–S6M). Conversely, following depletion of these TFs, these target sites showed a significant reduction in chromatin accessibility [\(Figures 6](#page-10-0)F–6I; Figures S6N–S6Q). Together with our previous observations, these data support the notion that Nfia, Atf3, and Runx2 directly target distal regulatory elements and induce their active state, as manifested by increased chromatin accessibility, to drive gene expression programs underlying the differentiation and maturation of astrocytes.

#### Nfia, Atf3, or Runx2 Overexpression Steers Neurogenic Radial Glia Away from Generating Neurons and Promotes Astroglial Progenitor Generation In Vivo

To corroborate our key observations on astrogliogenesis from ESCs, we employed primary cortical astrocytes from postnatal day 5 (P5) mice to analyze occupancy by the astrogliogenic TFs identified here and the occurrence of active chromatin landscape at defined target sites [\(Figure 7](#page-11-0)A). Primary astrocytes

<span id="page-11-0"></span>

#### Figure 7. Astroglial TFs Regulate a Similar Set of Sites in Primary Cultures, and Their Overexpression In Vivo Alters the Fate of Cortical **Progenitors**

(A) Schematic representation of primary astrocyte culture.

(B–E) ChIP-qPCR following H3K27ac ChIP in primary astrocytes derived from P5 mice to detect its enrichment at the Nfia and Atf3 co-occupied (B), Nfia (C), Atf3 (D), and Runx2 (E) motif sites in the candidate regulatory regions. Enrichments are plotted as a ratio of precipitated DNA (bound) to total input DNA and then further normalized to an intergenic region.

exhibited high levels of H3K27ac at the identified TF-target reg-ulatory elements ([Figures 7B](#page-11-0)-7E). Further analysis using ChIP assays for Nfia, Atf3, and Runx2 confirmed the occupancy by these TFs at their predicted sites [\(Figures 7](#page-11-0)F–7I).

Given that these TFs were found to function in activation of astroglial-specific regulatory elements but were not engaged in activation of similar elements in neurogenesis, we hypothesized that the expression of these factors might interfere with the execution of neurogenesis programs during embryonic development. We thus performed *in utero* electroporation (IUE) of mouse cortical progenitors on embryonic day (E) 15.5 (Figueres-Oñate [et al., 2016; Pataskar et al., 2016\)](#page-14-13) with plasmids encoding NFIA, ATF3, or RUNX2 or a mixture of these three TFs (MIX) (Figures S7A and S7B). The successful expression of the TFs was confirmed in HEK293T cells by immunoblotting (Figure S7C). At E18.5, cells electroporated with the control plasmid exhibited a typical distribution in the developing cortex that is characterized by migration to the cortical plate (CP) (54.8%  $\pm$  5.8% of red fluorescent protein (RFP)-positive cells in the CP,  $n = 3$  animals) ([Figures 7](#page-11-0)J and 7K). In contrast, most cells overexpressing each of the astrogliogenic TFs or their mixture failed to populate the CP ([Figures 7](#page-11-0)J and 7K). Instead, these cells were retained in the GLAST-positive progenitor zone [\(Figures 7J](#page-11-0) and 7K; Figure S7D; Table S6). Furthermore, to evaluate whether the cells ectopically expressing these TFs are prevented from acquiring a neurogenic fate, we analyzed the expression of SATB2, a marker of upper-layer neurons generated at this embryonic stage, in electroporated cells [\(Figure 7](#page-11-0)L; Figure S7E; Table S6). This analysis demonstrated that cells expressing any of these TFs alone or in combination fail to gain expression of SATB2, in line with their retention below the CP. Further analysis of the expression of the astroglial progenitor marker SOX9 in electroporated cells showed that the expression of these TFs causes neurogenic radial glia to generate a larger proportion of SOX9 positive progenitors [\(Figure 7L](#page-11-0); Table S6).

To characterize the influence of the TFs at a later stage of development, we performed *in utero* electroporation in E15.5 cortical progenitors using MIX and analyzed at the P5 stage. Consistent with our results at E18.5, TF-overexpressing cells were mostly retained in the progenitor zone at P5, although most control-electroporated cells generated neuronal cells that already populated the upper cortical layers ([Figure 7](#page-11-0)M). Moreover, we observed a significant increase in the number of SOX9-positive cells among TF-electroporated cells compared to control [\(Figures 7M](#page-11-0) and 7N; Table S6). In contrast, a converse pattern was observed for the mature neuronal marker NeuN [\(Fig](#page-11-0)[ures 7M](#page-11-0) and 7N; Table S6). Altogether, these results suggest an

increase in the generation of astroglial progenitors at the expense of neurons.

#### **DISCUSSION**

We aimed to identify key regulators of astrogliogenesis based on lineage- and stage-specific remodeling of the transcriptional and epigenetic landscapes during astrocyte differentiation. Using mouse ESC differentiation into astrocytes as a model system, we identified three transcriptionally and epigenetically distinct stages: (1) a progenitor stage (aNPC), (2) an early stage of astroglial differentiation (eA), and (3) a more advanced stage of astroglial differentiation (lA). We obtained evidence for epigenetic priming along the differentiation axis, indicating that astrogliogenesis follows a dynamic trajectory during which subsequent stages are being orchestrated by preceding ones. By searching for lineage- and stage-specific regulatory elements unique to each stage and inferring their most highly associated TF binding motifs, we were able to pinpoint drivers of the underlying differentiation trajectory: the TFs Nfia, Atf3, and Runx2. These TFs were found to be not only important for expression and maintenance of astroglial marker genes but also critically required for the epigenetic remodeling at the transition between distinct stages. Our molecular analysis of astrogliogenesis supports the notion that differentiation of neural precursors into astrocytes is not a one-step but rather a multi-phase process ([Kang et al.,](#page-14-4) [2012\)](#page-14-4). Furthermore, *in vivo* overexpression of these TFs together resulted in an increase in SOX9-positive progenitors at the expense of neurogenesis; however, because of the transient nature of the overexpression, a definitive cell fate could not be tracked. It is possible that the continuous expression of the TFs hampers proper astrocyte development and maturation, and using a system allowing temporally controlled TF expression could further drive the acquisition of an astrocytic phenotype.

A substantial body of prior work has provided compelling evidence for an important role of Nfia at the onset of astrogliogenesis, both in the developing spinal cord and in the cerebral cortex [\(Cebolla and Vallejo, 2006; Deneen et al., 2006; Kang et al.,](#page-14-14) [2012\)](#page-14-14). However, the precise role of Nfia remains largely elusive. Combined expression of Nfia with Nfib and Sox9 can induce conversion of fibroblasts into astrocyte-like cells ([Caiazzo](#page-14-15) [et al., 2015](#page-14-15)), supporting the notion of an instructive role in astrogliogenesis. The present study showed that Nfia is required for converting primed chromatin into a transcriptionally active one at the transition from aNPC to eA. Nfia also appears to play an important role in suppressing neuronal gene expression. Thus, Nfia may regulate distinct neuronal-fate-suppressing and

<sup>(</sup>F–I) ChIP-qPCR following Nfia (F and G), Atf3 (F and H), and Runx2 (I) ChIP in primary astrocytes derived from P5 mice, showing their occupancy. Enrichments are plotted as in B-E.

<sup>(</sup>J) Distribution of electroporated cells at the E18.5 stage upon *in utero* electroporation of E15.5 ventricular and/or subventricular progenitors using a control pCIDRE plasmid or pCIDRE plasmid encoding NFIA, ATF3, RUNX2, or a combination of all three factors (MIX) (revealed by RFP immunolabeling, shown as white cells).

<sup>(</sup>K) Quantification of (J).

<sup>(</sup>L) Percentage of SOX9- and SATB2-positive cells among RFP-positive cells from cortices electroporated with pCIDRE control or pCIDRE-TF plasmids (single TFs or in combination) and analyzed at the E18.5 stage.

<sup>(</sup>M) Tilescan images showing distribution of electroporated cells at the P5 stage following *in utero* electroporation as above in control or MIX condition (tagged with MYC). Representative images of SOX9, MYC, and NeuN immunofluorescent staining in control and MIX conditions.

<sup>(</sup>N) Quantification of SOX9 and NeuN in electroporated cells in the MIX condition. MYC was used to quantify the expression of SOX9 in electroporated cells in the MIX condition.

astrocyte-fate-instructing gene expression modules. The latter module may be co-regulated by Atf3, because there was significant overlap of genes, comprising putative shared direct targets, that were downregulated following knockdown of either TF. Moreover, Nfia became downregulated following Atf3 knockdown, indicating that Atf3 may function upstream of Nfia. Like Nfia, Atf3 was found to be crucial for the conversion of primed to active chromatin. However, in contrast to Nfia, knockdown of Atf3 resulted in the upregulation cell-cyclerelated genes, supporting a specific role of Atf3 in cell-cycle exit at the progenitor stage. So far, Atf3 has been largely recognized as a stress response gene ([Hai et al., 1999\)](#page-14-16). Stress response such as unfolded protein response (UPR) mediated by Atf4, itself known to induce Atf3, has been shown to regulate the stage-specific balance between direct and indirect neurogenesis from radial glia during cortical development ([Laguesse](#page-14-17) [et al., 2015](#page-14-17)), thus setting a precedent for ATF-mediated cellfate decisions. Finally, a study identified *Drosophila* Atf3 as a cell polarity response gene. While there is ample evidence for an important role for cell polarity in the regulation of neuronal-fate decisions [\(Donohoe et al., 2018\)](#page-14-18), virtually nothing is known regarding the acquisition of a glial fate. Our study warrants further investigation into the functional importance of UPR and cell polarity regulation as potential mechanisms in astrogliogenesis.

Virtually nothing is known about Runx2 function in the CNS [\(Wang and Stifani, 2017\)](#page-15-10). Runx2 plays a key role in osteoblast and chondrocyte differentiation ([Komori, 2018](#page-14-19)). Computational modeling of stem cell-fate decisions predicted Runx2 as a determinant of astrogliogenesis, and Runx2 overexpression induced astroglial differentiation in mouse neural stem cells ([Okawa](#page-15-11) [et al., 2016](#page-15-11)). One particularly intriguing observation was that knockdown of Runx2 resulted in the induction of genes associated with reactive astrocytes, suggesting that Runx2 may be important in repressing a reactive phenotype. Reactive astrogliosis represents in many respects a state of dedifferentiation and involves reacquisition of neural stem cell properties [\(Robel](#page-15-12) [et al., 2011\)](#page-15-12), which can eventually reactivate neurogenic potential [\(Magnusson et al., 2014; Nato et al., 2015\)](#page-15-13). It will be interesting to learn whether such dedifferentiation involves downregulation of Runx2.

Consistent with its known function in astrogliogenesis [\(Bonni](#page-14-3) [et al., 1997; Moon et al., 2002\)](#page-14-3), Stat3 was found to strongly influence astroglial marker expression following knockdown. Surprisingly, however, we failed to observe global changes in H3K27ac at its putative targets. Although we cannot exclude that Stat3 may induce chromatin remodeling via alternative mechanisms, such as sequence-independent recruitment of Stat3, our observation may genuinely indicate that Stat3 acts by transactivating already-accessible targets upon prior chromatin remodeling by pioneer TFs. Given its role in opening chromatin at the transition to the lA stage, Runx2 might be in a prime position to exert such a function. However, activation of Stat3 is also implicated in reactive astrogliosis [\(Herrmann et al., 2008\)](#page-14-20); hence, Runx2 and Stat3 may act partially antagonistically to each other.

Although our study revealed compelling evidence for dynamic transcriptional and epigenetic remodeling during astrogliogenesis, it also raises intriguing questions regarding the molecular mechanisms that account for the epigenetic writing of specific priming marks at gene loci that subsequently become specifically activated in astrogliogenesis. Furthermore, how the TFs driving astrogliogenesis are activated during this process remains to be investigated. Lastly, we lack molecular insights into how the TFs mediate transition of the chromatin state of the target regulatory elements from primed to active. Deciphering these mechanisms will lead us to a better understanding of how glial and neuronal lineage diverge during development. Our study represents a rich resource for the identification and characterization of further regulators of such key decisions during CNS development.

#### STAR+METHODS

Detailed methods are provided in the online version of this paper and include the following:

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#### SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures and seven tables and can be found with this article online at [https://doi.org/10.1016/j.stem.2018.09.008.](https://doi.org/10.1016/j.stem.2018.09.008)

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#### AUTHOR CONTRIBUTIONS

N.T. designed the study, performed experiments, analyzed data, and wrote the manuscript. A.P. developed computational pipelines, performed bioinformatics analysis, and wrote the manuscript. S.T. initiated the bioinformatics analysis, developed computational pipelines, and provided the first set of observations. S.P. performed *in utero* electroporation with the assistance of M.F.-O. and L.L.-M. S.K.S. helped set up astrocyte differentiation protocols. N.M. performed the calcium imaging experiments. V.K.T. and B.B. designed the study, analyzed data, and wrote the manuscript. All authors read and approved the final manuscript.

#### DECLARATION OF INTERESTS

The authors declare no competing interests.

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### <span id="page-16-1"></span>STAR+METHODS

#### <span id="page-16-0"></span>KEY RESOURCES TABLE











#### <span id="page-20-0"></span>CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resource/reagents should be directed to and will be fulfilled by the Lead Contact, Prof. Dr. Benedikt Berninger ([benedikt.berninger@kcl.ac.uk\)](mailto:benedikt.berninger@kcl.ac.uk).

#### <span id="page-20-1"></span>EXPERIMENTAL MODEL AND SUBJECT DETAILS

The handling of the mice and all experimental procedures were conducted in accordance with the European Union guidelines on the use and welfare of experimental animals (2010/63/EU). Experimental procedures were approved by the State of Rheinland Pfalz, Germany (23177-07/G17-1-067) or Spanish Ministry of Agriculture (RD 1201/2005 and L 32/2007) and the Cajal Institute, CSIC Animal Experimentation Ethics Committees and the Community of Madrid (Ref. PROEX 223/16). Animals were housed on a 12:12 hr lightdark cycle, with free access to food and water. None of the mice used were involved in previous procedure or drug treatment. Both male and female embryos were used and randomly allocated to experimental groups.

#### Cell culture

mESCs derived from C57/BL-6 mice were cultured in DMEM supplemented with 10% FBS, 2 mM Glutamax, 2 mM sodium pyruvate,  $2$  mM non-essential amino acids, Leukemia-inhibitory factor (LIF) and 50  $\mu$ M  $\beta$ -mercaptoethanol. The cell medium was changed every day, and the cells were passaged every second day using 0.05% trypsin and plated on 0.2% gelatin coated plates. Using the protocol proposed by Conti et al., the ESCs were further differentiated into aNPC ([Conti et al., 2005; Pollard et al., 2006](#page-14-8)). Briefly, the ESCs were passaged and grown in N2/B27 media containing 0.5x DMEM-F12, 0.5x N2 supplement, 0.5x B27 supplement, 0.5x Neurobasal media, 2 mM Glutamax and 2 mM Penicillin-Streptomycin for 7 days on 0.2% gelatin coated plates. The cells were trypsinized and grown on uncoated dishes for the following 7 days in NSA media containing 10 ng/ml FGF2 and 10 ng/ml EGF to form cell aggregates. The formed aggregates were further passaged and grown on 0.2% Gelatin coated plated to obtain the monolayer of the aNPC. To generate the astrocytes, the aNPC were differentiated in differentiation media containing DMEM, 1% FBS, 1x B27, LIF, 2 mM Glutamax and 2 mM Penicillin-Streptomycin for 1, 5 and 21 days. For differentiation of aNPC into neurons, cells were kept in neurobasal media, along with B27 for 2 weeks. 293T cells were cultured in DMEM, 10% FBS, Penicillin-Streptomycin and Glutamax.

#### Primary astrocyte culture

The primary astrocyte culture was prepared from the brains of P5-P7 C57/BL6 mice using a previously described protocol ([Heinrich](#page-14-21) [et al., 2011](#page-14-21)). Briefly, the cortices were dissected to remove the white matter and meninges and cultured in DMEM-F12 media supplemented with 1x B27, 10% FBS, 2 mM Glutamax and 2 mM Penicillin-Streptomycin.

#### <span id="page-21-0"></span>METHOD DETAILS

#### Immunofluorescence and Brdu labeling

The cells were grown on coverslips, fixed with 4% paraformaldehyde and permeabilized with 0.2% Triton X-100 for 5 minutes at room temperature. Subsequently, the cells were blocked with 5% BSA for 30 minutes, incubated with primary antibodies (mouse anti-GFAP-1:500, G3893, Sigma; rabbit anti-GFAP-1:200, Z0334(29-2), DAKO; rabbit anti-AQP4-1:200, sc-20812, Santa Cruz; mouse anti-S100B, ab11178, Abcam; rabbit anti-GLAST-1:200, AB\_2571715, Frontier Institute; rabbit anti-CX43-1:200, ab11370, Abcam; rabbit anti-OLIG2-1:200, AB9610, Milipore; rat anti-ASCL1-1:200, MAB256, R&D systems; rabbit anti-NES-1:200, 839801, BioLegend; anti-guinea pig anti-DCX-1:200, AB2253, Milipore; rabbit anti-SOX9-1:200, ab185966, Abcam and rat anti-CD44- 1:200, 550538, BD PharMingen) for 1 hour, and then incubated with a fluorochrome-labeled secondary antibody for 45 minutes at room temperature. The coverslips were counterstained with DAPI and imaged under a confocal laser-scanning microscope. The data were processed using Adobe Photoshop software. For the Brdu labeling, the aNPC were differentiated into astrocytes for 1 and 5 days, labeled with 30 µM Brdu for 8 hours. Cells were fixed in 70% ethanol and denatured by using 2N HCL/0.5% Triton X-100 for 30 minutes. HCL was neutralized by 0.1 M Na2B4O7 pH 8.5 buffer for 2 minutes. Cells were washed with 0.5%Tween-20/1% BSA/PBS buffer before proceeding for Brdu staining (1:200, B5002, Sigma) as above.

#### Immunoblotting

Immunoblotting was performed as described previously ([Tiwari et al., 2013\)](#page-15-19) using following antibodies: mouse anti-GFAP (1:1000, G3893, Sigma), rabbit anti-CX43 (1:000, ab11370, Abcam), rat anti-ASCL1 (1:1000, MAB2567, R&D systems), rabbit anti-NES (1:1000, 839801, BioLegend), rabbit anti-OLIG2 (1:1000, AB9610, Milipore), rabbit anti-NFIA (1:1000, ab41851, Abcam), rabbit anti-RUNX2 (1:1000, sc-10758 X, Santa Cruz), rabbit anti-GAPDH (1:1000, ab9484, Abcam) and mouse anti-FLAG (1:1000, F1804, Sigma). Briefly, cells were lysed using RIPA buffer (50 mM HEPES pH 7.5, 150 mM NaCl, 5 mM EGTA, 1.5 mM MgCl2, 1% Glycerol, 1% Triton X-100) and protein concentration was measured using Biorad reagent (Life Technologies) following manufacturer's protocol. 50 µg lysate was mixed with final 1x Laemelli buffer (5x Laemelli buffer: 10% SDS, 50% glycerol, 25% 2-mercaptoethanol, 0.02% bromphenol blue and 0.3125 M Tris HCl, pH approx. 6.8.), then run at 150 V and transferred at 100 V for 2-hours with BioRad Mini-PROTEAN and Mini Trans-Blot Cell tanks respectively. Subsequently, the membranes were blocked with 5% non-fat dry milk in TBST for 1 h at room temperature and incubated in primary antibodies diluted in 5% non-fat dry milk in TBST for overnight at 4°C. The membranes were washed three times for 5 minutes with TBST and incubated in horseradish peroxidase coupled secondary antibody (in 5% non-fat dry milk) for 1 hour at room temperature. After three 5 minutes washes with TBST, membranes were incubated with ECL solution (Amersham ECLTM Prime Western Blotting Detection Reagent) according to the manufacturer's instructions and imaged with the Peqlab Fusion-SL system using the fusion software version 15.16 by Vilber Lourmat.

#### $Ca<sup>2+</sup> Imaqina$

eA and IA were incubated with 10  $\mu$ M OGB1, diluted in culture medium, for 45 min at 37°C in a humidified atmosphere. After the incubation time, cells were washed with warm culture media and analyzed within 2 hr ([Lee et al., 2015\)](#page-14-27). Cells were placed in a recording chamber mounted on a fluorescent microscope (Zeiss Axio Imager A2) and superfused with artificial cerebrospinal fluid (ACSF) (in mM): NaCl, 125; KCl, 2.5; NaHCO<sub>3</sub>, 25; NaH2PO<sub>4</sub>, 1.25; CaCl2, 2; MgCl<sub>2</sub>, 1 and glucose, 25, saturated with 5% CO<sub>2</sub> and 95% O<sub>2</sub> (pH 7.4, 30-32°C). Cells were visualized with a 40x (0.7 NA) objective and movies (15 s) were acquired with a frequency of 8.9 Hz with a Hamamatsu Orca-0.3G camera and HCImage software ([Lee et al., 2015\)](#page-14-27). During acquisition, we tested the generation of  $Ca<sup>2+</sup>$  waves in astrocytes using mechanical stimuli with a glass pipette which was placed right above the membrane of cultured astrocyte (resistance of 5-10 M $\Omega$ , which corresponds to a tip opening of 1-2 mm) [\(Lee et al., 2015\)](#page-14-27). For quantification, we defined regions of interest (ROIs) in cells surrounding the site of mechanical stimulus. Intensity measurements for ROIs are expressed as the change of fluorescence relative to background fluorescence ( $\Delta F/F0$ ) and were calculated with ImageJ (ROI manager) [\(Lee et al., 2015\)](#page-14-27).

#### siRNA transfection and plasmid transfection

For the siRNA (Dharamcon) transfection, Lipofectamine RNAiMax (Invitrogen) was used according to manufacturer's instructions to knockdown the candidate TFs. The candidates were first knocked down in aNPC for  $-2$  days to reduce the basal level expression and were then re-transfected with siRNA at day 0 and kept in aNPC media for 1 day and changed to astromedia next day for 1 day experiment (Nfia and Atf3). On the other hand, for the 5 days experiment (Runx2), day 2 astrocytes were re-transfected with siRNA in astrocyte differentiation media. 293T cells were transfected with pCIDR-dsRed or pCIDR-TF-dsRed plamids by using Lipofectamine 2000, according to manufacturer's instructions.

#### RNA purification, cDNA preparation and Quantitative RT-PCR

RNA was extracted using the Tri-reagent from sigma, and cDNA was prepared according to the manufacturer's instructions (Applied Biosystems) using 1 µg of initial RNA material. The cDNA was used to quantify the gene expression on a StepOne plus real-time PCR machine. The primer sequences are provided in Table S7.

#### Chromatin Immunoprecipitation

The ChIP experiments were performed as previously described ([Tiwari et al., 2013\)](#page-15-19). In brief, crosslinked chromatin was sonicated to achieve an average fragment size of 500 bp. Starting with 60 µg of chromatin and 5 µg of the anti-H3K27ac (Abcam), anti-Nfia (Santa Cruz), anti-Atf3 (Santa Cruz) and anti-Runx2 antibodies (Santa Cruz), the samples were immunoprecipitated and then de-crosslinked to release the DNA fragments. In total, 1  $\mu$ L of the ChIP material and 1  $\mu$ L of the input material were used for the quantitative real-time PCR using specific primers covering the motifs of Nfia, Atf3 and Runx2 in eA and lA. Primers covering an intergenic region were used as a control. The efficiencies of the PCR amplification were normalized to the PCR product of the intergenic region. Primer sequences provided in Table S7.

#### Transcriptome analysis

The transcriptome samples under the astroglial differentiation (aNPC, eA and lA), control and TF knockdown conditions (GSE96539) were derived in triplicate for astroglial differentiation and in biological replicates for depletion studies. The RNaseq output in FASTQ format was subjected to an initial quality assessment using FASTQC v2.6.14. After the quality check, the files were used for the read alignment to the mouse genome (mm9) with UCSC annotations [\(Karolchik et al., 2014](#page-14-28)) using TopHat v.2.0.8 ([Trapnell et al., 2009](#page-15-15)); then, the uniquely mapped reads were retained in the output BAM file. SAMTOOLS v0.1.19 [\(Li et al., 2009\)](#page-14-22) was used for the file format conversions and sorting of the alignment file. The raw read counts per gene were calculated using HTSeq program v0.5.4p1 [\(Anders](#page-14-23) [et al., 2015](#page-14-23)), and the read count data matrix was then subjected to library-size normalization and differential expression analyses using DESeq ([Oshlack et al., 2010\)](#page-15-16). An FDR cutoff of 0.1 was used to call the differentially expressed genes. The transcriptomes from the *in vitro* astroglial differentiation were normalized together, while those from the control and TF KD were normalized independently. The GO term analysis of all gene lists (Gene Symbol) was performed using ToppGene [\(Chen et al., 2009\)](#page-14-24). Heatmaps depicting the gene expression were plotted using pheatmap R package. The analyzed microarray data from GSE9655 [\(Cahoy et al., 2008\)](#page-14-9) were directly downloaded from GEO, while the RNaseq data from GSE73721 ([Zhang et al., 2016\)](#page-15-7) were analyzed as described above and compared to the RNaseq data using a linear regression analysis. Gene lists derived from the RNaseq analysis are supplied in Table S1.

#### ChIP-seq and motif analysis

The ChIP-sequencing output in FASTQ format was subjected to a quality check using FASTQC v2.6.14 (Andrews). ChIP-seq was performed to detect the histone markers H3K27ac and H3K4me1 during each stage of astroglial differentiation (GSE96539) and H3K27ac under the control and shortlisted TF knockdown conditions. Bowtie v0.12.9 ([Langmead, 2010](#page-14-25)) was used to align the reads uniquely, i.e., each read was maximally aligned to one position, to mm9 genome with UCSC annotations [\(Karolchik et al., 2014](#page-14-28)). The alignment output files from two biological replicates were merged together after checking for correlations across the replicates using the SAMTOOLS v0.1.19 ([Li et al., 2009\)](#page-14-22) merge function. SAMTOOLS v0.1.19 [\(Li et al., 2009](#page-14-22)) was used for the alignment file format conversions and sorting of alignment output files. The WIGGLE files for the alignment files were generated using QuasR package ([Gaidatzis et al., 2015](#page-14-26)). The peaks were computed without providing input with MACS v2.0.10.20120913 [\(Zhang et al., 2008](#page-15-17)) using the default parameters. The enrichment was calculated using formula as previously described [\(Pataskar et al., 2016\)](#page-15-14). The heatmaps and the density plots representing the ChIP-seq normalized read distribution over the shortened genomic region were plotted using the ngs.plot.r program ([Shen et al., 2014\)](#page-15-18).

HOMER v4.7 was used to obtain annotation information for the peaks, including the distance to the nearest TSS and the nearest gene. The peaks were associated with their nearest genes if they were either intergenic and less than 50KB from the TSS or at promoters, intron and exons.

A motif analysis was performed by HOMER v4.7 using a GC% normalized background control on the shortlisted genome coordinate sets using the findMotifs.pl program. Motif scanning of the complete genome was performed using the scanMotifsGenome.pl program in HOMER v4.7.

#### ATAC

ATAC library preparation was performed exactly as described ([Buenrostro et al., 2013\)](#page-14-29). Briefly, 50,000 eA and lA cells were incubated with 0.1% NP-40 to isolate nuclei. Nuclei were then transposed for 30 min at 37°C with adaptor-loaded Nextera Tn5 (Illumina, Fc-121-1030). Transposed fragments were directly PCR amplified and used for qRT-PCR.

#### Cell growth assay

For growth curves, cells were seeded in each well of a 24-well plate and cell numbers were assessed over the time periods indicated by using a Neubauer counting chamber upon TF depletion. For cell cycle analysis, cells were trypsinized and fixed overnight in 70% ice-cold ethanol, washed twice with PBS, and resuspended in sodium citrate buffer with 5 µg/ml PI overnight. Stained cells were analyzed on a FACSCanto II using DIVA software.

#### Apoptosis assay

Apoptosis assays were performed using a Cy5 Annexin V antibody according to the manufacturer's instructions (BD Biosciences). Stained cells were filtered with a 40µm mesh and analyzed on a FACSCanto II flow cytometer using DIVA software.

#### In utero electroporation

E15.5 pregnant C57BL/6 mice were anesthetized with 1.5%–2.8% isoflurane (Isova vet, Centauro) in pure  $O_2$  and received an injection of enrofloxacin (5 mg/kg, s.c.; Baytril Bayer) and meloxicam (300 µg /kg, s.c.; VITA Laboratories)/ Rimadyl (carprofen; 5 mg/kg; Pfizer) before starting the surgical procedure. The surgical area was constantly maintained moistened with a physiological saline solution. The abdominal cavity was opened, and the uterine horns were carefully extracted. For each embryo, 1 µL of DNA solution was injected into the lateral ventricle using a glass capillary, and 5 consecutive electric pulses of 37 V (50 ms each, 950 ms intervals) were delivered through platinum electrodes (3 mm diameter) using a BTX electroporator (Holliston). The uterine horns were then replaced in the abdominal cavity, and the abdomen was sutured. The DNA consisted of a pCAG-IRES-dsRedexpress (pCIDRE;  $4 \mu$ g) [\(Pataskar et al., 2016\)](#page-15-14) plasmid as control, or encoding NFIA, ATF3 or Runx2.

#### Immunohistochemistry and image analysis

For the in utero electroporated brains analyzed at E18.5, the pregnant mice were sacrificed, and the embryos were extracted from the uterine horns. For analysis at postnatal stages, animals were perfused with a saline solution (NaCl 0,9% in PB 0,1M) followed by 4% paraformaldehyde (PFA). Harvested embryonic brains were placed in 4% PFA for 48 h for fixation and postnatal brains were postfixed in 4% PFA for 24 h. The brains were embedded in agarose (3%, in PBS) and cut into 50 or 70  $\mu$ m (for postnatal and embryonic brains, respectively) serial coronal sections using a vibrating microtome (Microm HM650V, Thermo Scientific). The sections were then stored in a cryoprotective solution (20% glucose, 40% ethylene glycol, 0.025% sodium azide, and 0.05 M phosphate buffer, pH 7.4). For immunohistochemistry, free-floating sections were incubated for 90 min at room temperature (RT) in a blocking solution (5% Donkey serum, 0.3% Triton X-100 in TBS 0.1 M, pH 7.6). The primary antibodies, which were diluted in blocking solution, were applied for 2 hours at RT and overnight at  $4^{\circ}$ C. The following antibodies were used: donkey anti-RFP (1:500, 600401379S, Biomol) to detect the pCIDRE-electroporated cells, rabbit anti-SOX9 (1:200, ab185966, Abcam), rabbit anti-SATB2 (1:200, ab34735, Abcam), mouse anti-TUBB3 (1:500, T8660, Sigma), rabbit anti-GLAST (1:300, AB\_2571715, Frontier Institute), mouse anti-MYC tag (1:200, 2276S, Cell signaling), and mouse anti-NeuN (1:500, MAB377, Milipore). The appropriate secondary antibodies were diluted in blocking solution and applied for 1 hour at room temperature. The sections were then counterstained with DAPI and mounted with Vectashield (Vector Laboratories) mounting medium. Image stacks were acquired under an SP5 confocal microscope (Leica) with a 20x dry objective (NA 0.7) or a 40x oil objective (NA 1.3). For [Figure 7](#page-11-0)M, tilescans showing the dispersion pattern of RFP +ve cells were acquired under an SP5 confocal microscope (Leica) with a 40x oil objective (NA 1.3). Output 2D images were generated in ImageJ using the Maximum Intensity Projection (MIP) method. The quantification of the distribution of the electroporated cells was performed in 3 animals/group ( $n = 3$ ) with 2 sections/brain.

#### <span id="page-23-0"></span>QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical significance was evaluated by One-Way ANOVA, followed by Dunnett's test for multiple group comparisons or by t test for 2-groups comparison, using the SPSS Statistics 22 software for [Figures 7](#page-11-0), S1, S5, and S7. P values represent \* < 0.05; \*\* < 0.01; \*\*\* < 0.001. The data are expressed as the mean ± SD. The number of experiments and biological samples used is specified in figure legends and [Star Methods.](#page-16-1) The bar plots were generated using Prism software.

#### <span id="page-23-1"></span>DATA AND SOFTWARE AVAILABILITY

All relevant data are available from authors. All the software used in this work are described in the relevant [Star Methods](#page-16-1) sections. RNA-seq and ChIP-seq data have been deposited (GEO: GSE96539).

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## Supplemental Information

## Stage-Specific Transcription Factors Drive

## Astrogliogenesis by Remodeling

## Gene Regulatory Landscapes

Neha Tiwari, Abhijeet Pataskar, Sophie Péron, Sudhir Thakurela, Sanjeeb Kumar Sahu, María Figueres-Oñate, Nicolás Marichal, Laura López-Mascaraque, Vijay K. Tiwari, and Benedikt Berninger







## **Supplemental Figure 4 (Related to Figure 4)**



nNPC eN lN aNPC eA lA

aNPC eA lA

## **Supplemental Figure 5 (Related to Figure 5)**



## **Supplemental Figure 6 (Related to Figure 6)**



## **Supplemental Figure 7 (Related to Figure 7)**







### **SUPPLEMENTAL INFORMATION**

### **Figure S1 (Related to Figure 1):**

(A) Bright-field images showing the morphology of ESC-derived astrocytes during different differentiation stages (aNPC, eA, lA\_1 and lA\_2). Scale bar is 50 µm. (B) qPCR depicting the expression, represented as fold change, of astroglial markers (Gfap, S100b, Nfia, Glast, Cx43, and Aldoc) in aNPC, eA, lA\_1 and lA\_2. (C) Western blot analysis to show the expression of astroglial markers (GFAP, CX43), progenitor markers (NES, OLIG2, NFIA) and Runx2 during astrogliogenesis. (D) Immunofluorescence staining of various astrocyte (GFAP, AQP4, S100B, GLAST, CX43, SOX9, CD44), progenitor (OLIG2, ASCL1, NES) and neuronal (TUBB3) markers in the ESC-derived astrocytes during different phases of astrogliogenesis, including the aNPC, eA and IA. The scale bar is 50  $\mu$ m. (E) Brdu +ve cells were quantified by incorporating Brdu in aNPC, eA and lA for 8 hours. (F) Propagation of calcium waves upon mechanical stimulation of eA and lA membrane in stimulated (top) and surrounding (bottom) cells. (G) Pictures depicting the propagation of calcium waves (arrowheads) in stimulated and surrounding cells upon mechanical stimulation in eA (upper panels) and IA (lower panels). ΔF/F0 analysis of ROIs (open circles) showed the time course of  $[Ca_{2}]$ i increase after mechanical stimuli application. (H) Immunofluorescence staining for GFAP and TUBB3 in aNPC cells kept in differentiation media for 2 weeks (I) Table depicting the quantification of the significantly differentially expressed genes in respective comparisons. (J-N) Normalized read counts showing the expression of genes in each cluster depicted in Figure 1C (O) Heat map visualizing the expression of commonly regulated genes in the glial network and our dataset during astrogliogenesis (P) Quality control plot depicting the number of H3K27ac reads aligned per sample compared to the total sequenced reads after merging the two biological replicates. (Q) Bar plot depicting the H3K27ac peak counts in aNPC, eA and lA. (R) Bar plots showing the number of H3K27ac peaks unique to a particular cell type, i.e., either aNPC, eA or lA. (S, T) ChIP-qPCR to analyze the enrichment of H3K27ac in the promoter (S) and distal regulatory region (T) of astrogliogenic genes in aNPC, eA and lA.

### **Figure S2 (Related to Figure 2):**

(A) Bar plot depicting the number of H3K4me1 reads aligned per sample compared to the total sequenced reads after merging the two biological replicates. (B) Left side: Bar plot depicting the H3K4me1 peak counts in aNPC, eA and lA. Right side: Bar plots showing the H3K4me1 peaks unique to a particular cell type, i.e., aNPC, eA or lA. (C, D) Bar graphs representing the ChIPqPCR validation of the non-promoter (C) and promoter (D) regulatory regions of representative

astroglial and neuronal marker genes. (E) Schematic representation of neuronal differentiation from ES cells. Scale bar is 100 µm. (F) Box plots depicting the ChIP-seq enrichment of H3K27ac during astrogliogenesis (green) and neurogenesis (red) and H3K4me1 enrichment during astrogliogenesis at genomic regions differentiated by the clusters shown in Figure 2J. (G) Same as (F) but a depiction of the expression of genes during astrogliogenesis (red) and neurogenesis (green) that are associated with genomic regions that are differentiated by the clusters shown in Figure 2J. (H, I) Bar and line plots depicting the top gene ontologies enriched with genes associated with the regions depicted in cluster 2 (H) and clusters 6 and 7 (I) in Figure 2J. Main xaxis shows gene count while alternate x-axis on top shows p-values.

### **Figure S3 (Related to Figure 3):**

(A) Principal component analysis plot depicting the distribution of the transcriptomes in ESC, aNPC, eA, lA\_1, lA\_2, nNPC (neural progenitors), eN (early neurons) and lN (late neurons). (B-D) Volcano plot showing the differentially expressed genes in the comparison of the transcriptomes (Red: downregulated, Green: upregulated; Grey: Not changing) between aNPC and nNPC (B), eA and eN (C) and lA and lN (D). (E-G) Top enriched gene ontologies for genes that are at least 10 fold or more expressed in aNPC compared to nNPC (E), eA compared to eN (F) and lA compared to lN (G) are represented as bar and line plots. (H-J) Heatmap depicting the expression of TFs that are at least 2 fold or more expressed in aNPC compared to nNPC (H), eA compared to eN (I) and lA as compared to lN (J). (K-M) Same as (H-J) but for the expression of epigenetic regulators. (N-P) Overlap of H3K27ac enriched sites during astrogliogenesis with neurogenesis (nNPC, eN and lN; N), CD4 (O) and liver (P)'s H3K27ac sites. (Q) Selected candidate TFs for each stage of astrogliogenesis. The one highlighted with red color did not give significant depletion in their expression upon siRNA treatment. Thus, they were excluded from the studies. (R-T) siRNA depletion of stage-specific selected candidates in their respective stages and analysis of progenitor and astroglial marker expression by q-PCR. (U) qPCR following the Nfia and Atf3 depletion in eA and the Stat3, Prdm9 and Runx2 depletion in lA to show their knockdown efficiency. (V) Normalized read counts showing the expression Nfia, Atf3, Stat3, Prdm9 and Runx2 during astroglio- and neurogenesis. (W, X) Immunofluorescence staining of DAPI, GFAP and candidate TFs in ESC-derived astrocytes during different phases of astrogliogenesis, including the progenitor stage (aNPC), D1 (early astrocytes; eA) and D5 (late astrocytes; lA). Scale bar is 100 µm.

## **Figure S4 (Related to Figure 4)**:

(A-C) ChIP-seq enrichment of H3K27ac during astrogliogenesis (A) and neurogenesis (B) and H3K4me1 during astrogliogenesis (C) at genomic locations marked by the Nfia motif in eAspecific H3K27ac peaks. (D-F) Same as (A-C) but at genomic locations marked by the Atf3 motif in eA-specific H3K27ac peaks. (G-I) Same as (A-C) but at genomic locations marked by the Lhx2 motif in lN-specific H3K27ac peaks. (J-L) Same as (A-C) but at genomic locations marked by the Runx2 motif in lA-specific H3K27ac peaks. (M-O) Same as (A-C) but at genomic locations marked by the Brn1 motif in lN-specific H3K27ac peaks. (P) Neuronal progenitor cells can either differentiate into astrocytes or neurons. During astroglial differentiation, the Nfia and Atf3 motif sites are specifically enriched among all H3K27ac peaks in eA-during astrogliogenesis and not during neurogenesis. These sites are already primed in aNPC by H3K4me1 mark. Runx2 sites show a gradual gain in H3K27ac enrichment during astrogliogenesis but they do not acquire this mark during neurogenesis. Interestingly, these sites also gain H3K4me1 during astrogliogenesis but its highest levels at Runx2 sites are achieved already at the eA stage. All representations are presented as boxplots with a fold enrichment above the input on the y-axis for H3K27ac during astrogliogenesis (left panels), H3K4me1 during astrogliogenesis (middle panels) and H3K27ac during neurogenesis (right panels) at motif instances of Nfia (eA), Atf3 (eA) and Runx2 (lA).

## **Figure S5 (Related to Figure 5):**

(A) Bright-field images showing the morphology of cells upon depletion of Nfia and Atf3 in eA and Runx2 in IA. Scale bar is 50 µm. (B) Immunofluorescence staining for GFAP and AQP4 upon depletion of Nfia and Atf3 in eA and Runx2 in lA. (C) siRNA ablation of Nfia, Atf3 in eA and Runx2 expression in lA for 1 day and quantification of cell numbers. (D) Rate of apoptosis as determined by Annexin-V staining and flow cytometery in eA transfected with Nfia or Atf3 and lA transfected with Runx2 for 1 day. (E) eA were transfected with Nfia or Atf3 and lA were transfecetd with Runx2 for 1 day and then stained with propidium iodide (PI), and the percentages of cells in G0/G1 and S-G2/M phases of the cell cycle were determined by flow cytometry. (F-H) Boxplot depicting the expression of the downregulated and upregulated genes upon the Nfia and Atf3 KD in eA (F,G) and the Runx2 KD in lA (H) during stages of astroglial and neuronal differentiation. (I-K) Gene set enrichment plot depicting the density of the occurrence of astrocyte genes (genes induced in astrocytes compared to those induced in neurons) and neuronal genes (vice-versa) in Nfia (I), Atf3 (J) and Runx2 (K) deregulated genes.

## **Figure S6 (Related to Figure 6):**

(A) Venn diagram depicting the overlap of genes significantly downregulated by the Nfia and Atf3 KD in eA and their comparison with that of the nearest genes to the H3K27ac peaks during

astrogliogenesis that are enriched with either the Nfia or Atf3 motifs. (B-I) H3K27ac enrichment (B, D-F) and mRNA level (C, G-I) enrichment at the putative Nfia, Atf3 and Runx2 targets sites. (J-M) qPCR following ATAC in aNPC, eA and lA at sites co-bound by Nfia and Atf3 (J) or bound by Nfia (K), Atf3 (L) and Runx2 (M) alone (N-Q). qPCR following ATAC in Nfia- (N, O), Atf3- (N, P) and Runx2-depleted (Q) cells in eA (N-P) and lA (Q) respectively at the putative target regulatory elements.

## **Figure S7 (Related to Figure 7):**

(A) Schematic representation of constructs used for IUE (B) Schematic representation of the IUE experimentation. (C) Immunoblotting after overexpressing the pCIDRE and pCIDRE plasmid encoding Nfia, Atf3 and Runx2 in 293T cells. Flag antibody is used for the immunoblotting. (D) Immunofluorescence of the E18.5 neocortex using the GLAST and TUBB3 antibody. (E) Staining for SATB2 at E18.8, an upper neuronal marker, following overexpression of astroglial TFs by IUE at E15.5. TFs-overexpressing cells are mainly devoid of SATB2 expression.