Polycomb Protein Eed is Required for Neurogenesis and Cortical Injury Activation in the Subventricular Zone

Bin Sun, Eunhyuk Chang, Anna Gerhartl and Francis G. Szele

Department of Physiology, Anatomy and Genetics, University of Oxford, Oxford OX1 3QX, UK

Address correspondence to Francis G. Szele, PhD, Department of Physiology, Anatomy, and Genetics, University of Oxford, South Parks Road, Oxford OX1 3QX, UK. Email: Francis.Szele@dpag.ox.ac.uk

Abstract

The postnatal subventricular zone (SVZ) harbors neural stem cells (NSCs) that exhibit robust neurogenesis. However, the epigenetic mechanisms that maintain NSCs and regulate neurogenesis remain unclear. We report that label-retaining SVZ NSCs express Eed, the core component of Polycomb repressive complex 2. In vivo and in vitro conditional knockout and knockdown show Eed is necessary for maintaining NSC proliferation, neurogenesis and neurosphere formation. We discovered that Eed functions to maintain p21 protein levels in NSCs by repressing Gata6 transcription. Both Gata6 overexpression and p21 knockdown reduced neurogenesis, while Gata6 knockdown or p21 overexpression partially rescued neurogenesis after Eed loss. Furthermore, genetic deletion of Eed impaired injury induced SVZ proliferation and emigration. These data reveal a novel epigenetic regulated pathway and suggest an essential role for Eed in SVZ homeostasis and injury.

Key words: adult neurogenesis, epigenetics, PRC2, subventricular zone

Introduction

The postnatal/adult subventricular zone (SVZ) harbors the largest population of neural stem cells (NSCs) in the brain (Ihrie and Alvarez-Buylla 2011). SVZ NSCs self-renew slowly throughout life and generate rapidly dividing transit amplifying progenitors (TAPs), which give rise to neuroblasts that migrate to the olfactory bulbs (OB) in the rostral migratory stream (RMS). SVZ lineage progresses along a gradual continuum (Llorens-Bobadilla et al. 2015) and cells can be distinguished by morphology, position, marker expression, and behavior (Doetsch et al. 1999; Mamber et al. 2013). The SVZ is an excellent system for uncovering developmental mechanisms and has therapeutic relevance, since progenitors migrate to injuries and provide neuroprotection (Pluchino et al. 2003; Young et al. 2011). SVZ NSCs can also become harmful when disruption of tightly regulated SVZ cell cycles leads to gliomagenesis (Sanai et al. 2005; Bardella et al. 2016). Therefore, understanding what mechanisms regulate activation of stem cell division and neurogenesis in SVZ cells is a fundamental developmental question which could yield therapeutic approaches.

Multiple cell cycle activators and inhibitors sustain SVZ NSC quiescence and facilitate a “ready for activation” state (Cheung and Rando 2013). The cyclin-dependent kinase inhibitor (CDKI) Cdkn1a encodes p21, which counteracts the cyclin-dependent kinases, Cdk1 and Cdk2. SVZ NSCs express p21, which is required for their quiescence (Kippin et al. 2005; Porlan et al. 2013). Cdkn2a encodes the CDKIs p16Ink4a and p19Arf (Kamijo et al. 1997), which also limit SVZ NSC self-renewal (Nishino et al. 2008). Epigenetic mechanisms regulate embryonic and adult NSC cell cycle length, stem cell activation and lineage progression via CDKIs. The epigenetic Polycomb repressive complex 1 (PRC1) component Bmi1 represses p16Ink4a, p19Arf and p21, to promote...
proliferation in the adult SVZ (Molofsky et al. 2003; Fasano et al. 2007). Since most SVZ NSCs are quiescent in vivo, these studies may have characterized PRC2 functions in TAPs rather than in NSCs. Yet, p21 is expressed in SVZ NSCs in vivo and is important for NSC quiescence and maintenance (Porlan et al. 2013). The question of how CDKI expression is balanced in different cells of the SVZ lineage is important but not yet resolved.

PRC2-mediated gene silencing controls developmental events and adult tissue maintenance through histone methylation (H3K27me3). PRC2 consists of the core protein Eed (Embryonic ectoderm development), and other subunits including the methyltransferase Ezh2 (Enhancer of zeste homolog 2), Suz12 (Suppressor of zeste 12 Homolog), and RbAp46/48 (Retinoblastoma protein associated protein 46/48) (Margueron and Reinberg 2011). PRC2 represses cell-type specific genes during development, and loss of Ezh2 in early embryogenesis disrupts the temporal order of cerebral cortex neuron generation and is necessary for adult neurogenesis (Pereira et al. 2010; Hwang et al. 2014). Although some biochemical features of Eed have been elucidated, its function in neural development and adult stem cells is unidentified. This is important, however, because Eed is an indispensable PRC2 subunit —when it is eliminated, PRC2 can no longer function. In contrast, the Ezh2 homolog Ezh1 can compensate for Ezh2 loss and removing Eed expression eliminates this confounding factor (Shen et al. 2008; Xie et al. 2013).

We first show that Eed is expressed by NSC, upstream in the SVZ lineage compared with Ezh2. We tested Eed function with conditional knockouts, knockdown and overexpression in vivo and in vitro and show it is necessary for SVZ NSC quiescence, maintenance, and neurogenesis. Unlike Ezh2, Eed suppresses Gata6, a transcription factor. Eed loss and increased Gata6 decreases p21 protein levels. These findings suggest that Eed regulation of Gata6 fine-tunes p21 levels and that this is essential for NSC activation and neurogenesis.

Materials and Methods

Mice

Glast-CreERT2 (Mori et al. 2006), Eedfl/fl mice (Xie et al. 2013), and Ezh2fl/fl (Su et al. 2003) were maintained on a C57BL/B6 background. For Cre induction in postnatal pups, 2 protocols were used: either 0.2 mg/g of tamoxifen (Sigma) was injected subcutaneously at P1 or pups were injected with 0.2 mg/g of tamoxifen at P1 followed with daily injections in the lactating mothers (150 mg/kg) for 5 days. For Cre induction in adult mice, 150 mg/kg/d tamoxifen was injected intraperitoneally for 5 days. Littermates or pups were injected with 0.2 mg/g of tamoxifen at P1 followed by daily injections in the lactating mothers (150 mg/kg) for another 2 days. For Cre induction in adult mice, 150 mg/kg/d tamoxifen was injected intraperitoneally for 5 days. Littermates were used as controls. To label fast proliferating cells, BrdU (Sigma, 50 mg/kg) was injected intraperitoneally 2 h before euthanasia. For label-retaining cells, BrdU (50 mg/kg) was injected every 12 h for 5 days and mice were euthanized 6 weeks after the last injection. All animal procedures were carried out with Oxford University Research Ethics Committee approval in accordance with the Animals (Scientific Procedures) Act of 1986 (UK).

Postnatal Electroporation

Electroporation was performed as published (Boutin et al. 2008; Chesler et al. 2008). Constructs for electroporation were prepared with Endofree Plasmid Maxi Kit (Qiagen). 0.1% Fast Green was added to plasmids for tracing ventilator injection. P0–P2 pups were anesthetized by hypothermia. A 1–2 pl plasmid solution was injected into the lateral ventricle via a pulled glass capillary. Pups were then subjected to electrical pulses (100 V, 50 ms ON with 850 ms intervals for 5 cycles) by tweezer electrodes coated with conductive gel. The dorso-lateral and striatal SVZ was primarily targeted. After surgery, pups were kept at 37°C in a warming box for recovery and then returned to the dam. Brains were collected at the indicated time points.

Neurosphere Culture

SVZ tissues were dissected and dissociated with Accutase (Sigma) at 37°C. Cells were then cultured in Neurobasal medium (Gibco) containing B27 (Gibco), N2 (Invitrogen), 1-glutamine (Invitrogen), Pen Strep (Invitrogen), EGF (20 ng/mL, Sigma), and FGF-2 (20 ng/mL, R&D). For bulk culture, cells were plated at 1 x 10^5/mL in 6-well plates (non-TC, BD). For clonal density culture, cells were seeded at 10 μL in 96-well plates (non-TC) and spheres were counted at 7–8 days. For differentiation, 1 x 10^5 cells were seeded per well without growth factors in 8-well chamber slides (Nunc) coated with poly–c–lysine (Sigma) and laminin (Invitrogen). Cells were fixed for immunocytochemistry 7 days later.

ChIP

ChIP was performed as described (Shen et al. 2008, 2009). Rabbit antibodies against H3K27me3 (Millipore 07-449) and mouse antibodies against Eed (Millipore 05-1320) were used for immunoprecipitation. DNA was purified with PCR purification kit (Qiagen) and subjected for quantitative PCR with SYBR Green. Relative enrichment was normalized to the input and the genomic background with the intron of actin as the internal control. The primer sequences are listed in Supplemental Experimental Procedures.

RT-PCR

Total RNA was obtained from samples with RNasey Mini Kit (Qiagen) following the manufacturer’s instructions. Genomic DNA was removed with on-column DNase set (Qiagen), and reverse transcription was performed by SuperScript III RT kit (Invitrogen). The cDNA was subjected to real-time quantitative PCR with SYBR Green PCR Master Mix (Kapa Biosystems or Invitrogen). The target mRNAs were normalized relative to 18S ribosomal RNA. The primer sequences are listed in Supplemental Experimental Procedures.

Cerebral Cortex Lesions

Cortical injuries were generated as described (Goings et al. 2002, 2004). Briefly, 2–3-month-old mice were anesthetized with Ketamine/Xyloazine. The left frontoparietal cerebral cortex from Bregma to 1.2 mm anterior to Bregma was aspirated with a fire-polished glass Pasteur pipette connected to a mild vacuum. Mice were monitored postsurgery and euthanised at the indicated time.

Constructs and Lentivirus Production

pCAG-Cre-IRES2-EGFP (26646), pCAG-IRES2-EGFP (pCAG-G, 11159), pLKO.1-puro (8453), pCAG-1-Gata6 (51929), and p21 shRNA (25868) were from Addgene (Liang et al. 2001; Stewart et al. 2003; Matsuda and Cepko 2004; Woodhead et al. 2006; Fasano et al. 2007). Eed shRNA construct was a kind gift from Prof. Neil Brockdorff (Cooper and Brockdorff 2013). siRNAs against firefly luciferase or Gata6 were purchased from Sigma. pLKO.1-based lentivirus was packaged as previously published (Tavares et al. 2012). Briefly, HEK293FT cells were transfected with pLKO.1, psPAX2 and pMD2.G constructs and fed with fresh packaged virus.
Supernatants were collected 48 and 72 h later and centrifuged at 25,000 rpm, 4°C for 100 min to concentrate the virus.

Western Blot

Proteins were run in 10% SDS-PAGE gel at 100 V and transferred into PVDF membrane at 100 V for 90 min. The membrane was blocked in 5% skim milk in PBST (PBS+0.1% Tween-20) and incubated with primary antibody at 4°C overnight. On the second day, the membrane was washed in PBST and incubated with the secondary antibody for 1 h. The membrane was washed again in PBST, and chemiluminescence was developed with Lumigen PS-3 kit. All the western blots have been repeated at least 3 times and representative images are shown in the figures.

Immunohistochemistry, Image Acquisition and Quantification

Thirty micrometer-thick free-floating sections were incubated with indicated primary antibodies overnight at 4°C. Alexa conjugated secondary antibodies (Invitrogen) were then used for immunofluorescence detection. For BrdU, Ezh2, and Eed immunostaining, sections were treated with 2 M HCl for 1 h at 37°C for antigen retrieval. Sections were then washed with PBS and processed for blocking. DAPI (Sigma) was used for nuclei staining before mounting. Primary antibodies used are listed in the Supplemental Experimental Procedures. Z-stack images were taken with confocal microscopy (Zeiss 710) to confirm colocalization. At least 4 layers with 2 μm intervals between images were acquired for analysis. The images were processed with Image J. For olfactory bulb quantification, evenly spaced...
10 sections per animal were analyzed to cover anterior to posterior OB. For percentage cell quantifications, at least 100 cells were counted but frequently several hundred were quantified.

**Results**

**Eed is Expressed in SVZ Stem and Progenitor Cells**

We first examined PRC2 component expression patterns. Western blots showed core PRC2 protein, Eed expression in several brain areas at P4 as well as in SVZ-derived neurospheres (Fig. 1A). In 6-week-old brains, Eed immunofluorescence was detected in the SVZ, the striatum and cerebral cortex (Fig. 1B). All DAPI+ nuclei examined in the SVZ expressed Eed, indicating it is expressed in all cells of the lineage. Importantly, both BrdU+ label-retaining cells (LRC) and Ki67+ actively proliferating cells expressed Eed in the SVZ (Figs 1C and S1A and S1B). The former result suggested quiescent NSCs express Eed. EGFR+/GFAP+ NSCs, EGFR+/GFAP– TAPs and EGFR-/GFAP+ niche astrocytes were all Eed+ (Fig. 1D, E). In the RMS and OB, Dcx+ neuroblasts and granular layer cells expressed Eed (Fig. 1F, G). Similar to Eed, PRC2 epigenetic mark H3K27me3+ nuclei were throughout the brain (Fig. 1H). GFAP+ astrocytes in the corpus callosum and SVZ (Fig. 1I, J), as well as S100β+ niche astrocytes and ependymal cells (Fig. 1K) and Dcx+ neuroblasts (Fig. S1C) were all H3K27me3+.

Unlike Eed, Ezh2 immunoreactivity was restricted to the SVZ (Fig. 1L) and RMS with few Ezh2+ cells elsewhere. However, only 6.45 ± 2.75% of GFAP+/Sox2+/SVZ NSCs were Ezh2+ (Fig. 1M). Similarly, only 7.58 ± 1.96% of SVZ LRC had Ezh2+...
nuclei (Fig. 1O). The majority \((97.37 \pm 1.96\%)\) of actively proliferating BrdU+ cells were Ezh2+ (Fig. 1N), which was confirmed with phospho-histone 3 (PHil3) and Ezh2 co-staining (Fig. S1D). Ezh2+ nuclei were mainly detected in doublecortin+ (Dcx) neuroblasts in the RMS and olfactory bulb (OB) (Fig. S1D). Our data showing Dcx+ neuroblasts express Ezh2 is consistent with the literature (Hwang et al. 2014). These data strongly suggest that Ezh2 is expressed in TAPs and neuroblasts but not in SVZ NSCs. The Allen Brain Atlas shows Ezh1 is expressed in cells or regions lacking Ezh2, which would explain the broad distribution of H3K27me3. Ezh1 mRNA was detected in multiple brain regions including the SVZ (Fig. 1P). Unfortunately there are no reliable Ezh1 antibodies for immunochemistry. However, the transcriptome of FACSorted SVZ cells shows an enrichment of Ezh1 in quiescent SVZ stem cells (type B cells) (Codega et al. 2014). This is consistent with the general idea that Ezh1 is expressed by quiescent stem cells while Ezh2 is expressed by active stem cells (Cheung and Rando 2013). Overall, our data suggests a model in which Eed is expressed in quiescent NSCs as well as after they become activated and enter the neurogenic program (Fig. 1Q). In contrast our data suggest Ezh2 is not expressed in SVZ NSC.

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Eed is required for postnatal OB neurogenesis. (A–D) Eed cKO schedule as in Fig. 2F. (A) Immunostaining for Dcx in the RMS. (B) Quantification of surface area occupied by Dcx+ cells in the RMS. \(N = 3-4\). (C) Co-staining for BrdU and NeuN in the granular layer of the olfactory bulb. (D) Quantifications of NeuN+/BrdU+ double labeled cells in the OB. \(N = 3-4\). (E) Diagram of postnatal electroporation schedule used in F–I. (F) GFP immunofluorescence in SVZ 5 days postelectroporation. (G–I) Confocal images and quantifications of GFP+ cells in the OB 5 days (G) or 14 days (H) postelectroporation. \(N = 5-6\). Data are shown as mean ± SEM and analyzed by two-tailed Student t-test: *\(P < 0.05\), **\(P < 0.01\), ***\(P < 0.001\). Scale bars represent 50 μm in (A, C), 100 μm (F–H).
Eed is Required for SVZ Neural Stem Cell Activation

We next examined Eed function in the SVZ. We crossed Glast-CreERT2 mice with Eed<sup>fl/fl</sup> mice (Mori et al. 2006; Xie et al. 2013) to remove Eed function in postnatal SVZ NSCs via tamoxifen (TMX) administration in P0–P1 pups (Materials and Methods). To boost Cre-recombination efficiency, we gave TMX to pups and to lactating dams. Genotyping and RT-PCR confirmed Eed cKO (Fig. S1E, F). At P14, the number of PH3+ cells was unchanged in Eed cKO SVZs (Fig. S1G). However, by P24 fewer GFAP+ cells were proliferating (Ki67+ (SVZ NSCs and possibly niche astrocytes) in Eed cKOs (Fig. 2A, B). Further analysis of Eed loss revealed 45% reduction in GFAP+/Sox2+NSCs that expressed Ki67 (Fig. 2A, C), suggesting fewer stem cells are entering the cell cycle. Ezh2 cKO SVZs did not exhibit such a decrease in GFAP+/Sox2+NSCs that expressed Ki67 (Fig. S2G), supporting the observation that NSCs do not express Ezh2.

Figure 4. Loss of Eed selectively activates PRC2 targets. (A–C) ChIP-qPCR from the P4 (A) or P56 (B) SVZ with anti-H3K27me3 or IgG antibody. N = 3. (D) qPCR analysis of PRC2 target gene expression in the P14 SVZ of littermate controls and Eed cKO mice. N = 3. (E) ChIP-qPCR using H3K27me3, Eed or IgG antibody of neurospheres cultured from postnatal SVZ. N = 3. (F–H) ChIP-qPCR for H3K27me3

This data is shown as mean ± SD (A–C, F–H) or mean ± SEM (D, E, J, K) and analyzed by two-tailed Student t-test. *P < 0.05, **P < 0.01, ***P < 0.001.
Stem cell activation defects or loss of stemness and acquisition of niche astrocyte phenotype could decrease NSC proliferation. The latter was assessed by coimmunostaining GFAP+ cells with S100β, a marker of niche astrocytes (and ependymal cells) (Raponi et al. 2007). Seventy-five percent more GFAP+ cells expressed S100β in the SVZ after Eed loss (Fig. 2D, E), suggesting loss of stemness and NSC differentiation into niche astrocytes. We next sought to determine if a longer survival time of P81 after Eed cKO would have similar results. For label-retaining analysis, BrdU was given from P35 to P39, and brains collected 6 weeks later (Fig. 2F). A smaller percentage of BrdU LRC cells in Eed cKO SVZ expressed immunodetectable H3K27me3.

Figure 5. Eed and Gata6 fine-tune p21 protein levels for neurogenesis. (A, B) Immunostaining and quantification of PHi3 in GFP+ cells derived from neurospheres transfected with either GFP or Gata6 cDNA. The arrows indicate PHi3+/GFP+ cells. N = 3. (C, D) Immunostaining and quantification of TuJ1 in differentiating cells from 7 DIV neurospheres transfected with either GFP or Gata6 cDNA. N = 3. (E, F) Immunocytochemistry and quantification of Dlx2 in GFP+ cells in SVZ of pups electroporated with either GFP or Gata6 cDNA. The arrows indicate Dlx2+/GFP+ cells. N = 3. (G, H) Immunostaining and quantification of Mash1 in GFP+ cells in SVZ of pups electroporated with either GFP or Gata6 cDNA. The arrows indicate Mash1+/GFP+ cells. N = 3. (I, J) Immunostaining and quantification of Ki67 in GFP+ cells in SVZ of pups electroporated with either GFP or Gata6 cDNA. The arrows indicate Ki67+/GFP+ cells. N = 3. (K, L) qPCR (K) and western blot (L) analysis of p21 expression in neurospheres transfected with either GFP or Gata6 cDNA. N = 3. (M) Western blot analysis of p21 in P15 SVZ tissues from littermate control or Eed cKO mice. N = 3. (N, O) p21 protein stability time course in neurospheres transfected with either GFP or Gata6 cDNA and treated with cycloheximide (CHX) added 48 h after nucleofection. N = 3. Data are shown as mean ± SEM and analyzed by two-tailed Student t-test. *P < 0.05. Scale bars represent 50 μm in (A, C); and 20 μm in (E, G, H).
Confirming loss of PRC2 function and presence of knockout cells (Fig. S1H, I). Although Eed loss did not significantly affect overall LRC cell numbers (Fig. S1J), more LRCs expressed S100b in Eed cKO mice. This suggests stem cells are less active and the data are consistent with them acquiring a niche astrocyte phenotype (Fig. 2G, H). Importantly, most S100b+/GFAP+ exhibited long basal processes, and these cells as well as S100b+/LRC cells are primarily within the SVZ body rather than lining the ventricle (Fig. 2D, G) indicating they are not ependymal cells. Examination of ependymal cells lining the ventricles shows that Eed cKO does not change S100b expression (Fig. 2G).

We next examined primary SVZ neurospheres, which can be formed by SVZ NSCs and TAPs. Lentiviral Eed knockdown reduced the number of neurospheres formed 4 days in vitro (div) (Fig. 2). Both P15 and P38 Eed cKO SVZs yielded 50% fewer clonal neurospheres (Fig. 2), K, L). Together, these data show Eed is required for NSC activation.

Eed is Required for Postnatal OB Neurogenesis

Next, we examined OB neurogenesis in P81 Glast-CreERT;EedLoxIE mice given TMX as neonates and BrdU between P35-39 (as in Fig. 2F). Eed cKOs had reduced RMS surface areas (Fig. 3A). Similar experiments in Glast-CreERT;Ezh2F/F mouse also resulted in decreased neurogenesis (Fig. S2 and Supplementary Results), comparable to a previous study (Hwang et al. 2014).

Glast-CreERT2-mediated recombination occurs in SVZ NSCs as well as in astrocytes outside the SVZ (Mori et al. 2006). Thus it was possible that Eed cKO in non-SVZ astrocytes affected the number of OB NeuN+/BrdU+ cells. For this control and to confirm our results, we postnatally electroporated SVZ cells in P1 pups with Eed shRNA or empty control constructs together with pCAG-IRES-GFP (Fig. 3J). Five days postelectroporation, GFP immunofluorescence in the SVZ revealed similarly large numbers of electroporated SVZ radial glia-like NSCs in control and Eed KD suggesting similar electroporation efficiency in the 2 groups (Fig. 3J). However, at this time point, the density of OB GFP+ cells decreased more than 35% in the knockdown compared with control mice (Fig. 3G, I). Neither oligodendrocyte differentiation nor cell death accounted for these differences (Figure S3A, S3B and S3C). By 14 dpe, GFP+ cells entered the OB granular layer and displayed mature morphology in controls (Fig. 3H). GFP-labeled cells were found in all layers of control OBs 14dpe but were predominantly in the granule layer (Fig. 3H). After Eed KD fewer GFP+ cells were seen in all layers. Eed knockdown caused a greater than 60% reduction in GFP+ cell density in the OB taken as a whole (Fig. 3H, I). The 2 complementary techniques of conditional knockdown and shRNA knockdown revealed that Eed promotes postnatal OB neurogenesis.

Loss of Eed Selectively Derepresses PRC2 Targets in Neural Stem Cells

Because Eed and Ezh2 are involved in histone 3 lysine 27 trimethylation (H3K27me3), we determined if the H3K27me3 repressive mark was associated with PRC2 targets in the SVZ. Chromatin immunoprecipitation (ChIP) analysis of P4 and P56 SVZ showed H3K27me3 enrichment on multiple classic PRC2 targets, including Gata6, Hoxa11, Ngxn1, Sox17, and Olig2 (Fig. 4A–C). Cdkn2a expresses 2 transcripts, p16nk4a and p19arf (Kamijo et al. 1997). PRC2 represses Cdkn2a during development (Pereira et al. 2010; Hidalgo et al. 2012; Xie et al. 2013; Hwang et al. 2014), and both Eed and Ezh2 conditional knockout (Materials and methods and Supplemental results) increased SVZ Cdkn2a expression (Fig. 4D, E). Previous work showed Ezh2 KO only derepressed p16nk4a but not p19arf in the SVZ (Hwang et al. 2014). In contrast, we found, that Eed loss increased both p16nk4a and p19arf expression (Fig. 4D). Eed cKO only increased expression of Gata6 among other PRC2 targets and Cdkn2a (Fig. 4D and S4A). In contrast Ezh2 cKO did not alter Gata6 expression in the SVZ (Fig. 4J).

SVZ heterogeneity implied the above results could have occurred in multiple cell types. We looked for PRC2 and Eed targets in NSC and TAP-enriched self-renewing neurospheres. ChIP analysis revealed H3K27me3 and Eed occupancy on Gata6, Hoxa11, Ngxn1, and Sox17 (Fig. 4F, G, H), similar to the SVZ. Eed KD correspondingly decreased H3K27 occupancy on these targets (Fig. 4F). Neurospheres cultured from EedLoxIE or Ezh2LoxIE pups were nucleoected with either GFP control or GFP-Cre plasmids, at 40–50% efficiency causing a 30–40% decrease in RNA (Fig. 4J, K). Gata6 expression increased more than 2-fold in the GFP-Cre neurospheres 7 days post transfection (Fig. 4J). Expression was selectively increased, Gata6, Cdkn2a, and Ngxn1 increased but Sox17 and Hoxa11 did not (Fig. 4J). In contrast to Eed cKO, Ezh2 cKO in vitro only increased expression of Cdkn2a but did not affect Gata6 or other PRC2 targets (Fig. 4K). Our in vivo and in vitro data indicate Eed selectively suppresses Gata6 in contrast to Ezh2, which did not affect expression of this transcription factor.

Gata6 Overexpression Limits Neurogenesis Both In Vitro and In Vivo

Gata6 is suppressed via CpG island hypermethylation in glioblastoma (Martinez et al. 2009; Skirud et al. 2012), suggesting it normally limits proliferation. The role of Gata6 in neural development was unknown, but our results also suggested it limits mitosis. Gata6 overexpression (OE) in cultured NSCs reduced the percent of nucleoected (GFP+) cells that were PH3+/BrdU+ (M phase of cell cycle) by 30% (Fig. 5A, B). Gata6 OE did not alter Caspase-3+/GFP+ cell numbers in vitro (Fig. S4B), suggesting no effect on cell death. Gata6 OE also had no effect on nestin or GAF immunostaining at 2DIV (Fig. S4C, D, E). In differentiation assays without growth factors Gata6 OE decreased the percent of cells that entered mitosis (Fig. S4F), together suggesting a negative effect on neurogenesis and/or neuronal differentiation. In contrast, differentiation assays showed the percent of cells that expressed GAF after Gata6 OE was not significantly different compared with controls (Fig. S4F), suggesting Gata6 does not affect fate choice in SVZ NSC.

To confirm that Gata6 limits SVZ neurogenesis, we coelectroporated Gata6 cDNA with GFP in –P1 pups. The majority of SVZ cells expressing Mash1 and Dlx2 are TAPs (Mamber et al. 2013). At 3dpe, fewer GFP+ cells in the Gata6 OE group expressed Dlx2 (Fig. S5E, F) or Mash1 (Fig. S5G, H) in the SVZ. Of note, neither Gata6 OE nor the loss of Eed directly affected Mash1 transcription (Fig. S4G, H) suggesting they inhibited lineage progression from NSC to TAPs rather than reducing marker expression. Similar to the reduction of proliferation in vitro, Gata6 OE reduced the percentage of electroporated cells that expressed Ki67 (Fig. S1J, J). These novel results show the transcription factor Gata6 is sufficient to restrict proliferation and neurogenesis in SVZ cells.
Elevated Gata6 Reduces p21 Protein Stability

p21 maintains SVZ NSCs (Kippin et al. 2005; Marques-Torrejon et al. 2013; Porlan et al. 2013), so we hypothesized that high levels of Gata6 expression limit SVZ neurogenesis through p21. Gata6 OE in vitro did not significantly alter p21 mRNA expression (Fig. 5K), but p21 protein levels were decreased (Fig. 5L). Eed cKO decreased p21 protein in vivo, further suggesting functional interaction with Gata6 (Fig. 5M). MicroRNA clusters (miR-224, miR-291, miR-303b, etc.) and RNA binding proteins (Msil, Fxr1, etc.) regulate p21 expression post-transcriptionally (Dolezalova et al. 2012; Freeman and Espinosa 2013), but Gata6 OE in neurospheres did not alter miRNA (8 examined) and RNA binding protein expression (11 examined) (Fig. S5A, B), suggesting they do not participate in this context. Stabilization and degradation of p21 protein is important in tumorigenesis (Abbas and Dutta 2009) and to determine protein stability, we examined p21 levels in neurospheres treated with the protein synthesis inhibitor (CHX). Forty-eight hrs after nucleofection, cells were treated with CHX and analyzed 0–2.5 h later. Gata6 OE significantly decreased p21’s half life after CHX treatment (Fig. 5N, O). The protease inhibitor MG132, increased p21 levels relatively more in Gata6 OE than control cells (Fig. S5C) suggesting Gata6 regulates p21 protein stability.

Eed Regulates SVZ Neurogenesis Partially Through Gata6 and p21

If Gata6 is necessary for Eed regulation of SVZ neurogenesis, Gata6 repression should rescue neurogenesis deficits caused by Eed loss. To test this, P1 pups were electroporated with an Eed

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**Figure 6.** Gata6 knockdown and p21 overexpression partially rescues Eed loss in NSCs. (A–C) Quantifications of Dlx2 (A) or Mash1 (B) or Ki67 (C) in GFP+ cells in SVZ of pups electroporated with different plasmids as listed in the figure. N = 3–5. (D) Western blot analysis of p21 in neurospheres after Eed and Gata6 double knockdown. (E, F) Immunostaining and quantification of β III-tubulin as detected with the Tuj1 antibody in NSC derived from neurospheres transfected with either empty vector or p21 shRNA. N = 3. (G) Quantification of Tuj1+ cells in NSC derived from neurospheres transfected with control, Eed KD, empty vector (EV), or p21 OE plasmids as listed. N = 3. (F) A schematic diagram of molecular pathways regulated by Eed and Enh2 in NSCs. Data are shown as mean ± SEM and analyzed by two-tailed Student t-test or one way ANOVA. *P < 0.05, **P < 0.01. Scale bar represents 20 μm in (E).
shRNA construct alone or along with either a luciferase siRNA (siFluc) as a negative control or a Gata6 siRNA that was confirmed in vitro (Fig. S5D). The percentage of GFP+ cells that were Dlx2+, Mash1+, or Ki67+ was comparable between the Eed shRNA and Eed shRNA plus siFluc groups (Fig. 6A–C), establishing a baseline for siGata6 comparison. 5dpe the percentage of electroporated GFP+ cells expressing Dlx2 decreased after Eed KD (Fig. 6A). When pups were co-electroporated with Eed shRNA and siGata6 constructs, the percentage of GFP+ cells that were Dlx2+ increased statistically compared with the siFluc group showing a partial but significant rescue (Fig. 6A). Similar to Dlx2, the percent of electroporated GFP+ cells expressing Mash1 decreased in the Eed shRNA group compared with controls (Fig. 6B). When pups were coelectroporated with Eed shRNA and siGata6 constructs, the percent of GFP+ cells that were Mash1+ increased significantly by 35% (Fig. 6B). Since we showed that Eed orchestrates p16/p19 and p21 cell cycle inhibitor expression in the SVZ (Figs 4 and 5), we examined whether Gata6 KD could rescue the decreased proliferation induced by Eed KD. Confirming our cKO results, Eed KD decreased the percent of electroporated GFP+ cells that were Ki67+ by 43% (Fig. 6C). However, double knockdown of Eed and Gata6, did not increase the percent of GFP+/Ki67+ suggesting Gata6 KD was insufficient to rescue proliferation after Eed KD.

Figure 7. Eed is required for injury-induced SVZ proliferation. (A) Immunostaining of GFAP and BrdU in the cortical lesion. (B) Immunostaining of GFAP and Eed in reactive astrocytes in the injured cerebral cortex. Arrows indicate astrocytes undergoing cell division. (C) Western blot of Eed expression in the ipsilateral and contralateral cerebral cortex. (D) Timeline of experimental design. (E–G) Immunostaining and quantifications of GFAP, Sox2, and Ki67 in the injured cerebral cortex. N = 3. (H, I) Immunostaining and quantifications of Ki67 in the SVZ. N = 3. (J–L) Immunostaining and quantifications of Dcx in the corpus callosum. N = 3. Data are shown as mean ± SEM and analyzed by two-tailed Student t-test or one way ANOVA. *P < 0.05, **P < 0.01. Scale bars represent 500 μm in (A), 10 μm in (B), 50 μm in (E), 30 μm in (H), and 100 μm in (J).
We next confirmed Gata6 is necessary for Eed loss-of-function induced p21 protein decrease. Eed KD in neurospheres reduced p21 protein, confirming our Eed cKO effects (Fig. 6D). Eed and Gata6 double KD in neurospheres restored p21 levels (Fig. 6D) indicating Gata6 was indeed necessary for the effect of Eed loss on p21. To confirm if Eed loss decreases neurogenesis via p21, we carried out p21 KD in neurospheres. p21 KD reduced the percentage of nucleofected cells expressing β-III tubulin compared with controls (Fig. 6E, F), which was consistent with the literature (Porlan et al. 2013) and suggested reduced neurogenesis and/or differentiation. Moreover, p21 OE rescued the percent of β-III tubulin+ cells in Eed KD of NSC cultures (Fig. 6G). These data establish a mechanism by which the PRC2/Eed complex regulates Gata6 and p21 to reneurogenic competence in NSCs, as schematized in Figure 6H.

**Eed is Dispensable for Parenchymal Reactive Astrocyte Proliferation After Injury but Necessary for Injury-Induced SVZ Activation**

Parenchymal astrocytes are generally postmitotic, but can become reactive after injury and become neurogenic (Buffo et al. 2008; Sirko et al. 2013). We tested Eed necessity for parenchymal astrocyte proliferation after a cerebral cortex injury (Fig. 7A) that activates SVZ neurogenesis and neuroblast emigration (Széle and Chesselet 1996; Sundholm-Peters et al. 2005). Peri-lesion GFAP+ reactive astrocytes were Eed+/− (Fig. 7B) but overall Eed levels were similar in ipsi- and contralateral cortex (Fig. 7C). To test Eed necessity for cortical astrogliosis after injury, we compared 5-week old Glst-CreERT2; Eedfl/fl (cKO) with Eed+/− (control) mice. Both groups received tamoxifen and then cortical lesions 4 weeks later (Fig. 7D). Peri-lesion astrocytes expressed high levels of GFAP and Ki67 and many expressed Sox2 (Fig. 7F). Control and cKO mice had similar numbers of Ki67+/GFAP+ proliferative astrocytes and Sox2+/GFAP+ cells (Fig. 7F, G), suggesting Eed is dispensable for parenchymal astrocytic proliferation.

To determine if Eed mediates SVZ function after injury, increasing proliferation, we compared Eed cKOs with controls. As expected, the SVZ ipsilateral to the lesion contained more proliferative cells (Ki67+) than the contralateral SVZ in control mice 5 days post injury (Fig. 7H, I). After Eed loss, both ipsilateral and contralateral SVZs had significantly fewer Ki67+ cells than controls. More importantly, cell proliferation was not statistically significantly increased after Eed cKO in the ipsilateral SVZ (Fig. 7H, I). Caspase-3 immunohistochemistry indicated this was not due to enhanced cell death (data not shown). SVZ neuroblasts migrate to the OB, but cortical injury redirects some into the corpus callosum and cortex (Goings et al. 2004; Saha et al. 2013). Fewer neuroblasts were found in the corpus callosum in Eed cKO compared with control mice (Fig. 7, K) consistent with the proliferation deficiency. After injury approximately 20% of all Dcx+ cells were found in the CC in Eed cKO compared with 44% in controls, suggesting Eed necessity for a robust response to cortical injury (Fig. 7J).

**Discussion**

The most important finding here is that Eed loss significantly reduces SVZ NSC activation and neurogenesis. It does so by increasing Gata6 expression, which reduces p21 protein levels. We confirmed these findings in vivo and in vitro with multiple knockdown, overexpression and rescue studies. Gata6, however, is not affected by EzH2, which suggests PRC2 works via multiple distinct molecular pathways. Eed is also necessary for SVZ proliferation but not for parenchymal astrocytosis.

**Eed Regulates SVZ NSC Proliferation and Neurogenesis via Gata6 and p21**

Several experiments demonstrated Eed loss reduces SVZ NSC proliferation and neurogenesis. We crossed floxed Eed (Xie et al. 2013) and Glst-CreERT2 mice (Mori et al. 2006) to knock-out Eed from SVZ NSC and their progeny. Though most NSC have Eed deleted after Tamoxifen, a few NSC likely “escaped” Cre recombination and thus retained normal Eed levels. This was consistent with incomplete loss of H3K27me3 in LRC-BrdU cells. Additionally, the SVZ is heterogeneous and some SVZ neural stem cells may not express GLAST. Nevertheless, conditional knockout of Eed significantly decreased the percent of proliferating SVZ NSCs. Interestingly, Eed conditional knockout also caused more GFAP+ and label-retaining cells to express S100β. RNA-Seq showed quiescent SVZ NSCs express >200-fold more S100β than active NSCs (Codega et al. 2014). The percentage of SVZ LRCs that express S100β significantly increased after Eed cKO, thus SVZ LRCs are likely to become quiescent after Eed loss. Finally, Eed conditional knockout and knockdown reduced newborn neuronal numbers in the OB suggesting reduced neurogenesis. We believe that the reduction in newborn neuroblasts in the OB after Eed loss is because fewer NSC become activated thereby inhibiting lineage progression to neuroblasts. Alternatively, Eed/Gata6/p21 may function directly in neuroblasts to reduce their numbers in the OB. Which stage of SVZ lineage does Eed affect? Eed knockdown with electroporation initially targets NSC (Boutin et al. 2008). Eed is expressed in all SVZ cells, and Eed and H3K27me3 expression were confirmed in label-retaining NSC, GFAP+/EGFR+/NSC, proliferative TAPs, and Dcx+ neuroblasts. This is consistent with Eed affecting NSC and then also functioning at subsequent steps in the SVZ lineage.

Eed loss increased Gata6 expression. Gata6 is a transcription factor essential for heart and primitive gut development (Burch 2005). Gata6’s role in mammalian neurogenesis was unknown but hypermethylation represses Gata6 in glioblastomas (Martinez et al. 2009; Skirute et al. 2012), suggesting Gata6 normally limits proliferation. Accordingly, Gata6 OE decreased SVZ proliferation in vivo and in vitro. Further studies showed Gata6 OE decreased TAP and neuroblast marker-positive cells. Gata6 OE could have increased Gata6 more than in Eed cKO and not reflected true physiological fluctuations. However, Gata6 OE may have a “ceiling effect”; the response to increased expression may have a threshold. These caveats aside, Gata6 knockdown suggest it is necessary for reduced neurogenesis after Eed loss—it partially rescued Eed loss effects. p16/p19 expression increased after Eed loss and this may have restricted general proliferation, limiting the extent of the rescue.

Eed loss-of-function increased p16/p19 (Cdkn2a) but did not significantly alter p21 (Cdkn1a) mRNA levels. In contrast, Eed cKO in vivo and KD in vitro reduced p21 protein levels. Similarly, Gata6 OE reduced p21 protein levels. MicroRNAs and RNA binding protein expression suggesting they do not affect p21 stabilization; therefore, how Gata6 destabilizes p21 is unclear. Nonetheless, Gata6 was required for Eed KD to decrease p21 protein levels since Gata6 loss blocked the effect. As well, Eed KD and p21 KD similarly reduced the number of neuroblast marker-positive cells and p21 OE rescued the Eed KD effect.
Thus, Eed normally represses Gata6, which maintains p21 protein. Together with Eed’s effect on p16/p19 this mechanism regulates SVZ NSC maintenance, proliferation and neurogenesis. Since post-transcriptional regulatory processes fine-tune p21 protein levels, p21 may have dose-dependent roles in the SVZ. Most knowledge of p21 function in the SVZ is from constitutive knockouts (Kippin et al. 2005; Marques-Torrejon et al. 2013; Porlan et al. 2013). Nonmutant approaches, such as p21 promoter modulation, and determination of p21 protein levels in stem cell lineages will clarify its functions.

Eed is Necessary for Injury-Induced SVZ Activation but dispensable for Parenchymal Astrocytosis

Increased SVZ proliferation and progenitor emigration towards lesions can improve histological and behavioral outcomes (Jin et al. 2010; Young et al. 2011). Brain injury induces both SVZ astrocyte-like NSC proliferation and parenchymal astrocytosis but the overlap of molecular mechanisms regulating these two processes is unclear. Since epigenetic mechanisms coordinate astrocytic and NSC-mediated brain repair, we examined Eed function after cortical injury. Eed loss inhibited lesion-induced increases in SVZ proliferation and the number of Dcx+ cells between the SVZ and the injury, showing Eed participates in injury-induced SVZ proliferation. As of now, it is unclear if the post injury effects of Eed on SVZ proliferation are Gata6 or p21 dependent. In contrast to its effects in the SVZ, Eed loss did not alter parenchymal astrocytosis after injury, suggesting Eed is dispensable for this process. Our data reveals a fundamental difference in post injury molecular regulation in SVZ astrocyte-like NSCs compared with parenchymal astrocytes.

Eed Versus Ezh2—Distinct Cellular Expression and Molecular Regulation

Eed was expressed by label-retaining cells and GFAP+/EGFR+SVZ cells, implying NSC expression (Mamber et al. 2013). In contrast, SVZ label-retaining cells and GFAP+/Sox2+SVZ cells did not exhibit Ezh2 immunofluorescence indicating it was not expressed by NSC (Mamber et al. 2013). However, Ezh2 may be expressed by NSC below the level of detection. Notably Eed, but not Ezh2, loss increased Gata6 expression. Eed and Ezh2 both repress p16/p19 (Cdkn2a)—our work is concordant with Hwang et al., 2014, showing Ezh2 loss increased p16. Eed and Ezh2 loss-of-function may regulate other distinct genes through selective H3K27me3 targets. Future RNA-sequencing studies could establish these global differences and help place Gata6 and p21 in the context of broad gene regulatory networks.

Conclusion

We show Eed is necessary for stem cell activation and neurogenesis. We delineated a functional role for Eed in regulating Gata6 and thereby maintaining p21 protein levels. Interestingly the PRC2 methyltransferase Ezh2 did not alter Gata6 levels suggesting divergence of Eed and Ezh2 function in the SVZ niche. Our study identifies Eed, Gata6, and p21 as possible molecular targets to manipulate stem cell cycles in injury and cancer.

Authors’ Contributions

B.S. designed and performed the experiments, analyzed data, and wrote the article; E.C. performed the lentiviral assay; A.G. helped with the immunohistochemistry and the relevant quantifications; F.S. designed experiments, analyzed data, and wrote the article.

Supplementary Material

Supplementary material is available at Cerebral Cortex online.

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Notes

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Supplemental results

Ezh2 cKO decreases neurogenesis

We crossed Glast-CreERT2 mice with Ezh2^{fl/fl} mice (Su IH et al. 2003; Mori T et al. 2006) to remove Ezh2 function in postnatal SVZ NSCs via tamoxifen (TMX) administration in P0–P1 pups (Materials and methods). Genotyping and RT-PCR confirmed Ezh2 cKO (Figure S2A and S2B). Ezh2 immunoreactivity was dramatically reduced at P14 in the SVZ of Glast-CreERT2;Ezh2^{fl/fl} mice, (Figure S2C) but we found no obvious change in H3K27me3 staining (data not shown).

We first examined the effect of Ezh2 cKO at P28 (Figure S2D). Ezh2 cKO SVZ had fewer Ki67+ (proliferating cells) and fewer Mash1+ (TAPs) at P28 (Figure S2E and S2F). However the percent of GFAP+/Sox2+ cells that expressed Ki67 was not significantly different after Ezh2 cKO (Figure S2G). Ezh2 CKO mice had slightly weaker Dcx immunostaining in the SVZ (Figure S2H). RT-PCR confirmed significant Mash1 and Dcx decreases at P28 in Ezh2 cKO and Id2 was also reduced (Figure S2I).

We next administered tamoxifen at P0, BrdU from P29 to P33 and analysed neurogenesis at P75 (Figure S2J). Ezh2 loss decreased the number of NeuN+/BrdU+ newborn neurons in OB granular and glomerular layers (Figure S2K, S2L and S2M). Dcx+ neuroblast surface area in the RMS also decreased (data not shown). Caspase-3+ immunoreactivity showed the decreased number of newborn neurons was not due to apoptosis in the RMS/SVZ (data not shown). By P75, the number of PHi3+ (Figure S2N) and Mash1+ (Figure S2O and S2P) SVZ cells in Ezh2 cKOs had returned to control levels. Finally, the total number of LRCs in the SVZ was unaffected in the Ezh2 loss-of-function studies (data not shown), suggesting Ezh2 is dispensable for SVZ NSC maintenance and long-term SVZ NSC label retention. These results are overall consistent with (Hwang WW et al. 2014), which showed loss of Ezh2 function reduces SVZ neurogenesis.
Supplemental Figure Legends:

**Figure S1. Generation of conditional knockout mice**

(A-B) High magnification image of Eed, BrdU and Ki67 costaining in label-retaining cells.

(C) Coimmunostaining of H3K27me3 and Dcx in the OB.

(D) Coimmunostaining of Ezh2, PHi3 and Dcx in the P42 SVZ.

(E) Genotyping for Eed transgene. Wildtype band is shorter than 200 bp and the floxed band is longer than 200 bp.

(F) qPCR analysis of SVZ tissue from P14 SVZ to confirm Eed knockout efficiency.

(G) Quantification of PHi3+ cells in P14 SVZ of control and Eed cKO mice. N=3.

(H-I) Immunostaining and quantifications of BrdU and H3K27me3 in mice for BrdU label-retaining analysis. White arrows indicate H3K27me3+/BrdU+ cells, while magenta arrows indicate H3K27m3-/BrdU+ cells. Time course as in Fig. 2F. N=3.

(J) Quantification of the number of label-retaining cells in the SVZ. N=3.

Data are shown as mean ± SEM and analysed by two-tailed Student t-test. **p<0.01. Scale bars represent 20 µm in (A), (B), (D) and (H), and 40 µm in (C).

**Figure S2. Ezh2 is dispensable for SVZ neural stem cell maintenance**

(A) Genotyping for Ezh2 transgene. The floxed band is 736bp and the deleted band is 625bp.

(B) qPCR analysis of SVZ tissue from P14 SVZ to confirm Ezh2 knockout efficiency.

(C) Immunostaining of Ezh2 or H3K27me3 in control or Ezh2 cKO SVZ (P14).

(D) Schematic of experimental outline used in E-J.

(E-G) Quantifications of Ki67+ (E), Mash1+ (F), and percent of GFAP+/Sox2+ cells that are Ki67+ (G) in P14 SVZ of control and Ezh2 cKO mice. N=3-4.

(H) Immunostaining of Dcx in P28 SVZ of control and Ezh2 cKO mice.
(I) qPCR analysis of P14 and P28 SVZ. Relative expression is normalised to littermate controls. N=3.

(J) A schematic diagram of the experimental design used in K-P.

(K-M) Costaining for BrdU and NeuN, and quantifications of double labelled cells in the OB. N=3-4.

(N) Quantification of PHi3+ cells in P75 SVZ. N=3-4.

(O-P) Immunostaining and quantification of Mash1+ cells in P75 SVZ. N=3-4.

Data are shown as mean ± SEM and analysed by two-tailed Student t-test. *p<0.05, ***p<0.001. Scale bars represent 20 µm in (C), 60 µm in (H), (O); and 100 µm in (K).

Figure S3. Control data for Eed knockdown in vivo

(A-B) Immunostaining and quantification of Olig2 among GFP+ cells in the RMS (5dpe). The arrows indicate Olig2+/GFP+ cells. N=5-6.

(C) Immunostaining of Caspase-3 among GFP+ cells in the RMS (5dpe). The arrow indicates a Caspase-3+/GFP+ cell. N=5-6.

Data are shown as mean ± SEM and analysed by two-tailed Student t-test. Scale bars represent 20 µm in (A) and (C).

Figure S4. The molecular regulation of Eed/Gata6 in SVZ NSC

(A) qPCR analysis of gene expression in P14 SVZ from control and Eed cKO mice. N=3.

(B) Quantification of Caspase-3 among GFP+ NSCs in culture (2 days post nucleofection). N=3.

(C) Immunostaining of Nestin among GFP+ cells in culture after control or Gata6 OE nucleofection.

(D-E) Immunostaining and quantification of GFAP among GFP+ cells in culture. N=3.

(F) Quantification of GFAP in differentiated NSC derived from neurospheres 7 DIV transfected with either GFP or Gata6 cDNA. N=3.
(G) qPCR analysis of Mash1 expression in NSCs 2 days after nucleofection with GFP (control) or Gata6 cDNA. N=3.

(H) qPCR analysis of Mash1 expression in Eed\(^{fl/fl}\) NSCs nucleofected with GFP or GFP-Cre. N=3.

Data are shown as mean ± SEM and analysed by two-tailed Student t-test. Scale bars represent 50 µm in (C), (D).

**Figure S5. Gata6 overexpression regulates p21 protein stability**

(A-B) qPCR analysis of miRNAs and RBPs in NSCs transfected with GFP or Gata6 cDNA (2 days in culture). N=3.

(C) Western blot of p21 in neurospheres transfected with either control GFP or Gata6 cDNA and treated with MG132 for 4hrs. N=3.

(D) Western blot to validate the knockdown efficiency of siGata6 in NSCs in vitro.

Data are shown as mean ± SEM and analysed by two-tailed Student t-test.
Supplemental Table 1. Antibodies for immunohistochemistry

| Antigen | Dilution | Host | Manufacturer Cat. |
|---------|----------|------|-------------------|
| BrdU    | 1:400    | mouse | Dako M0744        |
| BrdU    | 1:500    | sheep | Abcam Ab1893      |
| Caspase-3| 1:1000  | rabbit | Cell Signaling 9661 |
| Dcx     | 1:100    | goat  | Santa Cruz SC-8066 |
| Dlx2    | 1:500    | rabbit | Millipore AB5726  |
| Eed     | 1:100    | mouse | Abcam ab126542    |
| EGFR    | 1:200    | rabbit | Santa Cruz sc-03  |
| Ezh2    | 1:100    | mouse | BD Biosciences 612666 |
| GFAP    | 1:400    | rat   | Invitrogen 130300 |
| GFAP    | 1:5000   | chicken | Abcam ab4674    |
| GFP     | 1:500    | chicken | Aves GFP-1010    |
| H3K27me3| 1:250    | rabbit | Millipore 07-449 |
| Ki67    | 1:500    | rabbit | Abcam ab166667   |
| Mash1   | 1:100    | mouse | BD Biosciences 556604 |
| Nestin  | 1:200    | mouse | Millipore MAB353 |
| NeuN    | 1:400    | mouse | Millipore MA8377 |
| Olig2   | 1:1000   | rabbit | Millipore AB9610 |
| PHi3    | 1:400    | rabbit | Millipore 06-570 |
| S100β   | 1:200    | rabbit | Dako Z031129-2   |
| S100β   | 1:100    | mouse | Sigma S2657      |
| Sox2    | 1:500    | rabbit | Millipore AB5603 |
| Sox2    | 1:200    | goat  | R&D systems AF2018 |
### Supplemental Table 2. Primers for RT-PCR

| Gene     | Sequence (5’-3’)                                                                 |
|----------|---------------------------------------------------------------------------------|
| Gata6    | CTTGGGGCTCTATATGAACTCCAT                                                       |
|          | TAGAAGAAGGAGAGTGGAGAGTACGGACGACCA                                                |
| Sox17    | GCCAAAGACGAACGCAACGGAAGG                                                         |
|          | TTCTCTGCAAAGGTCAACGGCT                                                          |
| Hoxa11   | TCGGGCCACACTGAGGACAGAG                                                        |
|          | ACTCTGCTCAACGCTCTGGATCT                                                         |
| Olig2    | CGCAGCGACCTCAATCTCAA                                                             |
|          | CCCAGGGATGATCTAAGCTCTCGAA                                                        |
| Ngn1     | ATCACCACTCTCTGACACC                                                           |
|          | GAGGAAGAAGATTATGGATGGCTTTCCA                                                   |
| Ezh2     | TTACCTGCTGGACGGTCTGATGTG                                                          |
|          | TGCTCTGTTCTCATCCTGAGAAATAATCTCC                                                |
| Ezh1     | AATATGGGACGCAAGGGCTCTGTATGTG                                                    |
|          | CGCAGAATTTTCTTCCACCTCCATTGAGG                                                   |
| Eed      | GCACAGAGATGAAGTTCTGAGTGCT                                                       |
|          | ATAAGACTCTCTAAATTCATTACCTACCT                                                  |
| 18s      | GAAACGGTTGAAACCCATT                                                              |
|          | CCAATCCTAGCTAGAGGCG                                                            |
| Mash1    | GCAAACGGGCTCAAGTTGGG                                                           |
|          | GTGCAGGGATAGTGGGGGG                                                              |
| Id2      | ATGAAAGCCTCTCAGCTCCGTTG                                                         |
|          | ACAGAACTCATCGGCTGTG                                                            |
| Dcx      | CATTTTGACGAACGACACAAAGC                                                         |
|          | TGGAGTCCAATCCATCCTGGA                                                         |
| Bmp4     | CACTGTGAGGAGTTTCCACGGAAG                                                      |
|          | GGATGCCTGCTAGGGTGAAGAGGA                                                      |
| Bmp6     | GCCATCTGGTTCTTTAATTGATG                                                        |
|          | GTGGTTAAGGAGATGTTGTATG                                                         |
| Ngn2     | TCACCAGGGACACTGTATCT                                                           |
|          | CTGGAAATGAGAGTCCG                                                             |
| p21 (Cdkn1a) | CCTGTTAGTGCTCCGACCCTT                                      |
|          | CCATGAGGCATCAGCACATC                                                          |
| p27 (Cdkn1b) | TCAACGTGAGAGTGCTCAACG                                      |
|          | CCGGGCCAGAAGATTTCTG                                                            |
| p57 (Cdkn1c) | CGAGAGAGGGCGAGGAGGAGAATC                                              |
|          | GAAGAAGTGGTTCGCTGGGC                                                           |
| Cdkn2a   | CAGACCAGGCGGATAGCTCTCA                                                          |
|          | GGATTTAGCTCTGCTCTGGGGAGTTG                                                      |
| Bmp7     | TTCCCTGGAACAGGAGTCTG                                                           |
|          | CCTGAATCTCGGCGACTT                 |
Supplemental Table 3. Primers for ChIP-qPCR

| Gene             | Sequence (5’-3’)       |
|------------------|------------------------|
| Olig2            | GCCTGACGCTACAGTGACAA   |
|                  | GGCTAATTCCGCTCAATGAA   |
| Gata6            | CGGTCCTTCGTTTAAGAGTTGAGG |
|                  | ACACAGACCAGCAAAGATGAAGAAGA |
| Ngn1             | CATTGTGGCCGCGTA        |
|                  | GCGATCAGATCAGCTCCT     |
| Actin (promoter) | CCAACACACCTAGCAAATTGAACCAC |
|                  | CCTGGATTGAATGGACAGAGTCACT |
| Actin (intron I) | CGTATTTGTCATCTTGAGATGACAGTATT |
|                  | GCCATTGAGGCGATGATGCTAGC |
| Hoxa11           | GCTGGGAGAAGTGCTGAAACG  |
|                  | CGGTGGGTGAGGGATACCTCTCTGG |
| Sox17            | CCACTCACTCTGAGGCTGCTGTA |
|                  | CCAAACAGACCTGAGGCTGCAAA |

Supplemental Table 4. Antibodies for western blot

| Antigen | Dilution | Host  | Manufacturer Cat.   |
|---------|----------|-------|---------------------|
| p21     | 1:1000   | Rabbit| Abcam ab109199     |
| Eed     | 1:5000   | Mouse | Millipore 05-1320  |
| Gata6   | 1:200    | Rabbit| Abcam ab22600      |
| β-actin | 1:5000   | Rabbit| Abcam ab8227       |
Supplemental references

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Su IH, Basavaraj A, Krutchinsky AN, Hobert O, Ullrich A, Chait BT, Tarakhovsky A. 2003. Ezh2 controls B cell development through histone H3 methylation and Igh rearrangement. Nat Immunol 4:124-131.
A. Relative expression of miRNAs (miR-17, miR-93, miR-224, miR-291, miR-295, miR-322b, miR-322c) at 2DIV.

B. Relative expression of genes (Fxr1, P53, Msi1, Igf2bp2, Igf2bp3, Rmm2, Cnot7, Hnmp1, Hnmp4, Hnmt1, Raly) at 2DIV.

C. Neurospheres (3DIV):

|          | control | Gata6 OE |
|----------|---------|----------|
| WB:      | DMSO MG132 | DMSO MG132 |
| p21      |          |          |
| β-actin  |          |          |
| p21/β-actin | 0.57 | 0.77    | 0.19 | 0.44 |
| relative increase | 1 | 1.35 | 1 | 2.33 |

D. Neurospheres (2DIV):

|          | Gata6 OE + siFluc | Gata6 OE + siGata6 |
|----------|-------------------|-------------------|
| WB:      |                   |                   |
| Gata6    |                   |                   |
| β-actin  |                   |                   |

* indicates a significant increase.