SETD4 cells contribute to brain development and maintain adult stem cell reservoir for neurogenesis

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SUMMARY

Cellular quiescence facilitates maintenance of neural stem cells (NSCs) and their subsequent regenerative functions in response to brain injury and aging. However, the specification and maintenance of NSCs in quiescence from embryo to adulthood remain largely unclear. Here, using Set domain-containing protein 4 (SETD4), an epigenetic determinant of cellular quiescence, we mark a small but long-lived NSC population in deep quiescence in the subventricular zone of adult murine brain. Genetic lineage tracing shows that SETD4+ cells appear before neuroectoderm formation and contribute to brain development. In the adult, conditional knockout of Setd4 resulted in quiescence exit of NSCs, generating newborn neurons in the olfactory bulb and contributing to damage repair. However, long period deletion of SETD4 lead to exhaustion of NSC reservoir or SETD4 overexpression caused quiescence entry of NSCs, leading to suppressed neurogenesis. This study reveals the existence of long-lived deep quiescent NSCs and their neurogenic capacities beyond activation.

INTRODUCTION

Adult neural stem cells (NSCs) residing within the subventricular zone (SVZ) of the lateral ventricles and the subgranular zone (SGZ) of the hippocampus are known for their ability to generate new neurons that contribute to complex sensory and cognitive functions throughout the lifetime of mammals (Bond et al., 2015; Choi et al., 2018; Doetsch et al., 1999; Mobley et al., 2014; Seri et al., 2001). In the mature brain, these NSCs exist in either a quiescent or an active state (Furutachi et al., 2015; Lugert et al., 2010; Ming and Song, 2011). The state of quiescence has been considered to contribute to long-term maintenance of a reservoir of NSCs by preventing the exhaustion of their proliferation potential and permitting the avoidance of accumulation of damage to DNA, proteins, and mitochondria that might otherwise result in malignant transformation or senescence (Audesse et al., 2019; Cameron et al., 2019; Prozorovski et al., 2008). Upon aging, the higher fraction of NSCs identified in a quiescence state suggests a compensatory mechanism to avoid full depletion of the NSC compartment (Kalamakis et al., 2019). After brain injury, quiescent NSCs can be activated to produce multiple types of progenies to contribute to brain repair (Faiz et al., 2015; Llorens-Bobadilla et al., 2015; Otsuki and Brand, 2020; Delgado et al., 2021). Because of the overall heterogeneity of the NSC population and the lack of defining makers of NSCs that occur in a quiescent state, it is currently unknown how quiescent NSCs originate in the embryo. Their specification and maintenance in adulthood and into old age also remains unclear. Understanding of how quiescent NSCs react in the injured or aged brain and any related potential for therapeutic contributions towards regeneration in the human brain would represent significant advances.

Systemic and local signals seem to regulate NSC quiescence by modulating the expression or function of molecules acting within stem cells. These include transcription factors, cell cycle regulators, and metabolites, which in turn control NSC quiescence or activation (Cavallucci et al., 2016; Kippin et al., 2005; Knobloch et al., 2017; Sueda et al., 2019). Although several molecular players in the regulation of NSC quiescence have been reported (Ahn and Joyner, 2005; Chavali et al., 2018; Engler et al., 2018; Ganapathi et al., 2018; Kandasamy et al., 2014; Marqués-Torrejón et al., 2021; Sueda et al., 2019; Urbán et al., 2019), we know remarkably little about the direct determinants of quiescence and the mechanisms of the transition between active and quiescent states in NSCs.

Epigenetic studies have shown that factors such as chromatin remodeling and histone modification play critical roles in regulating stem cell behaviors, providing genome-wide transcriptional changes and long-lasting effects for downstream signaling pathways (Yao et al., 2016). As a lysine methyl-transferase of histone, SET domain-containing protein 4 (SETD4) has been reported to play a role in cell proliferation by controlling related gene expressions (Faria et al., 2013; Liao et al., 2021). Our previous report revealed that SETD4 regulates cellular dormancy or deep quiescence by H4K20ME3 catalysis in diapause embryos of Artemia, a quiescent model system (Dai et al., 2017). SETD4 was then also found to epigenetically regulate quiescence of cancer stem cells in human breast tumor and C-KIT cells in the murine heart by...
facilitating heterochromatin formation via H4K20ME3 (Ye et al., 2019; Xing et al., 2021). Thus, an evolutionarily conserved mechanism of cellular quiescence as modulated by SETD4 was proposed.

In this study, upon confirmation of SETD4 marking of a small NSC population in deep quiescence in the SVZ, we performed identification lineage tracing and conditional knockout and knockin of Setd4 in mice. We found that SETD4+ cells occurred in the neuroepithelium before neuroectoderm formation, contributed to the brain development in early embryo, and then entered a quiescent state in the late embryo. As a long-lived quiescent NSCs, postnatal SETD4+ cells preserve the stem cell reservoir for retaining their function of neurogenesis over extended periods and, upon activation, are able to function in the repair of the injured olfactory bulb (OB).

RESULTS

SETD4 marks a small population of long-lived quiescent NSCs in the SVZ after birth

To investigate whether SETD4+ cells occur in the stem cell niches of brain, we used RNAscope, an RNA fluorescence in situ hybridization technique, to reveal endogenous Setd4 expression in the adult SVZ and SGZ. Using RNAscope combined with immunostaining analysis we found that a small population of NSCs (GFAP, SOX2, and NESTIN) expressed the Setd4 gene in the adult (2-month-old) SVZ (Figure 1A). However, we did not observe any Setd4 expression in the adult SGZ (Figure S1A). We then generated inducible transgenic Setd4-CreERT2::mTmG lineage tracing mice in which CreER2 recombinase is under the control of the SETD4 promoter. Upon tamoxifen (TAM) induction, membrane-based green fluorescent protein (GFP) was expressed, allowing endogenous SETD4+ cells to be followed (Figure 1B). Beyond a series of TAM pulses and 3 day chases, RNAscope analysis confirmed that TAM-induced GFP accurately marked SETD4+ cells (Figure 1C). We identified a small population of GFP+ cells (1.78% ± 0.10%) in the SVZ of 2-month-old Setd4-CreERT2::mTmG mice (Figure 1D). Consistent with observations using the RNAscope assay, we did not observe any GFP+ cells in the SGZ of the dentate gyrus (Figure S1A). In the SVZ, these GFP+ cells existed along the lateral ventricular wall as single-cell clones and lacked any Ki67 or PCNA expression, markers of cell proliferation, indicating that they were in quiescent state (Figure 1E). To confirm this, an EdU incorporation assay was performed. We found that all TAM-induced GFP+ cells were EdU negative (Figure 1B). This was consistent with the former Ki67 and PCNA negative data. Moreover, the majority (more than 90%) of GFP+ cells expressed GFAP, SOX2, and NESTIN as NSC markers but lacked any expression of MASH1, DCX, NEUN, or OLIG2, markers of transit-amplifying cells (TAPs), and also lacked any immature migratory neuroblasts (NBs), mature neurons or oligodendrocytes (Figures 1F and S1B). These GFP+ cells exhibited the typical radial morphology of NSCs having minute apical endings to their ventricular surfaces and long basal processes (Figure 1G) (Mirzadeh et al., 2008; Obernier and Alvarez-Buylla, 2019; Delgado et al., 2021). We considered such data as confirmation of the SETD4+ quiescent NSC identity.

To identify whether SETD4+ cells also exist as quiescent NSCs in the aged SVZ, we also analyzed Setd4 expression in aged (20-month-old) mice by RNAscope. Results showed that the SETD4+ NSCs (GFAP+, SOX2+, and NESTIN+) existed in the SVZ of aged mice (Figure S1C). Similarly, 15-month-old Setd4-CreERT2::mTmG mice were induced by TAM (Figure 1B). RNAscope analysis showed that TAM-induced GFP+ cells could still accurately mark SETD4+ cells (Figure S1D). As observed in 2-month-old mice (Figures 1D–1G), a small population of GFP+ cells were also identified as quiescent NSCs in the aged SVZ, occurring along the lateral ventricular wall, showing positive for NSC markers, negative for Ki67 expression, and displaying NSC morphology (Figures S1E–S1H). We analyzed the whole-brain sections of 24 h TAM-induced Setd4-CreERT2::mTmG mice at early postnatal day (P7), adult (2 months), and aged (20 months) (Figure S1I). We found that SETD4-expressing cells existed only within the wall of the ventricular zone in the forebrain and expressed SOX2 but not Ki67 throughout these stages (Figures S1J–S1L). We concluded that SETD4 marks a small population of quiescent NSCs in the SVZ of the mouse brain.

As we observed SETD4+ NSCs existed in the SVZ of neonatal (P3) brain (Figure S2A), to explore SETD4+ NSCs properties after birth, lineage tracing was performed at early postnatal and adult. Setd4-CreERT2::mTmG mice were treated with TAM at P3, then analyzed at P4, P21, and P56 (Figure 1H). Almost all GFP+ cells expressed markers of NSCs but lacked the cell proliferation marker of Ki67, each maintained as a single-cell clone in the SVZ throughout all time points (Figures 1I, S2B, and S2C). These SETD4+ NSCs still showed typical radial morphology along the SVZ at P56 (Figure 1I). Previous reports have shown that NBs derived from SVZ NSCs migrate to the OB and distribute to the granule cell layer (GCL) and the glomeruli (Lim and Alvarez-Buylla, 2016). Here, we did not observe any progeny (GFP+) of SETD4+ NSCs in the OB, indicating that these SETD4+ NSCs were maintained in a quiescent state in the SVZ during postnatal development (Figures 1I and S2D). The long-term behavior of SETD4+ quiescent NSCs was also examined by lineage tracing in adult (2-month-old) Setd4-CreERT2::mTmG mice. After 1 and 12 months of TAM induction, GFP+ cells were observed as remaining as single-cell clones in the wall
Figure 1. SETD4 marks long-lived quiescent NSCs in the SVZ of adult brain

(A) Confocal images of in situ hybridization of Setd4 expression using RNAscope and combined with immunostaining for the NSC markers GFAP, SOX2, and NESTIN in the adult SVZ of C57BL/6 mice. The boxed area represents a higher magnification. The yellow dotted circle highlights a Setd4+ cell.

(B) Schematic representation of Setd4-CreERT2 lineage tracing mice. Adult lineage tracing mice were injected with TAM and chased after 3 days for the following analyses.

(C) Confocal images of in situ hybridization of Setd4 and egfp expression using RNAscope in the adult SVZ of Setd4-CreERT2::mTmG mice after 3 days of TAM induction. A higher magnification is showed in the boxed area. The yellow dotted circle highlights a Setd4+ cell.

(D) Confocal images of GFP+ cells in the whole SVZ of 2-month-old mice with higher magnification boxed area (right).

(E and F) Confocal images of cell proliferation markers (Ki67 and PCNA), EdU-incorporated signal (E) and the NSC markers GFAP, SOX2, and NESTIN (F) in GFP+ cells in the SVZ (top) of 2-month-old mice. A higher magnification is showed in the boxed area (bottom).

(G) Representative image of a SETD4+ cell in the SVZ of 2-month-old mice, with a representative minute apical ending to the ventricular surface and a long basal process.

(H) Experimental overview. Postnatal Setd4-CreERT2::mTmG mice were injected with TAM at P3 for lineage-tracing analysis at P4, P21, and P56. Adult Setd4-CreERT2::mTmG mice were injected with TAM at P56 for extended period lineage-tracing analyses at different time points.

(I) Confocal images of immunostaining for NSC markers in GFP+ cells in the SVZ at P56 (left) and the mature neuron marker NEUN in the OB (right). The yellow arrowhead signifies a single NSC (GFAP+SOX2+GFAP+) and its radial process.

(J and K) Confocal images of GFP+ cells consisting of a single NSC (GFAP+) in the SVZ and few progenies including oligodendrocytes (OLIG2+) in the striatum (top) and neurons (NEUN+) in the OB (bottom) after 12 months of TAM induction (J). Quantification of the number of GFP+ cells in lineage tracing experiments of adolescent and adult mice (K).

Nuclei were stained with DAPI (blue). LV, lateral ventricle; OB, the olfactory bulb. n = 4 mice were analyzed in each experiment.
A. Setd4-Cre::mTmG
- Setd4 expression
- Reporter expression
- Analysis
- E8.5
- E14.5
- P0
- P56

B. DAPI GFP
- LV
- E8.5
- E14.5
- P0
- P56

C. Setd4-CreERT2::mTmG
- TAM
- Analysis
- E7.5
- E8.5

D. DAPI GFP
- Percentage among all GFP+ clones (%)

E. Setd4-CreERT2::mTmG
- TAM
- Analysis
- E8.5
- E15.5

F. DAPI GFP KIR7
- LV

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of the SVZ (Figures 1J and S2E). At very low cell numbers, GFP+OLIG2+ oligodendrocytes in the striatum and GFP+NEUN+ neurons in the GCL of the OB were identified, showing that SETD4+ quiescent NSCs are long lived and have neurogenic capacity. We found that lineage tracing of adolescent and adult SETD4+ NSCs resulted in ∼8 GFP+ cells per SVZ examined, and the number of GFP+ cells did not change over time (Figure 1K). These results reveal that a small population of long-lived SETD4+ quiescent NSCs occur from embryo to the adult SVZ.

Embryonic SETD4+ cells contribute to brain development in the early embryo, then enter quiescence in the late embryo

As the earliest neural stem cells, radial glia cells (RGCs), derived from neuroepithelial cells (NECs), arise around embryonic day 10.5 (E10.5) (Kriegstein and Gotz, 2003; Kriegstein and Alvarez-Buylla, 2009). They then contribute to both embryonic and postnatal neurogenesis, giving rise to all the neuronal and glial lineages of the CNS (Anthony et al., 2004; Merkle et al., 2004). However, the early embryonic origins of NECs remain unclear. Before embryonic neurogenesis, the neuroectoderm is composed of a single layer of NECs, which form between E6.5 and E10.5 (Gotz and Huttner, 2005). Considering that the CNS originates from this, we generated Setd4-Cre::mTmG mice, which permitted observation of the descendants of SETD4+ cells without the requirement of TAM induction (Figure 2A).

We performed identification of GFP+ cells in the neuroepithelium at E8.5, the developing CNS at E14.5, and the neonatal brain at P0. We found that GFP+ cells, including SETD4+ and their daughter cells, constituted more than 36.40% ± 2.0% in the neuroectoderm at E8.5, which increased to more than 88.20% ± 4.83% and 96.40% ± 0.97% in the embryonic ventricular zone (VZ) at E14.5 and the SVZ at P0, respectively (Figure 2B). From E8.5 to P0, these GFP+ cells were identified via detection of NEStin and Sox2 to label the majority of NECs in the neuroectoderm, then RGCs in the embryonic VZ and NSCs in the SVZ (Figure S3A). TUJ1 detection confirmed that these GFP+ cells also largely contributed to neurons (TUJ1+) in the embryonic and neonatal brain (Figure S3A). Interestingly, beyond developmental stages, these GFP+ cells were also identified as distributed throughout the entire brain of adult Setd4-Cre::mTmG mice at P56, including the OB, striatum, cortex, SVZ, and SGZ regions (Figure S3B). Co-localization analysis via cytoplasmic-localized reporter (Setd4-Cre::Luc-EGFP) further validated our observation that almost all cells were indeed GFP positive at P56, including the radial glial-like cells in SGZ (Figure S3C). Flow cytometric analysis showed that approximately 98.33% ± 0.24% cells were GFP positive of the whole brain (Figure S3D). These results suggest that fetal SETD4+ cells occurring at very early embryonic stages were vital to the origin of subsequent brain development.

To further investigate fate specification of individual early embryonic SETD4+ cells, we next administered TAM to pregnant female Setd4-CreER<sup>+</sup>::mTmG mice at E7.5 and chased at E8.5 before the neuroepithelium was fully specified in the embryos. The GFP+ cells were identified in the neuroectoderm with NEStin positive as NECs (Figure 2C). The GFP+ clones consisted of single, double, or multiple cells, indicating that SETD4+ cells had occurred at E7.5 and contributed to NEC population in the neuroepithelium (Figure 2D). To examine whether this GFP+ population produces fetal RGCs and neurons during embryonic neurogenesis, we performed a single TAM pulse at E8.5 and chased at E15.5 (Figure 2E). As expected, these GFP+ clones were observed in cortical, striatal, and septal sides of the VZ in the forebrain. Upon immunostaining with SOX2, these GFP+ clones were further confirmed as consisting of RGCs (GFP+SOX2<sup>+</sup>), and their migrating differentiated progeny (GFP+SOX2<sup>+</sup>). These
A

Nestin-CreER<T>^{<tdTomato}

B

C

D

E

F

G

H

I

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results demonstrate that SETD4+ cells originate NECs, then generate RGCs that contribute to embryonic neurogenesis in the forebrain.

Previous research has demonstrated that RGCs can enter a quiescent state between E13.5 and E15.5, which can then persist for a long period into the adult SVZ (Fuentealba et al., 2015; Furutachi et al., 2015). To directly assess whether SETD4+ cells enter a quiescent state during embryonic development, we injected pregnant female Setd4-CreER<sup>T2</sup>.:mTmG mice with TAM at E13.5, E15.5, and E18.5 (Figure 2F). One day later, embryos were examined for the presence of GFP+ cells. The GFP+ cells showed the characteristic morphology of RGCs with long radial fibers on the ventricular surface at all time points. We examined the division state of these cells by immunostaining for Ki67. The percentages of Ki67+GFP+ cells decreased from 85.71% ± 9.71% at E14.5 to 40.00% ± 14.53% at E16.5 and then dropped to zero at E19.5, indicating SETD4+ cells had entered a quiescent state during that period. These results show that SETD4+ cells appear in the early embryo, contribute to brain development, and then enter quiescence before birth.

**Conditional knockout of Setd4 activates quiescent NSCs and rapidly promotes neurogenesis**

The functions of these long-lived quiescent NSCs could be examined only after their activation. Because of the NSC marker’s expression in SETD4+ cells, Setd4 conditional knockout (cKO) was constructed by generating Nestin-CreER<sup>T2</sup>.:Setd4<sup>flx/flx</sup>.:tdTomato (SETD4 cKO) mice (Figure 3A). To activate SETD4+ quiescent NSCs, we performed Setd4 cKO in adult SETD4 cKO mice (Figure S4A). Adult Nestin-CreER<sup>T2</sup>.:tdTomato reporter mice were used as a control, and the majority of the NESTIN+ NSCs were labeled with tdTomato (Td). After 3 days of TAM induction, the expression of SETD4 in the SVZ had been abolished, as indicated by western blot analysis, showing that SETD4 cKO mice had been successfully constructed (Figure S4A). Meanwhile, we observed a significant increase of active NSCs (Ki67+TdT+) in contrast to controls (Figure S4B). This indicated that deletion of SETD4 had resulted in the activation of quiescent NSCs, enabling entrance into the cell cycle. This progress was also supported by a significant increase of EdU-labeled NSCs (EdU+TdT+) after 2 h of EdU incorporation in SETD4 cKO mice (Figure S4C). At 3 days, the increased proliferation of NSCs had not affected the number of TdT+GFAP+ NSCs in the SVZ of the SETD4 cKO mice (Figure S4D). This suggested that deletion of SETD4 causes activation of SETD4+ quiescent NSCs.

Next, we extended the study to one month to identify the effect of SETD4 deletion on NSC activation and the differentiation of their progeny (Figure 3B). One month after deletion, the number of GFAP+TdT+ NSCs had significantly increased in the SVZ of SETD4 cKO mice compared with control mice (Figure 3C), in which the numbers of BrdU label-retaining NSCs (GFAP+BrdU+TdT+) and active NSCs (Ki67+TdT+) decreased and increased, respectively (Figures 3D and 3E). SETD4 deletion also led to significant increases of newborn TAPs (MASH1+TdT+) and NBs (DCX+TdT+) in the SVZ and progeny neurons (NEUN+TdT+) in the OB (Figures 3F–3H). These demonstrated a strengthening of neurogenesis of SETD4 cKO mice. We propose that one month of SETD4 deletion was sufficient to both activate the quiescent NSCs in the SVZ and then give rise to newborn neurons in the OB.

To validate this, we also performed Setd4 cKO in the SVZ of adult Setd4<sup>flx/flx</sup>.:tdTomato mice by injection of AAV9-GFAP-Cre (Figure S4E). As observed in the SETD4 cKO mice after one month of TAM induction, the number of SOX2+TdT+ NSCs had significantly increased after 6 weeks of SETD4 deletion, in contrast to controls (Figure S4F). Correspondingly, the numbers of active NSCs (Ki67+TdT+), TAPs (MASH1+TdT+), and NBs (DCX+TdT+) in the SVZ and their progeny neurons (NEUN+TdT+) in the GCL of the OB had also increased after SETD4 deletion (Figures S4G–S4J). These findings show that the deletion of SETD4 causes quiescence exit of quiescent NSCs and increased neurogenesis, resulting in more newborn neurons in the OB and strengthening the suggestion that SETD4 regulates the maintenance of quiescence of NSCs (Figure 3I).

**Persistent knockout of Setd4 results in NSC exhaustion and a decrease in neurogenesis**

Age-related decline in adult neurogenesis has been linked to NSC exhaustion (Signer and Morrison, 2013; Shi et al., 2018). Although over a shorter period of one month, we...
Figure 4. Long-term deletion of SETD4 results in NSC exhaustion and decrease of neurogenesis

(A) Experimental overview. Control and Setd4 cKO mice were pulsed with TAM injection, followed by BrdU injection after 3 months of TAM induction and analyses in (B)–(G) after 4 months of TAM induction.

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had observed that Setd4 cKO causes an increase in neurogenesis, we then examined the longer term consequences of SETD4 deletion. We extended TAM induction to 4 months in adult SETD4 cKO mice and labeled NSCs in the SVZ and newly generated neurons in the OB (Figure 4A). BrdU label-retaining NSCs, GFAP+SOX2+BrdU+ cells had significantly reduced in the SVZ after 4 months Setd4 cKO (Figure 4B). This was in line with the observation that the numbers of active NSCs (GFAP+KI67+) and GFAP+SOX2+ TAPs (MASH1+KI67+) and NBs (DCX+KI67+) and BrdU label-retaining NSCs had also decreased (Figures 4C and 4D). Long-term Setd4 cKO in mice also showed a significant reduction in TAPs (MASH1+KI67+) and NBs (DCX+KI67+) and BrdU label-retained newborn neurons (NEUN+BrdU+) in the GCL of the OB, compared with those of controls (Figures 4E-4G). We concluded that persistent deletion of SETD4 leads to depletion of the NSC reservoir and removal of their corresponding neurogenic abilities (Figure 4H).

In addition, we specifically knocked out Setd4 using a UBC-CreERT2 driver (Figure S5A). We observed that ablation of SETD4 in UBC-CreERT2::Setd4floxflox mice led to similar results to those of long-term Setd4 cKO in SETD4 cKO mice with the overall number of GFAP+SOX2+ NSCs, BrdU label-retaining NSCs active NSCs, TAPs and NBs in the SVZ, and newborn neurons in the GCL of the OB all being decreased in contrast to the controls (Figures S5B-S5G). In conclusion, SETD4 may function to sustain quiescent NSCs, thus protecting NSCs from exhaustion and retaining their neurogenesis potential in the adult.

**Beyond activation, SETD4+ quiescent NSCs promote regeneration in the injured OB**

To investigate whether, beyond their activation, SETD4+ quiescent NSCs are the instigators of regeneration, we injured right side of the OB of adult SETD4 cKO mice by N-methyl-D-aspartic acid (NMDA) injection, leaving the left side undamaged as a control. Examinations then began after one month of Setd4 cKO by TAM induction (Figure 5A). Following one week BrdU label-retaining and 3 week chase, BrdU signals were used to mark newborn neurons after NMDA induced injury. We found that the size of the injured OB recovered to the control levels after one month of injury (Figure 5B). As newborn neurons, the number of BrdU+Td+ cells significantly increased in the injured OB (Figure 5C). Moreover, along with the regeneration after Setd4 cKO, active NSCs (Td+GFAP+KI67+) markedly increased in the SVZ, an observation that may support neurogenesis of the injured OB (Figure 5C).

To assess such a contribution to neurogenesis in the injured OB by the activation of quiescent NSCs after Setd4 cKO, we also performed the OB injury in adult Nestin-CreERT2::tdTomato control mice without Setd4 cKO (Figure S6A). Although newborn neurons labeled by BrdU increased in the injured OB compared with control OB (Figure S6B), the level of newborn neuron numbers in control mice (without Setd4 cKO) was significantly lower than that in SETD4 cKO mice after one month of Setd4 cKO (Figure S6C). Notably, no discernible change in active NSCs (Ki67+GFAP+Td+) was observed in the SVZ in control mice after one month OB injury (Figure S6D). These results indicate that, one month beyond activation of Setd4 cKO, these quiescent NSCs were enabled to promote the regeneration of the injured OB.

However, if Setd4 cKO occurred a considerable period before OB injury, the previous positive effects were reversed. If OB injury was induced after a delay of 4 months beyond Setd4 cKO in SETD4 cKO mice, after a further 1 month, the size of the OB injured side showed a significant decrease in size and an atrophic appearance, in contrast to the left side control (Figure 5E). The number of newborn neurons (BrdU+Td+) in the OB labeled by BrdU and active NSCs (GFAP+Ki67+Td+) in the SVZ showed no further changes between injured-side OB and control-side OB, indicating that the regenerative capacity for injury repair had been compromised (Figures 5F and 5G). We propose that SETD4+ quiescent NSCs are critical to protect the NSCs from exhaustion and to retain neurogenesis capacity in the adult, in which SETD4 may also be active maintaining a stable quiescent NSCs number in the SVZ throughout the lifetime.

**Overexpression of Setd4 results in NSC quiescence entry and suppresses neurogenesis**

In contrast to the consequences of SETD4 deletion, we also performed overexpression (OE) of Setd4 in NSCs via generating Nestin-CreERT2::Rosa26-Setd4::tdTomato (SETD4 OE) mice (Figure 6A). Upon RNAscope analysis, we observed that more SVZ NSCs expressing Setd4 mRNA after 3 days of TAM induction compared with controls (Figures S7A and S7B). The expression level of SETD4 protein in the SVZ had also been up-regulated successfully (Figure S7C). This overexpression of SETD4 resulted in a significant (B-G) Confocal images and quantification of BrdU label-retaining NSCs (BrdU+SOX2*GFAP*) (B), active NSCs (Ki67*GFAP*) (C), total NSCs (SOX2*GFAP*) (D), TAPs (Ki67*MASH1*) (E), NBs (Ki67*DCX*) (F) in the SVZ, and newborn neurons (NEUN*BrdU*) (G) in the GCL of the OB in control (Ctrl) and Setd4 cKO mice.

(H) Summary of the effects of 4 month Setd4 cKO mice on NSCs in the SVZ and production of newborn neurons in the OB. Nuclei were stained with DAPI (blue). GCL, granule cell layer; OB, olfactory bulb. Values represent the mean ± SEM. **p < 0.01 and ***p < 0.001. n = 4 mice.
decrease in the number of active NSCs, as confirmed by detection of Ki67 and assay of EdU incorporation (Figures S7D and S7E), indicating that the self-renew ability of NSCs has been abolished. The overall number of GFAP"TdT" NSCs in the SVZ remained unchanged compared with those in control mice (Figure S7F). These data suggested that SETD4 conveys quiescence to SVZ NSCs.

We then investigated the effects of Setd4 OE after one month (Figure 6B). We observed that the number of TdT"GFAP" NSCs in the SVZ of one month SETD4 OE mice were significantly fewer than those in the control mice in normal physiological conditions, as the inhibition of NSC self-renews (Figure 6C). Meanwhile, the numbers of active NSCs (TdT"Ki67") were also decreased beyond Setd4 OE (Figure 6D). It is tempting to speculate that the normal neurogenesis progress was suppressed. In line with this, the numbers of TAPs (TdT"MASH1") and NBs (TdT"DCX") in the SVZ were reduced (Figures 6E and 6F). Importantly, the significant decrease of newborn neurons (NEUN"BrdU") in the GCL of the OB, and the link of this to the global decrease in these cells, indicated that overexpression of Setd4 induces NSCs to enter a quiescent state, leading to deprivation of their subsequent neurogenic capabilities (Figures 6G and 6H).

SETD4 epigenetically regulates the quiescence of NSCs via multiple signaling pathways

To investigate whether SETD4 epigenetically controls NSC quiescence via the previously noted conserved mechanism by which heterochromatin formation was facilitated via H4K20ME3 catalysis (Dai et al., 2017; Ye et al., 2019), we performed immunostaining for H3K9AC (a marker of
Figure 6. Overexpression of Setd4 results in quiescence entry of NSCs and a decrease of neurogenesis

(A) Schematic representation of Nestin-CreER<sup>T2</sup>:Setd4::tdTomato mice.

(B) Experimental overview. Control (Ctrl) and Setd4 overexpression (OE) mice were pulsed with TAM injection, followed by BrdU injection and analyses in (C)–(G) after one month of TAM induction.

(C–G) Confocal images and quantification of total NSCs (GFAP<sup>Td+</sup>) (C), active NSCs (Ki67<sup>Td+</sup>) (D), TAPs (MASH1<sup>Td+</sup>) (E), NBs (DCX<sup>Td+</sup>) (F) in the SVZ of lateral ventricles, and newborn neurons (NEUN<sup>BrdU+</sup>) (G) in the GCL of the OB in control and Setd4 overexpression mice.

(H) Summary of the effects of one month Setd4 overexpression mice on NSCs in the SVZ and the production of newborn neurons in the OB. Nuclei were stained with DAPI (blue). GCL, granule cell layer; OB, olfactory bulb. Values represent the mean ± SEM. **p < 0.01 and ***p < 0.001. n = 4 mice.
euchromatin) and H4K20ME3 (a marker of heterochromatin) to analyze the changes in the epigenetic state of adult NSCs after Setd4 OE. We observed that SETD4 overexpression induced an increase of H4K20ME3 and decrease of H3K9AC in NSCs (Td+), in direct contrast to the situation of active NSCs (Td+Ki67+) (Figure 7A). This was also

Figure 7. SETD4 epigenetically regulates quiescence of NSCs via multiple signaling pathways

(A) Confocal images and quantification of the level of H3K9AC and H4K20ME3 in NSCs in the SVZ of lateral ventricles in control and Setd4 overexpression mice after 3 day TAM induction. The white dotted circle line highlights the nuclear location of the higher magnification boxed area.

(B) Heatmap of gene expression in primary adult SVZ NSCs after Setd4 overexpression using RNA-seq.

(C) Volcano plot of total changed genes after Setd4 overexpression.

(D and E) Heatmap of differentially expressed genes from top 30-folded-changed genes (D) and representative gene expression for NSC quiescence-maintaining, activation, and cell cycle (E) after Setd4 overexpression.

(F and G) Gene set enrichment analysis (GSEA) (F) and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis (G) after Setd4 overexpression.

Nuclei were stained with DAPI (blue). LV, lateral ventricle. Data are represented as mean ± SEM. n = 3 mice were analyzed per experiment. Student’s t test.
consistent with our observation of high H4K20ME3 levels and low H3K9AC levels in SETD4+ (GFP+) cells dissociated from SVZ of Setd4-CreERT2::mTmG (Figure S7G).

Next, we aimed to reveal molecular mechanisms by which SETD4 controls the quiescence of NSCs in the adult SVZ. For this we performed RNA sequencing (RNA-seq) analysis in primary SVZ NSCs isolated from adult C57BL/6 mice that had been treated with either ADV-GFP (control) or ADV-SETD4-GFP (overexpression). Compared with the controls, 1,759 genes were found to be significantly up-regulated and 1,100 genes were down-regulated after Setd4 overexpression, with a false discovery rate (FDR) ≤ 5% and |log2(fold change)| > 1 (Figures 7B and 7C). Analysis of the differentially expressed genes showed that Setd4 overexpression had resulted in significantly up-regulated expressions of quiescence-maintaining related genes including Draxin, Nidrg2, Notch2, Id4, Foxa3, Apeo, and Bmp4 and down-regulated expressions of both activation-associated genes Akt1, Pi3kr6, Myc, Ctnnbip1 and cell cycle genes, such as Ccnd1, Ccnd2, Mcm2, Cdk1, Mki67, and Pena (Figures 7D and 7E). Gene set enrichment analysis and Kyoto Encyclopedia of Genes and Genomes orthology indicated that the NSC activation-associated signaling pathways of Wnt β-catenin, mTOR, PI3K-AKT, and MYC targets were all inhibited (Figures 7F and 7G). We conclude that SETD4 epigenetically controls quiescence of NSCs for sustaining a protected NSC population and maintaining a stem cell reservoir.

DISCUSSION

NSC quiescence does not simply occur in a uniform manner but is found to be heterogeneous. This diversity has been attributed to cells being at different depths of quiescence, forming a continuum of developmental stages along the quiescence to activity trajectory (Llorens-Bobadilla et al., 2015). Overall, it has been difficult to define NSCs in their quiescent state because of a lack of their specific markers. Identification of their exact function via tracing their lineages during brain homeostasis, damage repair and aging has likewise proved elusive. In this study, we found that SETD4+ NSCs enter quiescence before birth and persist into the SVZ even through to 15-month-old mice, indicating that they remain in a deeply quiescent state over an extended period. SETD4 was then successfully used to mark a long-lived and deeply quiescent NSC population in the SVZ of adult mice. Long-term lineage tracing over one year showed that SETD4+ NSCs maintain their quiescent state with a little or no progeny production in such a state of homeostasis. However, after one month activation by Setd4 knockout, formerly quiescent NSCs showed increased numbers in the SVZ, also with the production of large number of newborn neurons in the OB. This neurogenesis significantly contributed to repair of the damaged OB. We suggest that the quiescence of SETD4+ NSCs relating to brain injury or age-related senescence occurs firstly to protect the NSC reservoir from depletion and then to facilitate subsequent activation from quiescence for reparative and regenerative neurogenesis.

Results of Setd4 cKO and overexpression indicate that, as a determinative factor, SETD4 modulates the quiescence of NSCs. We suggest that SETD4 epigenetically controls NSC quiescence via the previously noted conserved mechanism by which heterochromatin formation is facilitated via H4K20ME3 catalysis. As noted in our previous report (Ye et al., 2019), SETD4 may regulate genome-wide transcription effecting many downstream signaling pathways with a conserved mechanism. In the present study, though deletion of SETD4 resulted in quiescence exit of NSCs in the SVZ and promoted neurogenesis in the OB during an initial period, persistent deletion led to NSC exhaustion and subsequent decrease of neurogenesis. We propose that SETD4 may also endow quiescent NSCs with the potential for asymmetric division for maintenance of a consistent cell number in the SVZ throughout the lifetime. Previous studies have manipulated the general neural cell precursors of various embryonic stages to examine the impact on the adult SVZ neural progenitor pool (Hu et al., 2017; Furutachi et al., 2015; Ohtsuka and Kageyama, 2021). During brain development, NSCs have been noted as contained within the lineages stemming from NECs in the fetal neuroepithelium to postnatal/adult B1 cells via RGCs (Kriegstein and Gotz, 2003; Kriegstein and Alvarez-Buylla, 2009). Here, using Setd4-Cre and Setd4-CreERT2 mouse lines, we found that SETD4+ cells appear before neuroectoderm formation, a very early stage of embryonic neurogenesis and generate NECs, the precursors of the earliest embryonic neural stem cells, RGCs. Combined with their capacities for contribution to almost the entire brain in adult Setd4-Cre mice, we suggest that these SETD4+ cells originate the brain by generating a precursor population that includes both NECs and RGCs, which then contributes to all lineages of neurons during embryonic development. We also found that SETD4+ cells in early embryos produced almost all postnatal NSCs both in the SVZ and SGZ (Figure S3D). However, we did not observe quiescent SETD4+ NSCs in the adult SGZ, suggesting other distinct mechanisms for NSCs reservoir maintenance likely occur in the SGZ.

This appearance at a very early embryonic stage may endow SETD4+ quiescent NSCs with high pluripotency or strong neurogenic capabilities. However, such pluripotency seems to be negated by Setd4 knockout, which led to depletion of the NSC reservoir and removal of neurogenic capabilities over an extended period. We thus
conclude that SETD4 controls NSC quiescence to preserve a stem cell reservoir for retaining the function of neurogenesis over extended periods and contributes to regeneration after injury. Our findings provide SETD4 as a primary regulator, active in the switch between quiescence entrance and exit for NSCs, that governs the rate of neurogenesis in homeostasis and in response to damage. Moreover, our study may pave the way for future pharmacologic developments aimed at targeting quiescent NSCs with SETD4, potentially applicable to many aspects of brain damage and neurodegenerative disease.

**EXPERIMENTAL PROCEDURES**

**Animals**

All animal procedures used in this study were performed following the animal care guidelines approved by the Institutional Animal Care and Use Committee of Zhejiang University. The transgenic animals constructed and used in this study are shown in the supplemental methods. All transgenic mice in this study were backcrossed for at least five generations and maintained on a C57BL/6 background.

**TAM treatment and administration of EdU or BrdU**

A TAM solution (20 mg/mL) was prepared for intraperitoneal injections by dissolving the powder (#T5648; Sigma-Aldrich) in a 9:1 solution of corn oil/ethanol at 37°C with occasional vortexing until dissolved. The labeling of cells progressing through to S phase was performed by intraperitoneal injections of EdU (50 mg/kg; #E10187; Invitrogen), and animals were analyzed 2 h later. For the BrdU label-retaining test, BrdU was administrated in drinking water (1 mg/mL in 0.1% sucrose; #B5002; Sigma-Aldrich) for 1 week followed by a 3 week “chase” period.

**Clonal analysis**

For labeling of SETD4+ cell clonal lineage analysis in embryos, the timed pregnancy was determined by identifying a vaginal plug (E0.5), and then a single dose of 80 mg/kg TAM was administered intraperitoneally to the pregnant females at the target embryonic day. A clone could be defined as a cluster of GFP+ cells located along the neuroepithelium at E8.5 with a cell-to-cell distance of <100 μm.

For labeling of SETD4+ cells lineage tracing in neonates, mice were injected with a single dose of 80 mg/kg of TAM at P3. For population fate mapping of SETD4+ cells in the adults and for the Setd4 conditional knockout or overexpression study, a single dose of 160 mg/kg of TAM was intraperitoneally injected into mice every 24 h and repeated 5 times.

**Immunofluorescent staining**

The immunofluorescent staining details and antibodies used in this study are shown in the supplemental methods.

**RNA sequencing**

Total RNA for RNA-seq was extracted from primary NSCs using TRIzol reagent (#15596018; Invitrogen) after 24 h infection with ADV-GFP (Ctrl) and ADV-SETD4-GFP (Setd4 OE), respectively. RNA-seq was performed at Tianjin Novogene Bioinformatics Technology Co., Ltd., using an Illumina HiSeq 4000 machine (see supplemental methods).

**Statistical analysis**

The studies were blinded during data collection and quantification. The data in figures reflect several independent experiments performed on different days. No data were excluded. Statistical analysis was performed with two-tailed unpaired Student’s t tests, as indicated in the text and figures. All statistical analyses were performed in GraphPad Prism 8.0 software. All data are presented as mean ± SEM. A p value of <0.05 was considered to indicate statistical significance (p > 0.05 was considered to indicate not significant [NS]), with the exact p values stated in the corresponding figures.

**Data and code availability**

The processed bulk RNA sequencing data were deposited in the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO) database under accession code GEO: GSE200159.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.stemcr.2022.07.017.

**AUTHOR CONTRIBUTIONS**

W.-J.Y. and S.-L.C. developed the concept and designed the study. S.-L.C. performed most of the research. W.-J.Y. supervised the research. W.-J.Y., S.-L.C., and Y.-S.Y. wrote the manuscript. Y.-S.Y., Y.-F.D., S.-H.Y., X.-Z.J., Y.-W.G., X.-T.H., and J.-S.Y., provided help in investigation and data analysis. C.W. provided advice during writing and proofread and revised the final manuscript.

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**CONFLICTS OF INTEREST**

The authors declare no competing interests.

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