Functional rejuvenation of aged neural stem cells by Plagl2 and anti-Dyrk1a activity

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The regenerative potential of neural stem cells (NSCs) declines during aging, leading to cognitive dysfunctions. This decline involves up-regulation of senescence-associated genes, but inactivation of such genes failed to reverse aging of hippocampal NSCs. Because many genes are up-regulated or down-regulated during aging, manipulation of single genes would be insufficient to reverse aging. Here we searched for a gene combination that can rejuvenate NSCs in the aged mouse brain from nuclear factors differentially expressed between embryonic and adult NSCs and their modulators. We found that a combination of inducing the zinc finger transcription factor gene Plagl2 and inhibiting Dyrk1a, a gene associated with Down syndrome (a genetic disorder known to accelerate aging), rejuvenated aged hippocampal NSCs, which already lost proliferative and neurogenic potential. Such rejuvenated NSCs proliferated and produced new neurons continuously at the level observed in juvenile hippocampi, leading to improved cognition. Epigenome, transcriptome, and live-imaging analyses indicated that this gene combination induces up-regulation of embryo-associated genes and down-regulation of age-associated genes by changing their chromatin accessibility, thereby rejuvenating aged dormant NSCs to function like juvenile active NSCs. Thus, aging of NSCs can be reversed to induce functional neurogenesis continuously, offering a way to treat age-related neurological disorders.

[Keywords: adult neurogenesis; aged brain; Ascl1; ATAC-seq; ChIP-seq; Dyrk1a; lentivirus; mouse; neural stem cell; Plagl2]

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analyses have been performed [Llorens-Bobadilla et al. 2015; Shin et al. 2015; Arregiani et al. 2017; Dulken et al. 2017; Hochgener et al. 2018; Urbán et al. 2019]. These studies as well as previous ones showed that the activation of Wnt signaling, inhibition of Bmp signaling, induction of Ascl1 oscillations, or lysosomal activation increases the number of active NSCs [Lie et al. 2005; Mira et al. 2010; Benraiss et al. 2013; Jang et al. 2013; Seib et al. 2013; Díaz-Moreno et al. 2018; Leeman et al. 2018; Kalamakis et al. 2019; Sueda et al. 2019]. Notch1 signaling is also required for maintenance of active NSCs, although Notch2 signaling induces quiescence [Nyfeler et al. 2005; Ables et al. 2010; Ehm et al. 2010; Imayoshi et al. 2010; Basak et al. 2012; Engler et al. 2018]. By modulating these signaling pathways, NSCs can be activated to produce some neurons in the young to middle-aged brain; however, this effect is limited in the aged brain (>18 mo of age for mice) [Berdugo-Vega et al. 2020]. Therefore, to restore active neurogenesis, it is important to rejuvenate both the proliferative and neurogenic potential of dormant NSCs.

To rejuvenate aged NSCs, many attempts to change cell-intrinsic properties have been performed. The expression of the cyclin-dependent kinase inhibitor p16INK4a linked to senescence increases during aging, thereby decreasing regenerative functions of NSCs [Molofsky et al. 2006]. However, inactivation of p16INK4a failed to rejuvenate aged NSCs in the hippocampus [Molofsky et al. 2006], and therefore it is not clear whether NSCs can be functionally rejuvenated in the aged brain. It was previously shown that the proneural gene Ascl1 is absolutely required for activation of NSCs and production of new neurons in the adult brain [Andersen et al. 2014], and that Id4 promotes the degradation of Ascl1 protein, leading to quiescence in NSCs [Blomfield et al. 2019; Zhang et al. 2019]. Indeed, inactivation of Id4 increases Ascl1 expression and activates NSCs in the adult hippocampal dentate gyrus; however, the effect on neurogenesis does not continue, probably due to compensation by other Id proteins [Blomfield et al. 2019]. It was shown that oscillatory expression of Ascl1 is important for efficient proliferation of NSCs [Imayoshi et al. 2013], and that induction of Ascl1 oscillations can activate NSCs to proliferate and produce new neurons in the adult brain [Sueda et al. 2019]. However, this activation is transient, and the effect on neurogenesis is limited [Sueda et al. 2019], suggesting that adult/aged NSCs are resistant to proliferation. Overexpression of the cell cycle regulators Cdk4 and cyclinD1 is able to activate proliferation of NSCs [Arregiani et al. 2011], but the effect on neurogenesis is also limited in the aged brain, probably because aged NSCs lose neurogenic potential [Berdugo-Vega et al. 2020]. Therefore, despite all these intensive efforts, the long-term robust activation or rejuvenation of the proliferative and neurogenic potential of NSCs has not been achieved in the aged brain.

Here, to rejuvenate aged dormant NSCs, we sought to change their intrinsic properties and searched for the genes that enhance neurogenesis from those differentially expressed between embryonic and adult NSCs. We found that inducing Plagl2 and anti-Dyrk1a activity rejuvenates NSCs to proliferate and produce new neurons continuously in the aged brain, thereby ameliorating age-related learning and memory deficits. We further analyzed the mechanism of how this treatment rejuvenates dormant NSCs and found that this activation results from up-regulation of key genes, such as Ascl1, whose chromatin structures become less accessible during aging.

Results

Screening for NSC-activating genes

To rejuvenate the proliferative and neurogenic potential of aged NSCs, we reasoned that the activity of multiple genes would need to be changed. As the majority of adult NSCs are quiescent, we compared the transcriptomes [DDBJ Bio-Project accession no. PRJDB9010] of G1/G0 NSCs from the ganglionic eminence and dorsal cortex of E14 mouse embryos with those of NSCs from the LV-SVZ and DG-SGZ of 2- to 3-mo-old mice to identify differentially expressed nuclear factors and their known modulators. We first focused on the top 80 nuclear factor genes expressed at a high level in embryos (“embryonic-high”) and examined their activity in NSC cultures. Each gene was overexpressed by using the Tet-ON system in NSC cultures, which were maintained in a quiescent state (EdU+) by bFGF and BMP [Fig. 1A; Supplemental Fig. S1A,B] by comparing the transcriptomes with those of NSCs from the LV-SVZ and DG-SGZ. The proportions of EdU-incorporating proliferating cells were measured 2 d after doxycycline treatment to identify genes that can activate the proliferation of cultured NSCs [Supplemental Fig. S1B]. The combination of proteins including Gsx2, Dmrt3, Cdk4, Plagl2, Sox21, Ascl1, Tgif2, Plagl1, and Hmgag2 induced the efficient activation of quiescent NSC cultures [Supplemental Fig. S1C,D] and were tested further in vivo. These nine genes were cloned individually together with mCherry into a lentivirus under the control of the Herpes Simplex virus [Supplemental Fig. S1E], which directs NSC- and astrocyte-specific expression [Lugert et al. 2010; Sueda et al. 2019]. Each lentivirus was injected into the hippocampal dentate gyrus of 6-mo-old mice, and brain sections were examined 1 wk later [Fig. 1A; Supplemental Fig. S1F]. Cdk4 was overexpressed with cyclin D1 (D1K4) [Supplemental Fig. S1G], because this combination reportedly activates NSCs in the adult brain [Arregiani et al. 2011]. Among the tested genes, Plagl2 most efficiently increased the number of MCM2+ proliferating cells [most likely, active NSCs and intermediate progenitor cells [IPCs]] in the hippocampal dentate gyrus of 6-mo-old mice compared with the control [Supplemental Fig. S1G]. The combination of Plagl2 with Plagl1, Hmgag2, Sox21, Dmrt3, Gsx2, or Ascl1 did not significantly increase the number of MCM2+ or DCX+ cells (data not shown).

We next performed knockdown of nuclear factor genes expressed at a high level in adult NSCs (“adult-high”) and their modulators by introducing siRNA targeting these genes in quiescent NSC cultures to identify those that could activate the proliferation of NSCs [Fig. 1A; Supplemental Fig. S2A]. Among 124 genes tested in knockdown screening in vitro, we selected 11 genes, Nr4a1, Stat6,
Figure 1. Screening for NSC-activating genes and analysis of the iPaD lentivirus. (A) Strategy for NSC-activating gene screening. Genes involved in in vitro NSC activation were searched for by lentivirus-mediated overexpression or siRNA knockdown. Top genes were introduced into the hippocampal dentate gyrus of 6- or 18-mo-old mice using lentivirus, and brain sections were examined. (B) Control lentivirus, lentivirus inducing Plagl2 expression, or lentivirus inducing Plagl2 expression and Dyrk1a knockdown (KD) was injected into the hippocampal dentate gyrus of 18-mo-old mice, and at 4 wk postinfection (wpi; at 19 mo of age), the hippocampal dentate gyrus was examined immunohistochemically. Virus-infected cells were mCherry+. (C, D) Quantification of DCX+dendrite+ immature neuron formation in the hippocampal dentate gyrus of 6-mo-old mice at 2wpi (C; n = 3) and 19-mo-old mice at 4wpi (D; n = 4). The upper and lower names along the horizontal axis indicate overexpressed and knockdown genes, respectively. [E, G, H] Control or iPaD lentivirus was injected into the hippocampal dentate gyrus of 18-mo-old mice, and 2 or 4 wk later, the hippocampal dentate gyrus was examined immunohistochemically. Yellow arrowheads indicate virus-infected mitotic cells [E, mCherry’BrdU’MCM2+], active NSCs [G; mCherry’BrdU’GFAP’Sox22+], and immature neurons [H; three examples of mCherry’BrdU’DCX’dendrite” cells], while white arrowheads indicate mCherry”BrdU” (non-virus-infected) cells. (F) Quantification of quiescent NSCs [MCM2−] and active NSCs or IPCs [MCM2+ or BrdU-incorporating] in the hippocampal dentate gyrus at 2 wk after control and iPaD lentivirus injection of 18-mo-old mice. BrdU was administered intraperitoneally for 7 d until sacrifice. At least three samples were examined for each condition. Each value represents the mean ± SEM. Scale bars: B, 50 µm; E, G, H, 20 µm.
Among them, examined 2 or 4 wk later (Fig. 1A; Supplemental Fig. S2G) 6- and 18- or 20-mo-old mice, and brain sections were examined 2 or 4 wk later [Fig. 1A; Supplemental Fig. S2G–J]. Among them, Cdktl, Prkc, and Dyrk1a shRNA similarly increased the number of MCM2+ active NSCs/IPCs in these mice [Supplemental Fig. S2G–J], but Dyrk1a shRNA also more efficiently increased the number of DCX+/dendrite+ cells than others in both 6- and 19-mo-old mice [Fig. 1B–D]. Compared with this combination, Dyrk1a knockdown alone did not effectively increase MCM2+ or DCX+dendrite+ cells [Fig. 1C, Supplemental Fig. S2I]. These data indicated that the combination of overexpression of Plagl2 [pleiomorphic adenoma gene-like 2], a zinc finger transcription factor, and knockdown of Dyrk1a [dual specificity tyrosine phosphorylation-regulated kinase 1a], a Down syndrome-related kinase, most efficiently activated neurogenesis in the hippocampal dentate gyrus of both 6- and 19-mo-old mice, and we named this combination iPaD [inducing Plagl2 and anti-Dyrk1a activity].

We first examined the short-term effects of lentivirus with iPaD under the control of the Hes5 promoter [iPaD lentivirus] on aged NSCs. Two weeks after the control or iPaD virus infection at 18 mo of age, very few were activated by the control virus, whereas more than half of the iPaD virus-infected cells were activated [MCM2+ and/or BrdU+] [Fig. 1E,F], which included Sox2+,GFAP+ active NSCs [Fig. 1G, arrowhead] and DCX+,dendrite+ immature neurons [Fig. 1H, arrowheads]. Many of the iPaD virus-infected MCM2+BrdU− [nonactivated] cells were astrocytes, which did not become active or proliferative by iPaD. When the iPaD lentivirus was injected into nonneurogenic brain regions, many astrocytes were labeled because the Hes5 promoter is active in astrocytes. However, none of the virus-infected cells expressed MCM2 or DCX [Supplemental Fig. S3D]. We also examined the effects of iPaD in the LV-SVZ and found that NSCs in this region were activated similarly to those in the DG-SGZ [Supplemental Fig. S3A–C]. These results suggest that iPaD can specifically activate NSCs.

Long-term effect of iPaD on neurogenesis

To examine the long-term effect of the iPaD lentivirus on neurogenesis in the aged brain, we introduced it into the hippocampal dentate gyrus of 18-mo-old mice. At this age, there were only a few proliferative cells [MCM2+], including activated NSCs and IPCs, and immature neurons [DCX+,dendrite+] in this region, and the control lentivirus did not affect their numbers [Fig. 2A,B [panel B1],C,D]. In contrast, the iPaD lentivirus infection activated NSCs efficiently and induced the formation of immature neurons 1 mo after infection [19 mo of age] [Fig. 2A,B [panel B2],C,D]. A similar effect was observed even 8 and 12 wk after infection [20 and 21 mo of age, respectively] [Fig. 2B [panels B3,B4],C,D], suggesting that neurogenesis is activated continuously by the iPaD lentivirus in the aged brain. Plagl2 alone also activated neurogenesis, but less efficiently than the iPaD lentivirus [Fig. 2A,C,D]. These results suggested that NSCs activated by the iPaD lentivirus proliferate continuously and are not exhausted even after 3 mo. In wild-type hippocampi, the number of MCM2+ or DCX+ cells was significantly decreased during aging and mostly disappeared at 18 mo of age [Fig. 2E]. However, the iPaD lentivirus increased the number of these cells in the hippocampal dentate gyrus of the aged brain [19 mo of age] to a level comparable with 9-mo-old or even younger wild-type mice [Fig. 2E], suggesting that neurogenesis in the aged brain can be reactivated effectively by the iPaD lentivirus.

Improved cognitive function by iPaD

To examine whether activated neurogenesis improves the cognitive function of aged mice, we analyzed spatial learning and memory with the Barnes maze test. Using this test, it was shown previously that spatial learning and memory are dependent on neurogenesis in the hippocampal dentate gyrus [Imayoshi et al. 2008], and that aged mice exhibit poorer performance than young mice, requiring a longer distance and latency to reach the target hole in the training session [Shoji and Miyakawa 2019]. We used two groups of 19-mo-old mice—one injected with the control lentivirus and the other injected with the iPaD lentivirus 1 mo before the test [Fig. 2F]. We found that those injected with the iPaD lentivirus reached the target hole via shorter distances, made fewer errors, and had shorter latencies than those injected with the control lentivirus [Fig. 2G–I; Supplemental Fig. S4A–C]. In the first probe test [1 d after training], iPaD mice stayed longer at the target hole than control mice [Fig. 2J, Supplemental Fig. S4D]. Furthermore, in the second probe test [7 d after retraining], iPaD mice still stayed longer around the target region than the other regions, whereas control mice did not [Supplemental Fig. S4E,F]. These data indicated that injection of the iPaD lentivirus improves the performance of aged mice in the Barnes maze test.

We next tested novel object recognition memory, which is dependent on hippocampal activity [Clark et al. 2000]. Mice from both groups were individually exposed to two copies of an identical object in the testing box for 10 min and then returned 24 h later to the same box containing one new but identical copy of the original object alongside one novel object [Supplemental Fig. S4G]. iPaD mice visited the novel object more often and spent a longer time investigating it than control mice [Supplemental Fig. S4H, I], suggesting that recognition memory is improved by injection of the iPaD lentivirus. Taken together, these data indicated that injection of the iPaD lentivirus activates neurogenesis in the aged brain and improves learning and memory ability.
Clonal analysis of NSCs with iPaD

The above results indicated that iPaD can activate functional neurogenesis in the aged brain. We next examined to what extent the proliferative and neurogenic potential of NSCs can be rejuvenated by iPaD. To this end, we performed a sparse labeling approach using a lentivirus expressing the tamoxifen-inducible Cre recombinase CreERT2 together with Plagl2 and anti-Dyrk1a activity under the control of the Hes5 promoter (iPaD-CreERT2 lentivirus) (Fig. 3A). As a control, we used a lentivirus expressing CreERT2 only under the control of the Hes5 promoter (control-CreERT2 lentivirus) (Fig. 3A). These viruses were introduced into the hippocampal dentate gyrus of 8- and 18-mo-old Ai14 mice, in which Cre recombinase labels infected not only NSCs (Fig. 3B) but also their progeny with tdTomato (Madisen et al. 2010). At day 5 after virus infection, a low dosage of tamoxifen was administered for 3 d for sparse labeling of NSCs, and brain sections were examined 3 wk later (Fig. 3A). To determine the virus-infected cell types, we first examined control-CreERT2 lentivirus-infected brains at 1 wk after tamoxifen treatment. We found that ~40% of tdTomato+ cells exhibited radial glia-like (RGL) morphology and expressed GFAP and Sox2, indicating that they were quiescent NSCs (Supplemental Fig. S5A,A′). The other tdTomato+ cells were mostly isolated single astrocytes (Supplemental Fig. S5A,A′). Similarly, 3 wk after tamoxifen treatment, 30%–50% of the tdTomato+ clones contained RGL cells (Fig. 3B), while the others were isolated single astrocytes. As the Hes5 promoter is also active in astrocytes, it is likely that those isolated single tdTomato+ astrocytes were infected directly with the virus rather than differentiating from virus-infected NSCs. Therefore, tdTomato+ single astrocytes were excluded from the clonal analysis.

To determine the composition of each clone derived from single NSCs, we examined virus-infected brains 3 wk after the last day of tamoxifen treatment (Fig. 3A),...

Figure 2. Long-term effect of the iPaD lentivirus on neurogenesis. (A) Schematic structures of the control, Plagl2, and iPaD lentiviruses. [B] The control [panel B1] or iPaD [panels B2–B4] lentivirus was injected into the hippocampal dentate gyrus of 18-mo-old mice. Immuno-histochemical analysis was performed at 8wpi of the control lentivirus (20 mo of age; panel B1) and at 4wpi (19 mo of age; panel B2), 8wpi (20 mo of age; panel B3), or 12wpi (21 mo of age; panel B4) of iPaD lentivirus. (C,D) Quantification of active NSCs/IPCs [MCM2+; C] and immature neurons [DCX2+dendrite+; D] in the hippocampal dentate gyrus after virus injection at 18 mo of age. Two-way ANOVA with Bonferroni’s post hoc test. Non-virus-infected cells were included for quantification, but virtually all MCM2+ or DCX2+dendrite+ cells at 20 and 21 mo of age were mCherry+. [E] Quantification of MCM2+ active NSCs/IPCs and DCX2+dendrite+ immature neurons in the wild-type hippocampal dentate gyrus at 6, 9, 12, and 18 mo of age. Quantification at 4wpi of iPaD lentivirus injection into 18-mo-old mice (19 mo of age) is shown at the right. Least four samples were examined for each condition. One-way ANOVA with Tukey HSD post hoc test was conducted. Each value represents the mean ± SEM. Scale bar, 50 µm. (F) Schedule of Barnes maze test. Control or iPaD lentivirus was injected into the hippocampal dentate gyrus of 18-mo-old male mice. N = 10 for control, N = 10 for iPaD male mice. (G–I) Quantification of the walking distance (G), latency (H), and errors (I) before entering the target hole. Error bars indicate SEM. (J) Time spent around the target hole was measured at the probe test. Error bars indicate SEM. [∗∗] P < 0.01, [∗∗∗] P < 0.001, one-way ANOVA with Tukey’s HSD test.
Figure 3. Clonal analysis of iPAd lentivirus-infected NSCs. (A) Strategy for clonal analysis using Ai14 mice. Schematic structures of the control-CreERT2 and iPAd-CreERT2 lentiviruses and the Rosa26 locus of Ai14 mice (bottom left), and the schedule of experiments for sparse labeling of NSCs (bottom right). (B) Quantification of clones containing RGL cells. (C) Immunohistological and morphological analyses of sparsely labeled cells (tdTomato+). (D) Immunohistological and morphological analyses of sparsely labeled cells (tdTomato+) from control-CreERT2 (top two panels) and iPAd-CreERT2 (bottom three panels) lentivirus-infected hippocampal dentate gyrus. Cell types were determined by morphology and the following markers (indicated by yellow arrowheads): RGL cells (GFAP+Nestin+), astrocytes (GFAP+S100β+), IPCs (MCM2+DCX+dendrite–), immature neurons (MCM2–DCX+dendrite+) and mature neurons (DCX–Prox1+). (E–H) Quantification of the clonal composition of sparsely labeled NSCs with the control-CreERT2 (E,G) and iPAd-CreERT2 (F,H) lentiviruses in 9-mo-old (E,F) and 19-mo-old (G,H) Ai14 mice. The number in parentheses indicates the number of each clone. (R) RGL cell, (P) IPC, (N) immature and mature neuron, (A) astrocyte. The number of clones per hemisphere was 38.3 ± 2.7 at 9 mo of age and 43.5 ± 3.6 at 19 mo of age. (I) Clonal sizes of sparsely labeled NSCs at 9 mo of age (left) and 19 mo of age (right). One-way ANOVA with Tukey HSD post hoc test was conducted. (J) Quantification of DCX+ IPCs/neurons and astrocytes per activated clone. Six samples were examined for each condition. [N] IPCs and immature and mature neurons, [A] astrocyte. Two-way ANOVA with Bonferroni’s post hoc test was conducted. Each value represents the mean ± SEM. Scale bars: C (top), 100 µm C (bottom), D, 20 µm.
which exhibited sparse labeling [Fig. 3C; Supplemental Fig. S5A]. We used immunohistochemical and morphological markers to identify RGL cells, IPCs, immature and mature neurons, and astrocytes [Fig. 3D; Supplemental Fig. S5D,E] and characterized a total of 93 and 91 clones from the control-CreER T2 lentivirus-infected and iPaD-CreER T2 lentivirus-infected brains, respectively, at 9 mo of age (at 4 wk postinfection of 8-mo-old brains). In control-CreER T2 lentivirus-infected brains, 68.8% [64 out of 93] of the clones consisted of one RGL cell [Fig. 3E], while only 31.2% [29 out of 93] of the clones consisted of two or more cells [Fig. 3E], indicating that the majority of the control virus-infected cells remained as quiescent NSCs. In agreement with this notion, the average clonal size was 1.5 cells/clone after 3-wk tracing. Among the clones derived from activated NSCs (n = 29), 13.8% [four out of 29] consisted of two RGL cells [R + R] [Fig. 3E], suggesting that symmetric cell division occurs occasionally at this stage. Among these activated clones (n = 29), only 13.8% [2P + N] + 2[R + N] + 4] contained DCX+ IPCs or neurons, while ~72.4% [10[A + A] + 11[R + A] + 21] contained astrocyte[s] [Fig. 3E]. These results indicated that wild-type NSCs exhibit low proliferative/neurogenic potential at 9 mo of age [Fig. 3E]. In contrast, in iPaD-CreER T2 lentivirus-infected brains, only 19.8% [18 out of 91] of the clones consisted of one RGL cell [Fig. 3F], while 80.2% [73 out of 91] of the clones contained two or more cells [Fig. 3F], indicating that the majority of the virus-infected NSCs were activated. In agreement with this notion, the average clonal size was 3.6 cells/clone after 3-wk tracing. Among the clones derived from activated NSCs (n = 73), 56.2% [2N + 5[P + N] + 2[N + A] + 21[R + P] + 9[R + N] + 2[R + N + A] = 41] contained DCX+ IPCs or neurons, while 34.2% [2[N + A] + 7[A + A] + 2[R + N + A] + 14[R + A] + 25] were astrogenic [Fig. 3F]. These results indicated that the iPaD lentivirus substantially enhances the proliferative (~2.4-fold increase) and neurogenic (~4.1-fold increase) potential of NSCs at 9 mo of age [Fig. 3J].

We next characterized a total of 78 and 97 clones from control-CreER T2 lentivirus-infected and iPaD-CreER T2 lentivirus-infected brains, respectively, at 19 mo of age (at 4 wk postinfection of 18-mo-old brains). In control-CreER T2 lentivirus-infected brains, 85.9% [67 out of 78] of the clones consisted of one RGL cell [Fig. 3G], indicating that the majority of the control virus-infected cells remained as quiescent NSCs. Furthermore, the other clones were mostly astrogenic, and no clones contained neurons, while very few [2.6%, two out of 78] contained IPCs [Fig. 3G]. The average clonal size was 1.2 cells/clone after 3-wk tracing, which is similar to that at 9 mo of age. These results indicated that NSCs exhibit very low proliferative and virtually no neurogenic potential at 19 mo of age [Fig. 3J]. In contrast, in iPaD-CreER T2 lentivirus-infected brains, only 26.8% [26 out of 97] of the clones consisted of one RGL cell, while 73.2% [71 out of 97] of the clones consisted of two or more cells [Fig. 3H], indicating that many virus-infected NSCs were activated. Among the clones derived from activated NSCs (n = 71), 71.8% [2P + 3[P + N] + 32[R + P] + 12[R + N] + 2[R + N + A] + 51] contained DCX+ IPCs or neurons, while 26.8% [8[A + A] + 2[R + N + A] + 9 [R + A] = 19] were astrogenic [Fig. 3H]. The average clonal size was 4.1 cells/clone for 3-wk tracing. These results indicated that the iPaD lentivirus endows nonproliferative, nonneurogenic NSCs with proliferative and neurogenic potential at 19 mo of age [Fig. 3J]. The clonal size of iPaD lentivirus-infected 19-mo-old NSCs was even larger than 4.1 cells/clone, if only activated NSCs are considered (~4.9 neurons/clone in 3-wk tracing) [Fig. 3J], and this value was comparable with or larger than that of wild-type activated [Ascl1+] NSCs at 2 mo of age, which generated, on average, 4.8 neurons in 2-mo tracing [Pilz et al. 2018]. These results suggest that iPaD can rejuvenate 19-mo-old NSCs to function like 2-mo-old or even younger NSCs.

Gene expression analysis of iPaD-dependent rejuvenation of aged NSCs

To characterize the rejuvenation ability of iPaD, we analyzed the senescence-associated factors β-galactosidase, p16Ink4a, and p19Arf (Molofsky et al. 2006; Nishino et al. 2008). Their expression was up-regulated in aged NSCs compared with young NSCs but repressed by iPaD, indicating that iPaD can reverse the aging process of NSCs [Fig. 4A–D]. It was previously shown that Plagl2 enhances the self-renewal of telencephalic neural progenitors and glioblastoma cells [Zheng et al. 2010; Adnani et al. 2018], while a proper dosage of Dyrk1a, a gene associated with Down syndrome, which accelerates aging [Francesi et al. 2019], is important for normal neurogenesis [Fotaki et al. 2002], but the mechanism by which iPaD rejuvenates aged NSCs is unclear. To investigate the mechanism, we next performed RNA-seq analysis of E13, young [1-mo-old], adult [9-mo-old], and aged [19-22-mo-old] NSCs infected with the control or iPaD virus [Fig. 4E; Supplemental Table S3]. Because of the technical difficulty in obtaining NSCs from DG-SGZ in the aged mice, we used NSCs from LV-SVZ. We confirmed that Plagl2 expression is down-regulated in aged NSCs but up-regulated by iPaD, while Dyrk1a expression is up-regulated in aged NSCs but down-regulated by iPaD [Supplemental Fig. S6A]. Principal component [PC] analysis showed a high reproducibility between biological duplicates or triplicates of each sample and dynamic transcriptomic changes caused by aging [Fig. 4F], which exhibited a high similarity to the previously published data [Supplemental Fig. S6B; Kalamakis et al. 2019]. Interestingly, the transcriptomic profiles of aged NSCs became very similar to those of 1-mo-old NSCs after infection with the iPaD virus, agreeing with the above observation that iPaD can rejuvenate the proliferative and neurogenic potential of aged NSCs in the DG-SGZ [Fig. 4F; Supplemental Fig. S6C]. The gene ontology [GO] terms of PC3, which represent iPaD-induced rejuvenation, included genes involved in development, cell fate commitment, and response to interferon-β [Supplemental Fig. S6D,E]. Analysis by c-means clustering revealed eight patterns of transcriptomic changes in NSCs from embryonic to aged stages [Fig. 4G; Supplemental Fig. S6F]. GO enrichment analysis indicated that genes involved in cell proliferation, such as Ccnd, Cdk4, and E2f1, are enriched in

Rejuvenation of aged neural stem cells

GENES & DEVELOPMENT 29
clusters 2 and 3 (Fig. 4H,J; Supplemental Fig. S6G), whose expression is down-regulated in aged NSCs compared with embryonic NSCs but up-regulated by iPAd (Fig. 4G). In contrast, genes involved in aging such as Tgfb2 and Igfbp2 are enriched in cluster 6 (Fig. 4I,J; Supplemental Fig. S6H), whose expression is up-regulated in aged NSCs but repressed by iPAd (Fig. 4G). These results suggest that iPAd can rejuvenate aged NSCs by up-regulating embryonic-high genes and repressing aged-high genes.

Further gene expression analyses showed that iPAd down-regulated quiescence-related genes such as Id4 [Blomfield et al. 2019; Zhang et al. 2019] and up-regulated activation-related genes such as Ascl1 [Fig. 4I; Supplemental Fig. S6I,J; Andersen et al. 2014; Sueda et al. 2019]. Ascl1 expression is mostly negative in the aged brain [Fig. 5A], but subsets of iPAd lentivirus-infected cells expressed Ascl1 protein after 4 wk of infection in 19-mo-old hippocampal dentate gyrus, whereas none of
the control lentivirus-infected cells expressed Ascl1 [Fig. 5B]. Furthermore, Ascl1 expression in cultured quiescent NSCs was up-regulated by iPaD but not by the control (Supplemental Fig. S7), indicating that iPaD can induce Ascl1 protein expression in quiescent NSCs. We next examined whether iPaD lentivirus-infected cells express Ascl1 in an oscillatory manner, a hallmark feature of activated NSCs (Imayoshi et al. 2013; Sueda et al. 2019). The iPaD lentivirus was injected into the dentate gyrus of 10-mo-old Ascl1 reporter mice, and 4 wk after injection (11 mo of age), time-lapse imaging of brain slices was performed. Ascl1 expression induced by the iPaD lentivirus was oscillatory [Fig. 5C], suggesting that iPaD is able to maintain active NSCs in the aged brain by inducing dynamic Ascl1 expression for an extended period of time.

Epigenetic analysis of iPaD-dependent rejuvenation of aged NSCs

Because Plagl2 is a zinc finger transcription factor, we next performed Plagl2 chromatin immunoprecipitation assays with sequencing (ChIP-seq) of HA-tagged Plagl2-overexpressing NSCs to identify its downstream targets. Motif enrichment analysis revealed the consensus sequence GGGGCCC recognized by Plagl2, which matched the previously characterized Plagl1/Zac1-binding sites [Fig. 6A; Hensen et al. 2002; Varrault et al. 2017], and these Plagl2-binding regions were identified in the promoter of 4731 genes [Supplemental Table S4]. GO terms involved in gene regulation and chromatin modifications were enriched in genes containing Plagl2-binding sites and up-regulated by iPaD [Fig. 6B]. Expression of chromatin-modifying enzymes and DNA methylation enzymes decreased with aging but were recovered by iPaD [Fig. 6C, D]. More than 70% of chromatin-modifying enzymes belong to clusters 2 and 3, and many of them contain Plagl2-binding regions around their promoters [Supplemental Fig. S8]. These results suggest that iPaD regulates rejuvenation by changing the chromatin accessibility.

To examine this possibility, we performed an assay for transposase-accessible chromatin by sequencing (ATAC-seq) to identify the age-associated changes in global chromatin accessibility and examined whether aged profiles can be reversed by iPaD. It was previously reported that chromatin accessibility profiles were distinct between embryonic and adult NSCs (Berg et al. 2019), raising the

**Figure 5.** Induction of Ascl1 expression by the iPaD lentivirus. (A) In situ hybridization of Ascl1 in the hippocampal DG-SGZ and the LV-SVZ of 1.5- and 17-mo-old mice. (B) The iPaD or control lentivirus was injected into the hippocampal dentate gyrus of 18-mo-old mice, and 4 wk later, the infected regions were examined immunohistochemically. Subsets of iPaD lentivirus-infected cells expressed Ascl1 (arrowhead), whereas none of the control lentivirus-infected cells did. (C) Bioluminescence imaging and quantification of Luc2-Ascl1 levels in iPaD lentivirus-infected cells. The iPaD lentivirus was injected into the hippocampal dentate gyrus of 10-mo-old Ascl1 reporter mice, in which Luc2-Ascl1 fusion protein was expressed from the endogenous Ascl1 locus. At 4 wk after injection (11 mo of age), time-lapse imaging of brain slices was performed. Luc2-Ascl1 expression occurred in iPaD virus-infected cells [arrowheads]. Three representative cells were quantified. Scale bars: A, 200 µm; B, 30 µm; C, 100 µm.
possibility that they could be further changed during aging. ATAC-seq analysis of embryonic (E14.5), adult (6-mo-old), and aged (19-20-mo-old) NSCs showed highly correlated signals between biological duplicates in embryonic, adult, and aged NSCs (\(r = 1.00\) in embryonic, \(r = 0.97\) in adult, and \(r = 0.99\) in aged NSCs, Pearson correlation coefficient) (Supplemental Fig. S9A). PCA also showed a high reproducibility between biological duplicates of each sample and revealed that chromatin accessibility profiles were significantly different between adult and aged NSCs, suggesting that the chromatin landscapes dynamically change during aging (Supplemental Fig. S9B). Analysis by \(k\)-means clustering revealed six patterns of chromatin landscape changes in NSCs from embryonic

Figure 6. The epigenetic mechanism of iPaD-dependent rejuvenation of aged NSCs. (A) Motif enrichment analysis of Plagl2 ChIP-seq. (B) GO analysis of genes up-regulated by iPaD and containing Plagl2-binding sites. (C,D) Changes of chromatin/DNA modification gene expression. (E) Analysis by \(k\)-means clustering of ATAC-seq signals around promoters of embryonic, adult, and aged NSCs revealed six clusters. (F) Proportions of Plagl2 and Ascl1 target genes in each cluster. (G) Changes of chromatin accessibility in aged NSCs by iPaD. In aged NSCs, closed genes containing Plagl2-binding sites exhibited higher accessibility by iPaD, while opened genes containing Plagl2-binding sites exhibited lower accessibility by iPaD. (H,I) RNA-seq, ATAC-seq, and Plagl2 ChIP-seq enrichment profiles of Nestin (H) and Gfap (I).
to aged stages [Fig. 6E]. Notably, 67.9% and 11.3% of the embryonic-high genes belonged to clusters A and B, respectively, whose chromatin structures are open in embryonic NSCs but mostly inaccessible in aged NSCs, suggesting that a total of 79.2% of the embryonic-high genes decrease chromatin accessibility during aging [Supplemental Fig. S9C,D]. Among these genes, Plagl2- and Ascl1-binding regions are enriched [Fig. 6F,G; Supplemental Fig. S9E], suggesting that Plagl2 and iPaD-inducible Ascl1 are involved in up-regulation of such embryonic-high genes whose chromatin structures are less accessible in aged NSCs. In agreement with this notion, iPaD preferentially up-regulated Plagl2 target genes compared with nontarget genes irrespective of their chromatin accessibility in aged NSCs [Supplemental Fig. S9F]. Furthermore, while iPaD increased chromatin accessibility (>1.5-fold) of 945 genes in aged NSCs, 628 and 168 (84.2%) of them belonged to clusters A and B, respectively. These results suggest that iPaD can reopen many genes whose chromatin structures become less accessible with aging. Indeed, the active NSC-specific gene Nestin exhibited high RNA expression and chromatin accessibility at an embryonic stage and very low RNA expression and chromatin accessibility at an aged stage, but chromatin accessibility of Plagl2 binding regions and RNA expression increased in aged NSCs by iPaD [Fig. 6H]. Similarly, in genes involved in proliferation of NSCs, such as E2f1, Tcf3, and Ascl1, RNA expression and chromatin accessibility decreased during aging but increased by iPaD [Supplemental Fig. S9G–I]. In contrast, in genes up-regulated during aging, such as Gfap, RNA expression and chromatin accessibility increased during aging but decreased by iPaD [Fig. 6I]. These results suggest that iPaD up-regulates many embryonic-high genes and down-regulates aged-high genes by controlling chromatin accessibility, thereby rejuvenating dormant NSCs.

**Discussion**

**Rejuvenation of aged NSCs by iPaD**

Aged NSCs are mostly dormant, and even when they are activated, they primarily produce astrocytes. Thus, aged NSCs lose their proliferative and neurogenic potential, leading to the cessation of neurogenesis. In this study, we showed that the iPaD lentivirus substantially rejuvenated the proliferative and neurogenic potential of NSCs in the aged brain. Clonal analysis by a sparse labeling approach as well as transcriptome analysis indicated that iPaD can rejuvenate aged NSCs [19–21 mo of age] to a level comparable with those at 1 or 2 mo of age and successfully improved cognition of aged mice. Once rejuvenated and activated by iPaD, aged dormant NSCs can generate, on average, 4.9 neurons but very few astrocytes in 3-wk tracing [Fig. 3J]. Furthermore, these activated NSCs were maintained for as long as 3 mo in the aged brain, suggesting that active neurogenesis continues for an extended period of time after iPaD treatment. Nevertheless, iPaD-activated neurogenesis gradually declined [Fig. 2D]. Furthermore, clonal analyses showed that 78.1% [9-mo-old] and 81.7% [19-mo-old] of iPaD-activated clones maintained RGL cells 4 wk after activation [Fig. 3F,H], suggesting that ~20% of activated NSCs are exhausted during this period. However, it is unknown whether this exhaustion is due to limitation of iPaD or loss of iPaD activity, and further analyses are required to answer this question.

A recent study showed that resting NSCs, those once proliferated but returned to quiescence [Urbán et al. 2016], are the major origin of active NSCs in the aged brain [Harris et al. 2021]. This population comprises only 3%–5% of the total NSCs, while the other major population is dormant NSCs, which have never proliferated [Harris et al. 2021]. Because the iPaD is able to activate 70%–80% of NSCs in the aged brain, it is likely that it mostly targets dormant NSCs, raising the possibility that the higher the infection efficiency of the iPaD virus, the more NSCs are activated to produce new neurons.

**The mechanism of NSC rejuvenation by iPaD**

During aging, the gene expression and accessible chromatin landscapes change dynamically in NSCs. Transcriptome analysis showed that there are eight clusters of genes showing different expression patterns. Among them, clusters 2 and 3, and 6 are of particular interest: In clusters 2 and 3, gene expression is down-regulated in aged NSCs compared with embryonic NSCs but up-regulated by iPaD, while in cluster 6, gene expression is up-regulated in aged NSCs but repressed by iPaD [Fig. 4G]. Genes involved in cell proliferation are enriched in clusters 2 and 3, while genes involved in aging are enriched in cluster 6. In addition, chromatin-modifying genes are also enriched in clusters 2 and 3. These results suggest that iPaD can rejuvenate aged NSCs by up-regulating embryonic-high genes and repressing age-associated genes via modulation of chromatin accessibility. To confirm this possibility, we also examined chromatin accessibility in aged NSCs. ATAC-seq analysis revealed that there are six clusters [A–F] of genes showing different patterns of chromatin landscape changes from embryonic to aged stages. The majority of embryonic-high genes belong to clusters A and B, whose chromatin structures are open in embryonic NSCs but mostly inaccessible in aged NSCs. Interestingly, many genes in clusters A and B contain Plagl2-binding sites as well as iPaD-induced Ascl1-binding sites, and therefore it is most likely that their expression is controlled directly or indirectly by iPaD. These results raised the possibility that iPaD up-regulates gene expression by increasing the chromatin accessibility in aged NSCs. In contrast, some aging-associated genes such as Gfap increased the chromatin accessibility in aged NSCs, but their transcription and chromatin accessibility are repressed by iPaD, suggesting that iPaD can also down-regulate gene expression by decreasing the chromatin accessibility in aged NSCs. Thus, iPaD can both up-regulate and down-regulate gene expression by modulating the chromatin accessibility, but the detailed mechanism by which iPaD differentially regulates the chromatin structures remains to be analyzed. In addition, while the chromatin/DNA modification gene expression patterns in
iPaD-treated NSCs are more similar to those of embryonic NSCs [Fig. 6C], their transcriptome is more similar to 1m quiescent NSCs [Fig. 4F], suggesting that additional factors, such as environmental conditions, may be required to further rejuvenate aged NSCs.

We also found that iPaD is able to induce Ascl1 oscillations, a hallmark feature of active NSCs [Imayoshi et al. 2013; Sueda et al. 2019], in the aged brain. We previously showed that Ascl1 has dual activities: It induces neuronal differentiation when its expression is continuous, whereas it activates proliferation of NSCs when its expression is oscillatory, indicating that Ascl1 oscillations are important for the active state of NSCs [Imayoshi et al. 2013; Sueda et al. 2019]. In aged NSCs, chromatin structures of Ascl1 are less accessible but become more widely accessible after iPaD treatment, suggesting that iPaD is involved in the chromatin accessibility of Ascl1. A single Ascl1-expressing cell was found in each mCherry+ cluster, which was most likely derived from a single iPaD lentivirus-infected NSC. This observation agrees well with the clonal analysis, which showed that most clusters derived from iPaD lentivirus-infected NSCs contained a single RGL cell [Fig. 3F,H], suggesting that these activated NSCs repeat an asymmetric cell division, in which one daughter cell remains as an NSC while the other starts neuronal or astrocytic differentiation.

It was previously reported that Plagl2 plays an important role in self-renewal of telencephalic neural progenitors [Adnani et al. 2018]. Indeed, we found that Plagl2 alone can increase cell proliferation [MCM2+] as efficiently as iPaD [Fig. 2C], suggesting that Plagl2 endows dormant NSCs with proliferative potential. Plagl2 alone generated fewer DCX+dendrite+ cells than iPaD but more than the control [Fig. 2D], suggesting that Plagl2 also has weak neurogenic as well as astrogenic activity. It was also reported that Plagl2 functions as a proto-oncogene by activating Wnt signaling in gliomas [Zheng et al. 2010]. However, we observed no tumor formation in the brain 3 mo after iPaD lentivirus infection, suggesting that iPaD cannot activate tumorigenesis. Furthermore, whereas the Wnt ligand gene Wnt6 was up-regulated by Plagl2 in p53-deficient NSCs [Zheng et al. 2010], it was not much changed in wild-type NSCs by the iPaD lentivirus. These results suggest that iPaD is not involved in tumor formation in the brain, and that the downstream events may be different, depending on the cellular context.

iPaD significantly increased the number of DCX+dendrite+ cells compared with Plagl2 alone [Fig. 2D], suggesting that inhibiting Dyrk1a endows NSCs with neurogenic potential, although its mechanism remains to be analyzed. Dyrk1a is known to prevent nuclear entry of the transcription factors NFATc by phosphorylation and thereby block NFATc activity [Arron et al. 2006], suggesting that iPaD can up-regulate NFATc activity. Interestingly, NFATc4 is included in the embryonic-high genes; however, overexpression of NFATc4 did not efficiently activate quiescent NSCs [Supplementary Table S1]. Another NFATc member, NFATc1, is included in the adult-high genes, but knockdown of NFATc1 did not efficiently activate quiescent NSCs [Supplementary Fig. S2C; Supplementary Table S2]. Thus, NFATc factors may not be involved in iPaD-induced neurogenesis, and further analyses will be required to identify the target proteins for Dyrk1a.

Possible application of iPaD to regenerative medicine

While the mechanism of aging has been intensively analyzed, how to reverse the aging process is still a difficult challenge to pursue [Negredo et al. 2020]. It has been shown that the activation of neurogenesis ameliorates many age-related neurodegenerative disorders, such as Alzheimer’s disease and Huntington’s disease [Benraiss et al. 2013; Choi et al. 2018; Díaz-Moreno et al. 2018], and our method may offer a new strategy to rejuvenate brain activity for treatment of neurodegenerative disorders. Since Dyrk1a increases amyloid-β production [Ryoo et al. 2008], it is possible that iPaD arrests progression of Alzheimer’s disease by inhibiting amyloid-β accumulation while replenishing neurons. Conflicting results about neurogenesis in the adult human hippocampus have been reported [Kempermann et al. 2018], yet even if most NSCs are quiescent/dormant in aged humans, our results suggest that such NSCs could be rejuvenated to produce many new neurons continuously, showing the applicability of our strategy to neuronal regeneration technology.

Materials and methods

Animals

All mice were handled in accordance with the Kyoto University Guide for the Care and Use of Laboratory Animals. The experimental protocols were approved by the Experimental Animal Committee of the Kyoto University. Wild-type mice (C57BL/6J) were purchased from SLC. Ai14 mice were obtained from Jackson Laboratory.

Lentivirus infection of mice

For screening and neurogenesis induction, 1 µL of lentivirus (3.75 × 10⁴ U/mL) was delivered with a flow rate of 0.125 µL/min stereotactically into the dentate gyrus bilaterally with the following coordinates: anteroposterior = −2.15 mm from bregma, lateral = ±1.85 mm, and ventral = 2.2 mm. For sparse labeling, 1 µL of lentivirus (5.0 × 10⁴ U/mL) was delivered. Brain sections were immunohistoologically examined.

Sparse labeling of NSCs

We administered 0.25 mL of tamoxifen (Tam; 20 mg/mL in corn oil, Sigma) by oral gavage 5 d after injection for three consecutive days to activate CreER²2 activity sparsely in Ai14 mice injected with the control-CreER²2 or iPaD-CreER²2 lentivirus. Brain sections were examined 1 or 3 wk later. To determine the virus-injected cell types, we first examined control-CreER²2 lentivirus-infected brains at 1 wk after tamoxifen treatment. We found that ~40% of tdTomato+ cells exhibited radial glia-like (RGL) morphology and expressed GFAP and Sox2, indicating that they were quiescent NSCs [Supplementary Fig. S5A,A′,B]. The other tdTomato+ cells were mostly isolated single astrocytes [Supplementary Fig. S5A,A′,C]. Similarly, at 3 wk after tamoxifen
treatment, 30%–50% of the tdTomato+ clones contained RGL cells (Fig. 3B), while the others were isolated single astrocytes. As the Hes5 promoter is also active in astrocytes, it is likely that those isolated single tdTomato+ astrocytes were infected directly with the virus rather than differentiating from virus-infected NSCs. Therefore, tdTomato+ single astrocytes were excluded from the clonal analysis.

**Time-lapse imaging of brain slices**

Time-lapse imaging of brain slices was performed as described previously [Sueda et al. 2019]. Coronal brain slices (150-µm thickness) were cultured in a medium (135 mM NaCl2, 5 mM KCl, 10 mM HEPES, 1 mM CaCl2, 1 mM MgCl2, 5% horse serum, 5% fetal bovine serum) containing 1 mM luciferin at 37°C in 5% CO2 and 80% O2. Bioluminescence was acquired using a CCD camera (iKon-M DU934P-BV, Andor).

**Senescence-associated β-galactosidase assay**

Activity of β-galactosidase was detected by senescence cell histochemical staining kit (Sigma CS00300). NSCs were cultured in an eight-well chamber (LAB-TEK 154534) with PLO coating. The schedule of lentivirus addition, medium change, and sample fixation was the same as transcriptome analyses. NSCs were fixed by 1× PFA solution for 7 min. To detect the activity of β-galactosidase, NSCs were incubated for 3 h at 37°C and reaction was stopped by replacing the reaction mixture with PBS. Twenty-month-old quiescent NSCs and 20-mo-old iPd NSCs were cultured in the same eight-well chambers but different wells to ensure that those isolated single tdTomato+ astrocytes were infected directly with the virus rather than differentiating from virus-infected NSCs. Therefore, tdTomato+ single astrocytes were excluded from the clonal analysis.

**Western blot**

Western blot was performed as described previously [Kobayashi et al. 2019]. NSCs in a 12-well plate were collected by plastic cell scraper and lysed by RIPA buffer with protease inhibitor cocktail (Merck 1169749801) for 30 min at 4°C. Protein concentrations of total cell lysate were measured by the DC protein assay, and 5 µg of protein from each sample was loaded for SDS-PAGE. Western blots were visualized by chemiluminescence using Amersham ECL (GE Healthcare) or ECL prime (GE Healthcare), quantified on an LAS3000 image analyzer (Fujifilm), and normalized against the corresponding intensity of α-Actin. The following primary antibodies were used for Western blotting: rabbit anti-Actin (Sigma), rabbit anti-p16INK4a (Abcam), and rabbit anti-p19ARF (Abcam). Secondary antibodies were HRP-conjugated antirabbit antibody (GE Healthcare).

**RNA in situ hybridization**

RNA in situ hybridization was performed using a digoxigenin-labeled Ascl1 antisense RNA probe, as described previously [Imayoshi et al. 2013].

Other detailed experimental procedures are in the Supplemental Material.

**Competing interest statement**

A national [Japanese] patent application (2021-033464) has been filed by Kyoto University.

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**Author contributions:** T. Kaise performed the experiments, analyzed the data, and wrote the manuscript. M.F. performed in vitro screening. R.P. performed Ascl1 expression analysis. W.P. and T. Kobayashi examined chromatin accessibility. M.Y. and I.I. analyzed the data. R.K. designed and supervised the project and wrote the manuscript.

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