Neural Stem Cell Grafting Counteracts Hippocampal Injury-Mediated Impairments in Mood, Memory, and Neurogenesis

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ABSTRACT

The hippocampus is vital for functions such as mood and memory. Hippocampal injury typically leads to mood and memory impairments associated with reduced and aberrant neurogenesis in the dentate gyrus. We examined whether neural stem cell (NSC) grafting after hippocampal injury would counteract impairments in mood, memory, and neurogenesis. We expanded NSCs from the anterior subventricular zone (SVZ) of postnatal F344 rat pups expressing the human placental alkaline phosphatase and grafted them into the hippocampus of young adult F344 rats at 5 days after an injury inflicted through a unilateral intracerebroventricular administration of kainic acid. Analyses through forced swim, water maze, and novel object recognition tests revealed significant impairments in mood and memory function in animals that underwent injury and sham-grafting surgery. In contrast, animals that received SVZ-NSC grafts after injury exhibited mood and memory function comparable to those of naive control animals. Graft-derived cells exhibited excellent survival and permissive migration, and they differentiated into neurons, subtypes of inhibitory GABAergic interneurons, astrocytes, oligodendrocytes, and oligodendrocyte progenitors. Significant fractions of graft-derived cells also expressed beneficial neurotrophic factors such as the glial cell line-derived neurotrophic factor, brain-derived neurotrophic factor, fibroblast growth factor, and vascular endothelial growth factor. Furthermore, SVZ-NSC grafting counteracted the injury-induced reductions of graft-derived cells also expressed beneficial neurotrophic factors such as the glial cell line-derived neurotrophic factor, brain-derived neurotrophic factor, fibroblast growth factor, and vascular endothelial growth factor. Furthermore, SVZ-NSC grafting counteracted the injury-induced reductions and abnormalities in neurogenesis by both maintaining a normal level of NSC activity in the subgranular zone and providing protection to reelin+ interneurons in the dentate gyrus. These results underscore that early SVZ-NSC grafting intervention after hippocampal injury is efficacious for thwarting mood and memory dysfunction and abnormal neurogenesis.

INTRODUCTION

The hippocampus is vital for cognitive and mood function [1–3]. It is also a brain area well known for postinjury plasticity [4]. Hippocampal injury associated with neurodegeneration can ensue through multiple causes, including head injury [5], ischemia [6], acute seizures [7], and severe stress [8]. Among hippocampal alterations in the early postinjury period, increased neurogenesis from neural stem cells (NSCs) and upregulation in the concentration of neurotrophic factors are conspicuous [9–13]. Although the implications of these plastic changes are still being examined, it is believed that these changes signify compensatory mechanisms to lessen the overall hippocampal dysfunction. Nonetheless, hippocampal injury leads to mood and memory impairments months after injury [14–16], which are allied with reduced NSC proliferation in the neurogenic subgranular zone (SGZ) of the dentate gyrus (DG) and aberrant hippocampal neurogenesis. Abnormal hippocampal neurogenesis is typified by both reduced incorporation of newly born neurons into the dentate granule cell layer (GCL) and abnormal migration of newly born neurons into the dentate hilus [10, 14]. These changes are associated with reduced concentration of multiple neurotrophic factors important for neurogenesis [10, 13] and loss of DG interneurons secreting reelin, an extracellular matrix protein that controls newly born dentate granule cell migration [17, 18].

From the above perspectives, interventions that are competent for averting the evolution of initial hippocampal injury into mood and memory impairments have significance. In particular, therapeutic strategies that have promise for maintaining the normal extent and pattern of neurogenesis in the injured hippocampus are of great interest. This is because hippocampal neurogenesis is considered to be vital for functions
such as mood and memory [2, 19, 20], and the aberrant neurogenesis that ensues after injury is believed to contribute to mood and memory dysfunction, as well as dentate hyperexcitability [21, 22]. In this context, NSC transplantation therapy appears to be a good candidate for ameliorating hippocampal injury-induced impairments, as these cells have the ability to survive, migrate, and engraft into brain regions exhibiting neuron loss [23]. Furthermore, NSCs can contribute new neurons, including the inhibitory GABAergic interneurons, introduce new astrocytes that are capable of secreting neurotrophic factors [24], and improve neurogenesis through stimulation of the proliferation of endogenous NSCs in the neurogenic SGZ [25]. Although NSCs can be obtained from a variety of sources, we chose subventricular zone (SVZ)-derived NSCs as donor cells because of the feasibility of their expansion in culture for extended periods without losing multipotency owing to their self-renewal ability [26, 27]. Moreover, harvesting of SVZ-NSCs from autopsied postnatal or adult human brains and live human brain is feasible [28–30].

We ascertained the efficacy of SVZ-NSC grafting into the hippocampus shortly after an injury for counteracting the injury-induced impairments in mood and memory function and neurogenesis. We expanded and characterized NSCs in vitro from the anterior SVZ of postnatal day 2 rats (F344) expressing the transgene alkaline phosphatase (AP). The AP+ NSCs exhibiting robust expression of multiple neurotrophic factors were then labeled with 5′-bromodeoxyuridine (BrdU) and grafted into the hippocampus of young adult F344 rats at 5 days after an injury inflicted through a unilateral administration of kainic acid (KA) into the posterior lateral ventricle. At 1.5 months postgrafting, animals were evaluated for functions such as mood, recognition memory, and spatial memory and compared with animals that received sham-grafting surgery after injury and age-matched control animals. Animals were then euthanized to measure the yield, migration, and phenotypic differentiation of graft-derived cells and the extent and pattern of hippocampal neurogenesis. To understand the potential mechanisms of grafting-mediated normalization of neurogenesis, both NSC proliferative activity in the SGZ and the survival of reelin+ interneurons in the DG were quantified. A flowchart summary of the experimental design is shown in Figure 1.

**Materials and Methods**

**Induction of Unilateral Partial Hippocampal Injury**

Four-month-old male F344 rats (Harlan Sprague-Dawley) were used in this study. A group of rats \( n = 6 \) received unilateral partial hippocampal injury followed by SVZ-NSC grafting into the injured hippocampus at 5 days postinjury (hereafter referred to as grafted animals). A second group of rats \( n = 6 \) received unilateral partial hippocampal injury followed by sham-grafting surgery at 5 days postinjury (sham-grafted animals). A third group of age-matched rats \( n = 6 \) served as naïve controls (control animals). An unilateral partial hippocampal injury was induced via KA (Tocris) administration into the right lateral ventricle, using procedures detailed in our earlier reports [12, 13, 31] and in the supplemental online data.

**Preparation of SVZ-NSC Suspension**

Anterior SVZ tissues were dissected from the forebrain, triturated, and expanded as neurospheres in vitro using standard NSC expansion procedures [32]. In order to label the AP+ neurosphere cells with a second marker, 0.25 μM BrdU was added to the proliferation medium. Neurospheres were mechanically triturated, and preparations having >75% cell viability were selected for further study. Samples from the NSC suspension were
incubated in culture dishes coated with poly-o-lysine and containing Dulbecco’s modified Eagle’s medium (DMEM) and Ham’s F-12 medium (F12) for 1 hour and processed for BrdU immunohistochemistry to calculate the BrdU labeling index [24, 32]. For grafting experiments, the live cells were adjusted to a density of 1.0 × 10^5 cells per microliter of culture medium containing brain-derived neurotrophic factor (BDNF) (200 ng/ml). The detailed procedures are described in the supplemental online data.

**Characterization of Neurotrophic Factors in SVZ-NSCs**

Samples from the NSC suspension were washed in DMEM and incubated in culture dishes coated with poly-o-lysine and containing the same medium for 1 hour and then processed for various immunofluorescence methods for detecting the presence of BDNF, glial cell line-derived neurotrophic factor (GDNF), vascular endothelial growth factor (VEGF), and fibroblast growth factor-2 (FGF-2) in NSCs. Details on these procedures are available in the supplemental online data.

**Phenotypic Differentiation of SVZ-NSCs in Culture**

Samples from the SVZ-NSC suspension were incubated in culture dishes coated with poly-o-lysine and containing DMEM, F12, and B-27 nutrient mixture for 7 days. The cultures were processed for various immunofluorescence methods to quantify fractions of neurons, interneurons, astrocytes, and oligodendrocytes. These procedures are described in detail in the supplemental online data.

**Grafting of SVZ-NSCs into the Injured Hippocampus**

Grafting was performed on postinjury day 5, using methods described in the supplemental online data. Three grafts (each containing 100,000 live cells in 1 µl of the culture medium including BDNF at a concentration of 200 ng/ml) were placed into the injured hippocampus along its septotemporal axis. Based on the BrdU labeling index (90% of all cells) in the cell suspension, this amounted to grafting of 90,000 BrdU+ cells per graft and 270,000 BrdU+ cells per hippocampus. The injured hippocampus of animals assigned for sham-grafting surgery received injections of the culture medium including BDNF (200 ng/ml) into three sites (1 µl/site).

**Analyses of Depressive-Like Behavior Through a Forced Swim Test**

Animals were subjected to a forced swim test (FST) at 1.5 months after the grafting/sham-grafting surgery. The forced swim test is one of the most widely used tests for assessing the extent of depressive-like behavior in rodents [33, 34]. A detailed description of this test is available in the supplemental online data.

**Measurement of Spatial Learning and Memory Function Using a Water Maze Test**

Animals were examined using a water maze test (WMT) after the completion of a NORT. A full description of the WMT used in this study is provided in our previous reports [24, 32] and the supplemental online data.

**Tissue Processing**

Following all behavioral tests (i.e., at ~2.5 months after grafting), rats were perfused with 4% paraformaldehyde, brain tissues were collected, and 30-µm-thick sections were cut coronally through the entire hippocampus using a cryostat and collected serially in 24-well plates containing phosphate buffer. A set of serial sections (every 15th section) through the entire hippocampus from animals belonging to different groups was processed for neuron-specific nuclear antigen (NeuN) immunostaining [39] to examine neurodegeneration.

**Quantification of the Yield of Graft-Derived Cells**

A set of serial sections (every 10th section) from grafted animals was first processed for BrdU immunostaining as described in our earlier report [40]. Cells positive for BrdU were then counted in serial sections through the entire anterior-posterior extent of the hippocampus using the optical fractionator counting method in a Stereo Investigator system (MicroBrightField Inc., Williston, VT, http://www.mbfbioscience.com). The counting procedure is detailed in our earlier reports [32, 40] and the supplemental online data. The yield of graft-derived cells in each hippocampus was expressed as the percentage of injected BrdU+ cells.

**Analyses of Graft Cell Differentiation and T Lymphocytes in the Host Brain**

We quantified the phenotype of graft-derived cells through dual immunofluorescence and confocal microscopy for AP and different neural cell antigens, and BrdU and different neural cell antigens. The neural cell antigens included markers of (a) mature neurons (NeuN), (b) inhibitory interneurons (GABA; the calcium-binding proteins parvalbumin [PV], calretinin [CR], and calbindin [CBN]); and the neuropeptides somatostatin [SS] and neuropeptide Y [NPY]), (c) mature astrocytes (S100β), (d) oligodendrocytes (2′,3′-cyclic nucleotide 3′-phosphodiesterase [CNPase]), and (e) oligodendrocyte progenitors (neuron glia proteoglycan 2 [NG2]). Additionally, we examined the presence of T lymphocytes using a CD4 antibody. The methods are described in the supplemental online data. Dual-labeled cells were quantified using z-section analyses using an Fv10i confocal microscope (Olympus, Tokyo, Japan, http://www.olympus-global.com). One hundred to 150 graft-derived cells from each of the grafted hippocampi (six sections per animal) were analyzed for every neural cell antigen examined.

**Characterization of the Expression of Neurotrophic Factors in Graft-Derived Cells**

We performed dual immunofluorescence on hippocampal sections passing through grafts using antibodies against BrdU or AP and antibodies for GDNF, BDNF, FGF-2, and VEGF. Using z-section analyses with a Fv10i confocal microscope, we then quantified the percentages of graft-derived cells expressing GDNF, BDNF, FGF-2, or VEGF. Approximately 200 graft-derived cells from each of the grafted hippocampus (using four to six sections per animal) were analyzed for every neurotrophic factor examined.
Measurement of the Extent and Pattern of Hippocampal Neurogenesis

A set of serial sections (every 15th section) through the entire hippocampus from all animals was processed for doublecortin (DCX) immunostaining [41]. Using these sections, numbers of newly born (DCX+) neurons in the SGZ-GCL were measured via the optical fractionator counting method using a Stereo Investigator system (MicroBrightField) as detailed in our earlier reports [25, 41]. We also measured percentages of DCX+ newly born neurons in the DG that are located in the dentate hilus, as described in our earlier report [40]. Furthermore, we examined the effect of SVZ-NSC grafting on the occurrence of abnormal basal dendrites [22, 40, 42] via measurement of percentages of relatively mature DCX+ neurons (i.e., DCX+ neurons with vertical dendrites projecting into the molecular layer) exhibiting basal dendrites, as described in our previous report [40].

Analyses of Proliferation of NSCs in the SGZ and Survival of Reelin+ Interneurons in the DG

We quantified fractions of putative NSCs in the SGZ (i.e., glial fibrillary acidic protein [GFAP]+ cells) that are positive for Ki67 (a marker of dividing cells). We first performed dual immunofluorescence on serial sections using antibodies against GFAP and Ki67 [11]. Next, using z-section analyses in a confocal microscope, we quantified the percentage of GFAP+ cells expressing Ki67. At least 200 GFAP+ cells in the SGZ of each hippocampus (using four to six sections per animal) were analyzed for Ki67 expression. Because none of the S100β+ mature astrocytes in the SGZ expressed Ki67 (supplemental online Fig. 1), the above quantification provided an indirect measure of the proliferation of NSCs in the SGZ. We measured the effects of NSC grafting on the survival of reelin+ interneurons via stereological quantification of reelin+ interneurons in the SGZ-GCL and the dentate hilus.

Statistical Analyses

All data are expressed as mean ± SEM. Data were analyzed using one-way analysis of variance followed by Student-Newman-Keuls multiple-comparison post tests.

RESULTS

SVZ-NSCs Express Neurotrophic Factors and Produce All Three Central Nervous System Phenotypes and Subtypes of GABAergic Neurons

The neurosphere cells generated from SVZ-NSCs displayed robust expression of BDNF, GDNF, VEGF, and FGF-2 (Fig. 2A1–2E3). Additional analyses demonstrated the ability of SVZ-NSCs to produce all three central nervous system cell types, including the GABAergic neurons (Fig. 2F–2K). Significant fractions of SVZ-NSCs differentiated into β-III-tubulin-positive (Tuj-1+) neurons (24%), GABA+ interneurons (22%), GFAP+ cells (70%), S100β+ mature astrocytes (53%), and oligodendrocytes positive for O4 (12%) and receptor-interacting protein (18%) (Fig. 2Q). The vast majority of Tuj-1+ neurons (90%) derived from SVZ-NSCs expressed GABA (Fig. 2F–2H), and the GABAergic neuronal population comprised subclasses of neurons expressing PV, CR, CB, SS, or NPY (Fig. 2L–2P).

Neurodegeneration After KA Administration

Neurodegeneration in the hippocampus ipsilateral to KA administration was typified by a partial loss of neurons in the dentate hilus and extensive loss of neurons in the cornu ammonis 3 (CA3) pyramidal cell layer spanning the CA3b and CA3c regions (supplemental online Fig. 2). Although neurons were spared in the CA3a region, some neuron loss was evident in the CA1 pyramidal...
cell layer. Furthermore, cell layers in the hippocampus contralateral to KA administration did not exhibit any sign of neuron loss (data not illustrated). This pattern of neurodegeneration was evident in both grafted and sham-grafted animals, which is consistent with our earlier studies using this injury model [31].

**SVZ-NSC Grafting Eases Hippocampal Injury-Mediated Mood Dysfunction**

Sham-grafted animals displayed increased depressive-like behavior, which was revealed by an increased time spent in immobility (or floating) in an FST [38] in comparison with control animals (Fig. 3A). Immobility time in these animals was 240% greater than the immobility time exhibited by control animals ($p < .001$; Fig. 3A). In contrast, grafted animals exhibited mood function that was equivalent to that of control animals and superior to that of sham-grafted animals ($p < .001$; Fig. 3A).

**SVZ-NSC Grafting Prevents Hippocampal Injury-Related Recognition Memory Dysfunction**

Sham-grafted animals spent an average of 35% of the total object exploration time with the novel object, which is 51% less than the novel object exploration time observed in control animals (Fig. 3B). In contrast, grafted animals spent 65% of the total object exploration time with the novel object, which is at par with control animals (Fig. 3B) and greater than sham-grafted animals ($p < .05$; Fig. 3B).

**SVZ-NSC Grafting Thwarts Hippocampal Injury-Mediated Spatial Memory Impairment**

The mean latency to reach the submerged platform over seven training sessions decreased progressively in all three animal groups ($p < .001$ to $p < .0001$; Fig. 3C1–3C3). However, a probe test conducted at 24 hours after the last training session revealed considerable memory retrieval dysfunction in sham-grafted animals but not in grafted animals. The sham-grafted animals spent less of the total probe test time in the platform quadrant and the platform area than control animals ($p < .01$, 37%–48% reduction; Fig. 3D1, 3D2), which is consistent with the spatial memory impairment observed in patients and animals with a unilateral hippocampal injury [43, 44]. In contrast, times spent in the platform quadrant and platform area by grafted animals were comparable to that of control animals ($p > .05$; Fig. 3D1, 3D2).

**Cells Derived from SVZ-NSC Grafts Survive and Exhibit Pervasive Migration**

Immunostaining for BrdU revealed the presence of graft cores in and around the injured CA3 region and migration of graft-derived cells into the DG and CA1 regions of the injured hippocampus (Fig. 4A1–4B4). Figure 4C depicts the migration of graft-derived cells into different regions of the hippocampus in representative serial sections along the septotemporal axis. Occasionally, a few graft-derived cells were also found outside the

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**Figure 3.** Grafting of SVZ-NSCs into the hippocampus at 5 days postinjury maintains normal mood and memory function. (A): Results of an FST, which used the time spent in floating as a measure of depression. Rats receiving sham-grafting surgery after hippocampal injury (red) displayed increased depressive-like behavior, whereas rats receiving SVZ-NSC grafts after hippocampal injury (green) exhibited mood function similar to that of naive control rats (blue), ***, $p < .001$. (B): Findings in a NORT, which used percentage of the exploration time spent with novel object as a measure of the object recognition memory. Rats receiving sham-grafting surgery exhibited impaired object recognition memory, whereas rats receiving SVZ-NSC grafts demonstrated object recognition ability comparable to that of naive control rats, *, $p < .05$. (C1–D2): Results of a WMT. Note that based on changes in the mean latency values to reach the hidden platform over seven learning sessions (C1–C3), rats in all three groups exhibited ability for spatial learning ($r^2 = 0.5–0.6$). (D1, D2): Comparison of memory retrieval function among the three groups, based on the amounts of time spent within the platform quadrant (D1) and the platform area (D2) in a probe test conducted at 24 hours after the seventh learning session. Note that rats receiving sham-grafting surgery after hippocampal injury exhibited impaired memory retrieval ability, whereas rats receiving SVZ-NSC grafts after hippocampal injury displayed memory retrieval ability similar to that of naive control rats, ***, $p < .01$; **, $p < .05$. Abbreviations: FST, forced swim test; NORT, novel object recognition test; SVZ-NSC, subventricular zone-neural stem cell; WMT, water maze test.
hippocampal area in the adjoining entorhinal cortex and the corpus callosum. Stereological quantification of the surviving graft-derived cells revealed that SVZ-NSC grafts gave rise to an average of 349,129 new cells (349,129/11006 46,303) in each injured hippocampus. We injected three grafts, each containing 100,000 live cells, amounting to 300,000 live cells per hippocampus. However, based on the 90% BrdU labeling index at the time of grafting, the number of BrdU cells injected per hippocampus was 270,000. Based on these injected and recovered numbers of BrdU cells, the overall yield is equivalent to ~116% of injected cells.

**SVZ-NSC Grafts Contribute Significant Numbers of NeuN+ and GABA+ Neurons**

Cells derived from SVZ-NSC grafts differentiated into NeuN+ neurons, GABA+ neurons, and subclasses of GABAergic neurons positive for CBN, NPY, and PV (Fig. 5A1–5E3). Neuronal differentiation was conspicuous in the graft core (Fig. 5A1–5A3). Quantification revealed that 29% of graft-derived cells differentiated into NeuN+ neurons, 25% into GABA+ neurons, 10% into CB1-interneurons, and 5% into PV+ interneurons (Fig. 5I). Extrapolation of the total yield of graft-derived cells with percentages of neuronal types suggested that SVZ-NSC grafting contributed ~101,247 NeuN+ neurons and ~88,330 GABA+ neurons into each injured hippocampus. Analyses of BrdU+ graft-derived cells revealed a similar trend (supplemental online Fig. 3). Additionally, 44% of graft-derived cells that migrated into the SGZ-GCL differentiated into NeuN+ neurons (supplemental online Fig. 3).

**SVZ-NSC Grafts Add Substantial Numbers of Astrocytes, Oligodendrocytes, and Oligodendrocyte Progenitors but Do Not Trigger Host Immune Response or Tumors**

Cells derived from SVZ-NSC grafts also differentiated into astrocytes, oligodendrocytes, and oligodendrocyte progenitors (Fig. S1–S5H). Forty-six percent of graft-derived cells differentiated into S100β+ astrocytes, 16% into CNPase+ oligodendrocytes, and 16% into NG2+ oligodendrocyte progenitors. Based on the total yield of graft-derived cells and percentages of glial cell types, this amounts to an addition of ~161,996 S100β+ mature astrocytes, ~52,369 oligodendrocytes, and ~52,369 oligodendrocyte progenitors into each injured hippocampus. Analyses

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**Figure 4.** Cells derived from the subventricular zone-neural stem cell grafts migrate profusely into different regions of the injured hippocampus. (A1, A2): Examples of injured hippocampi that demonstrated pervasive migration of 5'-bromodeoxyuridine-labeled graft-derived cells. (A2–A4, B2-B4): Respectively, graft-derived cells in magnified regions of the dentate gyrus and the CA3 and CA1 subfields from (A1) and (A2). (C): Tracings of every 15th section through the hippocampus (performed using Neurolucida [MicroBrightField]) to show the distribution of graft-derived cells in one of the grafted animals. Note that the graft core regions (solid pink areas) are located in the lesioned CA3 subfield, whereas the graft-derived cells migrated into all three regions of the hippocampus. Scale bars = 500 μm (A1, B1, C), 200 μm (A2–A4, B2–B4).

Abbreviations: CA, cornu ammonis; DG, dentate gyrus; DH, dentate hilus; GCL, granule cell layer.
Figure 5. Significant fractions of cells derived from the subventricular zone-neural stem cell (SVZ-NSC) grafts differentiate into different types of neurons and glia, and express several neurotrophic factors. (A1–H3): The AP-positive graft-derived cells (depicted in red) differentiated into neurons expressing NeuN (A1–A3), interneurons expressing GABA (B1–B3), calbindin (C1–C3), NPY (D1–D3), or PV (E1–E3); S100β+ astrocytes (F1–F3); CNPase+ oligodendrocytes (G1–G3); and NG2+ oligodendrocyte progenitors (H1–H3). The bar chart in (I) illustrates the percentages of different types of neurons and glia derived from SVZ-NSC grafts. (J1–M3): Cells derived from (BrdU+/AP+) SVZ-NSC grafts expressed GDNF (J1–J4), BDNF (K1–K3), FGF-2 (L1–L3), and VEGF (M1–M3). The bar chart in (N) illustrates the percentages of graft-derived cells expressing GDNF, BDNF, FGF-2, and VEGF. Scale bars = 10 μm (A1–H3, J2–M3), 100 μm (J1). Abbreviations: AP, alkaline phosphatase; BDNF, brain-derived neurotrophic factor; BrdU, 5′-bromodeoxyuridine; CBN, calbindin; CNPase, 2′,3′-cyclic nucleotide 3′-phosphodiesterase; FGF-2, fibroblast growth factor-2; GABA, γ-aminobutyric acid; GDNF, glial cell line-derived neurotrophic factor; NeuN, neuron-specific nuclear antigen; NG2, neuron glia proteoglycan 2; NPY, neuropeptide Y; PV, parvalbumin; VEGF, vascular endothelial growth factor.
of BrdU+ graft-derived cells revealed a similar trend (supplemental online Fig. 3). Furthermore, the injured hippocampus in both grafted and sham-grafted animals revealed only occasional CD4+ T lymphocytes. Importantly, graft areas did not attract CD4+ lymphocytes, implying that SVZ-NSC allografting does not trigger host immune response in the injured hippocampus. Additionally, none of the grafted hippocampi displayed tumors.

**Cells Derived from NSC Grafts Express GDNF, BDNF, FGF-2, and VEGF**

Considerable fractions of graft-derived cells expressed GDNF, BDNF, and FGF-2 (Fig. 5J1–5M3). Quantification revealed the occurrence of GDNF in 52% of graft-derived cells, BDNF in 40%, FGF-2 in 42%, and VEGF in 68% (Fig. 5N).

**SVZ-NSC Grafting Positively Influences Neurogenesis in Both Injured and Contralateral Hippocampi**

Neurogenesis in the injured hippocampus of sham-grafted animals exhibited 37% reduction at ~2.5 months postinjury compared with control animals (p < .05; Fig. 6A1–6D). However, the grafted injured hippocampus exhibited neurogenesis that was comparable to that of the control hippocampus (p > .05; Fig. 6A1–6D). We also measured the extent of neurogenesis in the dorsal and ventral segments of the hippocampus, as it is believed that neurogenesis in the dorsal segment is important for memory, whereas neurogenesis in the ventral segment is vital for mood function [45]. In comparison with the hippocampus of control animals, the dorsal segment of the injured hippocampus exhibited an ~40% decrease in neurogenesis in sham-grafted animals (p < .001; Fig. 6D) but no changes in grafted animals (p > .05; Fig. 6D). The ventral segment also showed a similar trend, although differences were not statistically significant (Fig. 6D). Furthermore, the hippocampus contralateral to the injury displayed increased neurogenesis in grafted animals but no changes in sham-grafted animals (details are given in supplemental online Fig. 4).

**SVZ-NSC Grafting Reduces Aberrant Pattern of Neurogenesis in the Injured Hippocampus**

Although the normal hippocampus exhibits apt migration of virtually all newly born neurons into the GCL (Fig. 6E1), the injured hippocampus exhibits abnormal migration of significant fractions of newly born neurons into the dentate hilus (Fig. 6E2). Consistent with this, the sham-grafted animals exhibited abnormal migration of ~23% of newly born neurons into the dentate hilus, in comparison with ~2% of newly born neurons exhibiting such migration in control animals (~12-fold increase, p < .001; Fig. 6E4). Interestingly, SVZ-NSC grafting after hippocampal injury reduced the aberrant migration of newly born neurons to ~13% (44% reduction, p < .01; Fig. 6E3, 6E4). Moreover, NSC grafting diminished the occurrence of basal dendrites in relatively mature newly born neurons (i.e., DCX+ neurons with vertically oriented dendrites projecting into the molecular layer; Fig. 6F1–6F3). This was evidenced by the occurrence of basal dendrites in only 14% of such neurons in grafted animals, in comparison with 36% of such neurons exhibiting basal dendrites in sham-grafted animals (62% reduction, p < .001; Fig. 6F4).

**SVZ-NSC Grafting Conserves Proliferative Behavior of NSCs in the SGZ**

We examined putative NSCs (GFAP+ cells) in the SGZ expressing Ki67 (Fig. 6G1–6I3). Proliferative activity of NSCs was ~17% in control animals, which reduced to ~7% in sham-grafted animals (56% reduction, p < .001; Fig. 6J). However, in grafted animals, proliferative activity of NSCs (~14%) was similar to that of control animals and greater than that of sham-grafted animals (p < .01; Fig. 6J).

**SVZ-NSC Grafting Preserves Reelin+ Interneurons in the SGZ-GCL and the Dentate Hilus**

Both SGZ-GCL and dentate hilar areas in sham-grafted animals displayed significantly reduced numbers of reelin+ neurons compared with control animals and grafted animals (25%–35% reduction, p < .05 to p < .01; Fig. 6K1–6K5). In contrast, both areas in grafted animals displayed numbers of reelin+ interneurons similar to those of control animals (p > .05; Fig. 6K1–6K5).

**DISCUSSION**

This study provides novel evidence that grafting of NSCs derived from the postnatal SVZ into the hippocampus is a highly efficacious approach for counteracting the hippocampal injury-induced mood and memory dysfunction. Preservation of normal mood and memory function in animals receiving SVZ-NSC grafts after hippocampal injury was associated with (a) robust survival and pervasive migration of graft-derived cells; (b) differentiation of substantial percentages of graft-derived cells into various subtypes of GABAergic interneurons, astrocytes, oligodendrocytes, and oligodendrocyte progenitors; (c) expression of the beneficial neurotrophic factors GDNF, BDNF, FGF-2, and VEGF in significant fractions of graft-derived cells; (d) normalization of the extent and pattern of neurogenesis in the injured hippocampus with maintenance of NSC proliferation in the SGZ at levels similar to those of the age-matched intact hippocampus; (e) protection of reelin+ interneurons in the DG of the injured hippocampus; and (f) increased neurogenesis in the hippocampus contralateral to injury. Additionally, SVZ-NSC grafting into the injured hippocampus did not trigger host immune response or tumor formation.

**Mechanisms Underlying SVZ-NSC Grafting Mediated Preservation of Mood and Memory Function After Hippocampal Injury**

On the basis of the differential structural plasticity of the hippocampus detected after injury with SVZ-NSC grafting or sham-grafting surgery, it is credible that SVZ-NSC grafting preserved mood and memory function through several mechanisms. Nonetheless, conservation of hippocampal neurogenesis in the injured hippocampus and enhancement of neurogenesis in the hippocampus contralateral to injury are the most perceivable possibilities. It is well known that a biphasic neurogenic response occurs in the injured hippocampus, which is typified by increased and anomalous neurogenesis in the acute phase and by decreased and abnormal neurogenesis in the chronic phase [10]. The aberrant neurogenesis is perceived from both migration of substantial fractions of newly born neurons into the dentate hilus and increased occurrences of basal dendrites from newly born neurons that are integrated into the dentate GCL [22, 42]. Our results validate that sham-grafting surgery after injury did not prevent such abnormal response. In contrast, early SVZ-NSC

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Figure 6. Grafting of SVZ-NSCs into the HPS maintains neurogenesis and neural stem cell (NSC) activity at levels comparable to those of the intact control HPS and preserves reelin+ interneurons in the dentate gyrus. *, p < .05; **, p < .01; ***, p < .001. (A1–D): Extent of neurogenesis measured through DCX immunostaining. (E1–F4): Pattern of neurogenesis. (G1–J): NSC activity in the SGZ examined through GFAP-Ki67 dual immunofluorescence. (K1–K5): Reelin+ interneurons in the SGZ-GCL and the DH examined through reelin immunostaining. Note that in comparison with naïve control rats (A1, A2, D; E1, E4; F1, F4; G1–G3, J; K1, K4, K5), rats receiving sham-grafting surgery after hippocampal injury exhibited (a) decreased neurogenesis (B1, B2, D); (b) greater fractions of newly born neurons migrating abnormally into the dentate hilus (E2, E4); (c) increased occurrences of aberrant basal dendrites from newly born neurons (F2, F4); (d) decreased NSC activity (H1–H3, J); and (e) decreased numbers of reelin+ interneurons (K2, K4, K5). However, in rats receiving SVZ-NSC grafts after hippocampal injury, the extent of neurogenesis (C1, C2, D), NSC activity (I1–I3, J), and surviving reelin+ interneuron numbers (K3, K4, K5) were comparable to those observed in naïve control rats. Additionally, both abnormal hilar migration of newly born neurons (E3, E4) and occurrences of aberrant basal dendrites (F3, F4) were greatly reduced in these rats. Scale bars = 200 μm (A1, B1, C1), 50 μm (A2, B2, C2, F1–F3), 100 μm (E1–E3), 10 μm (G1–I3), 200 μm (K1–K3). Abbreviations: DCX, doublecortin; DH, dentate hilus; GCL, granule cell layer; GFAP, glial fibrillary acidic protein; HPS, hippocampus; ML, molecular layer; SGZ, subgranular zone; SVZ-NSC, subventricular zone-neural stem cell.
grafting after injury preserved both the extent and the pattern of neurogenesis in the injured hippocampus at par with the control hippocampus. This was evinced by the following observations. Numbers of newly born neurons in the SGZ-GCL and numbers of proliferating NSCs in the SGZ of the injured grafted hippocampus were comparable to numbers observed in the respective regions of the control hippocampus. Furthermore, abnormal migration of newly born neurons into the dentate hilus and incidences of basal dendrites from newly born neurons were considerably reduced in the injured grafted hippocampus, in comparison with the injured sham-grafted hippocampus.

Our proposition that preserved neurogenesis in the injured hippocampus and enhanced neurogenesis in the hippocampus contralateral to injury underlie the conserved mood and memory function in grafted animals is centered on the following. First, a close association between neurogenesis and mood function has been recognized. For instance, a recent study demonstrates that induced neurogenesis deficiency results in an increased depressive-like behavior [20]. Besides, multiple previous studies have shown that recovery from mood dysfunction through antidepressant medications is intermediated by improved hippocampal neurogenesis. This was shown by findings that selective ablation of neurogenesis blocks behavioral responses to antidepressants [46, 47] and chronic antidepressant treatment enhances hippocampal neurogenesis [48, 49]. Second, even though a few earlier studies suggested uncertainties regarding the role of neurogenesis in certain memory functions [50, 51], a series of recent studies supports the idea that hippocampal neurogenesis plays a crucial role in maintaining both spatial and object recognition memories [27, 37, 52–54]. However, contributions from other factors may also be important. These may include the release of beneficial neurotrophic factors by graft-derived cells, protection of reelin+ interneurons, and addition of new GABAergic neurons.

Pertaining to neurotrophic factors, our analyses suggest robust expression of GDNF, BDNF, FGF-2, and VEGF in significant fractions of graft-derived cells. Besides their well-known ability for maintaining neurogenesis [55–57], these neurotrophic factors can directly impact mood and memory function. For example, studies show that major depressive disorder is associated with decreased concentration of BDNF [58], peripheral BDNF administration improves mood function [59], chronic intracerebroventricular FGF-2 treatment decreases depressive-like behavior [60], peripherally administered FGF-2 is highly effective for blunting anxiety [61], and chronic antidepressant treatment increases FGF-2 concentration in the hippocampus [62]. Furthermore, studies support the understanding that GDNF, BDNF, FGF-2, and VEGF have major roles in memory. These include the observations that expression of GDNF transgene in astrocytes ameliorates cognitive deficits in aged rats [63], mice lacking GDNF receptors exhibit significant memory dysfunction [64], forebrain-restricted deficiency of BDNF causes learning deficits [65], NSC grafts improve cognitive function in a mouse model of Alzheimer’s disease through BDNF [23], systemic FGF-2 administration enhances long-term memory [66], FGF-2 gene transfer into the hippocampus improves memory function in a mouse model of Alzheimer’s disease [67], and suppression of VEGF expression in the hippocampus by RNA interference causes memory impairments [68].

With reference to reelin+ interneurons, our results demonstrate that SVZ-NSC grafting greatly reduces the hippocampal injury-mediated loss of reelin+ interneurons in both SGZ-GCL and the dentate hilus. Reelin, a conserved extracellular glycoprotein, is believed to play an important role in both mood and memory function [69]. The role of reelin in mood function is evidenced by observations that (a) reelin gene expression is downregulated in postmortem brain samples of schizophrenia, bipolar disorder, autism, major depression, and Alzheimer’s disease patients [70]; (b) transgenic mice having reduced levels of reelin are more vulnerable to developing depressive-like behavior [71]; and (c) transgenic mice overexpressing reelin are protected against developing depressive-like behavior [72]. On the other hand, the significance of reelin in memory function is documented by findings that a single intracerebroventricular injection of reelin is sufficient to increase the activation of cAMP-response element-binding protein, hippocampal long-term potentiation, and spatial learning and memory [73].

As to the addition of new GABAergic neurons, our analyses showed that SVZ-NSC grafting added ~88,330 new GABAergic neurons into each injured hippocampus. Because hippocampal injury is associated with decreased numbers of GABAergic interneurons in the chronic phase [39], this addition is substantial and could also have positive effects on mood and memory function. This suggestion is substantiated by findings that major depressive disorders are associated with a reduced concentration of GABA in brain regions [74], GABA<sub>A</sub> receptor activation reverses memory deficits in an animal model of schizophrenia [75], and administration of GABA-enhancing drugs reverses memory deficits in aged rats [76]. Moreover, the addition of substantial numbers of new GABAergic neurons through grafts could also contribute to improved memory function through suppression of hippocampal hypexcitability that occurs in the chronic phase after hippocampal injury [77], as excess neural activity can impair memory function [76]. Collectively, our observations suggest that SVZ-NSC grafting preserved mood and memory function after hippocampal injury via several mechanisms.

**Potential Mechanisms Underlying the Beneficial Effects of SVZ-NSC Grafts on Hippocampal Neurogenesis**

One of the mechanisms likely involves the release of beneficial neurotrophic factors by graft-derived cells. This is supported by our findings that significant fractions of graft-derived cells express GDNF, BDNF, FGF-2, and VEGF. These neurotrophic factors are well known to enhance hippocampal neurogenesis through NSC proliferation and differentiation [56–58, 78]. Depleted concentration of most of these proteins in the chronic phase after hippocampal injury also supports this possibility [10, 12, 13, 79]. Addition of significant numbers of GABAergic neurons by SVZ-NSC grafts might also have contributed to normal NSC proliferation in the SGZ, as GABA has an important role in NSC proliferation and differentiation and integration of newly generated neurons [80]. Another possibility may be the addition of new NSCs to the SGZ of the injured hippocampus by SVZ-NSC grafts. However, this possibility is unlikely to be among the major mechanisms, as virtually all AP+/BrdU+ graft-derived cells that migrated into the SGZ differentiated into NeuN+ neurons, S100β+ astrocytes, or NG2+ oligodendrocyte progenitors. Among the potential factors that contributed to the maintenance of the normal pattern of neurogenesis in the injured hippocampus by SVZ-NSC grafts, considerable protection of reelin+ interneurons observed in the DG stands out.
This is because reelin is known to regulate the migration of newly born neurons into the GCL, and aberrant chain migration of newly born neurons into the dentate hilus after hippocampal injury has been suggested to be due to a reduced concentration of reelin occurring through a substantial loss of reelin + interneurons in the DG [18].

Suitability of SVZ-NSCs for Cell Therapy in Neurodegenerative Disorders

The postnatal SVZ-NSCs appear to be ideal for grafting into the hippocampus after injury, as placement of these cells shortly after injury results in excellent yield and differentiation of graft-derived cells and maintenance of mood and memory function and neurogenesis at normal levels. These cells may also be suitable for transplantation in neurodegenerative disorders, where restoration of mood and memory function and neurogenesis are some of the goals. Although NSCs can be obtained from a variety of sources, the selection of a particular type of NSC is challenging for clinical application [80]. Even though both pluripotent human embryonic stem cells and induced pluripotent stem cells are efficient for providing an unlimited supply of NSCs, their utility for routine clinical application is still unclear because of their potential for forming teratoma, as the presence of even a single pluripotent stem cell in the graft material can give rise to teratoma [81]. Therefore, rigorous testing of the therapeutic utility of NSCs with restricted fate potential (such as NSCs derived from the SVZ/hippocampus of fetal, postnatal, and adult brain) for promoting functional recovery in animal models of neurological disorders will be necessary.

CONCLUSION

The findings presented here demonstrate that grafting of NSCs expanded from the postnatal SVZ into the hippocampus early after an injury is highly efficacious for preventing injury-induced impairments in mood and memory function and hippocampal neurogenesis. Interestingly, the beneficial effects of grafting were associated with excellent survival and pervasive migration of graft-derived cells into different regions of the hippocampus. Furthermore, significant fractions of graft-derived cells differentiated into GABAergic interneurons, astrocytes, oligodendrocytes, and oligodendrocyte progenitors and expressed useful neurotrophic factors such as GDNF, BDNF, FGF-2, and VEGF. Thus, grafting of SVZ-NSCs into the hippocampus is a useful approach for alleviating mood and memory dysfunction in neurological disorders associated with hippocampal lesions or damage.

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AUTHOR CONTRIBUTIONS

B.H.: design, collection, assembly, analysis and interpretation of data, manuscript writing; A.K.S.: conception, design, collection, assembly, analysis and interpretation of data, manuscript writing, financial support.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST

The authors indicate no potential conflicts of interest.

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