Secretoneurin (SN), a neuropeptide derived from secretogranin II, promotes neurite outgrowth of immature cerebellar granule cells. SN also aids in the growth and repair of neuronal tissue, although the precise mechanisms underlying the promotion of brain tissue neuroprotection and plasticity by SN are not understood. Here, in a rat model of stroke and in ischemic human brain tissue, SN was markedly upregulated in both neurons and endothelial cells. SN-mediated neuroprotection rescued primary cortical cell cultures from oxygen/glucose deprivation. SN also induced expression of the antiapoptotic proteins Bcl-2 and Bcl-xL through the Jak2/Stat3 pathway and inhibited apoptosis by blocking caspase-3 activation. In addition, rats with occluded right middle cerebral arteries showed less cerebral infarction, improved motor performance, and increased brain metabolic activity following i.v. administration of SN. Furthermore, SN injection enhanced stem cell targeting to the injured brain in mice and promoted the formation of new blood vessels to increase local cortical blood flow in the ischemic hemisphere. Both in vitro and in vivo, SN not only promoted neuroprotection, but also enhanced neurogenesis and angiogenesis. Our results demonstrate that SN acts directly on neurons after hypoxia and ischemic insult to further their survival by activating the Jak2/Stat3 pathway.
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Introduction

Secretoneurin (SN) is a 33–amino acid neuropeptide produced by endoproteolytic processing of chromogranin/secretogranin family proteins, which are found in large dense-core vesicles in a wide variety of cell types of the endocrine tissue and nervous system (1, 2). In several recent reports, chromogranin/SN proteins have been found to be closely correlated with synaptic disturbance caused by neuronal/glial and inflammatory mechanisms in conditions such as Alzheimer disease (3–5). It has also been reported that SN can promote the neurite outgrowth of immature cerebellar granule cells (6). Furthermore, in a recent report, increased expression of SN was found in an animal model after transient forebrain ischemia (7). Although there is much evidence suggesting an important role for SN in the physiology and pathophysiology of the nervous system, its precise role in neuroprotection and neuronal plasticity has not been clarified.

Although several articles have reported a correlation between SN and neurological diseases including Alzheimer disease (3–5), Parkinson disease (8), and epilepsy (9, 10), little literature has examined the role of SN in stroke (7). Human stroke is a leading cause of death and disability worldwide (11), and as yet there is no effective treatment that enhances stroke recovery. One potential strategy for the treatment of stroke is transplantation of bone marrow stem cells (BMSCs) (12) leading to enhancement of neurogenesis and angiogenesis, which have been demonstrated to promote plasticity and assist in the recovery from stroke (13, 14). Recently, the role of bone marrow–derived circulating progenitor cells in postnatal angiogenesis and neurogenesis has been clearly demonstrated in hind-limb, myocardial, and cerebral ischemia (15–17). Due to the effects of SN on the induction of vasculogenesis through activation of the Akt signaling pathway (18), the mobilization of bone marrow–derived endothelial progenitor cells (19), and the increased SN expression seen in ischemic tissue (7), we hypothesized that SN might enhance neuroprotection and plasticity in the cerebral ischemic animal model. Furthermore, some growth factors may enhance the bone marrow–derived progenitor cells’ proliferation and angiogenesis via activation of the Jak2/Stat3 pathway (20, 21). Therefore, in the present study, we have examined the neuroprotective effects of SN against oxygen/glucose deprivation–induced (OGD-induced) neurotoxicity in primary cortical neurons and also analyzed the results of i.v. administration of SN on cerebral ischemic animals by measuring changes in the extent of induced cerebral infarction and neurological dysfunction. In addition, we also focused on the Jak2/Stat3 pathway to discern the possible molecular mechanism for the neuroprotective role of SN.

Results

Cerebral ischemia increases the immunoreactivity of SN in human and rat brains. In order to determine whether cerebral ischemia increases the expression of SN, levels of SN were measured by analysis of
SN-immunoreactivity (SN-IR). Brain samples from human stroke patients at 1, 3, and 7 days after ictus (n = 9) and experimental rats at 4 and 12 hours and 1, 3, and 7 days after undergoing cerebral ischemia (n = 20; 4 per time point) were examined for SN immunostaining. Significantly more SN-IR cells were found in the penumbral and hippocampal regions of the ischemic brain sample from human stroke patients 1 and 3 days after cerebral infarction than in those of controls (n = 3; Figure 1, A and B). In the animal model, greater numbers of SN-IR cells were detected 4 hours (n = 4) after cerebral ischemia than in the nonischemic control (n = 6). SN-IR cells were mainly detected in the ipsilateral cortex near the infarcted boundary region of the ischemic rat brains (Figure 1, C and D). In addition, there was a time-dependent increase in the number of SN-IR cells found around the penumbral region in ischemic rats; in comparison, nonischemic rats showed no increase (Figure 1D).

Cerebral ischemia enhances expression of SN in the ischemic brain and serum. In order to determine whether the expression of SN was increased in rats and humans, SN levels in brain samples from rats that had undergone cerebral ischemia (n = 20) and sera from human stroke patients (n = 20) were measured using ELISA. Samples of ischemic rat brains were taken from the cortical region and striatum. Brain samples from homologous areas of rats without middle cerebral artery (MCA) ligation were used as normal controls (n = 6). ELISA analysis of SN levels showed that cerebral ischemia in rat brains caused an increase in the expression of SN in a time-dependent manner (control, 1.1 ± 0.1 ng/g protein; 12 hours, 12.1 ± 2.2 ng/g protein; 1 day, 19.3 ± 3.2 ng/g protein; 3 days, 33.1 ± 5.2 ng/g protein; 7 days, 15.4 ± 3.3 ng/g protein; 14 days, 6.2 ± 2.7 ng/g protein; Figure 1E), which was in agreement with our immunohistochemical (IHC) studies of an ischemic rat’s brain that showed an increase in SN-IR (Figure 1A). In addition, sera from human stroke patients were also collected at different time points. Control serum samples were obtained from 6 healthy age-matched subjects. The level of SN in human stroke patients’ sera (12 hours, 3.1± 1.3 ng/ml; 1 day, 6.4 ± 2.1 ng/ml; 3 days, 13.3 ± 1.5 ng/ml; 7 days, 8.5 ± 1.4 ng/ml; 14 days, 4.1 ± 1.5 ng/ml) was significantly higher than that of the controls (0.12 ± 0.09 ng/ml; Figure 1F) and peaked at 3 days after stroke onset.

Immunoreactivity of SN colocalizes to neurons and endothelial cells after cerebral ischemia. In order to identify which cerebral cells expressed SN after cerebral ischemia, double immunofluorescence was performed on brain specimens of ischemic rats with laser-scanning confocal microscopy. Ischemic cortical areas of the rats revealed many SN-IR cells coexpressing a neuronal nucleus–positive (Neu-N+) neuronal phenotype (163 ± 26 cells/mm²; Figure 1G). Some SN-IR cells showing SMA+ vascular phenotypes were also found around the perivascular and endothelial regions (Figure 1G) of the ischemic hemispheres.

Neuroprotective effect of SN in primary cortical cell cultures. To evaluate the neuroprotective effect of SN in vitro, lactate dehydrogenase (LDH) activity and neuronal survival (positive microtubular-associated protein–2 [MAP-2] immunoreactivity) under OGD-induced neurotoxic conditions were measured in primary cortical cells (PCCs) with or without SN treatment (Figure 2, A–C). Treatment with 1 μg/l SN at 20 minutes prior to OGD significantly reduced LDH activity in cultures compared with the control group (Figure 2D). To analyze the effect of SN on neuronal survival after OGD, the number of MAP-2–immunoreactive cells was analyzed quantitatively. Pretreatment with 1 μg/l SN at 20 minutes prior to OGD significantly increased MAP-2–immunoreactive cell density in PCCs compared with the control group (Figure 2E). However, treatment with SN after OGD did not show a significantly neuroprotective effect in PCCs.

Neuroprotection by SN might be through an antiapoptotic effect in PCCs. In order to examine whether the neuroprotective effect of SN is caused by blocking an apoptotic pathway, an immunofluorescent study of caspase-3 under OGD-induced neurotoxic conditions was performed in PCCs with or without SN treatment. In the OGD environment, PCCs treated with SN for 12 hours showed significantly reduced caspase-3 activity using a fluorimetric method compared with the control group (Figure 2F). There was a significant reduction of caspase-3’ immunofluorescent cells in the SN-treated group compared with the control group (Figure 2G).

In addition, to investigate the molecular mechanism by which SN exerts an antiapoptotic effect on PCCs, we studied the expression of apoptosis-related proteins by Western blot analysis. Expression levels of antiapoptotic proteins such as Bcl-2 were lower when PCCs were subjected to OGD; this reduction was considerably inhibited by pretreatment with SN for 12 hours (Figure 2, H and I). SN activates the Jak2/Stat3 signal transductional pathway in PCCs. In order to examine whether SN exerts neuroprotective effects through specific cellular signals, the expression of activated protein kinases of both Jak2 and Stat3 in PCCs were measured after total protein was extracted. We found that the level of activated Jak2 and Stat3 after SN treatment increased in a time- and dose-dependent manner (Figure 2, J–M). The expression of Stat3 and Jak2 was not significantly different between SN-treated and control cells (Figure 2J). However, following the addition of 100 μM AG490, a specific inhibitor of activated Jak2, to OGD-treated cells, administration of 1 μg/l SN did not increase MAP-2–immunoreactive cell density in PCC (Figure 2N).

SN administration i.v. improves neurological behavior after cerebral ischemia. To evaluate the neuroprotective effect of i.v. administration of SN at 30 minutes after cerebral ischemia, body asymmetry trials and locomotor activity tests were used to assess neurological behavior in SN-treated and control stroke rats (n = 10 per group). From 14 to 28 days after treatment, rats treated with SN exhibited significantly reduced body asymmetry compared with control rats (Figure 3A). Locomotor activity such as vertical activity, vertical movement time, and number of vertical movements significantly increased in rats receiving SN treatment compared with control rats.
animals between 14 and 28 days after cerebral ischemia (Figure 3, B–D). Furthermore, measurement of improvement in grip strength was performed to examine the fore-limb strength for all experimental rats before treatment and at 28 days after each of the 2 treatments. A higher percentage of improvement in grip strength was found in the SN-treated group compared with the control group (Figure 3E). However, administration of AG490 or another Jak2 inhibitor, TG101209, to the SN-treated rats (n = 8 per group) blocked the recovery of all behavior modalities from neurological dysfunction after cerebral ischemia (Figure 3, A–E).

**SN administration i.v. reduces infarct volume after cerebral infarction.** To sequentially observe and quantify the volume of cerebral infarction showed mild infarction after cerebral ischemia. At 7 days after cerebral ischemia, the infarct volume was significantly less in SN-treated rats than saline-treated controls (73 ± 17 mm³ vs. 182 ± 16 mm³; Figure 3H). The area of largest infarction was significantly less in SN-treated rats than in control rats (9.4 ± 3.3 mm² vs. 19.7 ± 2.9 mm²; Figure 3H1). Infarcted slices were also significantly less in SN-treated animals than in control animals (3.1 ± 0.5 slices/rat vs. 6.7 ± 0.4 slices/rat; Figure 3H1).

**Enhancement of glucose metabolic activity in SN-treated stroke rats.** To verify whether i.v. SN administration enhances glucose metabolic activity, each experimental rat was examined by [18F]fluoro-2-deoxyglucose–PET (18FDG-PET). Glucose metabolism was measured by [18F] FDG-microPET 1 week after each treatment. The microPET image showed a striking increase of 18FDG uptake over the right cortex of the SN-treated group (Figure 3I). Semic quantitative measurement of relative glucose metabolic activity of the right hemisphere relative to the nonstoke hemisphere revealed significant enhancement in the SN-treated rats compared with controls (n = 8 per group; Figure 3, I and J).

**SN does not influence physiological parameters.** To demonstrate that the neuroprotective effect of SN did not occur as a result of changes to other physiological parameters, systemic physiological parameters were analyzed in 14 experimental rats at 1.5 hours after SN administration. Compared with vehicle control, i.v. administration of SN did not alter systemic blood pressure, blood gases, blood glucose, or serum electrolyte levels (n = 7 per group; Table 1).

**SN protects penumbral neurons from cerebral ischemic damage.** To verify the neuroprotective effect of SN in reducing neuronal injury in the penumbral region after cerebral ischemia (Figure 4, A and B), we analyzed ischemic brain tissues for neuronal survival using specific antibodies that recognize neuron-specific proteins (Neu-N conjugated with FITC and MAP-2 with Cy3; Figure 4, C–N). In the penumbra region surrounding the ischemic cores, the number of MAP-2- and Neu-N- cells significantly increased in SN-treated rats (n = 8; Figure 4, D and J) compared with vehicle control rats (n = 8; Figure 4, G and M). At 2 days after cerebral ischemia, ischemic brain tissue from animals injected with vehicle contained fewer MAP-2- or Neu-N- cells in the penumbral region than did that of SN-treated rats (Figure 4O).

**SN rescues neural tissue by blocking caspase-3 activity and increasing expression of antiapoptotic and signal-transduction protein.** To study the neuroprotective mechanism of SN in vivo, 12 rats (n = 6 per group) were euthanized 8 hours after MCA ligation and subjected to caspase-3 immunoreactivity tests. The penumbra surrounding the ischemic cores of the SN-treated rats only contained a few cells expressing activated caspase-3. Ischemic brain tissue from animals injected with the vehicle control, however, contained many cells positive for activated caspase-3 in both the penumbra and the ischemic core. Quantitatively, rats treated with SN showed fewer cells positive for activated caspase-3 after ischemia compared with controls (data not shown).

**To investigate the molecular mechanism underlying the antiapoptotic effect of SN, we examined the expression of apoptosis-related proteins and signal transduction proteins. Western blot showed significantly upregulated expression of antiapoptotic proteins such as Bcl-2 and Bcl-xL in SN-treated rats at 24 hours after cerebral ischemia compared with control rats (Figure 4, P and Q). In addition, significantly increased expression of phosphorylated Stat3 (p-Stat3) was found in the SN-treated ischemic rats compared with the vehicle control (Figure 4R). However, the increased expression of p-Stat3 was abolished by AG490 or TG101209 in the SN-treated rats (Figure 4R). Finally, the reduction of infarct volume after SN administration was also blocked by injection of AG490 or TG101209 (Figure 5, A and B). These data suggest that the antiapoptotic mechanism in cerebral ischemia related to SN treatment might involve inhibition of caspase-3 and upregulation of Bcl-2, Bcl-xL, and p-Stat3 expression.

**SN protects neural tissues from apoptosis.** Cellular apoptosis in ischemic rat brain was studied in SN-treated and control rats (n = 6 per group) using TUNEL analysis. The control animals, which did not receive MCA ligation, showed almost no TUNEL staining in their brain sections. The penumbral region surrounding the ischemic cores of SN-treated rats contained fewer TUNEL-positive cells than did that of vehicle controls (Figure 5C). Semic quantitatively, animals treated with vehicle had significantly more TUNEL-positive cells than those treated with SN (Figure 5C). TUNEL-positive cells were distributed mainly in the ischemic core of the cerebral cortex, with the labeling essentially found in the nucleus of neuronal cells.
**Figure 3**
SN administration i.v. in cerebral ischemic rats improves neurological dysfunction and reduces infarct size. (A) From 14 to 28 days after treatment, ischemic rats receiving SN i.v. showed significantly reduced body asymmetry after MCA ligation compared with controls. (B–D) Rats receiving SN showed significantly increased locomotor activities 14–28 days after treatment. (E) Final results of grip strength measurements showed a higher grip strength ratio in the SN-treated rats than in control rats. (F and G) Representative images of ischemic brain MRI; the white areas (white arrows) are the infarcted zones in the right cerebral cortices of the SN-treated and control rats on days 1, 7, and 28 after cerebral infarction. (H) Quantification of the infarct volume showed significant reduction in rats treated with SN on the seventh day after cerebral ischemia compared with controls. The area of largest infarction in the ischemic brain and the number of infarcted sections per rat were also reduced by SN treatment. (I) Representative 18FDG-PET (coronal view) of the right cortices (black arrows) of SN-treated and control rats. (J) Semiquantitative measurement showed that relative glucose metabolic activity in the right cortex, shown as the ratio of the ipsilateral hemisphere to the contralateral hemisphere, was much greater in the SN-treated group than in the control group. Data are mean ± SEM. *P < 0.05, **P < 0.01 vs. control.

**Table 1**
Physiological parameters were not altered by SN

<table>
<thead>
<tr>
<th>Parameter</th>
<th>SN (n = 7)</th>
<th>Control (n = 7)</th>
<th>P</th>
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<tr>
<td>pH</td>
<td>7.38 ± 0.041</td>
<td>7.33 ± 0.03</td>
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<td>PaCO2 (mmHg)</td>
<td>47.03 ± 1.11</td>
<td>51.4 ± 2.23</td>
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<tr>
<td>PaO2 (mmHg)</td>
<td>89.7 ± 3.15</td>
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<td>HCO3– (10–3 mol/l)</td>
<td>28.3 ± 1.56</td>
<td>24.8 ± 1.63</td>
<td>0.43</td>
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<td>Hematocrit (%)</td>
<td>45.13 ± 2.3</td>
<td>43.1 ± 3.21</td>
<td>0.28</td>
</tr>
<tr>
<td>Hemoglobin (10 g/l)</td>
<td>14.3 ± 0.61</td>
<td>15.4 ± 0.66</td>
<td>0.27</td>
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<td>Na+ (10–2 g/l)</td>
<td>138.4 ± 4.5</td>
<td>143.5 ± 2.3</td>
<td>0.65</td>
</tr>
<tr>
<td>K+ (10–3 mol/l)</td>
<td>4.1 ± 0.26</td>
<td>4.7 ± 0.42</td>
<td>0.79</td>
</tr>
<tr>
<td>Ca2+ (10–2 g/l)</td>
<td>4.1 ± 0.41</td>
<td>3.72 ± 1.2</td>
<td>0.54</td>
</tr>
<tr>
<td>Glucose (10–2 g/l)</td>
<td>149.4 ± 28.5</td>
<td>143.1 ± 15.2</td>
<td>0.53</td>
</tr>
<tr>
<td>MBP (mmHg)</td>
<td>78.9 ± 8.2</td>
<td>81.3 ± 6.4</td>
<td>0.55</td>
</tr>
<tr>
<td>HR (bpm)</td>
<td>391 ± 25</td>
<td>404 ± 19</td>
<td>0.66</td>
</tr>
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MBP, mean blood pressure; HR, heart rate. P values were determined by 2-tailed Student’s t test.
In agreement with previous studies, we found SN to have functions similar to those of glial cell–derived neurotrophic factor (GDNF), correlated to growth (18, 22–24) and repair (6, 25, 26) of injured neural tissue. Some literature has indicated that expression of SN is upregulated after cerebral ischemic insult (7). Similarly, TGF-β1 and GDNF mRNA levels have also been shown to increase after brain injury (27, 28). From this evidence, investigators have suggested that endogenous protective mechanisms could be activated after such insults (27, 28). In the present study, we have shown that increased expression of SN in the human stroke brain and in cerebral ischemic rats was induced to prevent further damage through endogenous genetic modulation following neuronal injury. Some reports have also demonstrated that a significant loss of SN-IR is found in the hippocampi of patients with Alzheimer disease and that a large part of the neuritic plaques in these patients are SN-immunopositive; the authors concluded that expression of SN in the hippocampus was closely correlated with synaptic loss in Alzheimer disease (3, 5).

In previous reports, SN has been considered to be an angiogenic cytokine like VEGF, which has mitogenic and antiapoptotic effects on endothelial cells (18, 25). These reports demonstrated that SN acts as a survival agent, as demonstrated by the reduction of annexin V–positive cells under serum starvation in HUVECs. In addition, these reports indicated that the mitogenic signaling pathway with SN is through activation of ERK1/2 and Akt pathways (18).

Figure 4
SN administration i.v. 30 minutes after cerebral ischemia rescued neurons from ischemic injury by directly inhibiting apoptotic signals. (A and B) Representative triphenyltetrazolium chloride–stained brain slices from rats injected with SN and vehicle. (C–N) Sections from the brain slices in A and B were immunostained to identify MAP-2+ neurons (C–H) and Neu-N+ neurons (I–N) in 3 different regions (corresponding with symbols in A and B). (O) The number of MAP-2+ and Neu-N+ immunoreactive cells in the penumbral area (shown in D, G, J, and M) was significantly increased in SN-treated rats compared with control rats. (P and Q) Significantly increased expression of Bcl-2 and Bcl-xL in SN-treated rats. (R) Increased expression of p-Stat3 in SN-treated rats, which was knocked down by AG490. Data are mean ± SEM. *P < 0.05 vs. control. Scale bar: 40 μm.

Figure 5
SN reduces apoptosis by activating the Jak2/Stat3 pathway. (A and B) Two days after cerebral ischemia, the SN-induced reduction of infarct volume was blocked by infusion of the inhibitor AG490. (C) Representative image of TUNEL (green) and Hoechst 33342 (blue) costaining showing cell death in penumbral of ischemic brains (arrowheads) from saline control and SN-treated rats. There were significantly fewer TUNEL-positive cells in SN-treated brain compared with saline control. Data are mean ± SEM. *P < 0.05 vs. control. Scale bar: 40 μm.
Although SN may exert a survival effect and induce immature cerebellar granular neurons to differentiate by promoting neurite outgrowth (6), few reports have focused on the neuroprotective role of SN in vitro and in vivo. We have demonstrated, for the first time to our knowledge, that SN exerts an antiapoptotic effect by reducing LDH activity, blocking activated caspase-3 activity, and upregulating expression of antiapoptotic proteins in primary cortical neurons in vitro. We also found that SN transduced an antiapoptotic signal in the ischemic penumbra, resulting in protection of cortical neurons and diminished infarct volume. In addition, i.v. administration of SN also increased the expression of Bcl-2 in primary cortical neurons and upregulated Bcl-2 and Bcl-xL in the penumbral region of cerebral ischemia. These in vivo effects of SN were conducted through induction of Jak2/Stat3, which was abolished by the specific inhibitors AG490 and TG101209. In summary, the neuroprotective effect of SN injection through direct inhibition of caspase-3 activation and enhancement of expression of antiapoptotic proteins (Bcl-2 and Bcl-xL) might be one of the more significant mechanisms involved in the rescue of injured neurons.

Taguchi et al. have recently shown that i.v. delivery of the CD34+ subpopulation of human umbilical cord blood cells can enhance angiogenesis, neurogenesis, and functional recovery in an animal stroke model (30). They further demonstrated that CD34+ cells stimulated angiogenesis with specific neovascularization around the cortical degeneration. Neurogenesis was in turn stimulated by these events, with subsequent migration of neuroblasts into the newly restored cortex, where these cells matured and contributed to functional recovery. Some reports have also assumed that enhancement of angiogenesis, and then neurogenesis, have a plasticity effect on the recovery of ischemic neurological insult (13, 14). In addition to the plasticity effect in the nervous system (31, 32), cytokines of neuronal origin (such as nerve growth factors) and...
several members of the neuropeptide group (such as neuropeptide Y; ref. 33) were reported to induce angiogenesis in injured areas of the brain. Furthermore, VEGF was shown to exert neuroplastic activity (34), by enhancing vascular supply of the nerves, and to improve neural function in ischemic and diabetic neuropathy (35, 36).

In the present study, we demonstrated that i.v. administration of SN 30 minutes after cerebral ischemia not only enhanced the incorporation of new vessels into the penumbral region to increase local cerebral blood perfusion, but induced neural differentiation of mobilized BMSCs and INPCs to reduce neurological dysfunction. Using a GFP-chimeric mouse model, we demonstrated that many GFP+ cells (from BMSCs), especially GFP+N Neu-N+ cells, migrated to the peri-infarcted area. A large amount of BrdU+N Neu-N+ and BrdU+GFAP+ cells were located in the subventricular and hippocampal regions in the SN-treated mice. In addition, some GFP+Musashi-1+BrdU+ and GFP+GFAP+BrdU+ cells were also found in the hippocampal, penumbral, and subventricular regions. Functional neural regeneration and neurogenesis after SN treatment contributed by both BMSCs and INPCs might be another important mechanism that induces improvement of neurological dysfunction following stroke. Therefore, SN may exert a significant neuroplastic effect because of efficient angiogenesis and neurogenesis in the penumbral region of the ischemic brain.

We know that circulating cytokines such as G-CSF, VEGF, stromal cell-derived factor-1 (SDF-1), statin, and erythropoietin (17, 37–40) enhance the mobilization of BMSCs. The mechanism of mobilization of BMSCs by these factors seems to be a receptor-mediated process involving CXCR4 and c-kit (41, 42), according to the SDF-1 concentration gradient theory (43). In previous studies, SN, an angiogenic cytokine, has been shown to chemoattract BMSCs, especially endothelial progenitor cells, from bone marrow to circulating peripheral blood (18, 25). However, the possible regulatory pathway by which SN mobilizes stem cells from bone marrow remains unknown. A previous investigation demonstrated that SN injection into mice caused significant proliferation of bone marrow cells positive for c-kit and Sca-1 (18). In addition, SN also mobilized and increased the number of Sca-1–positive cells in peripheral blood. In the present study, we demonstrated that SN enhanced the assembly of GFP+vWF+ and BrdU+vWF+ cells (from both BMSCs and INPCs) to form many vascular lumens over the penumbral, striatal, and hippocampal regions in GFP-chimeric mice. FITC-dextran perfusion of the ischemic brain and CD31+ blood vessel density revealed considerable new vessel formation in the SN-treated mice. At 7 days after cerebral ischemia, higher cerebral blood flow was detected in the SN-treated rats than in the controls.

In summary, we have shown that SN exerted neuroprotective effects in vitro by blocking activation of caspase-3, the downstream apoptotic enzyme, and upregulating the antiapoptotic proteins Bcl-2 and Bcl-xL. It was also neuroplastic in cerebral ischemic animals by means of instigating new neuronal and vascular formation within peri-infarcted regions of brain, attenuating tissue damage and subsequently reducing infarction volume and improving neurological function. We have disclosed, for the first time to our knowledge, a neuroplastic view of SN in a hypoxia and ischemia model. We believe that the SN treatment protocol presented here holds promise for the development of new therapeutic strategies, including gene therapy and that SN may possess utility as a small-molecular drug that can improve innate tissue regeneration and repair after cerebral ischemia to clinically significant levels.

Methods
IHC analysis of autopsy samples of human brain. We investigated autopsy brain specimens from 9 cases of fatal ischemic stroke (disease duration...
ranged from 15 hours to 7 days) treated at the Department of Neurology of Tzu-Chi General Hospital. Autopsy surgery was performed within a mean of 8 hours after death (range, 4–12 hours). Three patients who died of nonneurological causes served as controls (autopsies were conducted at 1, 2.5, and 4 days). The study protocol was approved by the Institutional Review Board of the Tzu-Chi General Hospital. Informed consent was obtained from relatives. Tissue sampling was based on individual infarct topography, which in each case was determined on the basis of cerebrovascular anatomy and the most recent MRI scan. On autopsy, brain areas with variable degrees of infarction were identified macroscopically, and cortical samples of about 1 cm$^3$, including subcortical white matter, were dissected and fixed with formalin prior to embedding in paraffin or frozen at –70°C until analysis, as described previously (44). SN immunostaining of brain samples was performed using specific antibody (1:200; Phoenix). Samples from corresponding areas of the contralateral or noninfarcted hemispheres and from the control brains were processed in a similar way. The extent of SN cell immunoreactivity was measured as number of cells per square millimeter.

Measurement of serum SN levels in acute stroke patients. We obtained serum samples at sequential time points (12 hours and 1, 3, 7, and 14 days) after onset of acute ischemic stroke. All patients’ protocols and consents were fully reviewed and approved by the Institutional Review Board of Tzu-Chi General Hospital. Levels of SN were measured with specific antibody using indirect ELISA techniques (Phoenix). Each serum sample was run in triplicate and compared with a standard curve. All samples were assigned a random number and run without knowledge of the donor’s disease or treatment status. Once the data were compiled, the sample classifications were revealed.

In vitro PCC preparation and OGD treatment. All animal research protocols were approved by the Academia Sinica Animal Care and Use Committee. PCCs were prepared from the cerebral cortex of gestation day–17 Sprague-Dawley rat embryos as previously described (45). Four days after isolation, the cultures were replenished with MEM (Invitrogen) containing 0.5 g/l BSA, 2% B27 supplement, 0.5 mM pyruvate, and antibiotics. Finally, the culture medium was changed to serum-free neurobasal medium containing 1 mM pyruvate, 1 mM glutamate, 0.5 g/l BSA, 2% B27 supplement, and antibiotics on the seventh day (45). For OGD treatment, the cells cultured with glucose-free Earle’s balanced salt solution were placed for 4 hours within a hypoxic chamber (Bug Box; Ruskinn Technology) and continuously flushed with 95% N$_2$ and 5% CO$_2$ at 37°C to maintain a gas-phase PO$_2$ less than 1 mmHg (OM-14 oxygen monitor; SensorMedics Corporation). Control cells were incubated in glucose-free Earle’s balanced salt solution in a normoxic incubator for the same time period. OGD was terminated by switching back to normal culture conditions.

Measurement of LDH activity and immunocytochemistry of MAP-2. In order to prove the concept of neuroprotection of SN, PCCs were prepared in 24-well plates and pretreated with 1 μg/l SN (Neosystems). After 20 minutes of SN pretreatment, PCCs were subjected to OGD in the hypoxia chamber for 4 hours, and then the culture media were collected for LDH activi-
bilateral common carotid artery (CCA) clamping as previously described. PCCs were pretreated with 1 μg/l SN and then placed in the hypoxia chamber for 2 hours. Fluorometric assays of caspase-3 activity were performed on the PCCs. PCCs treated as described above using commercial kits (Bio-Rad) according to the manufacturer’s instructions. For immunofluorescent study of activated-caspase-3, primary cortical neuron cultures were treated as described above and incubated with primary antibody against active fragment of caspase-3 (R&D Systems) as previously described (48). Caspase-3–positive cells were also quantified as previously described (48).

**Total protein extraction and Western blot assay.** Western blot analyses of p-Stat3, Stat3, p-Jak2, Jak2, Bcl-2, Bcl-XL, Bax, and Bad expression from PCC cultures were performed after SN treatment. Briefly, PCCs treated at a different time points (0.5, 1, 3, 8, and 12 hours) and with different doses (0.01, 0.1, 1 and 10 μg/l) of SN were lysed after treatment in a buffer containing 320 mM sucrose, 5 mM HEPES, 1 μg/ml leupeptin, and 1 μg/ml aprotinin. Lysates were centrifuged at 13,000 g for 15 minutes. The resulting pellet was resuspended in sample buffer (62.5 mM Tris–HCl, 10% glycerol, 2% SDS, 0.1% bromophenol blue, and 50 mM DTT) and subjected to SDS-polyacrylamide gel (4%–12%) electrophoresis. The gel was then transferred to a Hybond-P nylon membrane. This was followed by incubation with appropriately diluted antibodies to p-Stat3 (1:200; Cell Signaling), p-Jak2 (1:200; Cell Signaling), Stat3 (1:200; Santa Cruz), Jak2 (1:200; Santa Cruz), Bcl-2 (1:200; Santa Cruz), Bcl-XL (1:200; Transduction Laboratories), Bax (1:200; Santa Cruz), Bad (1:200; Transduction Laboratories), and β-actin (1:2,000; Santa Cruz). Specific Jak2 pathway inhibitor AG490 (100 μM; Calbiochem), which was used to pretreat cells, was applied to the PCCs to suppress enzyme binding in order to block the transcriptional signal of Jak2 as previously described (49). Membrane blocking, primary and secondary antibody incubations, and chemiluminescence reactions were conducted for each antibody individually according to the manufacturer’s protocol. The intensity of each band was measured using a Kodak Digital Science 1D Image Analysis System (Eastman Kodak).

**In vivo brain ischemia/reperfusion.** Adult male Sprague-Dawley rats (250–300 g) were used for this study. The rats were anesthetized with high-resolution small-animal PET (microPET Rodent R4; Concorde Microsystems Inc.). The system parameters were described by Visney et al. (57). In brief, 18F was produced by the 18O(p, n)18F nuclear reaction in a cyclotron at Tzu-Chi General Hospital and Tzu-Chi University, and 18FDG was synthesized as previously described (58) with an automated 18FDG synthesis system (Nihonkokan). Data were collected with high-resolution small-animal PET (microPET Rodent R4; Concorde Microsystems Inc.). The system parameters were described by Visney et al. (57). After 1 week of each treatment, animals anesthetized with chloral hydrate (0.4 g/kg i.p.) were fixed in a customized stereotactic head holder and positioned in the microPET scanner. The animals were then given an i.v. bolus injection of 18FDG (200–250 μCi/rat) dissolved in 0.5 ml saline. Data acquisition began simultaneously with injections.
and continued for 60 minutes in 1 bed position using a 3D acquisition protocol. The image data acquired from microPET were displayed and analyzed by IDL version 5.5 (Research Systems) and ASIPro version 3.2 (Concorde Microsystems) software. Coronal sections for striatal and cortical measurements represented brain areas between 0 and +1 mm from the bregma, while those for thalamic measurements represented areas between –2 and –3 mm from the bregma, as estimated by visual inspection of the unlesioned side. The relative metabolic activity in regions of interest of the striatum and cortex was expressed as percentage deficit, as previously described with modification (57).

Blood pressure, heart rate, blood glucose, and blood gas measurement. Physiological parameters were measured in 7 each of control and SN rats. The procedure has been described previously (59).

Activated caspase-3 immunostaining and Western blot analysis. Eight hours after ischemia, rats were anesthetized with chloral hydrate and perfused with 4% paraformaldehyde. Brain slices were incubated with primary antibody against caspase-3 (cleaved caspase-3 antibody, D175, dilution 1:500; Cell Signaling) conjugated with Cy3 (1:500; Jackson Immunoresearch) for 20 hours at 4°C, washed 3 times with PBS, and then observed by fluorescent microscopy (Axiovert 200M; Carl Zeiss). The extent of apoptosis was measured as the number of caspase-3+ apoptotic cells per square millimeter. In addition, apoptosis-related protein expression (Bcl-2, Bcl-xL, Bax, and Bad) and signal transduction protein (p-Stat3) in the right cortex and striatum region was also examined in the SN-treated and control rats using Western blot analysis as described previously (60).

TUNEL histochemistry. To detect cellular apoptosis, a TUNEL staining Kit (DeadEnd Fluorimetric TUNEL system; Promega) was used for the TUNEL assay. Twenty-four hours after ischemia, rat brains were fixed by transcardial perfusion with saline and immersed in 4% paraformaldehyde. After brains had been frozen on dry ice, a series of adjacent 10-μm-thick sections were cut in the coronal plane with a cryostat. The staining and semi-quantitating procedure was performed as described previously (61).

BrdU labeling and BrdU IHC. BrdU, a thymidine analog that is incorporated into the DNA of dividing cells during S-phase, was used for mitotic labeling (Sigma-Aldrich). The labeling protocol has been described previously (62). The BrdU immunostaining procedure with a specific antibody against BrdU (1:400; Boehringer Mannheim) and quantification of BrdU-immunoreactive cells have also been described previously (62). In brief, experimental rats’ brains were fixed by transcardial perfusion with saline, followed by perfusion and immersion in 4% paraformaldehyde. Subsequently, the brain samples were dehydrated in 30% sucrose. After brains had been frozen on dry ice, a series of adjacent 6-μm-thick sections were cut in the coronal plane with a cryostat, stained with H&E, and observed by light microscopy (E600; Nikon). For BrdU immunostaining, DNA was denatured by incubating each section in 20% formamide in 2x standard saline citrate at 65°C for 2 hours and then in 2N HCl at 37°C for 30 minutes, then rinsed in 0.1 M boric acid with pH 8.5. Sections were rinsed with Tris buffer and treated with 1% H2O2 to block endogeneous peroxidase. The immunostaining procedure was performed using the labeled streptavidin-biotin method (DAKO LSAB-2 Kit, Peroxidase; DAKO). Tissue slides were incubated with the appropriate diluted antibodies to BrdU (for nuclear identification, dilution 1:400; Boehringer Mannheim) at room temperature for 1 hour. After washing with Tris-buffered saline containing 0.1% Tween-20, the specimens were sequentially incubated for 10–30 minutes with biotinylated anti-rabbit and anti-mouse (1:200; R&D Systems) immunoglobulins and peroxidase-labeled streptavidin. Next, a pulse labeling method (n = 16) was used to observe the time course of cell proliferation in the brain after cerebral ischemia. Experimental rats were injected with BrdU (50 mg/kg i.p.) every 4 hours for 12 hours before sacrifice. A cumulative labeling method (n = 16) was used to examine the population of proliferative cells during 14 days of cerebral ischemia. Rats received daily injections of BrdU (50 mg/kg i.p.) for 14 consecutive days, starting the day after MCA ligation, and were sacrificed 14 days after the last injection.

Transgenic GFP-chimeric mouse preparation. In order to verify the enhancement of the INPC and BMSC mobilization and homing into brain by SN administration, a bone marrow sample was removed from the long bones of adult male donor mice as previously reported (63). Both ends of the femur and tibia were penetrated using a syringe with a 25-gauge needle, and the marrow was flushed out with sterile saline. Total marrow from 1 femur was diluted to 1 ml and then strained through 30-μm Spectramesh (Fisher Scientific). Before bone marrow transplantation, female recipient mice underwent whole-body gamma irradiation with 137Cs using a Gammacell 40 irradiator (MDS Nordion). A total dose of 9 Gy was administered to ablate the whole bone marrow. The mice received rescuing bone marrow transplantations within 24 hours of irradiation. Donor bone marrow was injected into the recipient animal’s tail as an 80-μl cell suspension containing 3 × 10^6 cells. At 3 weeks after transplantation, mice were anesthetized with chloral hydrate (0.3 g/kg i.p.) and subjected to right MCA ligation and bilateral CCA clamping for 60 minutes, as previously described with modification (50). At 60 minutes after arterial ligation, experimental mice were injected i.v. with recombinant human SN (20 μg in 200 μl saline; Neosystems) or vehicle (200 μl saline) through a 30-gauge syringe into the femoral vein. BrdU labeling was also performed for each mouse as described above.

Laser-scanning confocal microscopy for double-immunofluorescence analysis. To identify the coexpression of cell type–specific markers in SN-immunoreactive GFP+ and BrdU+ cells, immunofluorescent colocalization study with 3D images was performed to test for the expression of GFAP, α-SMA, vWF, MAP-2, Musashi-1, and Neu-N. The double-immunofluorescence technique with specific antibodies against BrdU (1:400; Mannheim), GFAP (1:400; Sigma-Aldrich), MAP-2 (1:200, Boehringer Mannheim), Nestin (1:400, Sigma-Aldrich), Neu-N (1:200, Chemicon), vWF (1:400, Sigma-Aldrich), Musashi-1 (1:100, Serotec) and SMA (1:100, BD Pharmingen) conjugated with Cy3, Cy5, or FITC (1:500, Jackson Immunoresearch) has been described previously (14). The tissue sections were analyzed with a Carl Zeiss LSM510 laser-scanning confocal microscope.

Angiogenic evaluation by FITC-dextran perfusion and CD31 IHC. In order to examine the blood vessels, cerebral microcirculation was analyzed by administering a fluorescent plasma marker (FITC-dextran; Sigma-Aldrich) to rats i.v. and observing them under fluorescent microscopy (Axiovert 200M; Carl Zeiss), as previously described (64). In addition, to quantify the cerebral blood vessel density and examine vascular remodeling by macrophage, experimental rats were anesthetized with chloral hydrate and perfused with 4% paraformaldehyde. Histological sections (6 μm) were stained with specific antibody to CD-31 (1:100; BD Biosciences—Pharmingen) conjugated with Cy3, Cy5, or FITC (1:500, Jackson Immunoresearch). The number of blood vessels was determined as previously described (30).

Measurement of CBF. Experimental rats were positioned in a stereotaxic frame, and baseline local cortical blood flow was monitored after cerebral ischemia with a laser Doppler flowmeter (LDF monitor; Moor Instruments) in anesthetized state (chloral hydrate) as previously described (65). In brief, CBF values were calculated as percentage increase over baseline local cortical blood flow.

Statistics. In this study, all observers were blind to the actual treatment result measurement. Sample sizes in animal treatment groups were monitored to ensure sufficient discriminatory power with Minitab, version 15, and were calculated to provide 80% power, sufficient to detect differences between each treatment. Results are expressed as mean ± SEM of all measurements in a treatment group. The behavioral scores have been evaluated...
for normality. Two-tailed Student’s t tests were used to evaluate significance of mean differences between the control and the treated group. Data lacking normal distribution were analyzed by nonparametric ANOVA.

P values less than 0.05 were taken as significant.

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