Implantation of olfactory ensheathing cells promotes neuroplasticity in murine models of stroke

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Murine olfactory ensheathing cells (OECs) promote central nervous system axonal regeneration in models of spinal cord injury. We investigated whether OECs could induce a neuroplastic effect to improve the neurological dysfunction caused by hypoxic/ischemic stress. In this study, human OECs/olfactory nerve fibroblasts (hOECs/ONFs) specifically secreted trophic factors including stromal cell–derived factor–1α (SDF-1α). Rats with intracerebral hOEC/ONF implantation showed more improvement on behavioral measures of neurological deficit following stroke than control rats. [18F]fluoro-2-deoxyglucose PET (FDG-PET) showed increased glucose metabolic activity in the hOEC/ONF-treated group compared with controls. In mice, transplanted hOECs/ONFs and endogenous homing stem cells including intrinsic neural progenitor cells and bone marrow stem cells localized with specific neural and vascular markers, indicating stem cell fusion. Both hOECs/ONFs and endogenous homing stem cells enhanced neuroplasticity in the rat and mouse ischemic brain. Upregulation of SDF-1α and CXCR4 in hOECs/ONFs promoted neurite outgrowth of cocultured primary cortical neurons under oxygen glucose deprivation conditions and in stroke animals through upregulation of cellular prion protein (PrPSc) expression. Therefore, the upregulation of SDF-1α and the enhancement of CXCR4 and PrPSc interaction induced by hOEC/ONF implantation mediated neuroplastic signals in response to hypoxia and ischemia.

Introduction

The olfactory mucosa (OM) contains unmyelinated olfactory nerves with olfactory ensheathing cells (OECs) and sheet-like processes of olfactory nerve fibroblasts (ONFs) (1, 2). The OECs guide growing axons from the neurons of the nasal cavity OM to the olfactory bulb to form synapses in the brain. Owing to their strong ability to guide axon outgrowth, OECs have been found to assist neuroregeneration in animal models of spinal cord injury and Parkinson disease (3, 4). Furthermore, OECs are pluripotent cells that can show Schwann cell–like properties (5). When transplanted into the demyelinated spinal cord, OECs can repair the defective myelin and restore conductance in remyelinated axons (6–8). Although much is known about OECs, few studies have focused on the characteristics of ONFs, which are flat and compact cells forming multiple layers with thin processes encircling OECs (2).

Neurotrophic factors support neuronal survival and axonal outgrowth (9). Many trophic factors secreted from human OECs/ONFs (hOECs/ONFs), such as brain-derived neurotrophic factor (BDNF), VEGF, and glial cell line–derived neurotrophic factor (GDNF), have been extensively studied to discover any potential they may have in promoting cerebral plasticity and enhancing functional recovery in animal stroke models (10–12). In recent reports, stromal cell–derived factor–1α (SDF-1α) and its receptor CXCR4 have been recognized as providing generalized trophic support to both embryonic and mature neurons and helping support neurons damaged by injury or inflammation (13–15). Furthermore, SDF-1α is a CXC chemokine produced by bone marrow stromal cells and is a potent chemotaxtractant for HSCs (16, 17) that is constitutively expressed by most tissues (18).

In the nervous system, cellular prion protein (PrPSc) is constitutively expressed in neurons and is abundant in regions such as the olfactory bulb, hippocampus, and synaptic neuropil in close spatiotemporal association with synapse formation. The physiological function of PrPSc remained obscure until several lines of prion gene–knockout mice were generated (19). Although these PrPSc-null mice were healthy throughout their life span, with normal development and resistance to scrapie infection (19), they were more susceptible to oxidative stress and apoptosis than wild-type mice (20). The localization of PrPSc in elongating axons suggests a role in axon growth (21, 22). Interestingly, interaction of PrPSc with the extracellular matrix protein laminin has been shown to promote neuritogenesis (23, 24).

In the present study, the cellular characteristics of hOECs/ONFs were studied in detail, including their immunophenotype, neuro-
trophic factor secretion, and signaling pathways under stress conditions. Using this cellular system, we intended to determine: (a) whether hOECs/ONFs could activate CXCR4 and produce SDF-1α in hypoxic and ischemic conditions; and (b) what role the interaction between soluble trophic factors such as SDF-1α and cell receptors including CXCR4 and PrP^C play in neuroregeneration in the ischemic brain. Specifically, transplantation of these olfactory accessory cells into stroke rats demonstrated that hOECs/ONFs enhance neuroplasticity, facilitate neurite outgrowth, and activate the stem cell homing process and thus promote reversal of the neurological deficit. In conclusion, this study demonstrates that hOECs/ONFs might potentially be of clinical benefit following autologous transplantation in stroke patients.

Results

Phenotypic characterization of hOECs/ONFs. hOECs/ONFs from surgical samples of nasal polyps were prepared and cultured on poly-D-lysine-coated chamber slides. They attached and grew slowly under standard culture conditions. The predominant cell morphology was spindle shaped, showing both a flattened fibroblast-like and an astrocyte-like pattern (Figure 1A). Immunocytochemical analysis consistently showed that at least 95% of cells expressed both low-affinity nerve growth factor receptor (p75) and S100 antigen and a variable percentage of cells (30%–70%) expressed fibronectin (FN) and glial fibrillary acidic protein (GFAP). Double immunofluorescence analysis demonstrated that the hOECs/ONFs coexpressed p75/GFAP, p75/S100, p75/FN, and GFAP/S100 (Figure 1B): 94% ± 2.8% of the cells expressed S100, 95% ± 3.3% of the cell population expressed p75, and 70% ± 2.1% expressed GFAP.

hOECs/ONFs secrete SDF-1α and upregulate CXCR4 under oxygen glucose deprivation treatment. In order to demonstrate the expression of SDF-1α and its receptor CXCR4, double immunofluorescence examination, ELISA, and Western blot analysis with specific antibodies were performed in the hOECs/ONFs. The hOECs/ONFs coexpressed SDF-1α and GFAP, SDF-1α and p75, CXCR4 and GFAP, and CXCR4 and p75 (Figure 1C). The level of BDNF, GDNF, and VEGF in the hOEC/ONF medium under oxygen glucose deprivation (OGD) conditions, as determined by ELISA, was higher than that in control (data not shown). Levels of SDF-1α (Figure 2A) and CXCR4 expression (Figure 2, B and C) also increased significantly 4 hours after OGD but fell to control levels over the next few hours. The corresponding cellular signaling pathways involved the activation of Akt and ERK1/2 one hour after OGD treatment (Figure 2, D and E), confirmed by the loss of increased SDF-1α expression following the addition of specific inhibitors of activated Akt (LY294002) or activated ERK1/2 (PD98059) to treated cells (Figure 2F). The expression of p38 and JNK was not significantly altered by OGD (Figure 2, D and E).

hOECs/ONFs enhanced neurite regeneration and survival of primary cortical cultures after OGD. To evaluate whether soluble factors secreted from hOECs/ONFs enhanced the neurite regeneration and survival of primary cortical cultures (PCCs) after OGD, neurite process elongation and number of neurons surviving were measured in PCCs cocultured with hOECs/ONFs. Following OGD, significantly enhanced neurite length (Figure 3, A and B) and significantly more neurite-bearing neurons (Figure 3B) were found in hOEC/ONF-cocultured PCCs compared with control.

To confirm the correlation between neurite regeneration and PrP^C expression, we performed Western blot and blocking antibody assays in a PCC and hOEC/ONF coculture system under OGD conditions. Western blot showed that expression of PrP^C in primary cortical neurons was significantly increased in PCCs cocultivated with hOECs/ONFs in comparison with PCCs alone (Figure 3C). Both the enhancement in neurite length and the increase in numbers of neurite-bearing neurons could be inhibited by addition of PrP^C-blocking antibody to the PCC coculture (Figure 3B).

PrP^C interacts with CXCR4 in vitro. In order to characterize the possible association between PrP^C and CXCR4, PCCs cocultured with hOECs/ONFs were analyzed by double immunofluorescence immunohistochemistry (IHC) and IP with specific antibodies. IHC showed that CXCR4 was distributed in the cytoplasmic aggregates and concentrated in the perinuclear caps of the PCC (Figure 3D). The PrP^C immunostaining appeared both in the cytoplasm and along neurites of the PCC (Figure 3D). Overlay of the IHC images for PrP^C and CXCR4 showed that they coincided in the neurite and perinuclear regions of the PCC (Figure 3D).

In order to verify whether CXCR4 binds to PrP^C, we carried out co-IP analysis with specific antibodies for CXCR4 and PrP^C. In the immunoprecipitated pellets, CXCR4 was seen to interact with PrP^C.
These experiments indicate that PrP<sup>C</sup> does interact directly with CXCR4 in the PCC and hOEC/ONF coculture system.

**hOEC/ONF transplantation improves neurological dysfunction after cerebral ischemia.** A body asymmetry trial and locomotor activity and grip strength assessments were used to assess whether implantation of hOECs/ONFs could improve neurological deficit after cerebral ischemia. Tests were conducted before and after middle cerebral artery (MCA) ligation in hOEC/ONF-treated rats and control rats. The behavioral measurement scores were all normalized to the baseline scores. Since unilateral cerebral ischemia causes imbalanced motor activity, all of the experimental rats developed significant body asymmetry, turning contralateral to the side of the induced ischemia on day 1 following cerebral ischemia. From 14 to 28 days after each treatment, rats with hOEC/ONF intracerebral transplants (n = 10) exhibited significantly reduced body asymmetry in comparison with control rats (n = 10) (Figure 4A). Locomotor activity was examined before and after cerebral ischemia in all animals. Vertical activity, vertical movement time, and the number of vertical movements significantly increased between 14 and 28 days after cerebral ischemia in rats receiving hOEC/ONF transplantation (n = 10) in comparison with control populations (n = 10) (Figure 4, B–D). Furthermore, measurement of forelimb grip strength for all experimental rats before and 28 days after treatment in each of the groups revealed a higher postischemia grip strength in the hOEC/ONF group (n = 10) than the control group (n = 10) (Figure 4E).

**Figure 2**

hOECs/ONFs upregulated the synthesis of soluble factors via the specific signaling pathway. (A–C) Under conditions of OGD, significantly increased expression of SDF-1α and CXCR4 was found by ELISA and Western blot analyses, respectively. (D and E) Expression of signal transduction proteins (p-Akt and p-ERK1/2) increased significantly by 1 hour after treatment. (F) The upregulation of SDF-1α could be blocked by specific inhibitors of p-Akt and p-ERK1/2 (LY294002 and PD98059). Data are expressed as mean ± SEM. *P < 0.05 and **P < 0.01 versus control.
Enhancement of glucose metabolic activity in intracerebral hOEC/ONF transplantation group after cerebral ischemia. To verify whether hOEC/ONF implantation could enhance glucose metabolic activity, experimental rats were examined by [18F]fluoro-2-deoxyglucose PET (FDG-PET) using microPET. At 4 weeks after each treatment, the uptake of FDG was strikingly greater in the right cortical region (ischemic area of model) of the hOEC/ONF-treated group (Figure 4F). Semiquantitative measurement of relative glucose metabolic activity of the right cortical region revealed significant enhancement in the hOEC/ONF-treated (n = 6) compared with the control group (n = 6) (Figure 4F).

Intracerebral transplantation of hOECs/ONFs increased expression of growth factor and antiapoptotic proteins in vivo. In order to demonstrate whether the improvement in neurological function was correlated with soluble factors and survival agents after cerebral ischemia in hOEC/ONF-treated animals, we examined the expression of growth factor and antiapoptotic proteins in the ischemic cortical area using ELISA, immunohistochemical analysis, and Western blot analysis. Consistent with our immunohistochemical evidence, ELISA revealed a significantly increased level of SDF-1α in the ischemic rats treated with hOECs/ONFs (n = 6) in comparison with vehicle-treated controls (n = 6) (Figure 4G). In addition, significantly upregulated expression of antiapoptotic proteins Bcl-2 and Bcl-xl, but not Bax and Bad, was found in hOEC/ONF-treated rats (n = 6) at 3 days after cerebral ischemia compared with control rats (n = 6) (Figure 4, H and I).

Intracerebral hOEC/ONF transplantation stimulates endogenous stem cell mobilization, homing, and engraftment into rat brain following cerebral ischemia. BrdU labeling was used to follow the growth of these homing stem cells in the brain to determine whether intracerebral hOEC/ONF transplantation could enhance homing and engraftment of endogenous stem cells (from host brain and peripheral blood) into the damaged regions of the brain following ischemia. Cumulative BrdU labeling in hOEC/ONF transplantation rats revealed BrdU-immunoreactive cells in the ipsilateral hemisphere near the penumbra area, the striatal region (Figure 5, A–C), and the subventricular region of the lateral ventricle (Figure 5, D–F).

Figure 3
In coculture, hOECs/ONFs promoted neurite regeneration and survival of PCC under OGD. (A) Representative images of neurite regeneration in PCC with or without hOEC/ONF cocultivation. Blo-Ab, blocking Ab. (B) Neurite length and the number of surviving neurons increased in the hOECs/ONFs cocultured with PCC compared with the control. The enhancement of neurite length and number of surviving neurons was inhibited by the neutralization peptide of PrP. (C) Upregulation of PrP expression was found in PCC cocultured with hOECs/ONFs. (D) In IHC, subcellular colocalization of CXCR4 and PrP was shown by costaining with CXCR4 and PrP antibodies in PCCs cocultured with hOECs/ONFs. (E) Using coimmunoprecipitation analysis, the cell lysates were immunoprecipitated (IP) with anti-CXCR4 or anti-PrP, and the signal of both proteins was detected by Western blotting (WB). Data are expressed as mean ± SEM. *P < 0.05 and **P < 0.01 versus control. Scale bars: 50 μm.
blood vessels of varying calibers (and also in endothelial cells of the vessel walls) (Figure 5, G–I) in the penumbral region. In BrdU pulse-labeling experiments, the number of BrdU-immunoreactive cells in the penumbral region of the ischemic cortex and striatum significantly increased in the hOEC/ONF-treated rats compared with saline-treated control rats (n = 8; P < 0.05) (Figure 5J). Intracerebral implantation of hOECs/ONFs could reconstruct the neural regeneration in the stroke model. To determine whether transplanted hOECs/ONFs and endogenous homing stem cells (from bone marrow stem cells [BMSCs] and INPCs) were responsible for the neuroplastic effect in the brains of hOEC/ONF-treated GFP-chimeric mice (n = 10) and BrdU-labeled mice (n = 10), the colocalization of cell type–specific markers at 28 days after transplantation was analyzed by double fluorescence IHC using laser scanning confocal microscopy (Figure 6A). There were significantly more BMSCs (GFP⁺ cells; green fluorescence) in the hOEC/ONF-treated GFP-chimeric mice than in controls (n = 10) (Figure 6A), and they were dispersed around the periphery of the transplanted hOECs/ONFs (Figure 6A). Moreover, there were also substantially more INPCs (BrdU ⁺ nestin⁺) in the hOEC/ONF-treated BrdU-labeled mice than in controls (n = 10) (Figure 6B). FISH analysis demonstrated that the implanted hOECs/ONFs labeled by bis-benzimide (blue fluorescence) were of human origin (Figure 6A). Interestingly, the implanted hOECs/ONFs and GFP⁺ cells showed evidence of cell fusion (or nuclear fusion) (Figure 6C). About one-third of the hOECs/ONFs that had fused with GFP⁺ cells (binucleated cells) (Figure 6C). In a colocalization study (3D image) of exogenous transplanted hOECs/ONFs and endogenous homing stem cells, some bis-benzimide-labeled cells and some GFP⁺ cells colocalized with microtubule associated protein–2 (MAP-2), vWF, and GFAP⁺ cells (red, neural cell–type specific markers) (Figure 6D) in the penumbra of hOEC/ONF-treated ischemic mouse brains. In addition, a few bis-benzimide-labeled hOECs/ONFs and GFP⁺ cells contained for SDF-1α immunoreactivity (Figure 6E).

Discussion

Although many studies have focused on OECs with regard to reversal of demyelination and axonal degeneration such as in spinal cord injury (3, 25–27), few reports have looked at the ability of OECs to repair ischemic neural injuries. In previous studies, it has been demonstrated that the olfactory epithelium (OE), which is highly vulnerable to injury, is endowed with a constitutive capacity for progenitor cell proliferation to reconstruct damaged olfactory neurons (28, 29). Furthermore, recent reports have shown that OE can induce simple neurogenesis following direct damage caused by exposure to methyl bromide gas (30, 31). This neurogenesis might be facilitated by factors including Mash1 (32) and Ngn1 (33) and enhance the proliferation of progenitor cells in the olfactory system. Therefore, in this report, we intended to re-verify the neuroplastic capacity of hOECs/ONFs using a different stress model of hypoxia/ischemia in both PCC and a rat stroke model. First, in view of the role of trophic factors in neuroprotection, the constitutive synthesis of many growth factors, such as BDNF, GDNF, and VEGF, has been shown to be protective under different stress models for both PCC and a rat stroke model. In our study, we also found that the level of BDNF, GDNF, and VEGF significantly increased in hOEC/ONF medium after OGD and we showed that SDF-1α was found both in the cultured hOEC/ONF medium after OGD and the hOEC/ONF-transplanted ischemic brain. The corresponding SDF-1α receptor, CXCR4,
was also highly expressed in the hOECs/ONFs. Since SDF-1α could be secreted from specific glial cells under certain conditions (13, 39), one can assume that hOECs/ONFs share features in common with glial cells (40) and can secrete SDF-1α under conditions of hypoxia/reoxygenation. Furthermore, there is also some evidence suggesting increased expression of SDF-1α and CXCR4 in cases of neuropathogenesis induced by many forms of injury, including trauma, stroke, and inflammation (14, 15, 41). Therefore, SDF-1α might not only provide generalized trophic support to both embryonic and mature neurons, but also help support neurons damaged by injury or inflammation. In addition, in a recent report, SDF-1α was found to play a critical role as a guidance molecule regulating neurite outgrowth in cultured cerebellar granular neurons (42). However, even if expression of SDF-1α was downregulated in the hOEC/ONF implantation, hOECs/ONFs could still produce other factors to rescue the injured neurons.

In this study, hOECs/ONFs transduced the neuroprotective signal through direct upregulation of antiapoptotic proteins (Bcl-2 and Bcl-xL) and enhancement of neurite regeneration, as well as via the known soluble trophic factors BDNF, GDNF, and VEGF. Therefore, SDF-1α, one of the major trophic factors secreted by hOECs/ONFs, might also play a significant role in neural survival and regeneration in our hypoxia/ischemia model.

In a recent report, Beites et al. concluded that neural stem cells resident in the OE could be regulated dynamically to induce neurogenesis during development and enhance regeneration after injury (43). In fact, one of the major components of OE stem and progenitor cells was OECs (43). Many reports have demonstrated that OECs can rebuild a demyelinated axon via production of a Schwann cell–like sheath in the spinal cord of an injured animal (3, 6, 44). Subsequent to the remyelination, OECs also replaced the myelin to regenerate disconnected axons and finally restored the neural conductance of the corticospinal tract (45, 46). Furthermore, OEC transplantation also stimulated neurite regrowth to recover body movement in spinal cord injury (47, 48).

In this report, we present the first evidence to our knowledge that implantation of hOECs/ONFs enhances homing of endogenous stem cells, from peripheral blood and host brain, into the ischemic brain. Recent reports indicate that SDF-1α/CXCR4 is a strong chemoattractant for stem cells and also plays a significant role in endogenous stem cell trafficking between peripheral circulation and bone marrow (49). A further study indicated that focal cerebral ischemia causes an upregulation in SDF-1α/CXCR4 expression over the peri-infarct area (13). This induction of upregulation of endothelial SDF-1α (50) and the appearance of increased CXCR4 expression in the ischemic hemisphere after ischemia explained in our previous report (51) together indicate that SDF-1α secreted from the implanted hOECs/ONFs in this study could be a chemoattractant for endogenous stem cells. Therefore, exogenous implanted hOECs/ONFs and endogenous homing stem cells might coordinate to exert a neuroplastic effect to repair the injured brain.

In recent reports, OECs have been proven to promote neurite regrowth in vitro (48), and secreted neurotrophic factors and cell-cell contact mechanisms are probably involved in the neurite regeneration (48). Although previous reports have demonstrated that growth factors secreted from OECs such as nerve growth factor (NGF), BDNF, GDNF, and neurturin (NTN) might indirectly...
facilitate axon regeneration (52), the actual molecules involved in the cell-cell contact mechanism that directly mediated the neurite outgrowth–stimulating effect of hOECs/ONFs are not completely known. Some investigations have found that cell adhesion molecules, such as L1 and N-cadherin, were associated with the neurite outgrowth (53, 54). Furthermore, it is known that PrP<sub>C</sub> plays an important role in the regulation of neurite regeneration (23). In a recent study, some proteins in the living brain involved in cell adhesion and neurite outgrowth were found in the presence of PrP<sub>C</sub> (55). Therefore, PrP<sub>C</sub> must interact with some intercellular matrix proteins to facilitate neurite regeneration. For example, PrP<sub>C</sub> participated in neurite adhesion through its interaction with laminin (24). In addition, the laminin receptor, which is a critical factor for cell differentiation and proliferation, was also identified as a cell-surface binding companion of PrP<sub>C</sub> (56). Since laminin contributes to axon growth and fiber tract formation, PrP<sub>C</sub> is particularly relevant for neurite formation during brain development (57). Although the SDF-1α/CXCR4 signaling pathway has also been shown to regulate axonal elongation (58) and guide the neurite growth cone (59, 60), few reports have investigated the interaction between CXCR4 and PrP<sub>C</sub> in neural regeneration (61). The combination of SDF-1α/CXCR4 with laminin might modulate cellular migration (62). In this study, we have demonstrated for the first time to our knowledge that coexpression of CXCR4 and PrP<sub>C</sub> might enhance the neurite growth in the cellular component of hOECs/ONFs cocultured with PCC under OGD conditions and in the brain of hOEC/ONF-implanted stroke animals. It is possible that these 2 proteins interact in trans with each other, that is, they facilitate an interaction between the cell surface of one cell and the extracellular protein matrix or the surface of an adjacent cell. Furthermore, investigation using blocking antibodies against CXCR4 and PrP<sub>C</sub> showed inhibition of neurite regeneration and even neurological function recovery.

In general, one of the major mechanisms by which bone marrow stem/progenitor cells repair tissue injury is cell/nuclear fusion (63–65). Terada et al. and Ying et al. investigated spontaneous fusion between embryonic stem cells (63) and neural stem cells (64). The frequency of cell/nuclear fusion ranged from 10<sup>-4</sup> per Figure 6

Biological mechanism of neuroplastic effects on the ischemic brain after intracerebral transplantation of hOECs/ONFs. (A) In a representative brain section of a GFP-chimeric mouse treated with or without hOECs/ONFs (white arrow indicates the injection site), GFP<sup>+</sup> cells are seen dispersed over the periphery of the transplanted hOECs/ONFs and were significantly increased in quantity in the hOEC/ONF-treated mice in comparison with controls. In FISH analysis (white arrow, 2 red spots), hOECs/ONFs were shown to be of human origin (inset square in left panel). (B) IHC of hOEC/ONF treatment in the BrdU-labeled mice. Many BrdU-nestin<sup>+</sup> cells were distributed around the transplanted hOECs/ONFs. (C) Interestingly, 1 cell with 2 nuclei (cell fusion) was found in the implanted hOECs/ONFs (white arrows, blue nucleus) and GFP<sup>+</sup> cells (white arrowheads, red nucleus). The nucleic dye TOTO-3 (red) was used to define the outline of all nuclei in the section. (D) In a colocalization study (3D image) some bis-benzimide–labeled cells and some GFP<sup>+</sup> cells colocalized with MAP-2<sup>+</sup>, vWF<sup>+</sup>, and GFAP<sup>+</sup> cells in the penumbra of hOEC/ONF-treated ischemic rat brains. (E) SDF-1α-immunoreactive cells colocalized with a few bis-benzimide–labeled hOECs/ONFs and GFP<sup>+</sup> cells. Data are expressed as mean ± SEM. *P < 0.05 versus control. Scale bars: 50 μm.
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cell plated (63, 64) to one-quarter of cells (66). They concluded that the phenomenon of cell/nuclear fusion was a frequent rather than a rare event. In our study, we demonstrated that cell/nuclear fusion occurred between hOECs/ONFs and BMSCs. Although a higher percentage of fused cells was found in this model than in previous investigations (66), the other 2 mechanisms also correlated well with stem cell–induced neuroplasticity in this study, including differentiation into tissue-specific phenotypes and secretion of many trophic factors.

**Methods**

**Separation and culture of hOECs/ONFs**: Human nasal polyp (hNP) samples (5 mm³, 0.5 g in weight) were collected in sterile boxes containing HBSS (Gibco-BRL; Invitrogen), for primary culture within 24 hours. Protocols for sampling hNP were approved by the Institutional Review Board of China Medical University and Hospital. Written informed consent was obtained from all patients. In brief, the donated tissue was dissected into small pieces under a dissecting microscope and placed in a phosphate-buffered solution at room temperature. The tissue was then ground with a dissection scalpel and transferred into 10 ml DMEM/F12 medium containing trypsin and EDTA and shaken at 37°C in a water bath for 5 minutes. It was then rinsed with DMEM/F12 solution and triturated with a fire-polished Pasteur pipette. The ground tissue explants were collected by centrifugation at 600 × g for 10 minutes. The resulting pellet was resuspended in DMEM/F12 medium containing B27 medium supplement, 10% FCS, and 1% (100 U/ml) penicillin/streptomycin at 3 × 10^6 cells/ml per culture medium. The tissue was placed in a 75-cm² flat flask and incubated in 5% CO₂ at 37°C. The tissue was left undisturbed for 5–7 days to allow for migration of the cells from the explants. These primary cells were passaged again once a week for 3–4 weeks.

**Immunocytochemical analysis and quantitative assessment of antigenicity in hOECs/ONFs**: Immunocytochemical studies of the hOECs/ONFs were performed using different antibodies: p75 (1:100; Millipore), GFAP (1:300; Millipore), FN (1:1,000; Molecular Probes; Invitrogen), S100 (1:1,000; Dako), oligodendrocyte marker 4 (04; 1:100; Millipore), SDF-1α (1:200; Torrey Pines Biolabs), CXCR4 (1:200; Torrey Pines Biolabs), neurofilament-200 (NF-200; 1:300; Sigma-Aldrich), βIII-tubulin (TuJ-1, 1:200; Millipore), MAP-2 (1:300; Millipore), and neuronal nuclear antigen (Neu-N; 1:50; Millipore). Cells were plated on a poly-L-lysine–coated chambered glass slide at a standard density and allowed to grow at 37°C in 5% CO₂ for 24 hours. The antigenicity of the cell population was quantitated as previously described with modification (67). In brief, a random field of view was selected at ×200 magnification with the UV filter. Images were captured and assessed using a spot digital camera (Diagnostic Instruments) and Northern Eclipse 5.0 software (Empix).

**Total protein extraction, Western blot assay, and ELISA in vitro**: For OGD treatment, the hOECs/ONFs were cultured with glucose-free Earle’s balanced salt solution, placed in a hypoxic chamber (Bugbox; Ruskinn) for 4 hours, and continuously flushed with 95% N₂ and 5% CO₂ at 37°C to maintain a gas-phase PO₂ of less than 1 mmHg (OM-14 oxygen monitor; SensorMedics). Control cells were incubated in glucose-free Earle’s balanced salt solution in a normoxic incubator for the same period. OGD was terminated by switching back to normal culture conditions. After OGD treatment, the cells were returned to a 37°C normoxic incubator (95% air and 5% CO₂) for different time periods (30 minutes, 1 hour, 4 hours, 12 hours, 24 hours, and 48 hours) of reoxygenation. The hOECs/ONFs were then collected at each time point and stored at -80°C for protein extraction. Western blot analyses of hOECs/ONFs were performed after hypoxia–reoxygenation treatment. Briefly, hOECs/ONFs were lysed 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-PAGE (4%–12%). Proteins in the gel were then transferred to a Hybond-P nylon membrane. This was followed by incubation with appropriately diluted antibodies against CXCR4 (1:100; Millipore), PrP<sup>β</sup> (1:300; M20; Santa Cruz Biotechnology Inc.), activated Akt (1:200; Calbiochem), activated ERK1/2 (1:200; Santa Cruz Biotechnology Inc.), activated p38 (1:200; Santa Cruz Biotechnology Inc.), activated JNK (1:200; Santa Cruz Biotechnology Inc.), and β-actin (1:2,000; Santa Cruz Biotechnology Inc.). Specific ERK1/2 inhibitor pathway inhibitor PD98059 (10 μM; Cell Signaling Technology) and Akt pathway inhibitors Wortmannin or LY294002 (10 μM; Calbiochem), which were used to pretreat cells, were applied to the hOECs/ONFs to suppress enzyme binding in order to block the transcriptional signal of ERK1/2 and Akt. 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. In addition, the total amount of growth factors in the medium was measured with a Quantikine ELISA kit (R&D Systems) according to the manufacturer’s instructions. Optical density was measured using a spectrophotometer (Molecular Devices), and standard curves were generated with the program SoftMax (Molecular Devices).

**In vitro PCC preparation and coculture with hOECs/ONFs**: PCC cells were prepared from the cerebral cortex of gestation day-17 embryos from Sprague-Dawley rats as described previously (68). In brief, pooled cortical tissues isolated from the brain were dissociated by mechanical trituration in HBSS. Then, cells were counted and plated at a density of 5 × 10⁶ cells in 24-well culture plates precoated with poly-L-lysine (0.02 g/l). Cells were grown in DMEM (Gibco-BRL; Invitrogen) with 10% heat-inactivated FBS, 1 mM pyruvate, 4.2 mM sodium bicarbonate, 20 mM HEPES, and antibiotics. Cultures were maintained at 37°C in a humidified atmosphere of 5% CO₂ and 95% air. Four days after isolation, the cultures were replenished with MEM (Gibco-BRL; Invitrogen) containing 0.5 g/l BSA and N-2 supplement, 0.5 mM pyruvate, and antibiotics. Finally, the culture medium was changed to a serum-free MEM containing 1 mM pyruvate, 1 mM glutamate, 0.5 g/l BSA, 0.3 mM KCl, and antibiotics on the seventh day of incubation.
During the coculture procedure, the culture medium was removed from hOEC/ONF culture and replaced with a 1.5-ml cell suspension containing 2 x 10^5 PCC cells. These cocultures were maintained for 3 days in the same culture condition. For OGD treatment, the cocultured cells were incubated with glucose-free Earle's balanced salt solution, placed in a hypoxic chamber (Bugbox) for 4 hours, and continuously flushed with 95% N2 and 5% CO2 at 37°C to maintain a gas-phase PO2 of less than 1 mmHg (OM-14 oxygen monitor; SensorMedics). After OGD treatment, the cocultured cells were returned to a 37°C normoxic incubator (95% air and 5% CO2) for different time periods (1, 3, and 7 days) of reoxygenation, before immunostaining and Western blot analysis.

**Immunocytochemistry and assessment of neurite regeneration.** For β-tubulin immunostaining, cell cultures were washed with PBS and fixed for 30 minutes at room temperature in 4% paraformaldehyde. After washing in PBS, the fixed cultures were treated for 30 minutes with blocking solution (10 g/l BSA, 0.03% Triton X-100, and 4% serum in PBS). Cells were incubated overnight at 4°C with an antibody against β-tubulin (1:200; Millipore; Millipore) for 3 hours, followed by a secondary antibody conjugated with FITC for 1 hour, and then rinsed 3 times in PBS. Finally, the slides were lightly counterstained with DAPI, washed with water, and then mounted.

The number of neurite-bearing cells and neurite length were assessed as previously described previously, with modification (69). In brief, cells in each treated group were plated after OGD, fixed, and immunostained for β-tubulin. For quantification, neurons with neurites were defined as those bearing a process greater than twice the cell body length. The length of the longest neurite of each neuron was measured from digitized images and quantified using the SigmaScan imaging analysis program (SigmaScan 4.01.003). All measurement data were calculated from experiments performed in triplicate.

To assess the correlation between neurite regeneration and PrP expression, we evaluated the neurite length and number of neurite-bearing cells in the presence of prion protein-blocking antibody (6H4; Prionics) under OGD conditions. The primary cortical neurons were pretreated with PrP-blocking antibody (6 μg/ml) and then cocultured with OECs for about 40 hours (70), before the cocultured cells received OGD treatment and neurite regeneration was assessed as outlined above.

**Communoprecipitation analysis.** The immunoprecipitation experimental procedures were as previously described (71). First, cells were managed with lysis buffer (50 mM Tris-HCl, pH 7.5, 1% NP-40, 150 mM NaCl, 0.5% sodium deoxycholate, and protease inhibitor). The lysate (300 μg) was incubated with protein A/G-agarose beads at 4°C for 6 hours. Then, CXCR4 (mAb 173; R&D Systems) and PrP C (C-20; Santa Cruz Biotechnology Inc.) (72) antibodies were added and reacted for 6 hours at 4°C. These immunocomplexes were incubated on protein A/G-agarose beads at 4°C overnight. After washing with lysis buffer 3 times, the immunocomplexes were examined by Western blot with anti-CXCR4 (Millipore) and anti-PrP C (M-20; Santa Cruz Biotechnology Inc.) antibodies.

**Animal brain ischemia/reperfusion model.** Adult male Sprague-Dawley rats were used in this study. Animals were subjected to 3-vehicle ligation, as previously described (31). All surgical procedures were performed using sterile/aseptic techniques in accordance with institutional guidelines. All animal experiments were approved by the Institutional Review Board of Animal Experiments, China Medical University Hospital.

**Intracerebral transplantation of hOECs/ONFs.** For cell labeling, cells were cultured in DMEM (Sigma-Aldrich) with 10% FCS at 37°C in a humidified atmosphere of 5% CO2/95% air and antibiotics. Prior to transplantation, the cells were incubated with 1 μg/ml bis-benzimide (labels nuclei with blue fluorescence) (Hoechst 33342; Sigma-Aldrich) for 5 hours at 37°C. The labeled cells were then collected and washed in PBS 5 times. Nucleated hOECs/ONFs were counted using a cytometer to ensure an adequate cell number for transplantation. One day after brain ischemia, the adult male Sprague-Dawley rats (weight, >300 g; age, 7–8 weeks) were anesthetized with chloral hydrate (0.4 g/kg, i.p.). Then they were injected stereotaxically with approximately 1 x 10^6 cells in 3–5 μl DMEM medium through a 26-gauge Hamilton syringe into 3 cortical areas adjacent to the right MCA, 3.0–5.0 mm below the dura as previously described (51). Cyclosporin A (CsA; 1 mg/kg/d, i.p.; Novartis) injections were given daily to each experimental rat from the day after cerebral ischemia for 3 weeks. An equal volume of CsA was injected to the transplantation group and saline control group, as previously described (73).

**Neurological behavioral measurement.** Behavioral assessments were performed 5 days before cerebral ischemia and 1, 7, 14, and 28 days after cell transplantation. The tests measured body asymmetry, locomotor activity, and grip strength (51, 74). The baseline scores were recorded in order to normalize those taken after cerebral ischemia, as previously described. Grip strength was analyzed using a Grip Strength Meter (TSE Systems) as previously described, with modification (74). In brief, the grip strength ratio for each forelimb was measured separately and was calculated as the ratio of the mean strength (n = 20 pulls) of the side contralateral to the ischemia to that of the ipsilateral side. In addition, the ratio of grip strength after treatment to that before treatment was calculated; the changes are presented relative to the pretreatment value.

**FDG-PET examination.** Since glucose metabolism is strongly correlated with functional plasticity of the brain, experimental rats were examined using microPET scanning of FDG to measure relative glucose metabolic activity, as previously described (75). In brief, 18F was produced by the 18O(p, n)18F nuclear reaction in a cyclotron at China Medical University and Hospital, and FDG was synthesized as previously described (76) with an automated FDG synthesis system (Nihonkoken). Data were collected with a high-resolution small-animal PET (microPET Rodent R4; Concorde Microsystems). The system parameters were described by Carmichael et al. (77). After 4 weeks of each treatment, animals were anesthetized with chloral hydrate (0.4 g/kg, i.p.), and the head was fixed in a customized stereotactic head holder and positioned in the microPET scanner. Then the animals were given an intravenous bolus injection of FDG (200–250 μCi/rat) dissolved in 0.5 ml saline. Data acquisition began at the same time and continued for 60 minutes in one 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 ASIPpro version 3.2 (Concorde Microsystems) software. FDG-PET images were reconstructed using a posterior-based 3D iterative algorithm (78) and overlaid on MR templates to confirm anatomical location (79). Coronal sections for striatal and cortical measurements represented brain areas between 0 and +1 mm from the bregma, while thalamic measurements were between –2 and –3 mm from the bregma, as estimated by visual inspection of the contralateral side. The relative metabolic activity in regions of interest of the striatum and cortex was expressed as percent deficit as previously described with modification (77).

**BrdU labeling and BrdU IHC.** BrdU (Sigma-Aldrich), a thymidine analog that is incorporated into the DNA of dividing cells during S phase, was used for mitotic labeling by a protocol described previously (80). Briefly, a pulse-labeling method was used to observe the time course of proliferative cells in the brain after cerebral ischemia. Experimental rats were i.p. injected with BrdU (50 mg/kg) every 4 hours for 12 hours before sacrifice. A cumulative labeling method 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. These rats were sacrificed 14 days after the last injection. The BrdU immunostaining procedure with a specific antibody against BrdU (1:400; Boehringer Mannheim) and quantification of BrdU immunoreactive cells have been described previously (80). In brief, experimental rats' brains were fixed by transcardial perfusion with saline, fol-
ollowed by perfusion and immersion in 4% paraformaldehyde. Subsequently, the brain samples were dehydrated in 30% sucrose. After brains were 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 first denatured by incubating each section in 50% formamide in 2x standard saline citrate at 65°C for 2 hours, then in 2 N HCl at 37°C for 30 minutes, and finally rinsed in 0.1 M boric acid with pH 8.5. Sections were then rinsed with Tris buffer and treated with 1% H2O2 to block endogenous peroxidase. The immunostaining procedure was performed using the labeled streptavidin-biotin (LSAB) method (LSAB-2 Kit, Peroxidase; Dako). Tissue slides were incubated with the appropriate diluted antibodies against BrdU (for nuclear identification; 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.

Preparation of transgenic GFP-chimeric mice. In order to verify the enhancement of the BMSC mobilization and homing into brain following hOEC/ONF implantation, transgenic GFP-chimeric mice generated as previously reported were used (81). In brief, 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, then strained through 30-μm Spectramesh (Fisher Scientific). Before bone marrow transplantation, female recipient wild-type (C57BL/6 mice) underwent whole-body gamma irradiation with 10 Gy using a Gammacell 40 (iradiator (MDS Nordic)). 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 x 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 common carotid artery clamping for 60 minutes, as previously described with modification (82). Then, 60 minutes after arterial ligation, experimental mice were implanted stereotactically with hOECs/ONFs (2 x 10^5 cells) or vehicle (200 μl saline) through a 30-gauge Hamilton syringe into 2 cortical areas, 2.0–3.0 mm below the dura. The approximate coordinates for these sites were 0.5–1.0 mm anterior to the bregma, 1.5–2.0 mm lateral to the midline, 1–1.5 mm posterior to the bregma, and 2–2.5 mm lateral to the midline.

Western blot analysis for expression of antipaptotic proteins and ELISA for growth factor in vivo. Experimental rats were anesthetized with chloridral hydrate (0.4 g/kg, i.p.) 3, 7, 14, and 28 days after initiation of the treatments. Ischemic cortical and striatal areas were evacuated on ice immediately. Subsequently, these brain tissue samples were homogenized with a plastic homogenizer, and growth factors were measured by ELISA analysis, as previously described with modification (83), with a Quantikine ELISA kit (R&D Systems) following the manufacturer’s instructions. Optical density was measured using a spectrophotometer ( Molecular Devices). The standard curves were obtained using the program SoftMax ( Molecular Devices). Western blot analysis was conducted with specific antibodies against β2 (1:200; Santa Cruz Biotechnology Inc.), β-actin (1:200; BD Biosciences), β-actin (1:200; Santa Cruz Biotechnology Inc.), Bax (1:200; BD Biosciences), and actin (1:2,000; Santa Cruz Biotechnology Inc.).

Immunohistochemical assessment of brain tissue. Animals were anesthetized with chloral hydrate (0.4 g/kg, i.p.) and their brains fixed by transcardial perfusion with saline, followed by perfusion with and immersion in 4% paraformaldehyde as previously described (51). FISH analysis. To demonstrate the xenografted human cells in the rats’ brain, FISH analysis was performed as previously described (63). In brief, cryostat sections 6 μm in thickness were fixed 3 times, for 10 minutes each, in Carnoy fixative. Tissue samples were digested with protease K (10 μg/ml; Sigma-Aldrich) for 5 minutes at 37°C, washed with water, and then rinsed in 2x SSC for 3 minutes and air dried again. Sections were denatured at 85°C for 2 minutes in preheated 70% formamide and 2x SSC buffer, pH 7.0, and then were quenched with ice-cold 70% ethanol for 1.5 minutes. The human X chromosome probe labeled with Cy3 (Star**FISH; Cambio) was heated to 72°C for 5 minutes, placed in a 37°C water bath for 3 hours to preanneal, and then applied to the sections at 45°C. Coverslips were placed on the sections for incubation overnight in a hydrated slide box at 42°C. The next day, the coverslips were removed in preheated 2x SSC buffer, pH 7.0, at 45°C. The sections were washed twice in preheated 50% formamide in 2x SSC buffer for 5 minutes each at 45°C and were then gently washed twice in preheated 0.1x SSC buffer for 5 minutes each at 45°C. Multicolored fluorescence staining of tissues was analyzed by a laser scanning confocal microscope (LSM510; Zeiss). Counting of FISH signal-positive nuclei was accomplished by systematically examining the FISH-stained tissue, field by field, under x630 magnification.

 Laser scanning confocal microscopy for immunofluorescence colocalization analysis. In order to identify whether cell-type-specific markers colocalized with endogenous stem cells in the ischemic rat brain, we labeled endogenous stem cells with BrdU, and exogenous cells (transplanted hOECs/ONFs) were labeled with bis-benzimide (blue fluorescence). Immunofluorescence colocalization analysis was performed with laser scanning confocal microscope. In immunofluorescence colocalization of hOEC/ONF cells and cell-specific marker studies, each coronal section was treated with cell-specific antibodies: WVF (for endothelial cells; 1:200; Dako), GFAP (for astrocytes; 1:300), Neu-N (for neuronal nuclei; 1:50), nestin (for neuronal dendrites; 1:50), and MAP-2 (for neuronal dendrites; 1:300), all from Millipore; SDF-1t (1:200) and CXCR4 (1:200) from Torrey Pines Biolabs; and PrPα (1:300; M20; Santa Cruz Biotechnology Inc.). The secondary antibodies in the subsequent visualization reaction were Alexa Fluor-conjugated donkey anti-mouse 555, anti-rabbit 555, or anti-rat 555 (1:500; Molecular Probes; Invitrogen) (51).

Assessment of neurite regeneration and Western blotting of PrPα and CXCR4 expression in vivo. Brain tissue samples were immunostained to measure neurite outgrowth. Measurement of neurite regeneration was performed as described previously (69). Briefly, brain tissue samples from each experimental rat were fixed and immunostained with specific antibody against β-tubulin (1:400; Sigma-Aldrich). For quantification analysis, neurons with processes greater than twice the cell body diameter were counted as neurite-bearing cells. The length of the longest neurite of each neuron was measured from digitized images and quantified using imaging analysis software (SigmaScan 4.01.003). Analysis of the expression of PrPα and CXCR4 was performed with specific antibody of PrPα (1:300; M20; Santa Cruz Biotechnology Inc.) and CXCR4 (1:300; Millipore) in a Western blot as described above. PrPα and CXCR4 activation was inhibited with PrPα-specific blocking antibody (10 μg/ml; 6H4; Prionics), CXCR4 neutralizing antibody (R&D Systems), and control human IgG (Sigma-Aldrich). The blocking protocol to inhibit PrPα activation involved pretreatment of the hOECs/ONFs (2 x 10^5 cells) with anti-PrPα blocking antibody for 24 hours as described previously with modification (84). In addition, the CXCR4 was neutralized by i.p. injection of CXCR4 neutralizing antibody (1 mg/rat) twice weekly for 2 weeks as described previously (85). Expression of PrPα and CXCR4, assay of neurite outgrowth, and neurological behavioral measurement (described above) were used to evaluate the outcome of the 4 treatment protocols (hOECs/ONFs; hOECs/ONFs with PrPα-blocking antibody; hOECs/ONFs with CXCR4-blocking antibody; and hOECs/ONFs with control human IgG).

Generation of PrPα-knockout mice. The PrPα−/− mice used in this study were wild-type C57BL/6 mice. PrPα−/− mice were a kind gift from...
Charles Weissmann, Institute of Neurology, London, United Kingdom, as previously described (86). Neurite regeneration after stroke was evaluated in the PrP+/+ and PrP−/− mice after hOEC/ONF (1 x 10⁵ cells) implantation as mentioned above.

**Statistics.** All observers in this study were blinded to the actual conditions of the experiment to minimize observer bias. Results are expressed as mean ± SEM. The behavioral scores were evaluated for normality, and for normally distributed data, 2-tailed Student’s t-tests were used to evaluate mean differences between the control and the treated groups. Data lacking normal distribution were analyzed by 1-way ANOVA. A value of P < 0.05 was taken as significant.

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