Neuroprotection by PlGF gene-modified human mesenchymal stem cells after cerebral ischaemia

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Intravenous delivery of mesenchymal stem cells (MSCs) prepared from adult bone marrow reduces infarction size and ameliorates functional deficits in rat cerebral ischaemia models. Placental growth factor (PlGF) is angiogenic to impaired non-neural tissue. To test the hypothesis that PlGF contributes to the therapeutic benefits of MSC delivery in cerebral ischaemia, we compared the efficacy of systemic delivery of human MSCs (hMSCs) and hMSCs transfected with a fibre-mutant F/RGD adenovirus vector with a PlGF gene (PlGF-hMSCs). A permanent middle cerebral artery occlusion (MCAO) was induced by intraluminal vascular occlusion with a microfilament. hMSCs and PlGF-hMSCs were intravenously injected into the rats 3 h after MCAO. Lesion size was assessed at 3 and 6 h, and 1, 3, 4 and 7 days using MR imaging and histology. Functional outcome was assessed using the limb placement test and the treadmill stress test. Both hMSCs and PlGF-hMSCs reduced lesion volume, induced angiogenesis and elicited functional improvement compared with the control sham group, but the effect was greater in the PlGF-hMSC group. Enzyme-linked immunosorbent assay of the infarcted hemisphere revealed an increase in PlGF in both hMSC groups, but a greater increase in the PlGF-hMSC group. These data support the hypothesis that PlGF contributes to neuroprotection and angiogenesis in cerebral ischaemia, and cellular delivery of PlGF to the brain can be achieved by intravenous delivery of hMSCs.

Keywords: angiogenesis; bone marrow transplantation; neural transplantation; regeneration; stroke

Abbreviations: BDNF = brain-derived neurotrophic factor; DWIs = diffusion-weighted images; ELISA = enzyme-linked immunosorbent assay; hMSCs = human MSCs; MCAO = middle cerebral artery occlusion; MOI = multiplicity of infection; MSCs = mesenchymal stem cells; PBS = phosphate-buffered saline; PlGF = placental growth factor; TTC = 2,3,5-triphenyltetrazolium chloride; TUNEL = terminal deoxynucleotidyltransferase dUTP nick-end labelling; VEGF = vascular endothelial growth factor

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Introduction

Transplantation of human mesenchymal stem cells (hMSCs) several hours after ischaemia onset can reduce infarction size and improve functional outcome in rodent cerebral ischaemia models (Chen et al., 2001a; Li et al., 2002; Iihoshi et al., 2004, 2005; Nomura et al., 2005; Honma et al., 2006). Under appropriate conditions, mesenchymal stem cells (MSCs) can differentiate into cells of neuronal and glial lineage (Prockop et al., 1997; Woodbury et al., 2000; Kobune et al., 2003, 2005; Nomura et al., 2005; Honma et al., 2006), but the beneficial effects of MSCs are not thought to result from neurogenesis. Rather, neuroprotective and angiogenic effects may be primarily responsible (Chen et al., 2001a).

There is low-level basal secretion from hMSCs of several neurotrophic factors in culture, but ischaemic rat brain extracts induce production of neurotrophins and angiogenic growth factors (Chen et al., 2002). One of these, brain-derived neurotrophic factor (BDNF), constitutively expressed at low level in primary hMSC cultures (Kurozumi et al., 2004; Nomura et al., 2005), increased in the ischaemic lesion following hMSC treatment in the rat middle cerebral artery occlusion (MCAO) model (Kurozumi et al., 2004; Nomura et al., 2005). Transplantation of BDNF gene-modified hMSCs resulted in stronger therapeutic effects with increased BDNF levels in the ischaemic lesion (Kurozumi et al., 2004;
Nomura et al., 2005). Thus, BDNF may play an important role in the therapeutic benefits on cerebral ischaemia following hMSC transplantation. The capacity of MSCs to release growth and trophic factors has been suggested to be the key to the beneficial effect in cerebral ischaemia (Chen et al., 2002).

In addition to neurotrophic factors that may confer neuroprotective effects, isolated cultured bone marrow-derived stromal cells secrete angiogenic cytokines, vascular endothelial growth factor (VEGF) and placental growth factor (PIGF) (Kinnaird et al., 2004). These factors promoted the growth of new and stable vessels in cardiac and limb ischaemia (Luttun et al., 2002b; Autiero et al., 2003). While the administration of VEGF is reported to consist of both neuroprotective and angiogenic effects in the ischaemic brain (Sun et al., 2003; Manoonkitiwongsa et al., 2004), the role of the VEGF family member, PIGF (Malione et al., 1991), on stroke therapy remains unclear. In this study, hMSCs and PIGF hypersecreting hMSCs (PIGF-hMSCs) were intravenously delivered at 3 h after induction of unilateral permanent cerebral ischaemia to investigate if cellular delivery of PIGF by hMSCs could influence structural integrity and functional outcome. We demonstrate that both cell types induce elevated PIGF levels in ischaemic brain and a concomitant reduction in lesion volume, increased angiogenesis and functional improvement. However, greater effects were observed with the PIGF-hMSC infusions. These results suggest that PIGF may be an important molecule in the therapeutic effects of hMSC treatment in cerebral ischaemia.

Material and methods

Preparation of human mesenchymal stem cells

hMSCs were processed for cell culture as described previously (Nomura et al., 2005; Honma et al., 2006). Briefly, human bone marrow from healthy adult volunteers was obtained by aspiration from the posterior iliac crest after informed consent was obtained; the subject’s consent was obtained according to the Declaration of Helsinki, and this study was approved by the Institutional Review Board at our university. Bone marrow mononuclear cells were isolated and were plated in 150 cm² plastic tissue culture flasks and incubated overnight. After washing away the free cells, the adherent cells were cultured in Mesenchymal Stem Cell Basal Medium (MSCBM, Cambrex, Walkersville, Maryland, USA) containing Mesenchymal Cell Growth Suplement (MCGS, Cambrex), 4 mM L-glutamine, in a humidified atmosphere of 5% CO₂ at 37°C. After reaching confluency, they were harvested and cryo-preserved as primary MSCs or used for gene transduction.

Adenoviral vectors

Adenoviral vectors carrying a human PIGF cDNA were constructed on the basis of the method of Kurozumi et al. (2004) with minor modifications. Briefly, human PIGF cDNA was cloned using the reverse-transcription polymerase chain reaction (RT–PCR) method using the total RNA extracted from primary MSC as the template. The identity of PIGF cDNA obtained in this manner was confirmed by sequencing and comparing it with the GenBank sequence NM_002632. The human PIGF primer sequence was forward, 5’-CGGAAATTCCACCATGCGTGGCATGCGTCCCT-3’, and reverse, 5’-CCAGATCTTACCTCCGCGGGGAAACATGC-3’.

The PIGF cDNA was inserted between the EcoRI site and the Bgl II site in the pCACC vector and the resulting plasmid was designated pCACCPIGF. The plasmid pCACCPIGF was digested with Cla I, and the fragment containing the PIGF cDNA expression unit was isolated by agarose gel electrophoresis. The adenoviral PIGF expression vector, pWEAxCAhPIGF-F/FGD, was prepared using LipofectAMINE 2000 (Invitrogen, Tokyo, Japan).

Before being used, the above viral vectors were evaluated for their viral concentration and titre, and viral stocks were examined for potential contamination with replication-competent viruses. To determine viral concentration [particle unit (pu)/ml], the viral solution was incubated in 0.1% sodium dodecyl sulphate and A₅₅₀ was measured. The viral titres of AxCAhPIGF-F/FGD were 1.0 × 10¹⁰ pu/ml.

Adenovirus infection

Adenovirus-mediated gene transfection was performed as described previously (Kurozumi et al., 2004, 2005). Briefly, the cells were seeded at a density of 2 × 10⁶ cells per 15 cm plate. hMSCs were exposed to the infectious viral particles in 7.5 ml DMEM at 37°C medium for 60 min; cells were infected with AxCAhPIGF-F/FGD or AxCAhPIGF-F/FGD at a multiplicity of infection (MOI) of 3.0 × 10³ pu/cell. The medium was then removed, and the cells washed once with DMEM and then re-cultured with normal medium for 24 h, after which transplantation was performed.

Phenotypic characterization of the hMSC and hMSC-PIGF cells

Flow cytometric analysis of primary hMSCs and PIGF-hMSCs was performed as described previously (Honma et al., 2006). Briefly, cell suspensions were washed twice with phosphate-buffered saline (PBS) containing 0.1% bovine serum albumin (BSA). For direct assays, aliquots of cells at a concentration of 1 × 10⁶ cells per millilitre were immunolabelled at 4°C for 30 min with the following antihuman antibodies: fluorescein isothiocyanate (FITC) conjugated CD45, CD34 (Immunotech, Marseille, France), and CD105 (SH-2) (AnCell, Bayport, MN, USA). As an isotype-matched control, mouse immunoglobulin G1–FITC (IgG1–FITC) (Immunotech, Marseille, France) was used. For indirect assays, cells were immunolabelled with antihuman CD73 (SH-3) (Alexis Biochemicals, San Diego, CA, USA). As the secondary antibody, goat anti-mouse IgG [heavy plus light (H plus L)]–FITC (Immunotech) was used. Labelled cells were analysed by a FACScalibur flow cytometer (Becton Dickinson, San Jose, CA, USA) with the use of CellQuest software. Dead cells were gated out with forward- versus side-scatter window and propidium iodide staining.

Cerebral ischaemic model

The rat MCAO model was used as a stroke model. We induced permanent MCAO by using a previously described method of intraluminal vascular occlusion (Longa et al., 1989). Adult male Sprague–Dawley rats (n = 151) weighing 230–250 g were initially anaesthetized with 5% isoflurane and maintained under anaesthesia with 1.5% isoflurane in a mixture of 70% N₂O and 30% O₂ with mechanical ventilation. Rectal temperature was maintained at 37°C with an infrared heat lamp. The left femoral artery was cannulated for measuring blood pH, pO₂, pCO₂ and blood pressure throughout the surgery. A length of 20.0–22.0 mm 4–0 surgical
dermalon suture with the tip rounded by heating near a flame was advanced from the external carotid artery into the lumen of the internal carotid artery until it blocked the origin of the MCA.

Transplantation procedures
Experiments consisted of three groups: Group 1, medium (MSCBM) alone (without donor cell administration) \( (n = 51) \), Group 2, hMSCs \( (1.0 \times 10^5) \) \( (n = 46) \), and Group 3, PlGF-hMSCs \( (1.0 \times 10^5) \) \( (n = 45) \). All injections were done 3 h after MCAO and cells were suspended in 1 ml of medium.

In some experiments, AxCALacZ-F/RGD adenovirus was used to transduce the LacZ gene into the hMSC \( (n = 8) \) and PlGF-hMSC \( (n = 16) \). Details of the construction procedures are described elsewhere (Niwa et al., 1991; Nakamura et al., 1994; Nakagawa et al., 1998; Takiguchi et al., 2000). For "in vitro" adenoviral infection, 1.0 \( \times 10^5 \) MSCs were placed with AxCALacZ-F/RGD at an MOI of 3.0 \( \times 10^5 \) pu/cell for 1 h and incubated at 37°C in a medium containing 10% foetal calf serum.

MRI studies and measurement of infarct volume
Rats were anaesthetized with ketamine (50 mg/kg) and xyrazine (6 mg/kg) intraperitonially. Each rat was placed in an animal holder/MRI probe apparatus and positioned inside the magnet. The animal’s head was held in place inside the imaging coil. All MRI measurements were performed using a 7-T, 18-cm-bore superconducting magnet (Oxford Magnet Technologies, Witney, Oxfordshire, UK) interfaced to a UNITYNOVA console (Oxford Instruments, Witney, Oxfordshire, UK and Varian, Inc., Palo Alto, CA, USA). T2-weighted images \( (T2-WI) \) were obtained from a 1.0-mm-thick coronal section with a 0.5 mm gap using a 30 mm \( \times \) 30 mm field of view, repetition time \( (TR) = 3000 \) ms, echo time \( (TE) = 37 \) ms, b-value = 0, and reconstructed using a 256 \( \times \) 256 image matrix. Diffusion-weighted images \( (DWIs) \) were obtained at the same condition as T2-WI except b-value \( (b \text{-value} = 1000) \). Accurate positioning of the brain was performed to centre the image slice 5 mm posterior to the rhinal fissure with the head of the rat held in a flat skull position. MRI measurements were obtained 3, 6, 24 h and 3, 4 and 7 days after MCAO.

The ischaemic lesion area was calculated from both T2-WI and DWI using imaging software (Scion Image, Version Beta 4.0.2, Scion Corporation, Frederick, Maryland, USA), based on the previously described method (Neumann-Haefelin et al., 2000). For each slice, the higher intensity lesions in both T2-WI and DWI where the signal intensity was 1.25 times higher than the counterpart in the contra-lateral brain lesion were marked as 

Detection of PlGF in vitro and in vivo
Forty-eight hours after hMSCs were transfected in vitro at various MOI \( (\text{pu/cell}) \), culture supernatants were collected for analysis. Furthermore, three and seven days after MCAO, rats were anaesthetized with ketamine and xyrazine intraperitonially, their brains were removed and coronal sections \( (30 \text{ mg}) \) from \(-0.3 \text{ to } 1.3 \text{ mm to bregma in the ischaemic hemisphere were dissected on ice and were stored at } \text{at } -80°C \text{ until use. Subsequently, each tissue sample was suspended in an equal weight of homogenate buffer (1 ml; } 137 \text{ mM NaCl, } 20 \text{ mM Tris, } 1% \text{ NP-40, 1 mM PMSF, } 10 \mu \text{g/ml aprotinin, } 1 \mu \text{g/ml leupeptin, } 0.5 \text{ mM sodium vanadate) and homogenized with a Dounce homogenizer. The homogenate was centrifuged (10000 g) for 10 min at 4°C, and the supernatant (1 } \mu \text{g/ul) was collected for analysis. Commercial PlGF ELISA (enzyme-linked immunosorbent assay) kits (R&D Systems Inc., Minneapolis, USA) were used to quantify the concentration of PlGF in each of the samples.

TTC staining and quantitative analysis of infarct volume
One week after transplantation, the rats were deeply anaesthetized with sodium pentobarbital \( (30 \text{ mg/kg, i.p.). The brains were removed carefully and dissected into coronal } 1 \text{ mm sections using a viva} \\

Detection of donor PlGF-hMSCs in vivo
Two weeks after transplantation, the LacZ-transfected PlGF-hMSCs in vivo were detected using laser scanning confocal microscopy. Brains of the deeply anaesthetized rats were removed, fixed in 4% paraformaldehyde in phosphate buffer, dehydrated with 30% sucrose in 0.1 M PBS for overnight and frozen in powdered dry ice. Coronal cryostat sections \( (10 \mu \text{m}) \) were processed for immunohistochemistry. To identify the transplanted cells, LacZ-labelled PlGF-hMSCs were visualized with the use of antibody to β-galactosidase (rhodamine-labelled polyclonal rabbit anti-β-galactosidase antibody, DAKO, Carpinteria, CA, USA). To identify the cell type derived from the donor PlGF-hMSCs, double-labelling studies were performed with the use of antibodies to neurons \( (\text{FITC-labelled monoclonal mouse NeuN, DAKO}) \), astrocytes \( (\text{FITC-labelled monoclonal mouse anti-glial fibrillary acidic protein (GFAP, Sigma, St Louis, MO, USA)}) \), oligodendrocytes \( (\text{FITC-labelled monoclonal mouse anti-O4, Boehringer-Mannheim, Mannheim, Germany)}) \), and endothelial cells \( (\text{FITC-labelled polyclonal rabbit anti-von Willbrand factor (vWF, DAKO)}) \). To excite the FITC fluorochrome \( (\text{green, a } 488 \text{ nm laser line generated by an argon laser was used, and for the Rhodamine fluorochrome (red, a } 543 \text{ nm laser line from a HeNe laser was used. Confocal images were obtained using a Zeiss laser scanning confocal microscope with the use of Zeiss software.})

TUNEL staining
Seven days after MCAO, rats were anaesthetized and transcardially perfused with PBS followed by 4% paraformaldehyde in PBS. Their brains were harvested and immersed in 4% paraformaldehyde in PBS for 2 days, after which 12 μm frozen sections were cut with a cryostat at \(-20°C. Cellular DNA fragmentation in the ischaemic boundary zone was detected using the TUNEL (terminal deoxynucleotidyltransferase dUTP nick-end labelling) method by means of an \textit{in situ} Apoptosis Detection Kit (TaKaRa Biomedicals, Shiga, Japan). Specifically, after proteinase digestion, the sections were incubated in a mixture containing terminal deoxynucleotidyltransferase and FITC-labelled dUTP. Sections were counterstained with propidium iodide. The number of TUNEL-positive cells was counted in three 651.5 μm \( \times \) 651.5 μm regions of the boundary zone.
Three-dimensional image acquisition for angiogenesis

Seven days after MCAO, rats were anaesthetized and intravenously perfused with FITC dextran (1 ml, Sigma). The brains were immediately removed, and were fixed in 4% paraformaldehyde at 4°C for 48 h. Coronal vibratome sections (100 μm) were analysed with a laser scanning confocal imaging system (LSM Pascal; Carl Zeiss, Jena, Germany). The sections were scanned in 512 × 512 pixel (651.5 μm × 651.5 μm) format in the x–y direction using a 5× frame-scan average (2.18 μm interval and 2.4 μm thickness along the z-axis) under a 20× objective. Vessel volumes were measured in the three dimensions using the software of Zeiss LSM.

Treadmill stress test

Rats were trained 20 min per day for 2 days a week to run on a motorized treadmill at a speed of 20 m/min with a slope of 0°C. Rats were placed on a moving belt facing away from the electrified grid and induced to run in the direction opposite of the movement of the belt. Thus, to avoid foot-shocks (with intensity in 1.0 mA), the rats had to move forward. Only the rats that had learned to avoid the mild electrical shock were included in this study. The maximum speed at which the rats could run on a motor-driven treadmill was recorded.

Limb placement test (LPT)

The rats’ four limbs were evaluated using the top and edges of a counter top (Ohlsson et al., 1995). Each test was scored as follows: 0, no placing; 1, incomplete and/or delayed (>2 s) placing; and 2, immediate and correct placing. For each body side, the maximum score from the tests used was 16. The forepaws were graded in all six tests; in tests 4 and 6, the hind limbs were also tested. During tests 1 through 4, the rat was held in a soft grip by the examiner. In test 1, limb placing was tested by slowly lowering the rat toward a table edge, the head of the rat was moved 45° upward while the chin was supported to prevent the nose and the vibrissae from touching the table. A normal rat places both forepaws on the table top. A rat with focal brain lesion may lose contact with the table with the paw contra-lateral to the injured hemisphere. In test 3, fore limb placement of the rat when facing a table edge was observed. A normal rat places both forepaws on the table top. Test 4 recorded fore limb and hind limb placement when the lateral side of the rat’s body was moved toward the table edge. For test 5, the rat was placed on the table and gently pushed from behind toward the table edge. A normal rat will grip on the edge, but an injured rat may drop the fore limb contra-lateral to the injured hemisphere. Test 6 was the same as test 5, but the rat was pushed laterally toward the table edge.

Statistical analysis

Data are presented as mean values ± standard deviation, and were statistically analysed. Differences among groups were assessed by ANOVA (analysis of variance) with Scheffe’s post hoc test to identify individual group differences. Differences were deemed statistically significant at P < 0.05.

Results

Characteristics of primary and PlGF gene-transduced hMSCs

Primary hMSCs were cultured as plastic adherent cells to subconfluency ~1 week (see Material and methods). A characteristic feature of hMSCs is a CD34-, CD45-, SH2*(CD105), SH3*(CD73) cell surface phenotype (Kobune et al., 2003). Both flattened and spindle-shaped cells can be recognized in the culture. PlGF-hMSCs showed similar flattened and spindle-shaped morphology. Flow cytometric analysis of the PlGF-hMSCs (Fig. 1B) was essentially identical to primary hMSCs (Fig. 1A).

Detection of immunoreactive human PlGF and quantitative analysis in vitro

Levels of PlGF in the supernatant of cultured hMSCs and PlGF-hMSCs with different MOI levels were studied. hMSC transfected with AxCAlPlGF/F-RGD (PlGF-hMSC) at an MOI of 300, 1000 and 3000 pu/cell secreted PlGF at a rate of 0.72 ± 0.33, 2.19 ± 0.36 and 7.57 ± 1.14 ng/10⁵ cells/48 h, respectively (n = 6). Non-transfected hMSC also produced PlGF protein (0.025 ± 0.002 ng/10⁵ cells/48 h) (n = 6). AxCALacZ/F-RGD at an MOI of 3000 pu/cell secreted PlGF at a rate of 0.02 ± 0.004 ng/10⁵ cells/48 h (n = 6). These data are summarized in Fig. 1C. The level of PlGF production from PlGF-hMSC transfected at an MOI of 3000 pu/cell was ~300-fold greater than that seen in non-infected or LacZ-infected hMSC.

Characterization of ischaemic lesion size by magnetic resonance image analysis

An estimate of lesion size was obtained using in vivo MRI (see Material and methods). The cells were intravenously delivered immediately after the 3 h MRI. DWI and T₂-WI are shown in Fig. 2A and B, respectively. These coronal forebrain sections were obtained at the level of caudato-putamen complex. Images obtained at 3 h (column 1), 3 days (column 2) and 7 days (column 3) are shown for control (medium) (n = 10), hMSC (n = 9) and PlGF-hMSC-injected groups (n = 8) (Fig. 2). The reduction in density in lesions on the right side of the brains that were subjected to ischaemic injury may be noted. Lesion volume (mm³) was determined by analysis of high-intensity areas on serial images collected through the cerebrum (see Material and methods).

As calculated from the DWI the lesion volume at 3 h after MCAO was similar in the control, hMSC and the PlGF-hMSC groups (Fig. 3A). In order to determine the efficacy of hMSC or PlGF-hMSC transplantation in modulating the ischaemic lesion volume, 1 × 10⁷ cells were delivered intravenously at 3 h after MCAO, and animals were re-imaged at 6 and 24 h, 3, 4 and 7 days later. In the sham control group, MRI-estimated lesion volume reached maximum at 3 days, and gradually decreased at Day 4, and at 7 days post-MCAO. Although lesion volumes estimated 3 h after MCAO were the same among the three groups, lesion volume was smaller 6 h after MCAO in the PlGF-hMSC-treated group (Fig. 3A). However, at 24 h, 3, 4 and 7 days, lesion volume was less in both hMSC and PlGF-hMSC groups as compared with control. The reduction in lesion volume was greater for the PlGF-hMSC group as compared with the hMSC group.
at 3, 4 and 7 days post-MCAO. These results are summarized in Fig. 3A. Lesion volume estimated from T2-WI showed similar results as with the DWI (Fig. 2B and 3B).

**Histological determination of infarction volume**

A second independent measure of infarction volume was performed. The brains were stained with TTC 7 days after MCAO (Fig. 4A). Normal brain (grey matter) tissue typically stains with TTC, but infarcted lesions show no or reduced staining (Bederson et al., 1986). TTC staining obtained 7 days after MCAO without cell transplantation is shown in Fig. 4A-1. The reduced staining on the lesion side may be noted. Lesion volume was calculated by measuring the area of reduced TTC staining in the forebrain (n = 5) (see Material and methods). As with MRI analysis there was a reduction in infarction volume with both hMSC (n = 5) (Fig. 4A-2) and PlGF-hMSC (n = 5) (Fig. 4A-3) treatment, but the effect of PlGF-hMSC was larger. Intravenous delivery of $1 \times 10^7$ PlGF-hMSCs resulted in very substantial reduction in lesion volume as estimated from TTC staining (Fig. 3C). We have shown in a previous paper (Kurozumi et al., 2005) that hMSCs transfected with AxAhEGFP-F/RGD directly injected into the brain of MCAO rats did not show increased neuroprotection as compared with cells transfected to express various cytokines.

**Identification and characterization of donor cells in vivo**

LacZ-transfected PlGF-hMSCs that had been intravenously administered ($10^7$ cells) 3 h after MCAO were identified in vivo (n = 16). The LacZ-expressing PlGF-hMSCs were found primarily in the lesion penumbra. Immunohistochemical studies were carried out to identify LacZ-positive cells in and around the lesion zone in animals transplanted with LacZ-transfected PlGF-hMSCs (Fig. 4B and C). The numbers of LacZ-positive cells were obtained from one section per rat, and four separate rats (n = 4) were analysed, demonstrating a large number of LacZ-positive cells in and around the lesion (74.00 ± 23.10 cells/mm², n = 4), although virtually no LacZ-positive cells were observed in the non-damaged hemisphere and non-infected group. A small number of the LacZ-positive cells expressed the NeuN (10.53 ± 3.43%, n = 4) (Fig. 4B), or GFAP (17.23 ± 4.00%, n = 4) (Fig. 4C), and very few of LacZ-positive cell population were O4-positive (n = 4). Moreover, only a small fraction of cells estimated from LacZ cell counts can reach the lesion in the brain, which is similar to previous studies (Chen et al.,
2001b; Li et al., 2001; Inoue et al., 2003). The density of LacZ-positive cells estimated from the immunohistochemical analysis was ~500 cells/mm³ in the ischaemic core and ~7400 cells/mm³ in the boundary zone. Infarcted volume ranged from 150 to 250 mm³. The number of intravenously injected cells was 10⁷ cells. Thus, the minimum percentage of intravenously injected cells accumulating in the ischaemic core was ~0.75%, and the maximum number in the boundary zone was ~18.5%.

**PIGF levels in vivo**

PIGF levels in focal brain tissue were measured by a sandwich ELISA 3 and 7 days after MCAO (see Material and methods). PIGF levels significantly increased in the ischaemic hemisphere of the hMSC-treated group (n = 10) compared with control group (n = 10) at 3 and 7 days after MCAO (Fig. 3D). In the PIGF-hMSC-treated rats (n = 10), PIGF increased in the ischaemic hemisphere to a greater extent than the hMSC-treated rats. It may be noted that the PIGF levels also significantly increased in the medium-treated group compared with the normal brain (n = 10). In the PIGF-hMSC groups, PIGF levels were similar at 47.7 ± 10.4 pg/mg on Day 3 versus 40.8 ± 3.5 pg/mg (n = 5, P = 0.22). Thus, PIGF secretion in the PIGF-hMSC group was above control and the hMSC group for at least a week after cell infusion. These results are summarized in Fig. 3D.

**Reduced nuclear DNA fragmentation in treated animals**

Nuclear DNA fragmentation was studied in the ischaemic boundary zone. There were significantly fewer TUNEL positive cells in animals treated with hMSC in the ischaemic...
boundary zone 7 days after MCAO (24.25 ± 5.78 cells/mm², n = 4; P < 0.01) (Fig. 5A-3) compared with those treated with medium (123.00 ± 33.25 cells/mm², n = 4) (Fig. 5A-2). The reduced apoptotic cell death is more obvious in the PlGF-hMSC-treated group (11.50 ± 4.44 cells/mm², n = 4; P < 0.05) compared with those in the hMSC-treated group (Fig. 5A-4). Figure 5A-1 indicated that there is no apoptotic cell in the normal rat brain (n = 4). These results were summarized in Fig. 5B.

**Angiogenesis introduced by the transplantation**

To examine whether the administration of hMSCs and PlGF-hMSCs induces angiogenesis, three-dimensional analysis of capillary vessels in the lesion was performed using Zeiss LSM5 PASCAL software. Figure 6A-1 shows the three-dimensional capillary image in the normal rat brain (n = 4). The capillary vascular volume in the boundary region 7 days after MCAO was increased in the hMSC-treated group (n = 4) (Fig. 6A-3) compared with the medium-treated group (n = 4) (Fig. 6A-2), and the angiogenesis was greater in the PlGF-hMSC-treated group (n = 4) (Fig. 6A-4). The capillary vascular volume was expressed as a ratio by dividing that obtained from the ischaemic hemisphere by that of the contra-lateral control hemisphere. The ratio was significantly higher in both the hMSC (0.48 ± 0.13; P < 0.05) and the PlGF-hMSC (1.37 ± 0.33; P < 0.001) treated groups as compared with the medium-treated group (0.25 ± 0.02). These results were summarized in Fig. 6B.

Immunohistochemical studies were carried out to show endothelial cell differentiation of LacZ-positive cells in the lesion zone in animals transplanted with LacZ-transfected PlGF-hMSCs (Fig. 7). The vWF-positive cells in the boundary region 7 days after MCAO were increased in the hMSC-treated group (n = 4) compared with the medium-treated
group \((n = 4)\), which was the greatest in the PlGF-hMSC-treated group \((n = 4)\) (Fig. 7C). Double-positive cells of vWF and LacZ were also increased in the PlGF-hMSC-treated group compared with the hMSC-treated group (Fig. 7D).

**Functional analysis**

To assess behavioural performance in the lesioned and transplanted animals, the LPT and the treadmill stress test were performed before ischaemia onset, 3 and 24 h, 3, 4 and 7 days after MCAO. Before MCAO, LPT neurological scores were similar among all animals. Three hours after MCAO, but before transplantation, there was equal deficit in limb placement score among control \((n = 7)\), hMSC-treated \((n = 7)\) and PlGF-hMSC-treated group \((n = 7)\) (Fig. 8A). LPT showed the lowest score in the sham-injected animals 24 h after MCAO with gradual improvement up to 7 days. In the hMSC transplantation group, the improvement in score was greater over the time course up to 7 days. PlGF-hMSC-treated group achieved a higher score at 3, 4 and 7 days as compared with hMSC-treated group. These results are summarized in Fig. 6A.

An independent behaviour study (treadmill stress test) was carried out. Before MCAO, there was no significant difference among control \((n = 7)\), hMSC-treated \((n = 7)\) and PlGF-hMSC-treated groups \((n = 7)\) (Fig. 8B), and at 3 h post-MCAO there was comparable impairment among three groups. Twenty-four hours after MCAO maximum velocity on the treadmill test was at its maximum deficit. An increase in maximum velocity was observed for the PlGF-hMSC group at 24 h. Both hMSC and PlGF-hMSC groups had greater maximum velocity from 3 to 7 days than sham control, but the PlGF-hMSC group attained a higher velocity than the hMSC group. These results are summarized in Fig. 8B.

**Discussion**

The present study demonstrates that intravenous infusion of either hMSCs or PlGF-hMSCs 3 h after permanent MCAO in the rat results in an increase in PlGF levels in the infarcted cerebral hemisphere, reduction in infarction volume and apoptosis, induced angiogenesis and improvement in behavioural performance. These results are consistent with previous studies showing beneficial effects of bone marrow cell transplantation in experimental cerebral ischaemic models (Chen et al., 2001b; Iihoshi et al., 2004; Nomura et al., 2005; Honma et al., 2006). While both hMSCs and PlGF-hMSC groups showed efficacy, the effects, including brain PlGF levels, were greater in the PlGF-hMSC group.

The PlGF gene was introduced into hMSC, and an elevated secretion of PlGF protein as compared with non-transgenic hMSC was confirmed in vitro using ELISA. This relatively high secretory rate was achieved by using Ad-F/RGD, which has a higher transfection rate (Nakamura et al., 2002; Tsuda et al., 2003). PlGF levels in the ischaemic brain lesion assayed by ELISA increased after either hMSC or PlGF-hMSC delivery, but levels were significantly higher in the PlGF-hMSC group. This suggests that the PlGF-hMSC could maintain high levels of PlGF in vivo during the critical post-ischaemic period and that this elevated PlGF secretion contributes to the enhanced neuroprotection.

Attenuation of PlGF production by viral dilution could occur through cell division in vivo. We estimate that several copies of adenovirus were transfected into a single hMSC. Doubling time of hMSC is \(\approx 5\) days. Therefore, in the present.
study, hMSC may have divided one or two times in vivo after transplantation. Only one copy of virus in an hMSC can produce significant levels of PlGF. Therefore, PlGF production is not severely weakened for a week in our experimental paradigm, which is supported by the in vivo data in Fig. 3D. Also, the adenovirus vector used in the present study is composed of a ubiquitously strong promoter based on the β-actin promoter (Niwa et al., 1991), which allowed strong expression in a variety of cell types. Neuroprotective and angiogenic effects that we expect from insertion of the PlGF gene require expression for perhaps several days. Prolonged expression of PlGF may cause unexpected side-effects. Therefore, transient expression may be sufficient and appropriate. Thus, even if some cell division occurs in vivo, it should not impede PlGF expression in the early critical period.

VEGF is an angiogenic and vascular permeability factor that exerts an acute neuroprotective effect in the ischaemic brain, and these effects operate independently (Sun et al., 2003; Manoonkitiwonga et al., 2004). Although PlGF is a member of the VEGF family of growth factors (Maglione

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**Fig. 5** Cells with DNA fragmentation were present in the ischaemic penumbra 7 days after MCAO. Fewer TUNEL-positive cells were detected in rats treated with PlGF-hMSCs (A-4) than those in hMSC-treated animals (A-3) and medium-treated animals (A-2) in the ischaemic boundary zone (yellow, TUNEL-positive; red, nuclear stain). Normal brain showed no TUNEL-positive cell (A-1). These results are summarized in B. Scale bar = 100 μm (A).

**Fig. 6** Seven days after MCAO, the angiogenesis in boundary zone was analysed using a three-dimensional analysis system. A-1 shows the three-dimensional capillary image with systemically perfused FITC-dextran in the normal rat brain. The total volume of the microvessels in the sampled lesion site decreased 7 days after MCAO (A-2), but was greater in the hMSC-treated group (A-3), and greater still in the PlGF-hMSC-treated group (A-4). These results are summarized in B. Scale bar = 150 μm (A).
et al., 1991), less attention has been devoted to the therapeutic potential of PlGF for angiogenic disorders (De Falco et al., 2002). However, recent work is focusing attention on PlGF (Luttun et al., 2002a). PlGF gene transfer into ischaemic myocardium and limb caused pre-existing vessels to enlarge, resulting in tortuous, thin-walled, pericyte-poor ‘mother’ vessels (Luttun et al., 2002a). The latter vessels remained enlarged and subsequently stabilized into mature, durable vessels by acquisition of a pericyte coat, occasionally, by deposition of a thin rim of perivascular collagen. These vessels remained functional and persisted for >1 year after PlGF gene transfer (Carmeliet et al., 2001). PlGF avoided

Fig. 7 Intravenously administrated PlGF-hMSCs differentiated into the endothelial lineage. Confocal images show the transplanted cells (LacZ in red: A-1, B-1) and endothelial cells (vWF in green: A-2, B-2). A-3 and B-3 confirm the co-labelling of LacZ/ vWF in the cells. B-4 also demonstrates z-axis projections of the image. The cell density of vWF-positive cells is shown in C. D demonstrates the cell density of vWF-positive cells co-localized with LacZ. Scale bar = 50 µm (A and B).

Fig. 8 Assessment of brain ischaemia-induced functional deficits using an LPT (A) and the treadmill stress test (B). Three hours after MCAO, but before intravenous cell delivery, there was no statistical difference in limb placement score among three groups. Twenty-four hours after MCAO, rats that received PlGF-hMSC achieved significantly higher limb placement scores compared with control rats. From 3 days on both hMSCs and PlGF-hMSCs showed greater functional improvement than medium alone, but the PlGF-hMSC group showed more improvement than the hMSC group. (B) The treadmill stress test demonstrates that the maximum speed at which the rats could run on a motor-driven treadmill was faster in the hMSC and the PlGF-hMSC-treated rats than control. Twenty-four hours after MCAO, rats that received hMSC-PlGF showed higher running speed compared with the medium alone, and the hMSC group. From three days on after MCAO, the hMSC and PlGF-hMSC groups attained higher velocities but the PlGF-hMSC group was greater. The highest level of functional improvement was attained at 7 days for both functional tests.
the harmful complications of VEGF such as oedema, fibrin deposition and growth of unstable vascular tangles and glomeruloid bodies in the impaired myocardium (Dvorak et al., 1979; Pettersson et al., 2000; Luttun et al., 2002a).

Although PlGF is reported to have angiogenic effects in the impaired myocardium, limb and skin (Dvorak et al., 1979; Pettersson et al., 2000; Carmeliet et al., 2001; Luttun et al., 2002b), its effects on ischaemic brain had not been studied. However, upregulation of endogenous PlGF in ischaemic brain tissue was reported on both the mRNA and protein level (Beck et al., 2002; Hayashi et al., 2003), suggesting that PlGF may play an important role in endogenous repair or neuroprotection in the ischaemic lesion. In the present study, elevated PlGF was observed in the infarcted brain after hMSC and PlGF-hMSC infusion, but PlGF levels were higher following infusion of the PlGF-hMSCs. PlGF induced angiogenesis and did not increase cerebral oedema as assayed by both in vivo MRI and histological analysis.

The therapeutic effect of PlGF-hMSC transplantation in the permanent rat MCAO model was observed even when applied 3 h after infarction. Thus, cellular delivery of PlGF by hMSCs or hypersecreting PlGF-hMSCs may have a therapeutic effect following cerebral ischaemia through neuroprotective and angiogenic mechanisms. These results do not rule out the potential beneficial effects of other trophic factors such as BDNF (Nomura et al., 2005) that is secreted by hMSCs. BDNF hypersecreting hMSCs induced comparable beneficial effects as PlGF-hMSCs. It will be interesting to determine if there is enhanced recovery by simultaneous administration of both cell types.

Cell-based therapeutic approaches are being considered for a number of neurological diseases. Improved neurological function in experimental autoimmune encephalomyelitis (EAE) has been reported following intravenous infusion of hMSCs (Zhang et al., 2005) and neurosphere-derived multipotent precursors (Pluchino et al., 2003, 2005). Suggested mechanisms include reduction of inflammatory infiltration and demyelination, and elevation of trophic factors that may be neuroprotective or stimulate oligodendroglialis. A cell-based therapy may have the advantage of exerting multiple therapeutic effects at various sites and times within the lesion, as the cells respond to a particular pathological microenvironment. As complex as these mechanisms are likely to be, our results indicate that PlGF contributes to neuroprotection and angiogenesis in cerebral ischaemia, and cellular delivery of PlGF to the brain can be achieved by intravenous delivery of hMSCs.

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