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Stem Cell-Based Therapies for Ischemic Stroke: Preclinical Results and the Potential of Imaging-Assisted Evaluation of Donor Cell Fate and Mechanisms of Brain Regeneration Peer-reviewed author version

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Stem Cell-Based Therapies for Ischemic Stroke: Preclinical Results and the Potential of Imaging-Assisted Evaluation of Donor Cell Fate and Mechanisms of Brain Regeneration

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Abstract: Stroke is the second most common cause of death and is a major cause of permanent disability. Given the current demographic trend of an ageing population and associated increased risk, the prevalence of and socioeconomic burden caused by stroke will continue to rise. Current therapies are unable to sufficiently ameliorate the disease outcome and are not applicable to all patients. Therefore, strategies such as cell-based therapies with mesenchymal stem cell (MSC) or induced pluripotent stem cell (iPSC) pave the way for new treatment options for stroke. These cells showed great preclinical promise despite the fact that the precise mechanism of action and the optimal administration route are unknown. To gain dynamic insights into the underlying repair processes after stem cell engraftment, noninvasive imaging modalities were developed to provide detailed spatial and functional information on the donor cell fate and host microenvironment. This review will focus on MSCs and iPSCs as types of widely used stem cell sources in current (bio)medical research and compare their efficacy and potential to ameliorate the disease outcome in animal stroke models. In addition, novel noninvasive imaging strategies allowing temporospatial in vivo tracking of transplanted cells and coinciding evaluation of neuronal repair following stroke will be discussed. © 2016 Wiley Periodicals, Inc. Med. Res. Rev., 00, No. 0, 1–47, 2016

Key words: ischemic stroke; mesenchymal stem cells; induced pluripotent stem cells; mechanisms of stem cell therapy; noninvasive imaging

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1. INTRODUCTION

Worldwide, stroke is the second single most common cause of death, accounting for 10–15% of deaths each year.^{1,2} Moreover, stroke is an important cause of adult disability as 90% of patients that survive from a stroke are left with a residual deficit.^{3,4} It might therefore be clear that stroke-related public and insurance costs constitute a major burden on healthcare systems worldwide.^{1,2} Combining the expectation that the amount of people over the age of 65 will double by 2030, and that the risk of suffering a stroke doubles for each decade over the age of 55, will even lead to a further increase in patient numbers with permanent disabilities and socioeconomic burden.^{2,4–7}

Despite this increased incidence, current available therapies are unable to sufficiently ameliorate the disease outcome or are even not applicable for subgroups of patients due to many contraindications as will be discussed below. Therefore, new therapeutic strategies are needed for treating and preventing stroke that can be applied to patients with distinct risk profiles and in a broader time frame, as time plays a crucial role in the treatment of acute ischemic stroke. In addition to clinical advances in stroke management, cell-based therapies have emerged as a potential candidate to promote functional recovery in patients suffering from stroke.⁸ Despite the promising results achieved with cell-based therapies in stroke, the host response, the precise mechanisms of action of these therapies, and the fate of the donor cells remain largely unknown.⁹ Therefore, noninvasive imaging modalities have been developed that are able to provide detailed temporospatial and functional information on the donor cell fate, the host microenvironment, and endogenous repair mechanisms,¹⁰ which will be discussed later.

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A. Pathophysiology of Stroke

The pathophysiology of stroke can be defined as a neurologic dysfunction of vascular origin with the sudden or rapid occurrence of symptoms and signs corresponding to the involvement of focal areas in the brain.¹¹ Two different types of stroke can occur: ischemic stroke (80–85%) and hemorrhagic stroke (15–20%). Ischemic stroke is most frequently caused by thromboembolisms while hemorrhagic stroke most often results from vessel wall pathology associated with hypertension and microaneurysms.¹² This review will only focus on ischemic stroke as the main pathology.

34 In ischemic stroke, the blood supply to certain brain areas is compromised due to vascular 35 occlusion thereby causing several changes at the (sub)cellular level and ultimately tissue damage. 36 These cellular and molecular processes start with energy depletion followed by glutamate 37 release leading to glutamate-induced excitotoxicity, ion channel dysfunction, and free radical 38 production. These processes in turn disrupt the cellular membrane, damage mitochondria and 39 DNA, generate an immune response, and trigger necrotic and apoptotic cell death (Fig. 1).¹³ 40 In the ischemic core, these cellular changes are irreversible.¹⁴ However, the tissue surrounding 41 the core, also termed as the ischemic penumbra, is functionally impaired but still viable.¹⁵ 42 This area "at risk" is therefore considered as the main target for therapeutic interventions that 43 are believed to exert a protective effect in intervening with the cellular processes discussed 44 above.^{16,17} Using noninvasive imaging methods, the ischemic penumbra has been divided in 45 additional border zones characterized by different grades of hypoperfusion and varying risk of 46 progressing toward lost infarcted tissue if a proper treatment is not initiated (Fig. 1).^{18,19}

When considering therapies that are aimed to salvage the ischemic penumbra by restoring perfusion, it is also important to take into account that restoring the blood flow in ischemic tissue by thrombolytic treatment can lead to secondary damage by reperfusion injury.^{13, 20} This reperfusion injury is mediated by leukocyte infiltration through local disruption of the blood-brain barrier (BBB) and accompanying matrix metalloprotease (MMP) production in



Figure 1. Areas at risk and pathophysiology of ischemic stroke. (1) Blood flow to focal areas of the brain is diminished by vascular occlusion by, for example, an embolism. The affected ischemic tissue can be divided 2.2 into the ischemic core (C) where tissue damage is irreversible, the salvageable ischemic penumbra (P), and 23 a zone of benign oligemia (B) where blood supply can be obtained by leptomeningeal collaterals. Additional 24 border zones with different grades of hypoperfusion and varying risk of progressing toward unsalvageable tissue 25 if a treatment is not initiated were identified with perfusion-weighted MRI. These areas are the core-penumbra border zone (BZ1) and the penumbra-benign oligemia zone (BZ2). (2) The cellular changes ultimately leading 26 to cell death initiate with ATP depletion due to ischemia, followed by depolarization of the affected neurons 27 that triggers (3) glutamate release. (4) Glutamate-induced excitotoxicity is mediated by an elevated sodium 28 and calcium influx that causes cell swelling, a depolarization wave that will lead to damage in neighboring cells, activation of a cascade of enzymatic reactions ultimately leading to membrane and mitochondrial damage 29 and ROS production, which will additionally damage mitochondria and DNA ultimately leading to cell death. 30 (5) Necrotic/apoptotic neurons secrete inflammatory mediators that activate resting microglia and enhance neutrophil and macrophage infiltration. The effects of activated microglia vary and include migration toward 31 and phagocytosis of damaged neurons and depending on the M1/M2 activation state of activated microglia, 32 proinflammatory and/or anti-inflammatory mediators are released. Image was created using Servier Medical Art. 33

addition to stimulation of reactive oxygen species (ROS) production, thereby damaging the
 reperfused environment.^{13, 14, 20–22} In turn, the reperfused ischemic stroke lesion can transform
 into a petechial hemorrhage that does not influence the prognosis or it can transform into an
 intracerebral hematoma, which is associated with a poor outcome.^{22–24}

Due to the complexity of the molecular processes that are involved in the onset of stroke, but also in ischemic reperfusion injury, multiple strategies are considered for treating stroke. These strategies include both acute and long-term approaches. Acute therapies aim to salvage the ischemic penumbra and limit reperfusion injury, while long-term therapeutic strategies aim to reconstitute the lost tissue from the ischemic core, as will be discussed later.

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B. Limitations and Potential Improvements of Available Therapies for Ischemic Stroke

Current therapies or approaches that have been proven to be effective in reducing the mortality
 rate and improving the functional outcome of acute ischemic stroke include the establish ment of a specialized stroke care unit (SCU),²⁵ thrombolysis with tissue plasminogen activator
 (tPA),^{26,27} aspirin administration,²⁸ and decompressive surgery following ischemic stroke.²⁹
 The most remarkable advance in stroke management that reduced the mortality and disability

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2 rate has been the establishment of a SCU, which is a separate physical space in general med-3 ical wards with specialized and specifically dedicated trained staff.²⁵ Despite these advances 4 in stroke management, they can only be applied in a short therapeutic window. The FDA-5 approved standard treatment for stroke, thrombolysis with tPA, is only applicable to less than 6 10% of stroke patients and ideally needs to be initiated within 3 hr after the onset of ischemia. 7 This low applicability rate of tPA is mainly associated with the higher risk of intracerebral 8 hemorrhage when tPA is administered longer than 3 hr after the onset of stroke symptoms.³⁰ 9 Moreover, patients often do not recognize stroke-associated symptoms and only show up at the hospital with advanced stroke symptoms³¹ or the symptoms are not immediately recognized by the hospital staff that can be counteracted by a SCU.²⁵ Therefore, the presenting stroke 11 12 patient has often exceeded the safe 3-hr time window to be considered for tPA treatment. This 13 time window can be extended to 4.5 hr if low-risk patients are selected together with providing 14 extensive care due to the higher risk of secondary damage and increased mortality due to reper-15 fusion injury.^{32,33} The major criteria to consider these low-risk patients included the absence 16 of cerebral hemorrhage or major infarction and they presented with acute ischemic stroke and 17 the symptoms started 3-4.5 hr before initiation of drug administration (for all inclusion and 18 exclusion criteria, see Table I in Ref. [34]). 19

Other approaches such as decompressive surgery and aspirin administration need to be started within 48 hr after the onset of stroke. Moreover, the benefits of aspirin administration are small while decompressive surgery is only applicable for patients whose stroke-associated infarct region is caused by middle cerebral artery-related pathology, combined with malignant space-occupying brain edema.³⁵

Various interventions are currently under investigation, which include extending the time window for thrombolysis with desmoteplase or alteplase,^{36,37} ultrasound-enhanced thrombolysis,³⁸ the creation of new thrombectomy devices,³⁹ stents, stent retrievers, and protective drugs.¹⁷ Asadi et al. provide an up to date, in depth overview of the different studies using endovascular treatments, which can be used as an important supplementary therapy to current intravenous thrombolysis.⁴⁰

30 However, when extending the therapeutic window for thrombolytic interventions, the is-31 chemic brain still needs to be protected from reperfusion injury. Therefore, reducing reperfusion 32 injury is also a route that is currently being investigated as an additional procedure to be imple-33 mented after thrombolytic therapy.²⁰ These approaches aim at reducing the local production of 34 ROS and BBB-damaging MMPs, or mediating the local immune response that would otherwise lead to secondary damage.⁴¹⁻⁴⁴ One of these approaches is therapeutic hypothermia⁴⁵ that has 35 36 been shown to decrease ischemic and reperfusion injury by influencing local excitotoxicity, 37 neuroinflammation, and ROS production.⁴⁶

While these therapies and novel interventions aim at mitigating the disease outcome, they can only be applied in the first few hours or days after the onset of ischemic stroke.^{32,47} Patients surviving stroke and not treated properly within this narrow time window are therefore often left with permanent disabilities, associated with the focal areas of the brain that are affected.³² In these patients, a therapy that can be applied weeks to months after stroke onset can be beneficial. These therapies aim at restoring the lost neural tissue or stimulating brain plasticity to improve the functional outcome but also muscle strengthening and physical conditioning has been shown to improve the quality of life of patients with permanent disabilities.⁴⁸ Stem cell based therapies have been shown to be a promising approach in achieving such results.⁴⁹

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C. Stem Cell Sources and Mechanisms of Action for Cell-Based Therapies

⁵⁰ When considering stem cells sources for a cell-based therapy in ischemic stroke, ex vivo ex-⁵¹ panded and manipulated neural stem cells (NSCs) or neural precursor cells (NPCs) would be

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NoSteppe. DescriptionInternational augroy $3 \times 10^{\circ}$ international $3 $		and/or treatment	Species	Occlusion	transplan- tation	location of transplantation	Fate of transplanted cells	Outcome	Reference
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No Wear rats 2 hr 3 hr yes, surgey 3 weaks are grifting, donor cells MUMC acclerate annolgie recorry aureany surgey 3 weaks are grifting, donor cells MUMC acclerate annolgie recorry and filteration integration 30 No SHR rats Permanent 1 weak post 5 × 10° in introvict 5 weaks after grifting, donor cells 70 71 70 No SHR rats Permanent 1 weaks post 5 × 10° in introvict 5 weaks after grifting, donor cells 70 71		hUMSCs and hUMSCs cultured in neuronal conditioned medium	Sprague- Dawley rats	90 min	24 hr post- surgery	2.5 × 10 ⁵ intracortical in two sites	36 Days survival, not quantified	Significant improvements in motor function, greater metabolic activity of cortical neurons, and better revascularization in the infarct cortex due to paracrine effects	71
No SHR rats Permanent Tasse poste Tasse poste <thtasse poste<="" th=""> <thtasse pos<="" td=""><td></td><td>No</td><td>Wistar rats</td><td>2 hr</td><td>24 hr post-</td><td>2×10^5 intracortical</td><td>5 Weeks survival, <3% expressing</td><td>hUMSC accelerate neurologic recovery after stroke by promoting angiogenesis</td><td>130</td></thtasse></thtasse>		No	Wistar rats	2 hr	24 hr post-	2×10^5 intracortical	5 Weeks survival, <3% expressing	hUMSC accelerate neurologic recovery after stroke by promoting angiogenesis	130
Noch-induced Sprague- bakiy Ihr 4-6 weeks post- rats 6 × 10 ⁴ intrastitatal bMSCs r-BMSCs show higher survival transport Improvement in beomotor and transport 137 No C57BL/61 1.5.th 1 week post- surgery 5 × 10 ⁶ in the stroke transport 1.45 week post- surgery 5 × 10 ⁶ in the stroke transport 1.47 ws. 7.0 ³ and differentiation of the BMSCs 142 No C57BL/61 1.5.th 1 week post- surgery 5 × 10 ⁶ in the stroke transport 1.47 ws. 7.0 ³ and differentiation of the PMSCs in munoludation by detrasting the presence of The I-H microfila and GFAP+ astroytes 143 Hypoxic C57BL/61 Fermanent 24 hr post- surgery 1.8 × 10 ⁶ intranstal 1.5 hr after admitstroin, door transport 140 Hypoxic C57BL/61 Fermanent 24 hr post- surgery 1.5 hr after admitstroin, door transport 150 1.5 hr after admitstroin, door transport 140 No Wistar rats 90 min 1.0 × 10 ⁶ intranstal 1.5 hr after admitstroin associated and significantly transport 140 No Wistar rats 90 min 1.0 × 10 ⁶ in the stroke 1.5 hr after admitstroke 1.5 hr after admitstroke		°N N	SHR rats	Permanent	l week post- surgery	7.5 × 10 ⁴ in three different cortical sites	neural markets 6 Weeks after grafting, donor cells expressed astrocyte, oligodendroglial, and neuronal markets. Functional integration was unlikely	atter stroke of profinoting augogenesis Improved functional outcome, mediated by paracrine factors that are produced by the surviving donor cells	70
No C57BL/64 1.5 th 1 week post- surgery 5 × 10 ⁶ in the stroke Large percentage hASCs express Cognitive recovery and decrease in infract 142 nice surgery lesion MAP2-Dowpercentage of GFAP expression size immunomodulation by decreasing the presence of Ibn-1 microglia and GFAP expression 142 No C57BL/6J Permanent 24 hr post- surgery 1.8 × 10 ⁴ above the Nigrulon after 1 week, toward Rehmin induces ASC-survisal GFAP expression 138 Hypoxic C57BL/6J Permanent 24 hr post- surgery 1.0 × 10 ⁶ intranasal 1.5 hr after administration, dolor Rehmin induces ASC-survisal differentiation into smooth maske cells 138 HPytoxic C57BL/6J Permanent 1.0 × 10 ⁶ intranasal 1.5 hr after administration, dolor 140 HPytoxic C57BL/6J Permanent 24 hr post- surgery 1.0 × 10 ⁶ intranasal 1.5 hr after administration, dolor 140 HPytoxic C57BL/6J Permanent 24 hr post- surgery 1.0 × 10 ⁶ intranasal 1.5 hr after administration into smooth miske cells 140 HPytoxic C57BL/6J Permanent 24 hr post- surgery 1.0		N otch-induced BMSCs	Sprague- Dawley rats	1 hr	4–6 weeks post- surgery	6×10^4 intrastriatal in three sites	r-BMSCs show higher survival (15% vs. 7%) and differentiation than h-BMSCs	Improvement in locomotor and neurological function. Reduced loss of striatal periinfarct cells.	137
No C57BL/6J Permanent 24 hr post- surgery 1.8 × 10 ⁴ above the surgery Migration after 1 week, toward Ischemia induces ASC-survival, migration 138 Hypoxic C57BL/6J Permanent 24 hr post- surgery 1.8 × 10 ⁴ above the surgery Migration after 4 weeks toward the lesion and microwssels 140 Hypoxic C57BL/6J Permanent 24 hr post- surgery 1.0 × 10 ⁶ intranasal 1.5 hr after administration, donor HP of BMSCs induced a higher expression 140 Protreatment mice cells were observed in the ischemic cortex. No long-term toward the lesion and microwssels 140 No Wistar rats 90 min 1, 6, 24, or augery 1.0 × 10 ⁶ intranasal 1.5 hr after administration, donor HP of BMSCs admit induces ASC-survival. migration 127 No Wistar rats 90 min 1, 6, 24, or augery 1 × 10 ⁶ into the could be detected 7 days non-HP BMSCs 127 No Wistar rats 2 hr post- surgery carotid artery could be detected 7 days non-HP BMSCs 127 No Wistar rats 2 hr post- surgery carotid artery could be		No	C57BL/6J mice	1.5 rh	1 week post- surgery	5×10^6 in the stroke lesion	Large percentage hASCs express MAP2, Low percentage of GFAP expression	Cognitive recovery and decrease in infarct size. Immunomodulation by decreasing the presence of Iba-1+ microglia and GFAP+ astrocytes	142
HypoxicC57BL/61Permanent24 hr post-1.0 × 10 ⁶ intransal1.5 hr after administration, donorHP of BMSCs induced a higher expression140pretreatmentmicesurgerycells were observed in theof migration associated and significantly140(HP)nicesurgerycells were observed in theof migration associated and significantly140(HP)NoWistar rats90 min1, 6, 24, or1 × 10 ⁶ into theq-dot nanocrystal marked BMSCsInjecting BMSCs1127NoWistar rats90 min1, 6, 24, or1 × 10 ⁶ into theq-dot nanocrystal marked BMSCsInjecting BMSCs1127NoWistar rats90 min1, 6, 24, or1 × 10 ⁶ into theq-dot nanocrystal marked BMSCsInjecting BMSCs1127NoWistar rats90 min1, 6, 24, or1 × 10 ⁶ into theq-dot nanocrystal marked BMSCsInjecting BMSCs1127NoWistar rats20 min1, 6, 24, or1 × 10 ⁶ into theq-dot nanocrystal marked BMSCsInjecting BMSCs127NoWistar rats2 hr post-carotid arterycould be detected 7 daysmon-HP BMSCs127NoWistar rats2 hr post-carotid arterypoststrokeeduction, and improvement of127NoWistar rats2 hr post-2 × 10 ⁶ into then/aBMSCs facilitate store indeciding and150NoWistar rats2 hr post-2 × 10 ⁶ into then/aBMSCs facilitate store indeciding and150		No	C57BL/6J mice	Permanent	24 hr post- surgery	1.8×10^4 above the corpus callosum	Migration after 1 week, toward vessels, 5% survival after 4 weeks	Ischemia induces ASC-survival, migration toward the lesion and microvessels, differentiation into smooth muscle cells	138
No Wistar rats 90 min 1, 6, 24, or 1 × 10 ⁶ into the 48 hr post- q-dot nanocrystal marked BMSCs Injecting BMSCs 24 hr after stroke had the most significant effect on graft 127 48 hr post- carotid artery could be detected 7 days Insecting BMSCs 24 hr after stroke had the surgery 127 80 most significant effect on graft surgery could be detected 7 days Insecting BMSCs 24 hr after stroke had the surgery 127 80 most significant effect on graft nost significant effect on graft nost significant effect on graft 127 80 most significant effect on graft nost significant effect on graft nost significant effect on graft 127 80 most significant effect on graft nost significant effect on graft nost significant effect on graft 127 80 most significant effect on graft nost significant effect on graft nost significant effect on graft 150 80 most significant effect n/a n/a BMSCs fabilitate should spouting and remotion should artery 150		Hypoxic pretreatment (HP)	C57BL/6J mice	Permanent	24 hr post- surgery	1.0×10^{6} intranasal	1.5 hr after administration, donor cells were observed in the ischemic cortex. No long-term follow up was performed	HP of BMSCs induced a higher expression of migration associated and significantly reduced infarct size and improved sensorimotor function compared to hon-HP BMSCs.	140
No Wistar rats 2 hr 2 hr post- 2 × 10 ⁶ into the n/a BMSCs facilitate axonal sprouting and 150 surgery carotid artery carotid artery boundary zone and corpus callosum		o	Wistar rats	90 min	1, 6, 24, or 48 hr post- surgery	1 × 10 ⁶ into the carotid artery	q-dot nanocrystal marked BMSCs could be detected 7 days poststroke	Injecting BMSCs 24 hr after stroke had the most significant effect on graft survival/integration, infarct size reduction, and improvement of neurological function. SDF-1 and bFGF were unregulated	127
		No	Wistar rats	2 hr	24 hr post- surgery	2×10^{6} into the carotid artery	n/a	BMSCs facilitate axonal sprouting and remyelination in the cortical ischemic boundary zone and corpus callosum	150

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	Reference	143	144	128	131	147	151	135	Continued
	Outcome	Magnetically labeled IA-delivered BMSCs could be detected with MR1 and high cerebral engraftment rates are associated with impeded cerebral blood flow after inicition	Localization of BMSCs in the brain but Localization of BMSCs in the brain but relocated to other organs 24 hr later. Increased radioactivity counts in the invilanceal stroke hemischere	Time dependent functional recovery and cell distribution around the lesion. Mechanisms of action: neuroprotection, angiogenesis and enhancing reactive astrocytes. downereolation of MMP9	Improvement of neurological deficits, migration of donor cells to lesion; attenuation of astroglial activity, inhibition of apoptosis, and promotion of cellular motification	The beneficial effects of cell transplantation persisted for at least 1 year. Donor cells survived, differentiated toward astrocytes or neurons or colocalized with microglia and endothelial cells. Reduction of	Improved functional outcome; reduction in neuronal cell death. No reduction in lesion size. VEGF and synaptofysin was upregulated and GFAP was downregulated in the treated groups. No difference was observed between the	IV-transplanted human MSCs induced IIV-transplanted human MSCs induced functional improvement, reduced infarct volume, and neuroprotection by providing IGF-1 and inducing neurotrophin expression in host brain	
	Fate of transplanted cells	Magnetically labeled BMSCs could be detected with MRI	Transient localization of engrafted cells in the host brain	Low survival, no expression of neuronal markers. Migration toward the lesion and secretion of BDNF	1.5% of surviving cells expressed NeuN; 1% survival of transplanted cells	Donor cells survive up to 1 year and preferentiate toward astrocytes	No migration/implantation of donor cells was observed	After 7 days no donor cells were detected	
	Cell dose and location of transplantation	1 × 10 ⁶ into internal carotid artery	1.1×10^{6} or 0.5×10^{6} into the external carotid artery	1 × 10 ⁶ into the carotid artery	2 × 10 ⁶ into the carotid artery	2×10^6 into the carotid artery	2 × 10 ⁶ h-ASCs or r-ASCs	3×10^6 into the jugular vein	
	Time of transplan- tation	30 min after reperfu- sion	24 hr post- surgery	24 hr, 4 days, and 7 days post- surgery	3 days post- surgery	24 hr post- surgery	30 min post- surgery	24 hr post- surgery	
	Occlusion	2 hr	90 mim	75 min	00 min	2 hr	Permanent	l hr	
	Species	Wistar rats	Wistar rats	Sprague– Dawley rats	Sprague– Dawley rats	Aged Wistar rats	Sprague Dawley rats	Wistar rats	
Continued	(Pre)diffe- rentiation and/or treatment	°Z	No	° Z	°N	°Z	No Z	Immortalized cells	
Table I.	Stem cell type	r-BMSCs	h-BMSCs	h-BMSCs	Autologous r-ASCs	r-BMSCs	h-ASCs and r-ASCs	h-BMSCs	

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n cell	(Pre)diffe- rentiation and/or		Occlusion	Time of transplan-	Cell dose and location of	لآمداء مراد بسميما معالية موالة	Orizonna	Daframen	
MSCs	No	C57BL/6J	30 min	24 hr post-	1×10^6 BMSCs or	n/a	Mice receiving EVs showed improved	129	
nd -BMSC- :Vs		mice	\mathcal{O}	surgery, Repeated after 3 and 5 days for hBMSC- EVs	E VS from 2 × 10° BMSC in the femoral vein		neurological function and long-term survival associated with improved angiogenesis and neurogenesis, which resembled BMSC responses.		
MSCs	No	Wistar rats	2 hr	30 min after reperfu- sion	1×10^{6} into the femoral vein	Magnetically labeled IV-delivered BMSCs could not be detected	Magnetically labeled IV-delivered BMSCs could not be detected	143	STEN
MSCs	BMSCs, PIGF gene transfected MSCs	Sprague- Dawley rats	Permanent	6 hr post- surgery	1 × 10 ⁷ intranvenously	LacZ-expressing PIGF-hBMSCs were found primarily in the penumbra and express NeuN (±10%) and GFAP (<17.23%)	hBMSCs and PIGF-transfected BMSCs improved angiogenesis, reduced the lesion size, and elicited functional improvement; the effect was more pronounced in PIGF-transduced BMSCs	136	I CELL-B
MSCs	BMSCs, CXCR4 gene transfected BMSCs, and siRNA-CXCR4 transfected BMSCs	Sprague- Dawley rats	2 hr	24 hr post- surgery	2 × 10° into the femoral vein	Increase in CXCR4-BMSCs surrounding the infarct compared to nontransfected and siRNA-CXCR4 transfected BMSCs	CXCR4-transfected BMSCs increased the periinfarct capillary bed, reduced the infarct size, and improved the functional outcome compared to nontransfected and siRNA-CXCR4-transfected BMSCs	141	ASED THEF
SCs	No	Sprague– Dawley rats	3 hr	0, 12, and 24 hr after stroke onset	2×10^{6} intravenously	Migration toward the lesion, questionable differentiation toward endothelial cells	Reduction of infarct region. Improvement in sensorimotor function, upregulation of CXCR4 and SDF-1. Decreased apoptosis in infarct region	132	RAPIES I
MCS and ASCs	No	Sprague- Dawley rats	60 min	30 min after reperfu- sion	2 × 10 ⁶ into the femoral vein	Migration of transplanted cells toward the lesion was not observed	BMSC and ASC administration improves functional recovery independent of reducing the infarct volume and cell migration. Treated groups show higher cell proliferation, oligodendrogenesis, synaptogenesis, and angiogenesis markers	6	FOR ISCHEM
3MSCs nd 1-ASCs	No	C57BL/6J mice	90 min	Immediately after reperfu- sion	1 × 10 ⁵ ASCs or BMSCs into the tail vein	n/a	ASC administration attenuated ischemic damage. Incomplete ASC incorporation in the brain. HGF and angiopoietin-1 expression was significantly increased in ASC-treated mice compared with the BMSC group	139	IC STROKE
MSCs	No	Sprague– Dawley rats	90 min	7 days post- surgery	$3.4 \pm 1.2 \times 10^{6}$ into the saphenous vein	Donor cells accumulate in the ischemic hemisphere, but also in the spleen and lungs	IV-injected 99mTc-HMPAO-labeled MSCs home to the ischemic lesion but also accumulate in the lungs and the spleen	145	•
								Continued	7

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Table I.	Continued							
Stem cell type	(Pre)diffe- rentiation and/or treatment	Species	Occlusion	Time of transplan- tation	Cell dose and location of transplantation	Fate of transplanted cells	Outcome	Reference
h-BMSCs	No	Sprague- Dawley rats	Hours, no details	60 days post- surgery	4×10^{6} in the jugular vein	Donor cells preferentially migrate to the spleen, up to 12 days postinjection	Significant reduction in striatal and periinfarct area. Reduced loss of hippocampal neurons, significant reduction in MHC-II activated inflammatory cells in gray and white matter. TNF-alfa expression in the spleen	134
r-BMSCs	° Z	Sprague Dawley rats	nim 06	1, 4, or 7 days post- surgery	3 × 10 ⁶ intravenously	Cells transplanted 1 day after stroke migrated toward the cortex, cells transplanted after 4 days or 7 days migrated to the striatum	was voterased Functional recovery (mNSS score) was highest when cells were transplanted I day after surgery. This was correlated with a time-dependent expression of SDF-1 and MCP-1 between ischemic	133
r-BMSCs	° Z	A ged Wistar rats	2 hr	1 month after surgery	3 × 10° intravenously	Preferential differentiation toward astrocytes (13%) over neurons (6%). Survival of donor cells was not quantified	regions Significant sensorimotor and general neurological recovery after cell compared with control animals BMSC treatment reduced scar thickness, and increased the number of proliferating cells and oligodendrocyte precursors. SDF-1 is upregulated in the ischemic boundary zone after stroke. BMSCs express CYCPA	146
r-BMSCs from SHR-SP rats	No	A ged SHR-SP rats	Permanent	30 days before stroke onset	5×10^5 into the tail vein vein	No direct transplantation, injected donor cells prior to MCAO	SHR-SP BMSCs transplantation increased microvasculature density in the periinfarct zone, reduced ischemic brain damage, and improved neurologic function. Rejuvenation of bone marrow from aged rats with young cells enhanced the ischemic response at the level of endotheial/vascular activation	152
h-UTCs	0 N	Aged Wistar rats	Permanent	24 hr post- surgery	1×10^7 cells/kg into the tail vein	Very few donor cells present at lesion site, no reactivity for MAP2 or GFAP	IV administration of hUTC improved neurological functional recovery without reducing infarct size, increased progenitor cell proliferation and vessel density in the ischemic boundary zone, and enhanced constructoresis	148
r-BMSCs, h-BMSC	No	Aged Sprague- Dawley rats	3 hr	6 hr post- surgery	1×10^6 cells/kg into the tail vein	1% migrates toward the lesion	Daily treatment with G-CSF improved neurological fundion. G-CSF + BMSC transplantation stimulated angiogenesis in the infarct core but did not further improve neurological function or infarct outwas size	149

IGF-1, insulin-like growth factor 1; HGF, hepatocyte growth factor; SHR, spontaneous hypertensive rats; SHR-SP, stroke-prone SHR; 99mTc-HMPAO, 99-technetium bound to hexamethylpropylene amine oxine; hUTCs, Human umbilical tissue derived cells; EV, extracellular vesicles; MCP-1, monocyte chemotactic protein-1; TNF-alfa, tumor necrosis factor alfa; prefix h, human; m, mouse; t, rat.

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STEM CELL-BASED THERAPIES FOR ISCHEMIC STROKE • 9

2 the ideal candidates to stimulate repair in the central nervous system (CNS) due to their neu-3 rogenic differentiation potential and predisposition.^{50–53} Promising results have already been 4 achieved with human NSCs in animal models of neurological disorders, including stroke.^{8,54} 5 However, there is a need for alternative stem cell sources with regenerative potential due to 6 ethical considerations with regard to the isolation of NSCs from embryonic or fetal tissue together with isolation and culturing complications of adult NSCs.^{55, 56} These alternative stem 7 8 cells need to be able to reconstitute the lost tissue or stimulate endogenous repair. The most 0 promising alternatives for NSC that are of nonembryonic or nonfetal origin are mesenchymal 10 stem cells (MSCs), induced pluripotent stem cells (iPSCs), and bone marrow mononuclear cells 11 (BMMNCs). These stem cell sources have shown to possess regenerative effects on the brain and allogeneic transplantation potential^{57–63} Moreover, these stem cell types can be obtained by 12 13 means of minimal invasive procedures thereby reducing donor site morbidity during isolation. 14 Although it remains a topic of debate whether MSCs possess NPC properties, several studies 15 have reported the ability of subtypes of MSCs to acquire neuronal features following exposure to the proper environmental stimuli.^{64–66} In addition to the discussion of which stem cell source 16 17 is most suitable for stroke research, different animal models such as the middle cerebral artery 18 occlusion (MCAO) model or photothrombotic stroke model are available to induce stroke in 19 an experimental setting, each with their own strengths and weaknesses.^{67–69}

20 Multiple mechanisms have been proposed for stem cell mediated therapies, including brain 21 protection, cell replacement, immunomodulation, and promoting both brain plasticity and angiogenesis in damaged brain regions (Fig. 2).⁴⁹ Interestingly, these mechanisms are mainly 22 23 thought to be mediated by the effect of the stem cell secretome on endogenous stem cells 24 and on the host microenvironment instead of directly replacing the lost cells,^{59,70,71} although 25 encouraging results have also been achieved with cell replacement studies.^{57,58} Therefore, the 26 transplanted cells can be seen as a vehicle for sustained growth factor delivery at the stroke 27 lesion, which can also respond dynamically to changes in the local microenvironment as will 28 be discussed next and into more detail in the following sections.

29 Stem cell mediated neuroprotective effects have been observed in in vitro and in vivo models of neurological disorders.^{72–74} These neuroprotective effects are mainly attributed to 30 31 the soluble factors secreted by the stem cells. In addition to the development of protective 32 therapies, interventions aiming at the directed recruitment and differentiation of NSCs to the 33 site of injury are considered. It is known that NSCs are present in the subventricular zone and dentate gyrus of the hippocampus in the adult brain.^{53,75,76} Moreover, following ischemic 34 35 stroke, endogenous NSCs differentiate into neurons and migrate toward the site of stroke injury and contribute to brain repair.^{77,78} A determining factor in the directed migration of neurons 36 37 is stromal cell derived factor $\alpha 1$ (SDF-1) and its receptor CXCR4.^{78,79} This SDF-1/CXCR-4 38 axis has been shown to act as an inflammatory mediator after acute cerebral ischemia^{80,81} but has also been shown to play an important role in CNS development,⁸² has a modulating effect 39 40 on different subsets of neurons,⁸² and has strong effects on cell migration, axon guidance, and angiogenesis in the postacute phase of stroke.79,82 41

42 Unfortunately, the endogenous repair by NSCs is insufficient to completely replace the 43 lost tissue. Therefore, in addition to exerting a protective effect on the brain, novel cell based 44 therapies are focusing on improving the recruitment of and repair by endogenous NSCs and 45 supporting cells.⁸ Direct cell replacement by neurons derived from stem cells themselves is also 46 a route that is being considered, although it is uncertain whether the transplanted stem cells are able to survive and adequately integrate into the host brain.^{57,59} It has been suggested that 47 48 damaged areas in the brain can only be successfully reconstituted by the equivalent homotopic 49 neurons, which stresses that adequate pretransplantation targeted differentiation of stem cells 50 grafts toward specific types of neurons is required for direct cell replacement by the stem cells 51 themselves.^{83–85} For example, it has been shown that grafting cortical donor tissue into the



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Figure 2. Mechanisms of action of cell-based therapies in ischemic stroke. The poststroke microenvironment 24 can be modulated by exogenously delivered stem cells by multiple mechanisms to trigger tissue repair. Stem cells 25 can contribute to poststroke recovery by stimulating the migration of endogenous NSCs toward the stroke lesion, where proliferation and differentiation toward replacement, neurons can be triggered. In addition, transplanted 26 stem cells are thought to be able to replace the lost neurons themselves in addition to stimulating host NSCs. 27 Moreover, the formation and attraction of new blood vessels toward the ischemic lesion and the stimulation of synaptogenesis and synaptoplasticity contributes to host repair. In addition to directly stimulating the formation 28 of new brain tissue, the degradation of existing cells in, for example, the ischemic penumbra is inhibited by 29 neuroprotective mechanisms such as ROS scavenging by the transplanted cells. Immunomodulatory effects 30 are also observed and include the inhibition of neutrophil activation and migration, effector T-cell and B-cell inhibition, reducing the activation and attraction of peripheral dendritic cells, and stimulating the M2 microglial 31 phenotype. These effects are predominantly caused by the soluble factors released by the stem cells, but also 32 cell-cell interactions appear to play a role. Image was created using Servier Medical Art. 33

35 damaged motor cortex reestablished cortical and even subcortical circuitry,⁸⁴ a feature that was 36 not observed when using heterotopical tissue such as occipital cortex.⁸⁶ More recently, a study 37 by Michelsen et al. demonstrated that in vitro differentiated mouse embryonal stem cell (ESC) 38 derived visual cortical neurons were able to reestablish connections with the damaged visual 39 cortex with recriprocal axonal projections and synaptic integration.⁸³ Interestingly, grafting 40 these cells in the damaged motor cortex or ESC-derived motor neurons in the damaged visual cortex did not lead to graft integration.⁸³ In addition, targeted differentiatiated iPSCs to pyra-41 42 midal cortical neurons have been shown to integrate in the host circuitry after transplantation into to the neonatal mouse brain⁸⁷ and have been used in preclinical stroke research,⁵⁷ as will 43 44 be discussed later. Similarly, it has been suggested that potential donor cells for Parkinson's 45 disease should be of the correct nigral dopaminergic neuron phenotype to improve functional engraftment with the appropriate targets.⁸⁵ Another theory in which cell-based therapies are 46 47 believed to improve the functional outcome in stroke is by directly inducing brain plasticity 48 after the ischemic insult. Although studies report the functionality of transplanted cells in the 49 endogenous neuronal circuitry, these effects are thought to be mainly mediated by promot-50 ing the formation of new synapses between existing neuronal cells and not by functionally 51 integrating into the host neuronal network.57,58,87,88

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A key concept in the regeneration of lost tissue is establishing adequate blood supply to the regenerating tissue. Without proper vascularization that provides oxygen and nutrients, the newly formed neuronal tissue will be unable to survive. Previously, it has been shown that stem cells can form vascular structures in vitro and secrete proangiogenic factors that can positively influence the growth of blood vessels in vitro and in vivo.^{89–91} Therefore, stimulating angiogenesis is another mechanism by which cell-based therapies can influence stroke outcome. Remarkably, revascularization appears to be the main mechanism by which BMMNCs are able to ameliorate the disease outcome.^{61–63}

10 In addition to protecting damaged neurons, restoring the lost neuronal circuitry and blood 11 supply, stem cells have been shown to be able to mediate the immune response.^{92,93} The mechanisms of these immunomodulating properties include influencing the activation state of mono-12 13 cytes, natural killer cells, B cells, T cells, and neutrophils. Stem cells were also shown to mediate 14 immunoglobulin release from plasma cells and upregulate the amount of regulatory T cells.^{92–94} 15 However, it is important to take into account that in ischemic stroke, one of the most common 16 causes of stroke-related morbidity is severe systemic immunosuppression, making patients 17 susceptible to infections.⁴¹ Therefore, additional systemic immunosuppression by cell-based 18 therapies could worsen stroke outcome. Fortunately, no adverse effects on systemic cytokine 19 levels were observed following stem cell transplantation in a rat model of stroke.94

Despite the promising results with BMMNCs in ischemic stroke from preclinical studies,^{61–63,95,96} in vitro evidence of the effect of BMMNCs on the above-mentioned mechanisms is scarce.^{97,98} Therefore, this review will focus on MSCs and iPSCs as readily available sources of stem cells and compare their efficacy and potential to ameliorate the disease outcome in animal models of ischemic stroke. In addition, novel imaging strategies allowing in vivo tracking of transplanted cells and noninvasive evaluation of brain repair following stroke will be discussed.

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2. MESENCHYMAL STEM CELLS AS A THERAPY IN STROKE

30 MSCs, initially discovered in the bone marrow stromal cells (BMSCs) by Friedenstein et al. 31 in the late 1960s,⁹⁹ were later found to be able to differentiate toward cells producing mesenchymal tissues including bone-forming osteoblasts, cartilage-producing chondroblasts, and 32 33 adipocytes.¹⁰⁰ In addition to bone marrow, MSCs have been isolated from a varying range of 34 other tissues including but not limited to adipose tissue (ASCs), Wharton's Jelly in the umbilical 35 cord (UMSCs), umbilical cord blood, and dental tissues.¹⁰¹⁻¹⁰⁵ Additional research into the 36 differentiation capacity of MSCs suggested that these cells were able to differentiate toward 37 hepatocytes,¹⁰⁶ cardiomyocytes,¹⁰⁷ and neuron-like cells.¹⁰⁸ The presence of MSCs in various 38 easily accessible and available donor tissues such as the dental pulp and adipose tissue makes 39 MSCs a promising cell type for stem cell based therapies. However, the main problem in the 40 extensive research with MSCs is the difficulty to compare study outcomes between different 41 research groups. Research groups often have their own methods of isolating, expanding, and characterizing the cells, leading to diverging criteria to define MSCs.^{101-103,109,110} MSCs are 42 43 a heterogeneous population that generally express the surface markers, CD29, CD44, CD90, 44 CD117, and CD146, while they do not express CD34 and CD45 although subpopulations of 45 CD45- and CD34-expressing MSCs were identified.¹¹¹

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A. In vitro Evidence for the Regenerative and (Neuro)protective Potential of MSCS on the Brain

⁵⁰ Despite interlaboratory differences in defining and culturing MSCs, researchers agreed on the ⁵¹ multilineage differentiation potential^{100, 101, 112} of these stem cells and subsequently investigated

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2 the ability of these cells to transdifferentiate into neuronal or neural-like cells in order to obtain 3 a cell source to replace the lost tissue after ischemic stroke. Early studies that investigated the 4 neurogenic differentiation potential of MSCs were performed using BMSCs, but also hASCs 5 and dental tissue and umbilical cord derived stem cells were successfully differentiated to cells 6 with a neuronal phenotype, expressing markers such as neuronal nuclei (NeuN), microtubuleassociated protein 2 (MAP2), neural cell adhesion molecule, and synapsin I.66, 108, 113, 114 Al-7 8 though a consensus was not found between the differentiation protocols, epidermal growth 9 factor and basic fibroblast growth factor (bFGF) are thought to play an important role in inducing MSCs toward a neuronal cell lineage.^{66,113} Subsequent maturation of the induced cells 11 was based on increasing intracellular cyclic adenosine monophosphate and protein kinase C signaling or by specific growth factor administration.^{66,108,113–115} However, few studies performed 12 13 electrophysiological measurements on the differentiated cells but were able to report both 14 voltage-gated sodium and potassium currents that could be reversibly blocked by tetrodotoxin and tetraethylammonium, respectively.^{66, 108, 113, 115} Of these studies, only the studies by Wislet-15 Gendebien et al.¹⁰⁸ and Gervois et al.⁶⁶ could demonstrate the ability of the differentiated cells 16 17 to generate a single action potential in neuronally differentiated BMSCs and human dental 18 pulp stem cells, respectively, demonstrating only incomplete neuronal differentiation.

19 In addition to the neuronal differentiation capacity of MSCs, researchers also investigated 20 the neuroprotective and regenerative potential of the MSC secretome. Hypoxia- and glutamate-21 induced excitotoxicity assays were used as in vitro models for ischemic stroke. It was shown 22 that the MSC protect SH-SY5Y neuroblastoma cells against hypoxia- and glutamate-induced 23 excitotoxicity both in coculture assays and assays using conditioned medium of MSCs, suggesting paracrine effects.^{72, 74, 116, 117} Although the influence of the MSC secretome on NSC survival 24 25 and/or differentiation is not evaluated in vitro, the MSC secretome has been shown to stimulate neurite outgrowth in dorsal root ganglia (DRG)^{118,119} and axotomized retinal ganglion 26 27 cells (RGCs),⁷³ and to enhance survival of these axotomized RGCs⁷³ and primary cortical¹²⁰ and dopaminergic neurons.¹²¹ Neurotrophins/growth factors that are secreted by MSCs include 28 29 glial-derived neurotrophic facto, neurotrophin-3 (NT-3), nerve growth factor (NGF), and brainderived neurotrophic factor (BDNF).^{66,73,118-122} These factors are believed to play an important 30 31 role in neurite outgrowth of DRGs^{118,119,122} and axotomized RGCs.⁷³ Moreover, these factors 32 are also suggested to protect RGCs from neurodegeneration after axotomization,⁷³ cortical 33 neurons from nitrix oxide exposure and withdrawal of trophic support,¹²⁰ and dopaminergic 34 neurons from 6-hydroxy-dopamine.¹²¹

35 As mentioned previously, another key concept in promoting brain regeneration after is-36 chemic stroke is stimulating revascularization of the regenerating tissue. Therefore, the influence 37 of the MSCs was not only investigated in (neuro)protective or neurite outgrowth assays but 38 also the ability of the MSCs to stimulate angiogenesis was evaluated. These studies showed 39 that MSCs are able to stimulate tube formation and endothelial cell migration, enhance wound 40 healing, and improve blood vessel formation in the chorioallantoic membrane essay.^{89, 123, 124} 41 These proangiogenic properties of MSCs were attributed to the soluble factors that are secreted 42 by the cells (Table I in Refs. [89] and [90]) Furthermore, it was shown that MSCs protect endothelial cells against hypoxia-induced cell death.¹²⁵ Several studies also suggest that MSCs not 43 44 only promote angiogenesis by paracrine effects but that these cells are also able to differentiate 45 into endothelial cells (Table II in Ref. [89]).

Although neurogenic differentiated MSCs express neuronal markers, differentiation of these cells toward mature neurons appears limited, as only immature electrophysiological profiles can be generated from these cells. Nonetheless, MSCs show great promise in vitro as shown by the neuroprotective and proangiogenic effects of the MSC secretome. Therefore, several studies have transplanted MSCs into animal models of ischemic stroke and evaluated the outcome, as will be discussed below.



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Table II.	Continued							
Stem cell type	(Pre)diffe- rentiation and/or treatment	Species	Occlusion time	Time of transplan- tation	Cell dose and location of transplantation	Fate of transplanted cells	Outcome	Reference
h-iPSCs	iPSC-derived It-NES cells	Aged Sprague- Dawley rats	30 min	48 hr post- surgery	1.5 × 10 ⁵ intracortical in two sites	49.2 % of the grafted cells survive 8 weeks posttransplantation, 30% of transplanted cells express DCX in the periphery, 91.3% express HuD, and 19.6% express GABA outside the graft core	Cell-grafted showed increased sensorimotor function compared to vehicle-treated rats. Transplanted It-NES cells expressed markers of neuroblasts, mature and GABAergic neurons. Microglia activation was diminished in It-NES grafted rats. Neuronal loss was diminished after transchation.	170
h-iPSCs	iPSC-derived NPCs	Sprague- Dawley rats	90 min	1 Week post- surgery	1×10 ⁵ intrastriatal	Survival not quantified, grafted cells express Sox2, nestin, Pax6, and extend MAP-2 expressing processes into the perilesional parenchym	Grafted iPSC-NPCs initially exert trophic effects on host brain structures, followed by iPSC-NPCs integration into the host brain	169
h-iPSCs	iPSC-derived NSCs	Sprague- Dawley rats	2 hr	Immediately after reperfu- sion	1 × 10 ⁶ intrastriatal	Transplanted cells survived and migrated into the damaged host tissue and express nestin (51.4%) and beta III tubulin (44.3%)	Engrafted cells survive, migrate, and differentiate toward neuronal cells Transplanted cells improved behavioral and sensorimotor function	164
h-iPSCs	iPSC-derived NSCs	C57BL/6J mice	1 hr	24 hr post- surgery	1 × 10 ⁵ intrahippocampal	Engrafied cells migrated toward the site of injury. Survival % not quantified	Improved motor and sensorimotor function in graft-receiving animals Mechanisms of action include a decrease in proinflammatory markers, adhesion molecules, and microglial activation BBB damage vas attenuated	172
h-iPSCs	iPSC-derived NPCs	Wistar rats	30 min	1 Week post- surgery	2.5 × 10 ⁵ intracerebral	Double amount of cells in the graft of which 41% were beta JIF tubulin/MAP2 positive; 5% of the grafted cells expressed GFAP	No migration of cells toward the lesion. No significant difference in behavioral recovery. Tumor formation was not present	171
h-iPSCs	Unclear	C57BL/6N and NSG-mice	Phototrom- botic stroke	1 Week post- surgery	 1 × 10⁵ iPSC in the stroke cavity with or without hyaluronic acid hydrogel 	38% versus 30% survival of cells with or without hydrogel 1 week after transplantation in NSG-mice. Grafted cells form DCX-positive neuroblasts	No assessment of functional recovery. Fumor formation was not evaluated	173
Lt-NES, long	e-term expandable neuroe	pithelial-like stem c	cells; NSG-mice,	NOD scid gamm	a immunodeficient mice;	prefix h, human; $m =$ mouse.	Ś	

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B. MSCS as a Therapy for Stroke In vivo

Due to the encouraging in vitro results of MSCs in protecting damaged neurons and stimulating revascularization in addition to secreting multiple soluble factors, the potential of different subtypes of MSCs to ameliorate stroke outcome after transplantation was evaluated in vivo (Table I). These studies evaluated the functional outcome after transplantation with a variety of behavioral tests. These tests include global neurological assessments to evaluate the disease severity such as the Bederson test and the modified neurological severity score (mNSS). In addition, specific motor, sensorimotor, and cognitive tests were performed. For detailed information on behavioral and disease severity tests in animal models of stroke, see Schaar et al.¹²⁶ Taken together, the studies that report an improvement in general neurological, sensorimotor, motor, or cognitive function are described in Table I.

The proposed underlying mechanisms responsible for the improvement in stroke outcome 14 suggested by the studies listed in Table I are diverse. One of the possibilities was that the trans-15 planted cells migrated toward the stroke lesion and differentiated locally toward neurons, estab-16 lishing new connections with the host environment, 127 although the majority of the in vivo stud-17 ies using MSCs as a therapy for stroke support paracrine mediated brain regeneration. 59, 128, 129 18 Regardless of the administration route, the fate of the transplanted cells was tracked using 19 markers such as DiI,^{60,130–133} DiR,¹³⁴ q-dot¹²⁷ or bromodeoxyuridine (BrdU)¹³⁵ incorporation prior to transplantation; LacZ¹³⁶ or GFP transduction;^{59,70,137–142} iron particle^{60,143} or radionu-20 21 clide labeling;^{144,145} in situ hybridization with the Y chromosome when male donors were used 22 in a female host146,147; or antibodies directed against human mitochondria or human nuclei 23 when human MSCs were grafted in a rodent stroke model.^{71,128,148,149} While the listed studies 24 could observe improvement of stroke outcome after transplantation, the amount of engrafted 25 cells that was present in the stroke lesion was limited. Pioneered by Zhao et al., intracranial 26 transplantation of MSCs showed that MSCs migrated toward the brain infarct region and 27 were able to survive in the host brain and promote functional recovery.⁷⁰ Additional studies showed that although they were present only in low numbers, ^{59,71,130,137,138} the transplanted 28 29 cells locally differentiated toward neural cells with a predisposition toward astroglial cells in 30 preference to neurons.^{59,130,137} Despite the local delivery of the transplanted cells in the stroke 31 lesion, functional integration and replacement of the lost neural circuitry does not appear to 32 be the mechanism of action of intracerebral transplanted MSCs to improve stroke outcome. 33 The results of these studies suggest that the soluble factors secreted by the MSCs are the main 34 actors in improving the functional outcome.^{59,70,130,137} After cerebral transplantation of the 35 MSCs, improved angiogenesis,^{71,130,138} increased neuronal activity,⁷¹ reduced loss of periin-36 farct cells,¹³⁷ and immunomodulatory effects¹⁴² were observed. After transplantation, the local 37 levels of soluble factors such as BDNF, vascular endothelial growth factor (VEGF), bFGF, 38 and angiopoetin-2 were elevated. 71,130 In order to increase the regenerative potential of in-39 tracerebral administered MSCs, several alternative research approaches were investigated. For 40 example, umbilical cord matrix stem cells were cultured in the presence of conditioned medium 41 obtained from a 5-day culture of rat-derived neural cells. However, this approach did not lead 42 to an additional improvement of stroke outcome.⁷¹ Alternatively, it was shown that hypoxic 43 preconditioning of MSCs promotes the survival, migration, and homing of the transplanted 44 cells toward the ischemic lesion compared to nonpreconditioned cells.¹⁴⁰ Transplantation of 45 these cells also leads to an additional functional improvement after intranasal administration, 46 which is assumed to be mediated by an increase in the expression of migration-related proteins 47 such as CXCR4 and MMPs in the hypoxic preconditioned stem cells.¹⁴⁰ 48

Intracerebral administration of MSCs is an invasive procedure and can lead to iatrogenic damage. Therefore, systemic administration of the transplanted cells via the arterial or venous route was considered. Intraarterial administration (IA) of MSCs in stroke has shown beneficial

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2 effects in animal stroke models. Interestingly, IA-transplanted cells are able to cross the BBB 3 after ischemic stroke and migrate toward the stroke lesion. Even though MSCs were found 4 in the core and periinfarct zone and expressed both astrocyte and neuronal markers, ^{128, 131, 147} 5 paracrine mechanisms of action of IA-delivered MSCs are thought to be responsible for the 6 enhanced stroke outcome, although integration into the host brain was observed but not 7 functionally confirmed.¹²⁷ At the ischemic boundary zone, MSCs are only present in low 8 numbers but were found to enhance axonal sprouting and remyelination,¹⁵⁰ angiogenesis, 9 and BDNF production while decreasing MMP-9 levels and suppressing microglial activity.¹²⁸ 10 Furthermore, BMSC administration into aged rats showed a sustained effect and donor cell 11 survival up to 1 year after transplantation.¹⁴⁷

Another route of MSC transplantation is via the venous system. Remarkably, MSCs 12 13 that were intravenously (IV) delivered, improved the functional outcome after stroke with-14 out MSCs being observed in the ischemic brain or in lower numbers than in IA or intracerebral transplantation.^{60,135,139,148,149,151} Despite being almost absent in the ischemic brain, 15 16 IV transplantation of MSC induced an increase in periinfarct zone microvasculature density and the expression of proangiogenic factors^{60, 135, 139, 148} and improved oligodendrogenesis and 17 synaptogenesis.^{60, 148, 152} Remarkably, when BMSCs were transplanted IV in a chronic stroke 18 19 model 60 days after surgery, the beneficial effects of the graft were attributed to their im-20 munomodulatory effects on the stroke lesion and on the spleen where the cells preferentially homed toward.¹³⁴ Another successfully applied therapeutic approach was to transduce BM-21 SCs with an adenoviral vector for PIGF¹³⁶ or a lentiviral vector for CXCR4.¹⁴¹ which further 22 23 improved the functional outcome after stroke compared to nontransduced BMSCs. The SDF-24 1/CXCR4 interaction in chemotaxis was found to be stimulated after intravenous administra-25 tion of MSCs and plays an important role in MSC-homing toward the stroke lesion.^{133,141,146} 26 In accordance with the other routes of administration, the paracrine effects of the transplanted 27 cells appear to be responsible for the posttransplantation effects on stroke outcome. This was 28 supported by a recent study by Doeppner et al., who showed that repeated IV administration 29 of extracellular vesicles produced by BMSC led to a similar improvement as injecting the stem 30 cells themselves.¹²⁹ Moreover, this study also showed reduced poststroke immunosuppression 31 after EV transplantation. 32

This overview shows a variety of studies that were performed using subtypes of MSCs and 33 different administration routes. Although similar results were achieved with different subtypes 34 of MSC, it remains debatable which is the most suitable source for MSC. Moreover, MSCs 35 showed great promise in replacing the neuronal tissue after transplantation due to their in 36 vitro neurogenic differentiation potential. However, cell replacement could not be proven in 37 vivo and it is assumed that paracrine factors are responsible for the beneficial effects after 38 MSC injection, which mainly stimulate angiogenesis and protect the host environment against additional damage without adequately stimulating endogenous neurogenesis or replacing lost 40 neurons. Therefore, additional stem cell sources are considered that are able to differentiate to functional mature neurons in vitro and can potentially improve stroke outcome more effectively than MSCs.

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3. INDUCED PLURIPOTENT STEM CELLS AS A THERAPY IN STROKE

In 2006, Takahashi and Yamanaka successfully transformed murine and in 2007, human fibroblasts to pluripotent stem cells by retroviral transduction with the transcription factors Oct3/4, Sox2, Klf4, and c-Myc, the so-called Yamanaka factors, allowing the formation of a patient-specific source of pluripotent stem cells.^{153,154} These iPSCs are able to differentiate toward tissues from all three germ layers in vitro and in vivo as proven by teratoma formation upon subcutaneous transplantation of iPSCs.

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The ability of iPSCs to differentiate into tissues of all three germ layers offers numerous potential therapeutic approaches. However, the main drawback in using iPSCs for transplantational studies in stroke research is that these cells, such as human ESCs, form teratomas when injected in an undifferentiated pluripotent state, with little to no improvement of the disease outcome.^{155,156} Therefore, iPSCs are irreversibly predifferentiated in vitro in order to minimize tumor formation and improve the functional outcome as only the undifferentiated iPSCs form teratomas. However, a recent study by Choi et al. showed that iPSC- derived NPCs reactivate the silenced exogenous retroviral genes caused by a downregulation of DNA methyltransferases during differentiation and can return toward their pluripotent and thus tumorigenic state.157 Moreover, nondifferentiated iPSCs can remain present within an iPSC-derived progenitor cell pool, which showed teratoma formation after subcutaneous transplantation. This teratoma formation was not observed when fully committed iPSC-derived cells were transplanted.¹⁵⁶ This study by Liu et al. was supported by Fu et al., who demonstrated that residual nondifferentiated iPSCs could not be eliminated by extended cell differentiation.¹⁵⁸ Therefore, adequate full neuronal commitment monitoring prior to transplantation is advised and studies that used nondifferentiated iPSCs in stroke research will be left out of this overview.

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A. In vitro Evidence for the Regenerative and (Neuro)protective Potential of IPSCS on the Brain

23 The multilineage differentiation potential of iPSCs makes these cells an attractive alternative for 24 NSCs as a source for cell-based therapies in neurological disorders. Neurogenic differentiating 25 iPSCs follow similar developmental principles as hESC-derived neurons, although the neural 26 differentiation efficiency can differ between different iPSC lines.¹⁵⁹ Targeted differentiation of 27 iPSCs toward neuronal subtypes has been achieved by various research groups, using different approaches to generate a variety of neuronal cells including medium spiny neurons, 160 dopamin-28 ergic neurons,¹⁶¹ motor neurons,^{159,162} nociceptors,¹⁶³ and pyramidal cortical neurons.^{57,87} Furthermore, it was shown that these neuronally differentiated cells are capable of repeated action 29 30 potential firing, suggesting advanced maturation. 57, 87, 159, 161, 163 More importantly, this targeted 31 32 differentiation of iPSCs toward specific neurons is of high importance and provides additional 33 regenerative potential as it has been suggested that damaged brain areas can only be successfully repaired by the neurons corresponding to the damaged area, as discussed earlier.^{83–85} Similar to 34 35 MSCs, various differentiation protocols were developed to induce neurogenic differentiation of 36 iPSCs. The protocols that were most successful are based on retinoic acid and sonic hedgehog 37 signaling or based on SMAD inhibition using Noggin, which blocks SMAD signaling by the transforming growth factor beta superfamily of signaling proteins.^{87, 159, 163, 164} 38

39 In contrast to MSCs, where the paracrine effects of the stem cell secretome have been in-40 vestigated thoroughly, there is a lack of stroke-related in vitro evidence of the (neuro)protective, 41 regenerative, and angiogenic properties of iPSCs. Among the few studies that investigated the 42 secretome of iPSCs, it was shown that iPSCs locally enhanced the production of proangiogenic 43 factors. However, these studies did not include an in vitro secretome analysis or an in vitro evaluation of the angiogenic properties of the iPSC secretome.^{165,166} Although the paracrine effects 44 45 of the iPSC secretome on angiogenesis were not evaluated in vitro, several studies have reported endothelial cell differentiation of iPSCs,^{91,167,168} which opened up the possibility that trans-46 47 planted iPSCs can directly contribute to establish new blood vessel formation in the damaged 48 brain.

Despite the lack of in vitro evidence for paracrine-mediated regeneration, the successful
 differentiation of iPSCs toward endothelial cells, neuronal precursors, and mature neuronal
 subtypes encouraged the use of iPSCs in animal models of ischemic stroke.

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B. Neurogenic Predifferentiated IPSCS in In vivo Stroke Models

Contrary to MSCs, iPSCs that were transplanted in animal stroke models were neuronally differentiated prior to engraftment in order to avoid teratogenicity (Table II). Moreover, no reports are available in which iPSCs are administered IV or IA. Information about mechanisms of action of iPSCs in ameliorating stroke outcome is therefore only available for intracranially transplanted neurogenic predifferentiated iPSCs.

When considering the overall results of the behavioral tests performed after iPSC transplantation after stroke, improvements in general neurological score,¹⁶⁹ motor,^{57, 58, 164, 169, 170} sensorimotor,^{57, 58, 164, 169, 170} and cognitive function¹⁶⁴ were observed. On the contrary, Jensen et al. did not observe an iPSC-mediated improvement in behavioral function.¹⁷¹

Underlying mechanisms of action of the transplanted iPSCs include migration and local 13 neuronal maturation of the transplanted iPSCs, ^{57, 58, 164, 170} synaptic integration into the host 14 brain^{57, 58} but also paracrine effects are considered to play a role.¹⁶⁹ The transplanted iPSCs were 15 predifferentiated toward long-term expandable neuroepithelial-like stem cells, 57,58,170 fated to-16 ward cortical neurons⁵⁷ or toward neuronal precursor cells,^{164,169,171} Transplanted iPSCs were 17 traced with antibodies directed against human nuclei, cytoplasm or mitochondria, 57, 58, 169-172 18 Dil labeling,¹⁶⁴ or by using GFP positive iPSCs.^{57,58,173} The engrafted cells survived up to 10 19 weeks after transplantation but the percentage of surviving cells varied considerably between the 20 different studies, which can be caused by several factors. For example, the host strain (i.e., nude 21 rats vs. immunocompetent rats) and species^{174–177} can have an influence on the stroke outcome 22 and cell survival rates.58 Moreover, the altered stroke microenvironment in aged animals might be less favorable for cell transplantation.^{170,178} Another factor that can influence cell survival 24 and amelioration is the time after stroke onset; the transplantation is performed as cell trans-25 plantation at later time points exposes the cells to the established immune response,¹⁷⁹ whereas 26 cell transplantation early after stroke onset or too close to the ischemic core can expose the 27 cells to limited blood supply, oxidative stress, and trophic factor deficiency.¹³ Remarkably, both 28 engrafted fated cortical neurons derived from long-term expandable neuroepithelial-like stem 29 cells and their nonfated counterparts, which were transplanted 48 hr after stroke onset, could be 30 detected 2 months posttransplantation in the damaged rat cortex in the study by Tornero et al.⁵⁷ 31 The transplanted cells differentiated locally toward neuronal cells as shown by the expression 32 of doublecortin (DCX) at the periphery of the graft. The expression of HuD and NeuN was 33 found at the core of the graft, 57, 58, 170 which was increased by differentiating the long-term 34 expandable neuroepithelial-like stem cells toward cortical neurons prior to transplantation.⁵⁷ 35 A study by Lam et al. demonstrated that 1-week survival of the iPSC grafts is enhanced— 36 although not significant—in photothrombotic stroke by delivering the cells in a hyaluronic 37 acid hydrogel. However, despite claiming to use iPSC-derived NPCs, the neuronal character-38 istics were not described, which might have had an impact on the survival rate. Additionally, 39 transplantation of the cells in the hyaluronic acid hydrogel favored DCX-positive neuroblast 40 formation 1 week after transplantation.¹⁷³ The study by Yuan et al. also observed the expres-41 sion of nonmature neuronal markers of the engrafted cells such as beta tubulin and nestin.¹⁶⁴ 42 Moreover, a low fraction of cells differentiated toward astrocytes, as shown by GFAP expres-43 sion. Nonetheless, this study showed a preferential neuronal differentiation of engrafted iPSC-44 derived neuronal precursors after transplantation,¹⁶⁴ which is supported by the study of Jensen 45 et al.¹⁷¹ 46

To examine whether the transplanted iPSCs contributed to the synaptic network in the host brain, various methods to determine synaptic integration were applied. One method that was used to determine synaptic integration was by retrograde tracing of fluorogold in the ipsilateral globus pallidus, 9 weeks after the injection of iPSCs.⁵⁸ In this study by Oki et al., it was shown that a small fraction of the transplanted iPSCs were fluorogold positive,

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2 meaning that striatal transplanted iPSCs extended projections toward the globus pallidus. Other 3 strategies to identify axonal projections of transplanted iPSCs used antibodies directed against donor specific cytoplasmic⁵⁷ or surface markers.¹⁶⁹ The study by Polentes et al. observed iPSC-4 5 derived axonal projections from the site of engraftment that was in the lesion cavity located 6 in the lateral quadrant of the striatum to the striatum and globus pallidus after 1 month and 7 these projections extended into the substantia nigra 1 month later. A few fibers were found 8 in the corpus callosum.¹⁶⁹ The study by Tornero et al. that compared long-term expandable 0 neuroepithelial-like stem cells and long-term expandable neuroepithelial-like stem cells fated 10 toward cortical neurons observed a higher density of projections extending from the site of 11 engraftment in the cortex over the corpus callosum in fated cells compared to the nonfated 12 cells.⁵⁷ In addition to axonal projections, the engrafted cells were shown to be functionally active 13 up to 6 months after transplantation, as was shown by whole-cell patch-clamp recordings in 14 acute brain slice preparations.57,58

15 In addition to cell-replacement mechanisms and synaptic integration, paracrine-mediated 16 improvement of stroke outcome is another mechanism by which transplanted iPSCs can exert 17 their effect. The study by Oki et al. showed that functional recovery was independent of long-18 term engraftment, suggesting a beneficial effect early after transplantation.⁵⁸ Furthermore, 19 they showed that VEGF reactivity was upregulated following transplantation in astrocytes and 2.0 the blood vessel wall of the damaged brain as early as I week after transplantation. How-21 ever, 9 weeks after transplantation, when animals receiving the cell graft showed significant 22 functional improvement, no difference in vessel length density and immunoreactivity for the 23 endothelial marker CD31 was observed between animals that were injected with iPSCs com-24 pared to vehicle-injected animals. An additional study by this research group detected only 25 weak expression of VEGF in the grafted cells and blood vessel walls of the damaged brain 26 8 weeks posttransplantation. VEGF-expressing astrocytes were not observed.¹⁷⁰ This suggests 27 that VEGF signaling is important in early recovery after stroke and can trigger long-lasting 28 effects in brain plasticity. In addition, it can also be postulated that VEGF signaling alone is not 29 sufficient to clarify the improved functional recovery after iPSC transplantation.¹⁸⁰ Moreover, 30 Tatarishvili et al. observed functional improvement from 1 to 4 weeks after transplantation, 31 making it inconceivable that the behavioral improvement is due to cell-mediated neuronal 32 replacement.170

33 Another possible mechanism by which transplanted iPSCs can influence functional recov-34 ery after stroke is by modulating the immune response. Microglia were not found to be more 35 prominent or more activated in vehicle-treated animals compared to animals that received 36 the iPSC graft 8 weeks after transplantation. However, microglia in the vehicle-treated group 37 showed a more round/amoeboid morphology compared to animals that received the iPSC 38 graft.¹⁷⁰ There is no information on how the number and activation status of the microglia 39 was affected in the early time points after transplantation. Previous studies have shown that 40 NSCs, transplanted in the cortex or striatum after stroke, can reduce the number of microglia in both early as late time points after engraftment.^{54,181} In addition to these results, Chen 41 42 et al. previously reported an upregulation of antiinflammatory cytokines and a downregula-43 tion of proinflammatory cytokines after nonpredifferentiated iPSC administration in the stroke 44 brain.¹⁸² More recently, a study by Eckert et al. showed that intrahippocampal transplantation 45 of iPSC-derived NSCs 24 hours after stroke onset reduced the expression of proinflammatory 46 markers, microglial activation, and adhesion molecules while attenuating BBB damage, leading 47 to a significant improvement in motor and sensorimotor function within the first week after 48 transplantation. These data support the influence of early transplantation on the host immune 49 response, leading to an improved stroke outcome.¹⁷²

These results show that functional integration of transplanted iPSCs in the host model is achievable. However, the timing of graft-induced improvement in behavioral tests suggests

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that early amelioration of stroke symptoms is mediated by paracrine mechanisms and may be mediated by the early influence of the graft on the host immune system. In addition, functional improvement caused by the integration of graft-derived neuronal cells is expected to take longer than the 8-9 weeks follow-up in the presented studies. The presented studies by Oki et al.⁵⁸ and Tornero et al.⁵⁷ followed the engrafted cells for up to 6 months after transplantation and were able to observe neuronal differentiation and functionality of the cells, but did not examine the underlying mechanisms into further detail.

One of the major issues in determining the fate of the transplanted cells and determining the exact onset of functional recovery is that most studies described above used traditional histopathological techniques to determine the fate of the transplanted cells and to visualize possible methods of action of the transplanted cells. Therefore, as will be discussed next, novel techniques are being developed to allow real-time and longitudinal noninvasive imaging and tracking of the transplanted cells. Furthermore, additional imaging methods can be applied in the same animals to get real-time information on revascularization and reinnervation as will be discussed next.

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4. CURRENT CHALLENGES AND FUTURE PROSPECTS OF STEM CELL BASED THERAPIES IN STROKE: VISUALIZING DONOR CELLS AND THE INJURED BRAIN

Traditional histopathological techniques can only be applied ex vivo. The most frequently used cell trackers include the use of GFP-transduced or BrdU-labeled stem cells in addition to antibodies directed against human-specific epitopes. However, it has been shown by Burns et al. that thymidine analogs such as BrdU may not be a suitable marker to track donor cells due to label transfer to phagocytizing cells.¹⁸³ Similar to cell tracking, morphological changes such as revascularization and reinnervation, evoked by the transplanted cells, are investigated by using tissue-specific antibodies or other ex vivo methods. Fortunately, noninvasive, quantifiable imaging methods have been developed to track the fate of the transplanted cells and evaluate the host environment, which can also be applied in humans. These methods will provide more insight into the exact timing of behavioral improvements following stem cell transplantation by correlating the presence or absence of the engrafted cells with morphological adaptations such as revascularization and reinnervation at the injured site.

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A. Visualizing Engrafted Cells in the Injured Brain

Noninvasive imaging methods of donor cells are based on magnetic resonance imaging (MRI), 38 positron emission tomography (PET), single photon emission computed tomography (SPECT), and optical methods such as bioluminescence imaging (BLI) and fluorescence imaging (FLI). 40 The advantage of MRI compared to the other imaging methodologies is its high spatial resolution. However, PET-, SPECT-, BLI-, and FLI-based imaging hold the advantage of a higher sensitivity, although the last two are only sensitive in preclinical research. Prior to injection, 43 donor cells can be labeled either directly or indirectly. For indirect labeling, imaging reporter genes are introduced into the host cells that encode for proteins or molecules that will lead to 45 the accumulation of a specific substrate or ligand within the cells in which the reporter is expressed. Direct labeling involves the (stable) attachment or incorporation of reporter molecules 47 into the cells after in vitro incubation (extensively reviewed in Ref. [184]). Direct pretransplan-48 tation donor cell labels such as superparamagnetic iron oxide (SPIO) particles^{60, 185-187} and 49 radionuclides^{144,145} have also been used to track donor cell fate, but as will be discussed below, are also subjected to several disadvantages.¹⁸⁸ The different labeling strategies are illustrated in 50 51 Fig. 3.



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Figure 3. Direct and indirect labeling strategies to track stem cell fate in vivo. Direct labeling methods such as SPIOs and radionuclides have the advantage that they have been shown to have minimal effects on stem cell properties. Furthermore, biosafety risks are minimal and the labeled cells can be easily monitored with noninvasive imaging techniques such as FLI, MRI, PET, and SPECT. The disadvantages of using direct labeling strategies are the decreasing signal over time and the possibility of monitoring a nonspecific signal. These disadvantages can be circumvented by using indirect labeling methods in which reporter genes are incorporated into the donor cell genome of which the reporter protein can be visualized in a substrate-dependent manner, allowing spatiotemporal control of cell tracking. The major disadvantage of using this labeling strategy is the labor-intensive design of reporter gene constructs and biosafety issues regarding the use of viral components to transduce the donor cells. Image was created using Servier Medical Art.

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SPIO particles are commonly used tracers that allow direct labeling of the donor cells for use with T2-weighted MR images without a significant effect on stem cell biology and differentiation potential of the labeled cells.^{60, 185–187} Despite the promising use of SPIOs in noninvasive cell tracking with MRI, this labeling method has several limitations.¹⁸⁹ It is impossible to discriminate between viable and dead cells as the SPIO particles remain present after the cells have died. Furthermore, macrophages and microglia present at the lesion sit may phagocytize the cell fragments of dead SPIO-labeled cells, which can lead to the occurrence of nonspecific signal not originating from transplanted cells. In proliferating cells, the SPIO signal decreases after transplantation, which is aggravated by asymmetrical replication of the donor cells.¹⁸⁹

42 Another approach to directly label stem cells is with radionuclides that are detectable 43 with the nuclear imaging methods PET or SPECT. These radionuclides can bind to the cell membrane (i.e., hexadecyl-4[¹⁸F]fluorobenzoate)¹⁹⁰ or can be taken up by the cell via passive 44 45 diffusion through the cell membrane or via ion channels, transporters, and pumps.¹⁹¹⁻¹⁹³ The 46 most widely used PET-compatible radionuclide is 2-deoxy-2-¹⁸F-fluoro-D-glucose (¹⁸F-FDG). ¹⁸F-FDG has been successfully used to monitor human autologous BMSC and peripheral 47 hematopoietic stem cell homing after myocardial infarction in a human study.^{194, 195} Survival, 48 49 proliferation, and differentiation of radiolabeled MSCs is maintained after radionuclide labeling, suggesting minimal radiotoxicity in these cells.^{196–198} The two most frequently used 50 51 SPECT-compatible radionuclides are oxine-bound Indium-111 (¹¹¹In-oxine) and technetium

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bound to hexamethylpropylene amine oxine (^{99m}Tc-HMPAO). ¹¹¹In-oxine and ^{99m}Tc-HMPAO have a half-life of 2.8 days and 6 hr, respectively, allowing long- and short-term imaging. ^{99m}Tc-HMPAO has been used as an effective tracer in a human study using labeled intraarterial-injected autologous bone marrow mononuclear cells after acute ischemic stroke¹⁹⁹ and shows minimal radiotoxic effects. ¹⁴⁵ Although ¹¹¹In-oxine is an FDA-approved tracer and has been used in animal models of stroke and myocardial infarction^{144,200} and human clinical studies,²⁰¹ it has been reported that ¹¹¹In-oxine binding is reversible, and that cell function and viability are impaired due to radiotoxic effects.^{202,203}

Similar to SPIOs, direct labeling of stem cells with radionuclides also has several disadvantages, such as tracer leakage into the extracellular space that induces labeling of nontarget cells.^{10,188} Furthermore, the tracer leakage to nontarget cells can be homogeneously distributed to these cells that leads only to a diminishable background signal compared to the initially targeted cells. More important is that the application window of these tracers is highly dependent on the half-life of the used radionuclide, reducing the use of these radionuclides for longitudinal follow-up of the labeled cells.¹⁰ Radiotoxicity does not appear to be a major hurdle when using low doses of ¹⁸F-FDG or ^{99m}Tc-HMPAO, but becomes a problem when ¹¹¹In-oxine is used as a tracer and is highly dependent on the dose that is used.¹⁹⁷

19 To minimize the problem of label transfer from donor to host cells or to cope with the 20 decreasing signal after transplantation of the donor cells, other noninvasive imaging methods 21 are available. One of the possibilities is using reporter genes of which the proteins are only 22 encoded in viable cells. Well-known examples are green fluorescent protein,²⁰⁴ which can be detected by FLI, and firefly luciferase (fluc),²⁰⁵ which catalyzes a light-emitting reaction that 23 24 can be detected by BLI upon oxidation of exogenously delivered luciferin. BLI-compatible 25 reporter genes have been successfully transduced in MSCs,^{206,207} and iPSCs¹⁵⁶ and have been 26 successfully used to monitor both donor cell fate²⁰⁷ as endogenous NSCs.²⁰⁸ Although these 27 optical imaging methods provide a highly sensitive technique^{209,210} to monitor cell survival and 28 proliferation, these methods are limited by a loss of spatial resolution due to light scattering and 29 by the anatomical depth of the signal-generating engrafted cells²¹¹ making them only useful for 30 preclinical research in small animals.

31 To improve the spatial resolution and to gain additional information of the engrafted cells. 32 MRI-detectable reporter gene methods have been developed, which are listed in Patrick et al. 33 (Table S1 in Ref. [212]) and reviewed in detail by Vandsburger et al.²¹¹ MRI reporter genes are 34 based on intracellular iron accumulation, enzymatic reactions, membrane-bound proteins such 35 as the biotin-streptavidin interaction, and chemical exchange saturation transfer (CEST).²¹¹ 36 It should be noted that noninvasive MRI reporter gene methods that are based on visualizing 37 iron accumulation cannot distinguish between living and dead cells, once the transduced cells 38 have bound or accumulated iron.²¹¹ Although MRI reporter genes based on cell membrane 39 interactions and enzymatic reactions were developed, the most promising MRI reporter method 40 is based on CEST. CEST is based on compounds containing exchangeable protons that resonate 41 at a different frequency than bulk water protons. These protons can be selectively saturated with 42 a radiofrequency pulse and are transferred to the bulk water molecules after proton transfer, 43 attenuating the signal of the water signal.²¹³ However, to date, no stem cell tracking experiments 44 have been performed with CEST. 45

In addition to MRI-detectable reporter genes, nuclear reporter genes that can be detected with PET or SPECT have been developed. These highly sensitive techniques allow for repeated visualization of migration and function of donor and host cells.²¹⁴ Imaging reporter genes can be subdivided into three main categories: enzymes, receptors, and transporters.¹⁰

The herpes simplex virus type 1 thymidine kinase gene (HSV1-*tk*) encodes the viral protein HSV1-TK, an enzyme that can phosphorylate nucleoside analogs that are subsequently

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negatively charged and become entrapped in the cells.²¹⁵ HSV1-TK can phosphorylate isotopelabeled pyrimidine analogs that can subsequently be used as reporter probes in PET and SPECT imaging.¹⁹¹ Unfortunately, the use of viral proteins can potentially cause an immune response and more specific for stroke research, none of the HSV1-TK compatible tracers is able to cross 6 the BBB.^{216,217} These disadvantages of HSV1-TK compatible tracers can be overcome by using reporter genes encoding for receptors such as the dopamine D2 receptor (D2R)²¹⁸ and human somatostatin receptor subtype 2 (hSSTr2)²¹⁹ or by reporter genes coding for transporters such as the human sodium iodide symporter (hNIS) that can transport all radioactive forms of I⁻ 10 as well as other isotopes (i.e., Technetium-99m).²²⁰ The advantage of using D2R, hSSTr2, and 11 hNIS is that they can be labeled with tracers able to cross the BBB and, since they are of human origin, are not likely to elicit an immune response.¹⁹¹ However, D2R and hSSTr2 have not been 12 13 used for longitudinal tracking of engrafted cells. hNIS has already been successfully used to 14 track MSCs²²¹ and iPSCs²²² in vivo but to date, no studies have been performed to trace donor 15 stem cells after transplantation in stroke with hNIS.

16 In order to combine the high spatial resolution and anatomical precision of MRI with the 17 sensitivity of radionuclide or BLI, dual-modality probes have been developed to exploit the characteristics of each imaging method ²¹²,²²³ (reviewed in Ref. [224]). For example, Patrick 18 19 et al. used Oatp1a1 as a reporter.²¹² This molecule is able to mediate the cellular uptake of 20 several small molecules, including gadolinium-based contrast agents that enhance T1-weighted 21 images in MRI and the radionuclide ¹¹¹In, which can be detected with SPECT. Dual-modality 22 imaging has also been used in stroke research to track transplanted NSCs²²⁵⁻²²⁷ and MSCs.¹⁴³ 23 In these studies, donor NSCs were transduced with *fluc* and labeled with SPIO particles to allow 24 BLI and MRI imaging,^{226,227} or were imaged with ¹⁹F MRI after transduction with *fluc*.²²⁵ In 25 a study by Walczak et al., SPIO-labeled MSCs were monitored with MRI combined with laser 26 Doppler flow.¹⁴³ 27

Over the past decade, several advances have been made in molecular imaging to facilitate stem cell tracking after transplantation. As will be discussed next, some of these methods are not only applicable to track the fate of the donor cells but can also be used to acquire additional information on the endogenous repair mechanisms following ischemic brain injury.

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B. In vivo Imaging of the Recovering Brain

34 Similar to tracking the fate of donor cells, noninvasive imaging modalities based on MRI, 35 radionuclide imaging, or optical methods can be applied to monitor the physiological and/or 36 functional properties of the host environment. These methods allow visualization of neurovascular processes and neurological function but can also be used to study endogenous stem cells 38 responses to treatment.

Optical methods are mainly based on FLI or BLI. In vivo two-photon FLI can be used to 39 40 monitor blood flow, synapse formation, and neuroinflammation.^{228–230} Two-photon imaging 41 requires fluorophores, although label-free imaging has been successfully used to visualize the 42 mouse brain.^{231,232} FLI can also be used to directly monitor neuronal activity by using voltage 43 sensitive dyes and proteins, which change their fluorescent properties in response to changes 44 in transmembrane voltage.^{233,234} Unfortunately, penetration depth is limited and a cranial window is required to visualize subcortical structures and repair processes.^{233,235} BLI-based 45 optical imaging methods are also limited by penetration depth, although it has been shown 46 that the BLI signal can be detected through the intact skull.²⁰⁸ BLI-based methods have been 47 used to track endogenous neuronal stem cells and neurogenesis after stroke.²⁰⁸ In addition, 48 49 fluc under the control of the VEGFR2 receptor promoter has been used to evaluate poststroke angiogenesis with BLI²³⁶ and when put under control of the toll-like receptor 2, the response 50 51 of microglia could be observed after photothrombotic stroke.²³⁷

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Another approach is to use PET or SPECT compatible radionuclides such as¹⁸F-FDG, which has been used in animal models of stroke to evaluate stroke outcome with PET after transplanting iPSCs and ESCs.²³⁸ This study by Wang et al. was able to correlate the ¹⁸F-FDG PET signal with functional improvement as shown by a decrease in mNSS score in animals 6 that received a nonpredifferentiated iPSC or ESC graft. Although the donor cells were not predifferentiated, tumor formation was not observed up to 4 weeks posttransplantation. A similar study was performed by Daadi et al. where NSCs were labeled with SPIO particles 9 and transduced with HSV1-tk to allow PET with the reporter probe [¹⁸F]-FHBG prior to transplantation in a rat stroke model.²³⁹ In accordance with the study of Wang et al., Daadi et al. demonstrated an increased ¹⁸F-FDG PET signal after NSC transplantation and correlated the ¹⁸F-FDG PET signal and the presence of NSCs as shown by [¹⁸F]-FHBG detection. Increased glucose uptake might be indicative of enhanced neuronal function but caution needs to be taken when interpreting ¹⁸F-FDG PET results as it is unclear which mechanisms are responsible for the increased ¹⁸F-FDG uptake in the injured brain.²⁴⁰ More recently, a study by Zinnhardt et al. combined a multitracer PET study with MRI to link the spatiotemporal relationship of MMP and microglial activation after transient MCAO providing more detailed information on early and delayed endogenous stroke responses.²⁴¹

MR-based methods have the advantage that they can couple physiological information with anatomical characteristics of the region of interest. Therefore, several MR-based modalities are used to gain additional information of the host environment in stroke and will be discussed next.

Blood oxygenation level dependent functional MRI (BOLD-fMRI) depends on the hemodynamic response to neuronal activity, which are directly related to the energy demand of the studied brain areas.²⁴² In small animal stroke research, BOLD-fMRI after electric forepaw stimulation has been successfully used to assess functional recovery and electric brain activity.²⁴³ As stated previously, current therapies for ischemic stroke aim to salvage the ischemic penumbra. Therefore, it is important to have an adequate monitoring tool to evaluate the size of the penumbra before and after therapeutic intervention. By using diffusion- and perfusionweighted MRI, the size of the ischemic penumbra can be determined based on the area of diffusion/perfusion mismatch.²⁴⁴ While the diffusion/perfusion ratio is mainly used clinically, it has also been successfully applied in a rat model of ischemic stroke.^{245,246}

33 In addition to providing a ways of evaluating the size of the penumbra, also the architectural 34 information of the affected brain region can be visualized with MR modalities. Diffuse tensor 35 imaging (DTI) images the anisotropy of water molecules in different tissues and is mainly used 36 in stroke research to study and visualize white matter tracts.²⁴⁷ Although its use in experimental 37 stroke in animal models is limited, DTI has been used to study the MRI evolution of stroke 38 macaques.²⁴⁸ While DTI can provide architectural information of white matter organization 39 after stroke, manganese-enhanced MRI (MEMRI) can be used to image synaptic connectivity 40 and assess changes in neuroarchitecture, anterograde axonal transport, and demarcates active regions of the brain independent of hemodynamic contrast compounds.^{249,250} MEMRI uses T1 41 42 contrast-enhancing Mn^{2+} ions that can enter the cell via Ca^{2+} channels. As Ca^{2+} plays a crucial 43 role in neuronal activation Mn^{2+} is transported toward the synaptic cleft, where it can be taken 44 up by other neurons in the circuit.²⁵¹ Neuronal connectivity during stroke and recovery has 45 been evaluated in MCAO models using this technique.^{252,253}

46 Reperfusion plays a key role in repairing the ischemic lesion and can be examined with 47 arterial spin labelling (ASL) scans in order to quantify the absolute amount of tissue perfusion in 48 different brain areas evoked by therapeutic interventions, including stem cell based therapies.²⁵⁴ 49 ASL measurements in small animal models of ischemic stroke were able to visualize perfusion 50 in the ischemic brain, allowing the potential longitudinal follow-up of therapeutic interventions 51 that aim to enhance reperfusion.^{255,256}

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5. DISCUSSION AND PERSPECTIVES

4 Because of the promising preclinical results of stem cell based therapies in in vivo models 5 of ischemic stroke, small-scale human trials were performed using IV delivered autologous MSCs.^{257–259} The outcome of these studies showed that MSC transplantation improved the 6 7 disease outcome but stress that the underlying mechanisms of action need to be determined 8 to provide a more directed approach, although it was reported that the SDF-1 levels in the serum of MSC-transplanted patients were associated with the clinical outcome.²⁵⁹ Therefore, it 0 10 is important that in the preclinical phase the potential of stem cell based therapies is thoroughly 11 investigated in animal models of ischemic stroke. Each model and route of administration has 12 its own advantages to investigate specific mechanisms of action of the donor cells. IA- and 13 IV-delivered donor cells would be the preferred method of administration in human applica-14 tions, but this administration route requires a substantial amount of donor cells compared to 15 intracranial delivered stem cells and are hindered by several limitations. These include high morbidity and cells ending up in the spleen after IA delivery^{143,260} and pulmonary obstruction 16 after IV delivery of donor cells.^{145,261} However, several reports are available that state that 17 18 donor cell migration toward the spleen is a possible mechanism of action of stem cell mediated 19 regeneration following stroke. Acosta et al. demonstrated that IV-delivered BMSCs end up in 20 the spleen, but that BMSC migration to the spleen inversely correlated with a reduced infarct 21 size, periinfarct size, and the number of MHC-II positive activated cells in the striatum.¹³⁴ 22 Although the influence of MSCs themselves on poststroke immunosuppression was not investi-23 gated, Doeppner et al. reported that poststroke immunosuppression was attenuated after MSC 24 extracellular vesicles were injected in stroke mice.¹²⁹ Supporting this hypothesis, Vendrame 25 et al. investigated the immunomodulatory effects of the mononuclear cell fraction of umbilical 26 cord blood cells.²⁶² In this study, it was demonstrated that IV transplantation of these cells 27 diminished spleen reduction and rescued CD8⁺ T-cell counts in addition to a reduction in 28 brain damage. Moreover, it was shown that the cell transplant increased II-10 and interferon 29 gamma mRNA expression and decreased tumor necrosis factor alfa mRNA expression.²⁶² 30 Donor cells applied to the host circulation migrate toward and integrate in low numbers into 31 the brain lesion and ameliorate the disease outcome (See Table I) presumably by neurotrophic effects as the cerebral level of neurotrophins was found to be elevated in some studies.^{132, 135, 139} 32 33 Moreover, IA or IV delivery of donor cells allows the interaction of the BBB with donor cells to 34 be studied that can contribute to knowledge of the neuroimmunological response after ischemic 35 stroke.¹²⁸ Intracranial delivery of donor cells is the most invasive route of transplantation but 36 a thorough meta-analysis of preclinical data by Vu et al. showed that intracranial delivery of 37 MSCs provided greater clinical benefit, although this mode of administration is less favorable 38 for human applications due to the highly invasive nature of the transplantation procedure.²⁶³ 39 Nonetheless, intracranial transplantation of fetal nigral tissue to treat patients with Parkinson's 40 disease has been performed although the clinical benefit remained debatable.²⁶⁴

Donor cells remain detectable at the site of injury and ameliorate the disease outcome as shown by behavioral results and postmortem tissue analysis. However, functional integration of the engrafted MSCs is based on marker expression instead of electrophysiological recordings. The latter has only been performed for iPSC-derived cortical differentiated longterm expandable neuroepithelial-like stem cells, demonstrating functional integration of the engrafted cells.^{57,58} IA or IV delivery of iPSC-derived neuronal precursor or committed cells has not been performed to date.

When considering the most suitable source of stem cell as a potential therapy in stroke research, comparative studies are needed to highlight differences in therapeutic potential of stem cells from different sources in an analogous experimental setup. For example, studies comparing different subtypes of MSCs showed that ASCs are more suitable as a candidate

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2 MSC than BMSCs¹³⁹ while another study compared the same subsets of MSC and could not 3 observe any difference.⁶⁰ Moreover, the studies listed in Tables I and II describe both xenogenic 4 transplantation of human-derived MSCs and iPSCs as allogeneic-derived MSCs in animal 5 models for ischemic stroke. Studies that compare xenogenic and allogeneic MSCs in ischemic stroke are limited to the study by Yasuhara et al.¹³⁷ and Balseanu et al.,¹⁴⁹ who compared 6 7 human and rat BMSCs, and Guttiérrez-Fernández et al. who compared human ASCs with 8 rat-ASCs¹⁵¹ In the study by Yasuhara et al., it was shown that the allogeneic BMSC graft 9 showed a higher survival rate and a higher number of neurogenic differentiated cells. In both 10 engrafted groups, improvement in locomotor and neurological function and a reduced loss of 11 striatal periinfarct cells was observed.¹³⁷ Although Balseanu et al. used both rat and human 12 BMSC in their study, no direct comparison was made between the xenograft and allograft.¹⁴⁹ 13 Preclinical studies with autologous-derived stem cells are scarce and were only described in 14 Jiang et al.'s¹³¹ study who transplanted autologous rat ASCs in ischemic rats, whereas for other 15 subtypes of MSCs, preclinical studies with autologous stem cells are not available. This study 16 by Jiang et al. did not compare the outcome of the transplantation between autologous and 17 allogeneic ASCs, which would provide additional information on extrapolability of their results. 18 Although xenogenic stem cell transplants are not likely to be used in human studies, preclinical 19 studies using xenogenic grafts provide insight into the potential of human stem cell sources in 20 ischemic stroke.

21 To date, no studies have been performed that compare the therapeutic potential of iPSCs 22 with a subtype of MSCs. Although the most thoroughly studied stem cell type in stroke re-23 search are MSCs, the potential of iPSCs and iPSC-derived neural progenitor cells is currently 24 being intensively investigated with several very promising results when iPSCs are predifferenti-25 ated toward neuronal precursor cells or committed cortical neurons.^{57,58} Nonetheless, adequate 26 screening of fully neuronal committed cells preceding engraftment is recommended for several 27 reasons. As stated previously, a recent study by Choi et al. demonstrated the ability of the 28 iPSC-derived NPCs to return toward their pluripotent and thus tumorigenic state by transgene reactivation during differentiation.¹⁵⁷ This statement was supported by Liu et al.¹⁵⁶ and Fu 29 30 et al.¹⁵⁸ who demonstrated that nondifferentiated iPSCs remain present in an iPSC-derived 31 progenitor pool. Another problem with iPSCs, and more specifically with retroviral trans-32 duced iPSCs, is the retroviral gene integration in the host, which promotes tumorigenicity.²⁶⁵ 33 Therefore, additional approaches have been developed to generate iPSCs with a lower risk for 34 tumorigenicity.^{266, 267} Moreover, it has been shown that iPSCs retain an epigenetic memory 35 related to the somatic donor tissue, leading to spontaneous redifferentiation to the cells of the 36 tissue of origin, ^{268–270} although it has been shown that this epigenetic memory and redifferenti-37 ation rate can vary between the somatic cells of origin with different tumorigenic propensities 38 between the somatic donor cells.^{269,270} The tumorigenicity of human iPSCs (and ESCs) was thoroughly reviewed by Ben-David and Benvenisty.²⁷¹ As an alternative to iPSCs in which the 39 40 Yamanaka factors are transduced, exogenous gene-free iPSCs can be used.²⁷² Another option 41 is to additionally engineer iPSC-derived cells to express suicide genes to eradicate the cells, an 42 approach that was successfully used in ESCs and BMSCs.^{273,274}

43 It should be noted that in addition to various MSC sources and iPSCs, encouraging results 44 have been achieved by using BMMNCs in animal models of ischemic stroke where this subset 45 of cells was found to stimulate endogenous angiogenesis^{61,62} and neurogenesis by improving the NPC-vascular niche⁶³ or modulate the immune system.⁹⁵ Moreover, it was shown that 46 47 after ischemic stroke, the amount of CD34⁺ blood cells that migrate from the bone marrow 48 to peripheral blood is increased, which is associated with a better clinical outcome and is be-49 lieved to be mediated by granulocyte colony-stimulating factor (G-CSF).^{275,276} In addition to 50 its effect on BMMNC recruitment, this factor has previously been shown to possess neuropro-51 tective and neuroregenerative effects²⁷⁷ and is also believed to mobilize BMSCs and possess

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2 immunomodulatory properties.²⁷⁸ BMMNCs can also be quickly isolated from peripheral 3 blood by Ficoll-Paque density gradient centrifugation just before administration, circumvent-4 ing the additional cell culture period, which is required for MSC- or iPSC-based therapies.⁶³ 5 As was a hurdle with MSC-based therapy, also the administration route of BMMNCs remains 6 a topic of debate. Although study by Kamiya et al.⁹⁶ showed superior results after IA-delivered 7 BMMNCs over IV-delivered cells, this was contradicted by Yang et al. who showed that IA 8 delivery was not superior to IV-delivered BMMNCs.²⁷⁹ Subsequently, these cells have been 0 used in several clinical trials. IA administration of these cells was safe and showed an improved 10 clinical outcome.^{280,281} IV delivery of these cells also appeared safe,²⁸² although no clinical 11 improvement was observed in the study by Prasad et al.,²⁸³ whereas Savitz et al. were able to show an improved functional outcome.²⁸² Despite these encouraging results, in depth in vitro 12 13 evidence on the underlying mechanisms of action is largely unknown. The scarce in vitro data 14 on the effect of BMMNCs on regenerative processes showed that BMMNCs exerted protective 15 effects on rat hippocampal brain slices subjected to oxygen and glucose deprivation⁹⁸ and that 16 the BMMNC secretome induced neuronal differentiation of SH-SY5Y neuroblastoma cells.⁹⁷ 17 Nonetheless, additional in vitro data supporting the mechanism of action of BMMNC-based 18 therapy for ischemic stroke are required.

19 As stroke is a disease that mainly affects the elderly,⁷ it is important to take into account the 20 effect of the aged microenvironment, age-related comorbidities, and the aged immune system 21 on the outcome of stem cell based therapies.^{230,284,285} Although it does not appear from clinical 22 studies that the aged brain microenvironment is detrimental for stem cell based therapies, dif-23 ferences exist between the young and aged brain. For example, the formation of the glial scar is 24 accelerated after stroke that hinders functional repair in aged rats.²⁸⁶ Moreover, comorbidities 25 such as hypertension, hyperlipidemia, and diabetes mellitus appear to play a role in age-related 26 stroke severity.^{287,288} As described previously, angiogenesis is a key concept in establishing brain 27 repair. Buga et al. compared the transcriptome and immunochemistry of young and aged stroke 28 rats and poststroke patients. Remarkably, although the upregulation of proangiogenic genes 29 associated with processes such as vessel sprouting, tube formation, and maturation was delayed 30 in aged rats, angiogenesis in the aged brains was similar to their younger counterparts. In ad-31 dition, an upregulation of proinflammatory and scar-promoting genes was found in the aged 32 rats compared to the younger brains, supporting the accelerated scar formation and increased neuroinflammation in aged stroke subjects.²⁸⁹ Of the studies listed in Table I, two studies by 33 34 Shen et al.^{146,147} and studies by Taguchi et al.,¹⁵² Balseanu et al.,¹⁴⁹ and Zhang et al.¹⁴⁸ used 35 aged rats to perform MSC transplantation studies in ischemic stroke. Although these studies 36 did not directly compare the outcome of their transplantation study between young and aged 37 rats, several encouraging results were found in these studies. These included, but are not limited to, enhanced functional recovery,^{146, 148, 149, 152} a reduction of the glial scar thickness,^{146, 147} and improved angiogenesis.^{148, 149, 152} Remarkably, Balseanu et al. who used a G-CSF treatment, 38 39 40 which is believed to possess multiple regenerative effects,²⁷⁸ or a combination of G-CSF and 41 a single BM-MSCs dose to improve the functional outcome in aged stroke animals, observed 42 that the functional improvement was not increased in this combination treatment.¹⁴⁹ Simi-43 larly, Buga et al. who used a G-CSF and a combined G-CSF-BMMNC therapy to improve 44 the functional outcome after transplantation in aged stroke animals were also unable to ob-45 serve an increased functional improvement by using the combination treatment.²⁹⁰ The use 46 of iPSCs in aged rats was described by Tatarishivili et al. who showed that almost 50% of 47 the engrafted iPSC-derived long-term expendable neuroepithelial cells survived 8 weeks post-48 transplantation and caused functional improvement in the aged rats.¹⁷⁰ Moreover, these cells 49 differentiated toward neuroblast-like cells, compared to the BM-MSCs described in Refs. 146 50 and 147 where the few surviving cells predominantly differentiated toward astrocytes. These stud-51 ies support the use of aged animals for in vivo stroke studies in which they were able to observe Q8

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cell-mediated improvements in brain regeneration and functional recovery. Nonetheless, a direct comparison of stroke outcome and the molecular effect after cell transplantation between young and aged animals for the various age-related differences in (brain) microenvironment could provide additional information on which processes are mainly responsible for stem cell mediated brain repair. For example, the previously mentioned study by Buga et al., which described the transcriptome in young and aged rats after stroke, provided several new target pathways that can provide additional insight into age-related stroke pathology and therapeutic opportunities.²⁸⁹

The evaluation of the disease outcome in animal models for ischemic stroke is based on 11 behavioral testing, for which various tests can be applied to investigate specific aspects of neuronal recovery.¹²⁶ As stroke symptoms are largely dependent on the brain area involved, 12 13 different behavioral tests are applied to evaluate general, motor, sensorimotor, and cognitive 14 recovery and each test has its strengths and weaknesses. For example, the Bederson test is 15 easy to perform but reliable neurological ratings on this Bederson scale are limited because 16 of their subjective nature, a common feature of all behavioral tests that are based on human 17 observation. An overview with critical comments on the different behavioral tests that are often used in stroke research is provided by Schaar et al.¹²⁶ Nonetheless, functional assessment of 18 19 stroke outcome should include tests to cover all aspects of the disease outcome. 20

As indicated above, preclinical stroke research is facing several important issues that need to be resolved prior to more elaborate testing of the clinical potential of stem cell based therapies for ischemic stroke. Noninvasive imaging methods can aid in the longitudinal follow-up of stem cell fate and effect after transplantation via various administration routes. However, with the exception of a few studies,^{238,253} most of the studies that used noninvasive imaging to monitor the poststroke microenvironment or stem cell migratory pathways focused on the proof of principle of the imaging technique and did not link their results to functional recovery,^{206,208,236,246,256} which will most likely be the next step to be performed with these highly promising imaging modalities.

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6. CONCLUSION

Although multiple clinical advances have been made to improve the clinical diagnosis and outcome after acute ischemic stroke, beneficial long-term or delayed interventions are currently not available. Stem cell based therapies with MSCs and iPSCs have shown great promise in in vivo models of ischemic stroke through various administration routes. Nonetheless, the mechanisms of action of the transplanted cells remain poorly understood and are highly dependent on administration route, pretreatment, and full neuronal predifferentiation when using iPSCs. Although postmortem cell tracking methods provide detailed spatial information on the donor cell fate and host microenvironment, they are unable to deliver dynamic information on these subjects. A longitudinal follow-up with noninvasive imaging methods allows donor cell fate and changes in host microenvironment to be linked with behavioral and functional improvements, which can lead to additional insight into the mechanisms responsible for functional recovery in stroke after donor cell transplantation.

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CON	NFLICT OF INTEREST	
The	authors declare that they have no conflict of interest.	
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