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Mesenchymal and neural stem cells labeled with HEDP-coated SPIO nanoparticles: in vitro characterization & migration potential in rat brain

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ABSTRACT

Mesenchymal stem cells (MSC) may transdifferentiate into neural cells in vitro under the influence of matrix molecules and growth factors present in neurogenic niches. However, further experiments on the behavior of such stem cells remain to be done in vivo. In this study, rat MSC (rMSC) have been grafted in a neurogenic environment of the rat brain, the subventricular zone (SVZ), in order to detect and follow their migration using superparamagnetic iron oxide (SPIO) nanoparticles. We sought to characterize the potential effect of iron loading on the behavior of rMSC as well as to address the potential of rMSC to migrate when exposed to the adequate brain microenvironment. 1-hydroxyethylidene-1.1-bisphosphonic acid (HEDP)-coated SPIO nanoparticles efficiently labeled rMSC without significant adverse effects on cell viability and on the in vitro differentiation potential. In opposition to iron-labeled rat neural stem cells (rNSC), used as a positive control, iron-labeled rMSC did not respond to the SVZ microenvironment in vivo and did not migrate, unless a mechanical lesion of the olfactory bulb was performed. This confirmed the known potential of iron-labeled rMSC to migrate toward lesions and, as far as we know, this is the first study describing such a long distance migration from the SVZ toward the olfactory bulb through the rostral migratory stream (RMS).

Section: cellular and molecular biology of nervous system

Keywords: mesenchymal stem cells; SPIO; cell migration; subventricular zone; olfactory bulb; magnetic resonance imaging

Abbreviation list: BrdU: bromodeoxyuridine; BSA: bovine serum albumin; DMEM: Dulbecco’s modified Eagle’s medium; DPBS: Dulbecco’s phosphate buffered saline; EDX: energy dispersive X-ray; FBS: foetal bovine serum; HEDP: 1-hydroxyethylidene-1.1-bisphosphonic acid; HG: high glucose; IHC: immunohistochemistry; LG: low glucose; MIAMI: marrow isolated adult multilineage inducible; MRI:
magnetic resonance imaging; MSC: mesenchymal stem cell; OB: olfactory bulb; PB: Prussian blue; PFA: paraformaldehyde; RMS: rostral migratory stream; rMSC: rat mesenchymal stem cell; rNSC: rat neural stem cell; RT: room temperature; SEM: scanning electron microscopy; SPIO: superparamagnetic iron oxide; SVZ: subventricular zone; TEM: transmission electron microscopy
1. INTRODUCTION

Stem cells are often described as the best candidates for cell therapy studies due to their self-renewal capacity and their large differentiation potential. Among them, mesenchymal stem cells (MSC) remain easy to isolate and expand. They may exhibit immunodepressive characteristics which make them less sensitive to rejection by the host immune system [25,30,34]. Moreover, MSC allow autologous grafts to be performed in cell therapy protocols. Bone marrow MSC typically differentiate into connective tissue cell types [12,21], but various laboratories have also reported the transdifferentiation potential of MSC into a neuronal-like phenotype [5,39,46]. We previously showed that a subpopulation of human MSC, marrow-isolated adult multilineage inducible (MIAMI) cells may transdifferentiate in vitro in a neurotrophin-dependent manner into neuronal-like cells. These cells express neuronal markers and present electrophysiological characteristics similar to those observed in mature neurons [44]. Moreover, a fraction of MSC transplanted in adult rat brains may respond to microenvironmental cues and transdifferentiate into neuronal-like cells [20,23,52]. Using different brain lesion models, it has been shown that implanted MSC may be involved in functional improvement, either directly or indirectly by their ability to produce various growth factors [9,27,28,51]. In addition, a damaged environment resulting e.g. from ischemia or from the presence of a tumor is known to stimulate the migration of transplanted MSC [20,28,41] as well as of neuronal precursors [1,22]. MSC may thus be considered as potential candidates for cell therapy studies in the central nervous system. However, the possible use of MSC for brain repair studies still requires an evaluation of their behavior, migratory dynamic and fate in vivo. Moreover, as only a fraction of the transplanted cells may respond to the stimuli of the microenvironment, highly sensitive procedures will be required in the future.

Magnetic resonance imaging (MRI) is a non-invasive tool that has demonstrated a high sensitivity for cell tracking after systemic or in situ injection of cells having incorporated magnetic tags. Indeed, using phagocytic cells, the detection of a single cell in mouse brain has been obtained with this technique [19]. This strategy spares laboratory animals and ultimately may be used for human stem cell
therapy studies. Toward this end, new magnetic tracers readily and specifically taken up by stem cells need to be formulated. Superparamagnetic iron oxide (SPIO) nanoparticles stand as promising tools to label and track various cell types in vivo by MRI [6,31,43]. We developed SPIO nanoparticles coated with 1-hydroxyethylidene-1.1-bisphosphonic acid (HEDP) [38], which are interesting due to their possible functionalization allowing the targeting and uptake of a specific cell type. In order to be able to translate this tool into the clinic, the innocuousness and the non-interference of these nanoparticles with the response of stem cells to their microenvironment have to be demonstrated.

In this study, we sought to characterize the potential effects of labeling rat MSC (rMSC) with these HEDP-coated SPIO nanoparticles on their viability and functions in vitro. Toward this end, rMSC iron uptake was characterized in vitro by Prussian blue (PB) staining and MRI. Iron-labeled rMSC viability and ultrastructure were also studied, as well as their osteogenic and neuronal differentiation potentials. Furthermore, we assessed the migratory potential of these iron-labeled MSC in vivo in response to the brain neurogenic stimuli. Indeed, it has been shown that neural stem cells (NSC) migrated from the subventricular zone (SVZ) to the olfactory bulb (OB), via the rostral migratory stream (RMS), where they differentiate into post-mitotic interneurons [11]. Therefore, we studied the ability of iron-labeled rMSC to migrate in a similar fashion than iron-labeled rat neural stem cells (rNSC), when transplanted into the SVZ of the lesioned or non-lesioned rat brain. This study was performed by bromodeoxyuridine (BrdU) immunohistochemistry (IHC) and PB staining. Finally, a double staining with PB/CD11b, specific for macrophage/microglia, was used to confirm the fate of the grafted cells and of the SPIO nanoparticles.
2. RESULTS

Iron uptake and cell characterization

Prussian blue staining, after incubation of the rMSC with the iron-oxide nanoparticles, demonstrated that the intensity and the percentage of labeled cells increased with the iron concentration used. Forty eight hours incubation with an amount of nanoparticles corresponding to 25 µg iron/mL resulted in ca. 10% of Prussian blue positive cells, 50 µg/mL efficiently labeled more than 90% of the cells whereas 100 µg iron/mL did not increase the percentage of positive cells compared to 50 µg iron/mL (fig. 1A, B, C respectively). For rNSC, PB staining revealed that 24 hours of incubation with 50 µg/mL iron gave the most efficient labeling (fig. 1D). rNSC were expanded as floating clusters called neurospheres but incubation in the iron containing medium was performed on adherent progenitors as the 3D structure of the neurospheres prevented an efficient labeling of all the cells.

Spectroscopic iron titration confirmed that the iron content per rMSC varied with the nanoparticle concentration used during labeling. Over the concentration range used (from 25 to 100 µg iron/mL), the intracellular concentration increased from ca. 3.5 pg/cell to 10 pg/cell (table 1). The differences in iron content per cell were also observed using in vitro MRI. Indeed, fig. 1E shows typical T2*-weighted images of an agarose gel containing iron-labeled rMSC. As the iron concentration increased in the incubation media, the T2* effects induced by the cells increased. Thus, cells incubated with 50 µg iron/mL induced a global T2* reduction of ca. 50% compared to unlabeled cells.

Trypan blue counting and MTS assay showed that cell viability was not affected one day post-incubation when using 25 or 50 µg iron/mL, whereas it decreased to 70% of control with 100 µg iron/mL. Consequently, 50 µg iron/mL incubated for 48 hours was chosen as the best trade-off in terms of iron content, percentage of marked cells and cell viability. It resulted in an average concentration of 5.6 ± 1.6 pg iron/cell as determined by spectroscopic iron titration (table 1). It should be noted that the facilitating Fugene® reagent was tested but did not significantly enhance iron uptake (data not shown). It was consequently not used in this study.
Under these conditions, iron-labeled cells did not exhibit morphological differences with unlabeled cells (fig. 2A, B). Moreover, transmission electron microscopy (TEM) demonstrated that iron-labeling did not affect rMSC ultrastructure and that the SPIO nanoparticles were taken into the cells rather than adhering to the exterior of the membrane. They were located in endo-lysosomal vesicles with a mean diameter of 355 nm (fig. 2C). Observations by scanning electron microscopy (SEM) confirmed the intracytoplasmic and perinuclear distribution of the nanoparticles. The small white dots observed on back-scattered SEM images (fig. 2D) were correlated to a difference in molecular composition and an energy dispersive x-ray (EDX) spectrum confirmed their iron nature (fig. 2E).

To confirm that iron-labeled rMSC maintain their function in vitro, differentiation studies were performed. Iron-labeled rMSC were still able to undergo an osteogenic differentiation. Moreover, a similar proportion of cells responded to this induction when compared to the control, unlabeled cells, as shown by Alizarin red staining of hydroxyapatite associated calcium mineral deposits (fig. 3A, B). In addition, rMSC developed features of neuronal cells in response to published protocols [49]. Most importantly, there was no significant difference between unlabeled and iron-labeled rMSC responses to the neuronal induction in vitro. Flat, uninduced rMSC expressed low levels of nestin and β3-tubulin (table 2). When exposed to the neuronal induction medium, some cells detached but most of them exhibited a narrower morphology that evolved towards a neuronal-like morphology, ranging from simple bipolar to branched cells (fig. 3C to H). Moreover, rMSC with a neuronal-like morphology acquired the expression of the neural-related markers nestin and β3-tubulin. Up to 50 % and 65 % of rMSC were positive for nestin and β3-tubulin, respectively, 24 hours post-induction. β3-tubulin expression remained stable 48 hours post-induction whereas nestin expression decreased to 5-35 % positive cells (table 2). The neuronal marker NeuN was already expressed in expanding cells and at a similar level at every step of the neuronal induction (fig. 3G, H).

Labeling for histochemical detection
Different in vitro labeling techniques (Hoechst 33342, PKH26, GFP transduction and BrdU) were tested to co-localize the iron-labeled cells after transplantation and determine their fate in vivo. PB staining of the labeled cells in vitro demonstrated that none of the methods used significantly interfered with iron uptake. However, some of these techniques were not completely satisfactory. Co-culture experiments with GFP-positive rMSC and Hoechst 33342-positive rMSC demonstrated that Hoechst fluorescent dye diffused into GFP-positive rMSC. PKH26 was readily absorbed by the rMSC and did not diffuse to the surrounding cells but the correct visualization of the cells was impaired in vivo due to a weak fluorescent signal. Lentiviral transduction yield of GFP was around 80% but resulted in a decreased cell proliferation rate, as well as a lower response of the cells to neuronal induction in vitro (data not shown).

Cell detection by PB staining and BrdU IHC was the most suitable labeling strategy, in our hands, to study the in vivo distribution of iron-labeled rMSC in rat brains. A minimum incubation time of 72 hours was used to label 90% of the iron-labeled rMSC in vitro with BrdU (fig. 4A). Indeed, an incubation time of 48 hours resulted in only 40% of BrdU positive rMSC (fig. 4B). The rapidly dividing iron-labeled rNSC were efficiently labeled with BrdU after only 6 hours incubation (fig. 4C). These incubation times were then used to load the cells with BrdU prior to transplantation.

**In vivo detection of rNSC and rMSC**

In the non-lesioned brain model, rMSC grafted into the SVZ did not respond to the stimuli of the neurogenic microenvironment and were only found in the grafting site (fig. 5A). Preliminary results indicated that, 5 days after grafting, rMSC close to the injection site stained positive for nestin as well as slightly for β3-tubulin (data not shown). We demonstrated that a mechanical lesion of the OB strongly induced the migration of rMSC to the OB through the RMS. This migratory behavior was confirmed by the visualization of the cells on adjacent brain slices both by BrdU IHC and PB staining 4, 14 and 21 days after transplantation, with a fainter staining after 21 days. The pathway followed by the cells from
the injection site to the OB was approximately 7 mm long. Time after grafting was a critical parameter as only a few migrating cells were observed 4 days after transplantation whereas the highest number was observed after 14 days. A fraction of cells were observed near the injection tract, but also all along the RMS and many of them reached the OB and were dispersed in this region (fig. 5 & table 3). Prussian blue staining performed on control rats was negative; therefore confirming that PB staining in the grafting site as well as in the lesioned OB was specific for iron-labeled cells and did not result from endogenous iron deposit. No strong inflammatory reaction was observed with CD11b staining, a specific marker for macrophage/microglia. Moreover, PB/CD11b double staining revealed a co-localization of this marker with only a few PB-positive cells, but not with the entire PB-positive cell population (fig. 5K, L).

However, even though PB staining revealed a large number of iron-labeled rMSC in the lesioned OB region after the mechanical lesion (fig. 5D), this pool of migrating cells was not accessible to MRI as the lesion itself was responsible for a large hyposignal and labeled cells were only efficiently visualized at the grafting site (fig. 6A, B).

In non-lesioned rat brains, rNSC were found in the grafting site, in the RMS and in the OB 14 days after transplantation (fig. 5F, G & table 3). Thus, iron-labeled rNSC migrated in a similar way than endogeneous rNSC in the absence of lesion.
3. DISCUSSION

Potential adverse effects of iron nanoparticles is a topic currently under investigation [35], and the means to diminish this side effect still remain to be explored. Many authors described the use of facilitating agents [2] such as lipofectamin [16], conjugation with Tat peptide [26] or with internalizing antibodies [7] to increase the uptake efficiency while diminishing cell toxicity. However, these methods are also currently under debate [4] and transfecting agents may not be approved for clinical use [29]. In our study, we demonstrated an efficient and non toxic in vitro uptake of native HEDP-coated SPIO nanoparticles by rMSC without using any transfecting agent. In addition, the bisphosphonate (HEDP) coating offers the possibility to functionalize the nanoparticles, therefore increasing the range of potential applications.

Forty-eight hours incubation with 50 µg iron/mL of these SPIO nanoparticles was the best trade-off in terms of labeling efficiency and cell viability. Under the conditions used, more than 90 % of the rMSC contained enough iron allowing their detection with PB staining; amount that is in the same range than what is usually described e.g. with D-mannose-coated nanoparticles where 80 % of the cells were labeled [43]. An efficient uptake is indeed required as a simple adhesion on the cell surface could be detrimental for the cell response to the microenvironment in vivo [43]. In this regard, we confirmed the endocytosis of the iron nanoparticles by SEM and TEM studies. In our study, the average iron content per cell (5.6 ± 1.6 pg) was low compared to what is described with polycation-bound SPIO nanoparticles (38 pg iron/cell) [43], but in the same range to what is obtained when labeling human MSC with the ferucarbotran Resovist® [29].

In addition to an absence of acute toxicity, the HEDP-coated nanoparticles had no significant side effects on rMSC behavior in vitro. The ultrastructure was conserved and their differentiation potential toward the osteogenic as well as neuronal lineages did not exhibit significant differences with unlabeled rMSC. MSC differentiation toward a neuronal-like lineage is now well-documented [5,39,44,46] and the fraction of differentiating rMSC (50 to 65 %) confirmed results published with this
protocol [49]. Interestingly, NeuN was always detected in the rMSC used in this study. In this regard, the expression of neuronal markers in expanding, uninduced MSC has already been described [45]. These results complement previous studies which described a conserved adipogenic and osteogenic differentiation potential [3,14] even if chondrogenic differentiation was sometimes impaired with Feridex® nanoparticles [24]. Altogether, these results confirmed the possible use of the HEDP-coated nanoparticles for cell tracking as it does not seem to interfere with the in vitro rMSC biology. Moreover, trials made to label a subpopulation of human MSC, the MIAMI cells [12] showed that they readily absorbed HEDP-coated iron nanoparticles, even with a higher efficiency than rMSC (data not shown). As MIAMI cells present a large differentiation potential, including the ectodermal lineage, this may render possible their future use for MRI tracking in human clinical cell therapy applications [44].

In vivo, a double staining PB/CD11b (specific for macrophage/microglia) demonstrated that CD11b-positive cells did not, for the most part, co-localize with PB staining and therefore gave the proof that uptake by the host immune system is limited. Similar results with PB/ED1 double staining were obtained with rMSC containing dextran-coated Endorem® nanoparticles in a spinal cord injury model [41]. To our knowledge, this phenomenon has not been previously described in other cell types, suggesting that this event may be cell dependent. Therefore, the fate of iron nanoparticles will have to be addressed in more details in future studies using MSC. On the other hand, a possible transfer of BrdU to the host cells 3 weeks after transplantation has been described [8], but as BrdU labeling co-localized nicely with PB in our study, it suggests that the majority of detected cells were iron-containing rMSC and not host cells.

Iron-labeled rMSC did not respond to the microenvironmental stimuli when stereotactically implanted into the SVZ and did not migrate toward the OB via the RMS. However, when transplanted in the SVZ of a rat brain presenting a mechanical injury of the OB, the iron-labeled rMSC extensively migrated from the grafting site to the OB through the RMS, therefore covering a distance of approximately 7 mm. Indeed, rMSC were easily detected by BrdU IHC and PB staining all over the
migratory pathway 14 days post-transplantation when a sufficient amount of cells (80x10^3) was injected. The fainter staining observed by IHC after 21 days may result from a possible cell proliferation, and therefore from an iron dilution, as was described for MSC transplanted in a brain infarct model 3 months after transplantation [50]. Using MRI, the fraction of rMSC migrating toward the OB was not detected as the mechanical lesion induced a very important background hyposignal. Therefore, precautions will have to be taken in studies seeking to visualize iron-labeled cells in similar lesioned environments. It is of importance to note that we did not observe any adverse effect on the behaviour of the animals even 21 days post-transplantation. Moreover, no signs of toxicity were detected by use of current histological staining (hematoxilin/eosin), therefore confirming the safety of SPIO nanoparticles for in vivo applications [33].

Taken together, these in vivo results demonstrate that rMSC were not able to migrate in a similar fashion than rNSC when transplanted into the SVZ of a normal brain, but did confirm their known potential to be attracted by lesions, even after iron-loading [20,42,50]. It underlines the importance of the microenvironment on rMSC migratory behavior. Indeed, without lesion, in opposition to rMSC, neurosphere-derived iron-labeled rNSC, used as a positive control for cell migration, did respond to the SVZ niche stimuli and demonstrate an important migratory potential toward the OB through the RMS. This confirmed previous studies using labeled rat NSC [10], unlabeled NSC [13,17] as well as labeled human NSC [18]. The migratory response of MSC to appropriate adhesion or inflammatory molecules and to chemokines present in damaged tissues is now well documented [36,40]. Indeed, many chemoattractive molecules present in a lesioned environment, including for example the monocyte chemoattractant protein-1 (MCP-1), the macrophage inflammatory protein 1α (MIP-1α) and IL-8 [47], are known to promote cell migration via an interaction with MSC cell surface receptors. Therefore, we assumed it was the underlying mechanism leading the migration of rMSC observed in the presence of an OB lesion. However, the migration in a normal adult brain of a MSC subpopulation expressing receptors known to regulate neural cell activity and migration in brain has been reported [37]. However, MSC are
a heterogenous population so these receptors, including Neo1, Nrp2 and Robo1 & 4, may not be present on the rMSC used in this study as we did not observe any migration in an undamaged brain. Similarly to Phinney et al. [37], we may hypothesize that MSC migrating in an inflammatory environment are a different subpopulation from those that migrate in response to guidance cues modulating neural cell migration.

As a conclusion, HEDP-coated SPIO nanoparticles were used to efficiently label rMSC without impairing cell viability and structure, as well as their differentiation potentials toward the osteogenic and neuronal lineages. In vivo, iron-labeled rMSC migration toward the OB was not induced by the SVZ microenvironment. However, their well-known capacity to migrate toward lesions was confirmed as they extensively migrated toward a mechanically injured OB. Moreover, a double staining for macrophage/microglia and iron-containing cells confirmed that the majority of detected cells were not host cells. To the best of our knowledge, this was the first time that a long distance migration from the SVZ toward a lesioned OB through the RMS was described for iron-labeled rMSC. This conserved ability to migrate in vivo when stimulated by a lesion, in addition to their differentiation potential, underlines once again the possible benefits of using mesenchymal stem cells for brain cell therapy.
4. EXPERIMENTAL PROCEDURE

All animal experiments were conducted in accordance with the “Direction des Services Vétérinaires”, the “Ministère de l’Agriculture” of France and with the European Communities Council Directive of 24 November 1986 (86/609/EEC).

Isolation and culture of MSC

To obtain rMSC, tibias and femurs of 250 g female Sprague Dawley rats (Charles River, l’Arbresle, France) were dissected. Bone marrow was harvested by flushing the medullar cavities using a 26-gauge needle with 4 ml of Dulbecco’s modified Eagle’s medium high glucose (DMEM-HG) (Lonza, Levallois-Perret, France) containing 2 mM L-glutamine and antibiotics (100 U/mL penicillin, 0.1 mg/mL streptomycin, 0.25 µg/mL amphotericin B, Sigma, St Quentin Fallavier, France). Cells were plated at a density of 2x10^6 cells/cm² in DMEM low glucose (DMEM-LG) (Lonza) containing 20% foetal bovine serum (FBS) (Lonza) and antibiotics. All cell cultures were performed at 37°C in a humidified 5% CO₂ atmosphere. After 24 h, MSC were selected by removing the non-adherent cells. Fourteen days later, adherent cells were detached with 0.25% trypsin-EDTA (Sigma) and plated at 2000 cells/cm². Medium was changed twice a week and cells re-plated when at 70% confluency.

Isolation and culture of rNSC

Cortex from E14-15 Sprague Dawley rat embryos were dissected, cut into small pieces and finally triturated using a firepolished Pasteur pipette. Neurospheres were formed by culturing in 3v DMEM-HG, 1v Ham’s F12 (Lonza) supplemented with B27 (GIBCO, Cergy-Pontoise, France), 20 ng/ml EGF, 20 ng/ml bFGF (R&D Systems Europe, Lille, France), 5 µg/ml heparin (Sigma) and antibiotics. Cells were fed every 2 or 3 days and split every 5 days at 1:3 dilution. Large neurospheres were cut with micro-scissors. Adherent neural progenitors were cultured with the same media in poly-D-
lysine (Sigma) coated plates (100 µg/mL poly-D-lysine in Dulbecco's phosphate buffered saline, DPBS, Lonza).

**Synthesis of HEDP-coated SPIO nanoparticles**

SPIO nanoparticles were prepared and purified as previously described [38]. Briefly, bare iron cores were generated under agitation by the coprecipitation of an aqueous solution of 100 µL Fe₂Cl₂ (0.314 M) and 100 µL Fe₃Cl₃ (0.666 M) in 2.5 mL of a 1 M tetramethylammonium hydroxide solution (all from Sigma). After 3 minutes, the 4 nm bare iron particles were incubated for 20 min at 20°C with 48 µM 1-hydroxyethylidene-1.1-bisphosphonic acid (HEDP) (Sigma). The pH of the solution was lowered to 6.0 using 3 M tetraethylsulfamide (Sigma) to allow for HEDP adsorption on the bare cores. After incubation, the nanoparticles were purified on a 20 cm Biogel P6 column (BioRad, Marnes-la-Coquette, France) at room temperature (RT). This resulted in a solution of HEDP-coated SPIO nanoparticles with a hydrodynamic diameter of 20 nm and a final iron concentration of 0.8 g/L. Magnetic relaxivities of those nanoparticles at 7 T were: r₁=1 mM⁻¹s⁻¹ and r₂=50 mM⁻¹s⁻¹ [32].

**Nanoparticle uptake and cell characterization**

*Iron labeling.* Nanoparticles were added to the culture medium at a concentration of 25, 50 or 100 µg iron/mL for 24 or 48 hours after the rMSC reached 60 % confluency. Fugene® (Roche, Meylan, France) transfection agent was tested to decrease the incubation time with nanoparticles: 1 µL/mL Fugene® was added to DMEM containing nanoparticles for 30 min before incubation with the cells in culture.

*Cell viability.* It was estimated by Trypan blue (Sigma) exclusion and by an MTS assay kit (Promega, Charbonnières-les-Bains, France). Briefly, after washing the cells with DPBS and incubation with diluted (1:5) MTS reagent for 4 h, samples were read at 490 nm by use of a Multiskan ascent spectrophotometer (Thermo Fisher scientific, Cergy Pontoise, France).
**Prussian blue (PB) staining.** Cells were fixed in 4 % paraformaldehyde (PFA) (Sigma) for 15 min at 4°C, washed with DPBS and incubated 20 min with Pearl's reagent. Reagent was extemporaneously prepared: 1 % potassium ferrocyanide in 3 % HCl (Sigma), heated for 20 min at 60°C and filtered. Cells were counterstained with 1 % eosin, washed with 1 % acetic acid (Sigma) before dehydration and mounting (Eukitt) (Labonord, Templemars, France). Cells were examined under bright field using an Axioscop microscope (Carl Zeiss, Le Pecq, France), a CoolSnap ES camera (Photometrics, Tucson, Arizona) and Metavue analysis software (Roper Scientific, Evry, France).

**Quantitative intracellular iron content assessment.** The average iron content of rMSC was assessed after incubation using inductively coupled plasma optical emission spectroscopy. 5x10⁵ cells were incubated with 65 % (v/v) nitric acid for 2 h. Cell extracts were diluted in water and the total iron content was measured at 260 nm using a Jobin-Yvon 238 ICP-OES spectroscope (Ultrace Instruments SA, Longjumeau, France).

**TEM.** Iron intracellular localization was confirmed by TEM. Cells were fixed for 30 min in 2 % glutaraldehyde in DPBS, then washed with DPBS, post-fixed in 2 % osmium tetroxide in DPBS for 1 h, and washed again before dehydration with ethanol (Sigma). After exposure to propylene oxide (Sigma), samples were infiltrated in 100 % epon resin and polymerized in a 60°C oven for 48 h. Samples were sectioned using a diamond knife on an Ultracut-S ultramicrotome (Leica Microsystems, Rueil-Malmaison, France). 70 nm sections were picked up on copper grids, contrasted with uranyl acetate (Merck, Fontenay Sous Bois, France) and Pb nitrate (Merck) solutions and coated with carbon. Sections were examined with a Jeol 2011 transmission electron microscope at 140 kV (Jeol, Croissy sur Seine, France).

**SEM.** Cells were fixed with 2 % glutaraldehyde (Sigma), dehydrated with increasing concentrations of ethanol and desiccated with hexamethyldisilazane (Carl Roth, Lauterbourg, France). Samples were finally coated with carbon and examined with a Jeol 6301F scanning electron microscope.
at 11 kV for backscattered images. EDX analysis was performed to confirm the iron nature of the observed nanoparticles.

*In vitro MRI.* After incubation with the SPIO nanoparticles, cells were washed, trypsinized and centrifuged. Cells were then embedded in a low melting-point 0.5 % agarose type VII (Sigma) gel at a concentration of 1 cell/voxel. MRI experiments were performed on a Bruker Avance DRX 300 system (Bruker Biospin, Wissembourg, France) equipped with a 150 mm vertical super-widebore magnet operating at 7 T, an 84 mm i.d. shielded gradient set capable of 144 mT/m maximum gradient strength and a standard 64 mm diameter birdcage resonator, operating on a Paravision software platform (version 2.1.1, Bruker Biospin). A 3D spoiled gradient-echo sequence (TR=110 ms; FOV 30*30*15 mm; matrix 128*128*32, $\alpha=20^\circ$), modified to a multi-echo sequence (TE=5 ms, IE=5 ms, nEchos=6) was used for quantitative T2* mapping [15].

*Osteogenic differentiation.* rMSC were plated at a density of 5500 cells/cm$^2$ in DMEM-HG containing 10 % FBS, 10 mM $\beta$-glycerol-phosphate, 50 mM ascorbate-2-phosphate, 100 nM dexamethason and antibiotics (Sigma). Medium was changed twice a week during 21 days. Cells were washed 3 times with DPBS and fixed with ice-cold 70 ° ethanol for 1 hour at 4°C. Staining was performed by 10 min incubation with 40 mM Alizarin Red-S (Sigma), pH 4.2 at RT. Cells were finally washed with DPBS and samples examined using an Axiovert 40 CFL bright-field microscope (Carl Zeiss).

*Neuronal differentiation.* Neuronal induction was performed according to a previously described protocol [49]. Briefly, rMSC were plated at $10^3$ cells/cm$^2$ on 0.5 µg/cm$^2$ fibronectin (Sigma) coated coverslips in DMEM-HG containing 20 % FBS and antibiotics. Neuronal pre-induction medium, consisting of DMEM-HG containing 20 % FBS and 1 mM $\beta$-mercaptoethanol (Sigma), was added for 24 hours. Afterwards, cells were washed with DPBS and transferred to neuronal induction media composed of DMEM-HG, 2 % dimethylsulfoxide (DMSO) (Sigma) and 200 µM butylated
hydroxyanisole (Sigma). Uninduced cells and cells exposed to the induction medium for 24 hours and 48 hours were stained for nestin, β3-tubulin and NeuN to assess the extent of neuronal differentiation. After washing with DPBS, cells were fixed with 4 % PFA at 4°C for 15 min, permeabilized with 0.2 % Triton X-100 (Sigma) for 5 min and blocked with DPBS, 10 % normal goat serum (Sigma), 4 % bovine serum albumin (BSA) (Sigma) at RT for 45 min. After washing with DPBS, the cells were then incubated overnight at 4°C with 5 µg/mL anti-nestin (BD Biosciences, Le Pont De Claix, France), 3.8 µg IgG2b/mL anti-β3-Tubulin (Sigma) or 20 µg/mL anti-NeuN (Chemicon, St Quent en Yvelines, France) monoclonal mouse antibodies in DPBS, 4 % BSA, 0.2 % Triton X-100. After washing with DPBS, incubation with 2.5 µg/mL secondary biotinylated anti-mouse antibody (Ref BA2001, Abcys, Paris, France) in DPBS, 4 % BSA, 0.2 % Triton X-100 was performed for 1 hour at RT. Finally, after washing with DPBS, incubation with 20 µg/mL streptavidin FITC (Ref F0422, Dako, Trappes, France) in DPBS for 40 min at RT was made before mounting (Fluorescent mounting medium, Dako) and observation under fluorescence microscopy (Axioskop, Carl Zeiss). Isotype controls were performed.

**Labeling methods for histochemical detection**

*Hoechst 33342 labeling.* After washing, cells were resuspended in 1 ml DMEM-HG containing 10 mg/mL Hoechst (Sigma), left for 30 min at 37°C and finally washed again 3 times.

*PKH26 labeling.* Cells were washed and resuspended in 500 µl DMEM-HG before adding 1.5 µL PKH26 (Sigma) for 3 min. Excess label was washed away with FBS, then washed again with DPBS 4 times. Co-cultures of GFP+ cells and Hoechst or PKH26-labeled cells were performed for 24-72 hours in order to evaluate the diffusion of the fluorescent dye to neighbouring cells. Cells were finally washed with DPBS and samples examined by fluorescence microscopy.

*Lentiviral transduction of GFP.* TEFLYGA (Gabon ape) human cell line containing a simian retrovirus encoding GFP with a CMV promoter was used for virus production (kind gift from Dr JC.)
Infection was performed with 5 µg/ml Polybrene® (Sigma) and GFP\(^+\) cells were selected with 500 µg/ml Neomycin G418 (Sigma) for 7 to 10 days.

**BrdU labeling.** rMSC and rNSC were incubated with 1 µL/mL BrdU from a BrdU Immunofluorescence Assay Kit (Roche Diagnostics, Meylan, France) on a time scale ranging from 6 to 96 h. For the rapidly dividing rNSC, shorter incubation times of 6 and 24 hours were tested and the uptake efficiency was compared between floating neurospheres and adherent neural progenitors. BrdU uptake was assessed by immunofluorescent staining following the Immunofluorescence Assay Kit instructions. After BrdU labeling, rMSC and rNSC were seeded on fibronectin and poly-D-lysine-coated coverslips respectively, washed with DPBS and fixed with 4 % PFA in DPBS for 10 min at 4°C. After washing, cells were incubated with the primary anti-BrdU mouse antibody (1:20) (Clone BMG 6H8) for 30 min. The secondary anti-mouse fluorescein-conjugated antibody (1:20) was added for 30 min at 37°C. Mounting and observations were performed as described in the previous section.

**Cell transplantation**

A total number of 50 ten weeks old Sprague-Dawley female rats of 220 g were anesthetized with 16 mg/kg Xylasin (Rompun®, Bayer, Puteaux, France) and 80 mg/kg Ketamine (Clorketam 1000®, Vétoquinol, Lure, France). Prior to transplantation, rMSC were incubated for at least 72 hours with BrdU and the SPIO nanoparticles were added to the medium during the last 48 hours of incubation. rNSC (adherent neural progenitors) were incubated for 48 hours with the SPIO nanoparticles and BrdU was added during the last 6 hours of incubation. After labeling, 40x10\(^3\) to 100x10\(^3\) rMSC or rNSC were injected in a final volume of DMEM-HG ranging from 2 to 5 µL using a 10 µl microsyringe (Hamilton, Reno, Nevada). The injection was performed at a rate of 2 µL/min and the cannula was kept in situ for an additional 4 min before withdrawal. Control rats received an equivalent volume of DMEM-HG only or unlabeled cells. Injection coordinates were chosen in the SVZ, close to the beginning of the RMS: +0.9 mm rostral to Bregma, -1.2 mm lateral to the midline, -3.9 mm ventral to the dura and the tooth bar
was set at -3.3 mm [48]. For some rats, an OB lesion was performed just after grafting the cells, ipsilaterally to the injection site at 6.5 mm from Bregma, using a 3 mm diameter drill.

**In vivo immunohistochemical tracking**

Four, 14 and 21 days after transplantation, rats were anesthetized by CO$_2$ inhalation. 0.9 % NaCl and 4 % PFA were used sequentially for intracardial perfusion and brains were left in PFA containing increasing amounts of sucrose (from 2.5 to 20 %) (all from Sigma). Brain sections of 14 µm were made on a CM 3050S cryotome (Leica Microsystems). IHC was used in vivo for the detection of BrdU$^+$ cells and a mouse anti-CD11b antibody (specific for macrophage-microglia) was used to assess the inflammatory reaction. For BrdU staining, sections were washed with DPBS and incubated for 2 hours at 65°C in SSC buffer (1v sodium chloride-sodium citrate pH 7.4, 1v formamide), 30 min with 2 N HCl in DPBS at 37°C and finally 10 min in 0.1 M sodium borate, pH 8.5 at RT (all from Sigma). Quenching of peroxidases was made with 3 % H$_2$O$_2$ (Sigma) in DPBS at RT for 15 min. For BrdU and CD11b staining, blocking was performed overnight at 4°C or 40 min at RT, respectively. BrdU blocking buffer: DPBS, 3 % BSA, 0.1 % Tween 20 (Sigma). CD11b blocking buffer: DPBS, 10 % normal goat serum, 4 % BSA, 0.2 % Triton-X 100. Sections were incubated with the rat anti-BrdU antibody (1:100) (Ref OBT0030, clone BU1/75 ICR1) for 4 hours at RT and with the mouse anti-CD11b antibody (1:50) (Ref MCA275G, clone MRC OX-42) overnight at 4°C (both from Abd Serotec, Cergy Saint-Christophe, France). After washing, sections were incubated at RT for 2 hours with the secondary biotinylated anti-rat (1:200) (Ref BA4001) or for 1 hour with the anti-mouse antibodies (1:200) (Ref BA2001). BrdU incubation buffer: DPBS, 1 % BSA, 0.1 % Tween 20. CD11b incubation buffer: DPBS, 4 % BSA, 0.2 % Triton-X 100. Incubation with Vectastain® ABC reagent in 0.1 % Tween 20 in DPBS was made at RT (1.5 hours for BrdU and 1 hour for CD11b) (all from Abcys). Sections were washed and revealed with 0.03 % H$_2$O$_2$, 2 mg/mL diaminobenzidine (DAB) (Sigma) (supplemented with 0.16 % nickel chloride hexahydrate (Sigma) for BrdU), in DPBS for 10 min at RT and washed before mounting.
For PB/CD11b double staining, sections were fixed again with 2% PFA in DPBS for 5 min at 4°C after CD11b staining. Staining with Pearl’s reagent was then performed as previously described and samples were examined by bright field microscopy.

**In vivo MRI tracking**

*In vivo*, MRI was performed under the conditions defined for *in vitro* study. A TE = 10 ms was chosen as it allowed to depict microscopic susceptibility inhomogeneities induced by SPIOs without jeopardizing the overall image quality by macroscopic susceptibility artefacts. During the *in vivo* imaging process, animals were anaesthetized using a mixture of isoflurane/O₂ (1.5%, 2 L/min) and body temperature was maintained at 36.5-37.5°C using a feedback-regulated heating pad.
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LITERATURE REFERENCES


Fig. 1. In vitro Prussian blue assessment of iron uptake & MRI. The intensity and the percentage of PB positive cells (in blue) increased with the iron concentration used. rMSC incubated for 48 hours with 25, 50 & 100 µg/mL iron (A, B & C respectively). rNSC incubated for 24 hours with 50 µg/mL iron under adherent conditions (D). In vitro T2*-weighted images (TE = 15 ms) of an agarose gel containing rMSC incubated for 48 hours in culture medium containing HEDP-coated nanoparticles (E). From left to right, iron concentration in the media was 0, 25, 50 and 100 µg /ml.
Fig. 2. Iron-labeled rMSC microscopical analysis. Morphology of unlabeled rMSC (A) was identical to rMSC labeled with 50 µg iron/mL during 48 hours (B). TEM pictures showing the endo-lysosomal localization of the nanoparticles, iron in black (C). Back-scattered SEM picture (11 kV), iron in white (D). EDX spectrum confirming the iron nature of the white dots observed by SEM (E). Nu: nucleus.
Fig. 3. rMSC in vitro differentiation. When induced toward the osteogenic lineage, unlabeled rMSC (A) as well as iron-labeled rMSC (B) produced hydroxyapatite calcium mineral deposits that stained red with Alizarin. After treatment with the neuronal induction media, flat rMSC acquired a narrower, spindle shaped morphology and exhibited an increased expression of the neural-related markers nestin and β3-tubulin. The neuronal marker NeuN was already expressed in expanding cells and at every steps of the neuronal induction. Most importantly, unlabeled and iron-labeled rMSC responded similarly to the neuronal induction. Nestin, β3-tubulin and NeuN expression 48 hours post-induction was identical for unlabeled (C, E & G respectively) and iron-labeled rMSC (D, F & H respectively).
Fig. 4. BrdU immunofluorescent staining of iron-labeled rMSC and rNSC.

Ninety percent of rMSC were BrdU positive after 72 hours incubation (A) whereas only 40 % were positive after 48 hours (B) (white arrowheads point out BrdU negative cells). Six hours incubation under adherent conditions was sufficient to label rNSC (C).
Fig. 5. rMSC migratory potential assessment by BrdU immunochemistry and PB staining 14 days after transplantation. Schematic cell migration is presented on top right corner. Grafted rMSC were detected in the grafting site (A), in the RMS (B, C) and in the OB (D, E) after grafting into the SVZ. rNSC migrated from the SVZ to the OB through the RMS (F, G). In sham-operated rats, injected with medium only, no PB staining was detected in the grafting site and in the OB (H & I respectively). The BrdU isotypic control shows no background staining (J). In the grafting site, a fraction of Prussian blue positive cells co-localize with CD11b (K, black arrowheads), but not the entire rMSC population (K, white arrowheads). IgG2a isotype/Prussian blue staining shows no background (L). Cc: corpus callosum, Cr: cerebellum, Fmi: forceps minor corpus callosum, Lv: lateral ventricule.
Fig. 6. MRI visualization of grafted cells. A focal deposit of rMSC was clearly detected at the grafting site 14 days after transplantation on axial (A) and sagittal (B) planes (white arrowheads).