A dual gold nanoparticle system for mesenchymal stem cell tracking

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Stem cell-based therapies have demonstrated improved outcomes in preclinical and clinical trials for treating cardiovascular ischemic diseases. However, the contribution of stem cells to vascular repair is poorly understood. To elucidate these mechanisms, many have attempted to monitor stem cells following their delivery in vivo, but these studies have been limited by the fact that many contrast agents, including nanoparticles, are commonly passed on to non-stem cells. Specifically, cells of the reticuloendothelial system, such as macrophages, frequently endocytose free contrast agents, resulting in the monitoring of macrophages instead of the stem cell therapy. Here we demonstrate a dual gold nanoparticle system which is capable of monitoring both delivered stem cells and infiltrating macrophages using photoacoustic imaging. In vitro analysis confirmed preferential labeling of the two cell types with their respective nanoparticles and the maintenance of cell function following nanoparticle labeling. In addition, delivery of the system within a rat hind limb ischemia model demonstrated the ability to monitor stem cells and distinguish and quantify macrophage infiltration. These findings were confirmed by histology and mass spectrometry analysis. This work has important implications for cell tracking and monitoring cell-based therapies.

Introduction

Stem cell therapy shows great potential to treat a large variety of diseases, including cardiovascular diseases, which are the number one cause of death globally. In particular, bone-marrow derived mesenchymal stem cells (MSCs) are advantageous in that they possess angiogenic properties, are easily obtained in large numbers and expandable in culture, and are part of the ischemic response. Numerous preclinical and clinical trials have investigated the therapeutic benefits of stem cell therapy for cardiovascular diseases. However, advances in the field of stem cell therapy have been limited by the inability to track administered cells, which would provide information concerning cell engraftment and persistence, mechanisms of vascular regeneration, and the role of MSCs in vascular repair.

Conventional methods for assessing the biological mechanisms underlying disease states and potential therapies rely on postmortem histology, which only offers endpoint measurements and requires a large number of animals to be sacrificed in order to produce statistically significant results. A more ideal cell tracking method would involve using non-invasive longitudinal imaging to monitor cells. Towards this end, many contrast agents are currently being investigated to label cells for cell tracking purposes, including reporter genes, radionuclides, fluorescent probes, and nanoparticles. Nanoparticles, such as quantum dots, iron oxide nanoparticles, and plasmonic nanoparticles (gold and silver), offer many advantages over other contrast agents. For example, nanoparticles can be optimized to promote cellular uptake through shape, size, and surface coating modification and allow for long-term monitoring of cells. However, viable and non-viable cells cannot be distinguished using nanoparticle contrast agents. As a result, it is not possible to detect if a cell is dead and has been endocytosed by macrophages, leading to a transfer of contrast agent from the labeled cells to macrophages. Other investigators have found nanoparticle transfer to macrophages, resulting in the monitoring and tracking of macrophages instead of the stem cell therapy.

Thus, the goal of this study is to develop a nanoparticle system which is capable of tracking stem cells following delivery in vivo and also monitoring macrophage infiltration and transfer of contrast agents from stem cells to macrophages as a result of macrophage endocytosis. Macrophages are known to have key roles in wound healing and vascular regeneration and to be influenced by and exert paracrine effects on stem cells, including MSCs. The nanoparticle system will consist of gold plasmonic nanoparticles. Gold nanoparticles can be synthesized in various shapes and sizes and their absorption characteristics can be tuned to maximally absorb in the near-infrared region, where the absorption from endogenous tissue is the lowest. Gold nanoparticles are also non-toxic to cells in...
evaluate this nanoparticle system, various our lab. formation within the gels, as demonstrated by previous work in delivering within a 3D PEGylated and gold nanospheres to label macrophages. This system is nanoparticle system consisting of gold nanorods to label MSCs acoustic imaging. acoustic imaging experiments were performed to assess the nanoparticle labeling. In addition, photoacoustic imaging results. This study has important implications for cell tracking and the role of MSCs and photoacoustic imaging when they are endocytosed by macrophages and gold nanospheres will only be detected using photoacoustic light region (520 nm), but plasmon coupling following nano-particles maximally absorb in the near-infrared region. On the other hand, gold nanospheres maximally absorb in the visible light region (520 nm), but plasmon coupling following nanosphere endocytosis by cells leads to peak broadening. Thus, the gold nanospheres will only be detected using photoacoustic imaging when they are endocytosed by macrophages and imaged within the tissue optical window of 650–900 nm. To evaluate this nanoparticle system, various in vitro assays were performed, including labeling of cells with the nanoparticles and the assessment of cell function and viability following nanoparticle labeling. In addition, in vitro and in vivo photoacoustic imaging experiments were performed to assess the feasibility of monitoring the two cell types. Histological analysis and mass spectrometry were also performed to verify the in vivo photoacoustic imaging results. This study has important implications for cell tracking and the role of MSCs and macrophages in vascular regeneration.

Experimental section

Cell culture

Rat mesenchymal stem cells were isolated from the bone marrow of Lewis rats (200–300 g). The femoral marrow cavity was flushed and adherent cells were collected and cultured in Dulbecco’s Modified Eagle Media (DMEM) (Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum (FBS), 1% glutamax, and 1% penicillin–streptomycin. Non-adherent cells were removed after 24 hours by replacing the media. The media was then changed every two days and the cells were passaged once reaching 80% confluency. The cells were cultured under standard cell culture conditions (37 °C, 5% CO₂). Passage 3–7 cells were used in all studies.

Murine macrophages (J774A.1, ATCC) were cultured under standard cell culture conditions (37 °C, 5% CO₂) in DMEM with 10% FBS. The media was changed every two days and passaged once reaching 80% confluency. Passage 10–20 cells were used in all studies.

Nanoparticle synthesis

Spherical gold nanoparticle seeds (20 nm in diameter) were synthesized by heating 100 mL of distilled (DI) water to 100 °C and adding 1 mL of 10 mg mL⁻¹ gold(III) chloride hydrate (HAuCl₄). Then, 5 mL of 10 mg mL⁻¹ sodium citrate dissolved in DI water was added and the reaction was allowed to cool to room temperature. Gold nanoparticles 50 nm in diameter were synthesized by mixing 7.5 mL of 10 mg mL⁻¹ HAuCl₄, 15.61 mL of 0.2 M ammonium hydroxylamine, 750 mL of DI water, and 50 nm of seed solution while stirring. The surface of the gold nanoparticles was coated with methoxy-polyethylene glycol-thiol (mPEG–SH) at a concentration of 0.1 mM and allowed to react for 30 minutes on a shaker, followed by centrifugal filtration (Amicon ultra-15, Millipore) to remove residual PEG.

Gold nanorods (NRs) were synthesized via seed mediated growth as previously described. Briefly, the seed solution was synthesized by mixing 5 mL of cetyl trimethylammonium bromide (CTAB) (0.2 M) with 5 mL of H₃AuCl₄ (0.5 mM) while stirring vigorously. Then 0.6 mL of ice cold sodium borohydride (NaBH₄) was added to the solution. The growth solution was prepared by mixing 380 mL of 0.2 M CTAB, 8 mL of 0.01 M silver nitrate (AgNO₃), and 40 mL of 0.01 M H₃AuCl₄ while stirring at 30 °C. Then 4.4 mL of 0.1 M ascorbic acid (C₆H₈O₆) was added drop wise, resulting in a color change from yellow to colorless. To produce NRs, 0.958 mL of gold nanoseeds was added to the growth solution and allowed to stir for 3 minutes. The solution was allowed to age overnight, centrifuged twice at 14 000 rcf for 45 minutes, and resuspended in ultrafiltrated (18 M2 cm, Thermo Scientific Barnstead Diamond water purification systems) deionized water.

The NR surface was modified by replacing CTAB with mPEG–SH through ligand exchange. Briefly, an equal volume of mPEG–SH (0.2 mM) was added to the gold NR solution, sonicated for 1 minute, and allowed to react overnight. Excess PEG was removed by centrifugal filtration (Amicon ultra-15, Millipore) at 2000 rcf for 10 minutes. Finally, a layer of silica was deposited on the surface of the PEG modified NRs through a modified Stöber method, as previously described. Under vigorous stirring, 2 mL of PEGylated NRs was added to 3 mL of isopropanol, followed by 1.2 mL of tetraethyl orthosilica (TEOS) (3%) in isopropanol and 1.2 mL of ammonium hydroxide (8.4%) in isopropanol. The reaction was allowed to mix for 2 hours, followed by centrifugal filtration at 500 rcf for 15 minutes, twice. The NRs were then coated with a layer of poly-l-lysine (5 kDa; Sigma-Aldrich) at a concentration of 0.833 mM and allowed to
react for 2 hours, followed by centrifugation at 3000 rcf for 10 minutes and then 5000 rcf for 7 minutes.

The size of the nanoparticles was characterized using transmission electron microscope (TEM) analysis and dynamic light scattering (DLS), and the surface charge was characterized using zeta potential measurements. For TEM, an FEI Tecnai transmission electron microscope fitted with a top mount AMT Advantage HR 1k x 1k digital camera and operating at 80 kV was used. A drop of nanoparticle solution was placed on carbon coated copper 300 mesh grids (Electron Microscopy Sciences, Hatfield, Pa) which had been glow discharged. ImageJ analysis was used to quantify the size of the NRs and thickness of the silica coating \((n = 100)\). A DelsaNano (Beckman Coulter, Inc., Brea, CA) was used for DLS analysis of spherical nanoparticles. Nanoparticle solutions were loaded into a cuvette and readings were taken at a temperature of 25 °C. For size measurements, 5 repetitions were performed for each nanoparticle solution with 60 readings per repetition. A Zetavizer Nano ZS (Malvern) was used to measure the zeta potential of nanoparticle solutions. A total of 3 measurements, with 15 repetitions, were collected at a temperature of 25 °C.

### Nanoparticle labeling of cells

Gold nanosphere media was made following PEGylation of 50 nm nanospheres. The nanospheres were sterile filtered using a 0.22 µm filter, centrifuged in a centrifugal filter tube (Amicon ultra-15, Millipore) at 3000 rcf for 5 minutes, and resuspended in cell culture media at a concentration of approximately \(10^{12}\) NPs mL\(^{-1}\). Gold NR media was made following PLL coating of silica NRs. The NRs were sterile filtered using a 0.22 µm filter and centrifuged at 3000 rcf for 10 minutes. The supernatant was then centrifuged at 5000 rcf for 7 minutes. The NRs were resuspended in cell culture media at a concentration of approximately \(10^{12}\) NRs mL\(^{-1}\). Macrophages were incubated with nanosphere media (200 µL cm\(^{-2}\)) and MSCs were incubated with NR media (200 µL cm\(^{-2}\)) for 24 hours.

Bright field and dark field microscopy was used to qualitatively analyze nanoparticle uptake by macrophages and MSCs. Following nanoparticle incubation, the media was removed and the cells were washed with phosphate buffered saline (PBS) to remove excess particles. The cells were then incubated with DAPI (1 µg mL\(^{-1}\)) for 15 minutes, followed by PBS washes. A Leica DMI2000B microscope, equipped with a Leica DFC290 camera, was used to obtain bright field and dark field images. Controls consisted of cells not incubated with nanoparticles.

Inductively coupled plasma mass spectrometry (ICP-MS) was used to quantitatively analyze nanoparticle uptake by macrophages and MSCs in 2D, as well as by macrophages in 3D PEGylated fibrin gels. The PEGylated fibrin gels were synthesized as described below and contained 50 000 cells per mL and \(10^{12}\) NPs mL\(^{-1}\). Following nanoparticle loading in 2D, the cells were collected using trypsinization and counted. To collect the cells in 3D, the gels were digested by incubation with 500 µL of bovine pancreatic trypsin (5 mg mL\(^{-1}\) in 0.9% sodium chloride) per 1 mL of gel for 1 hour. The reaction was quenched with serum containing media and the solution was filtered with a 70 µm and 40 µm cell strainer and centrifuged at 600 rcf for 7 minutes to collect the cells. To prepare the samples for ICP-MS, all of the solutions were incubated in 300 µL of 70% nitric acid at 60 °C overnight. The solutions were then diluted to 3.7% nitric acid using ultrapure water prior to running the samples on the ICP-MS machine. The samples containing MSCs incubated with 50 nm gold nanospheres were prepared for ICP-MS analysis by heating the samples at 150 °C and bringing them to dryness. The samples were then resuspended in 200 µL of aqua regia and further diluted with 2% HCl. Nanoparticle loading was quantified using a standard curve. Standard solutions were made by diluting AAS gold standard (Sigma-Aldrich, St. Louis, MO) in the same background solution as the samples. The number of nanoparticles per cell was calculated based on the NP dimensions and cell number.

MSCs and macrophages were co-cultured in a 3D PEGylated fibrin gel to investigate the nanoparticle system in a 3D environment. PEGylated fibrin gels were synthesized as described below and contained 25 000 MSCs mL\(^{-1}\); 50 000 macrophages per mL; and \(10^{12}\) NPs mL\(^{-1}\). The gels were made in 12 well transwell plates with 8.0 µm pore sizes and were incubated at 37 °C in MSC growth media.

### Cell viability

Cell viability was assessed following nanoparticle labeling using LIVE/DEAD viability staining and an MTS proliferation assay. LIVE/DEAD staining was performed by incubating the cells with a working solution of calcein AM (4 µm) and ethidium homodimer-1 (1 µm) for 45 minutes at 37 °C. The cells were washed with PBS and imaged using fluorescence microscopy (Leica DMI2000B microscope equipped with a Leica DFC290 camera).

Cell proliferation following nanoparticle labeling was assessed using MTS. Cells were incubated with DMEM media containing 20% MTS solution for 4 hours at 37 °C. The supernatant was collected and the absorbance value measured at 490 nm \((n = 3)\). A blank sample containing no cells was subtracted from all measurements.

The maintenance of cell function in 3D PEGylated fibrin gels was assessed after laser irradiation by examining cell morphology and tubular network formation. Difunctional succinimidyl glutarate PEG (8 mg mL\(^{-1}\) in PBS without calcium; NOF America) was added to fibrinogen (80 mg mL\(^{-1}\) in PBS without calcium; Sigma) in a 1 : 1 volume ratio, and the reaction was allowed to take place at room temperature for 3–5 minutes. An equal volume of MSCs as the fibrinogen and PEG mixture was then added at a concentration of 200 000 cells per mL. Gels with either NR labeled MSCs or unlabeled MSCs were made. The reaction then underwent gelation by adding an equal volume solution of thrombin \((25 \text{ U mL}^{-1})\ in 40 \text{ mM CaCl}_2; \text{CalBiochem})\). The final concentrations in the gel were 10 mg mL\(^{-1}\) of fibrinogen; 1 mg mL\(^{-1}\) of SG-PEG-SG; 50 000 cells per mL; and 12.5 U mL\(^{-1}\) of thrombin. The gels were made in 4 well chambered coverglass slides (Laboratory-Tek II; Nalge Nunc) and incubated at 37 °C in MSC growth media.

The gels were irradiated at various time points (day 1, 4, and 7) using a 800 nm laser with a fluence of \(10 \text{ mJ cm}^{-2}\) for 100
pulses. Control conditions consisted of the same gel formulations without laser irradiation. At day 7, the cells were stained with calcine AM (10 μM) and 3D z-stacks were taken using a Leica SP2 AOBs confocal microscope (10× magnification; 512 × 512 resolution). The images covered approximately 1 mm with a step size of approximately 3 μm. The 3D images were reconstructed using ImageJ.

**In vitro photoacoustic imaging**

A tissue mimicking phantom with varying ratios of NR labeled MSCs and nanosphere labeled macrophages was prepared for combined ultrasound and photoacoustic (US/PA) imaging. The bottom layer of the phantom consisted of gelatin (8% by weight; Sigma-Aldrich) with 15 μm diameter silica particles (0.2% by weight; Sigma-Aldrich) for ultrasonic scattering. The cell inclusions (20 μm weight; Sigma-Aldrich) for ultrasonic scattering. The cell to deliver the laser pulses. RF signals were acquired for laser wavelengths ranging from 680–970 nm and with a laser fluence of 5–15 mJ cm⁻². The transducer was mechanically scanned along the tissue mimicking phantom to obtain 3D photoacoustic information. The signals exported from the imaging system were post-processed for beamforming, laser fluence compensation, image reconstruction, and signal quantification. For spectral analysis, the quantified photoacoustic signals were used to calculate the ratio of the NR labeled MSCs and the nanosphere labeled macrophages in each voxel by comparison to weighted sums of their optical absorption spectrum. Then, the estimated ratios in the voxels were averaged to calculate the overall cell ratio for whole inclusions.

**Ischemic injury and *in vitro* ultrasound and photoacoustic imaging**

Animal handling and care followed the recommendations of the National Institutes of Health (NIH) guidelines for the care and use of laboratory animals. All protocols were approved by the Animal Care Committee at the University of Texas at Austin. Lewis rats (11 weeks) weighing 250–300 g were used. A femoral artery ligation was performed in Lewis rats (11 weeks, male) to induce hind limb ischemia. Through a small incision on the medial side of the thigh, the femoral artery of a single hind limb was separated from the nearby nerve and vein and ligated immediately distal to the inferior epigastric artery and proximal to the branch point of the popliteal and saphenous arteries using Prolene 5-0 sutures. The ligated segment was then excised and the skin incision closed with interrupted sutures. The animal was allowed to recover overnight and the following day (about 24 hours later), MSCs were injected intramuscularly into the gastrocnemius muscle of the ligated limb. PEGylated fibrin injections were prepared by combining difunctional succinimidyl glutarate PEG (4 mg mL⁻¹ in PBS without calcium; NOF America) to fibrinogen (40 mg mL⁻¹ in PBS without calcium; Sigma) in a 1 : 1 volume ratio. An equal volume of MSCs labeled with gold NRs was mixed with the PEGylated fibrinogen solution in a 1 : 1 volume ratio at a concentration of 13 × 10⁶ cells per mL. Next, 50 nm PEGylated gold nanospheres was added to the cell/PEGylated fibrin solution at a concentration of 10¹⁴ NPs mL⁻¹. The solution was then loaded into a 23 G needle syringe, followed by an equal volume of thrombin (25 U mL⁻¹ in 40 mM CaCl₂). The solution was mixed thoroughly within the syringe and the gel solution (300 μL) was injected into the gastrocnemius of the rat. The final concentrations in the gel were 5 mg mL⁻¹ of fibrinogen; 0.5 mg mL⁻¹ of SG-PEG-SG; 3.33 × 10⁶ cells per mL; and 12.5 U mL⁻¹ of thrombin. Prior to delivery, MSCs were fluorescently labeled with CellTracker CM-DiI (Invitrogen). The cells were incubated with CM-DiI (15 μM) at 37 °C for 8 minutes and then 4 °C for 15 minutes, washed with PBS, and resuspended in DMEM.

Ultrasound and photoacoustic signals were captured using the Vevo LAZR (VisualSonics, Inc.) system. A 40 MHz array transducer incorporated with a fiber bundle was submerged in a water bath, and the rat hind limb was coupled through a transparent film at the bottom of the water bath. RF signals were acquired for laser wavelengths ranging from 680–970 nm and with a laser fluence of 5–10 mJ cm⁻². Post-processing was the same as that for the *in vitro* photoacoustic imaging. The processed photoacoustic and spectroscopic images were overlaid with ultrasound images using a user-defined threshold of the photoacoustic signals.

**Histology**

At the terminal endpoint of the study, animals were sacrificed and the gastrocnemius muscles were isolated, embedded in optimal cutting temperature (OCT) compound, submerged in liquid nitrogen-cooled isopentane, and stored at −80 °C until further processing. The samples were cut into 12 μm thick sections using a cryostat and placed onto positively charged microscope slides. The tissue sections were fixed in 10% formalin for 15 minutes. Immunohistochemical and histochemical staining were subsequently performed.

For immunostaining, the slides were incubated in 0.125% trypsin solution for 20 minutes at 37 °C. They were then permeabilized in 0.5% Triton X-100 in TBST for 30 minutes and blocked in 10% normal goat serum for 1 hour. The slides were incubated overnight at 4 °C in the primary antibody (1 : 50 dilution in 2.5% normal goat serum) (ED1; Millipore), followed by incubation with a fluorescence-conjugated secondary antibody (1 : 200 in 2.5% normal goat serum) (Alexa Fluor 488 Goat Anti-Mouse Immunoglobulin G; Life Technologies). As a negative control, normal immunoglobulin G was used instead of the primary antibody. The samples were counterstained with DAPI (5 μg mL⁻¹) for 15 minutes, mounted, and viewed using a fluorescence microscope (DM2000B; Leica).

TUNEL staining was performed with the APO-BrdU TUNEL Assay Kit (Invitrogen). The slides were incubated in 0.125% trypsin for 20 minutes at 37 °C, followed by permeabilization in 0.5% Triton X-100 for 30 minutes. The slides were then incubated with TdT and BrdU for 2 hours at 37 °C. Alexa Fluor 488
Dye-labeled anti-BrdU antibody was applied for 30 minutes. The slides were then counterstained with DAPI (5 μg mL⁻¹) for 15 minutes, mounted, and viewed using a fluorescence microscope (DMI2000B; Leica).

Fluorescence-activated cell sorting of macrophages

Seven days following intramuscular injection of nanorod labeled or unlabeled MSCs and nanospheres within PEGylated fibrin, animals were sacrificed and the gastrocnemius muscle was isolated. The muscle samples were finely minced and incubated in 1% collagenase in DMEM for 1 hour at 37 °C under constant agitation. Samples were then filtered through cell strainers (40 μm and 70 μm), washed with PBS, and blocked for 30 minutes in fluorescence-activated cell sorting (FACS) buffer (1% BSA in PBS + 1% sodium azide). Samples were then incubated with primary antibody (1:50; ED1; Millipore) in FACS buffer for 45 minutes, washed, and incubated in secondary antibody (1:50; Alexa Fluor 488 Goat Anti-Mouse Immunoglobulin G; Life Technologies) for 45 minutes. Next, samples were fixed in 4% neutral buffered formalin for 15 minutes, washed, and resuspended in PBS. Samples were then sorted for macrophages using a FACSAria flow cytometer. Following sorting, samples were digested for ICP-MS analysis. The cells were collected by centrifuging at 270 × g for 5 minutes and resuspended in a small volume of aqua regia solution. The samples were heated at 150 °C and brought to dryness. They were then resuspended in 250 μL of aqua regia and further diluted with ultrapure water to give an HCl concentration of 5%. A standard curve was used to quantify the amount of gold in the cells by diluting AAS gold standard (Sigma-Aldrich, St. Louis, MO) in the same background solution as that of the cell solutions. The data was quantified by calculating the mass of gold per macrophage.

Results

Nanoparticle characterization

The size, shape, and surface charge of gold NRs and nanospheres were characterized using ultraviolet-light spectroscopy (UV-vis), TEM analysis, DLS, and zeta potential measurements. The absorbance spectra of gold NRs had a peak around 750 nm, which red shifted 20 nm following silica coating and an additional 8 nm following poly(L-lysine) (PLL) coating (Fig. 2A). Zeta potential measurements confirmed the sequential coatings were deposited on the NRs (Fig. 2A), as the surface charge was +40 mV for CTAB coated NRs, −0.61 mV for PEG coated NRs, −30.37 mV for silica coated NRs, and +26 mV for PLL coated NRs. Using TEM analysis, the size of the NRs was measured as 55.63 ± 9.14 × 15.59 ± 3.45 nm, with a silica coating thickness of 25.04 ± 3.06 nm (Fig. 2A). The 50 nm spherical nanoparticles had an absorbance peak around 520 nm (Fig. 2B), with a red shift of a few nm following PEG coating (data not shown). TEM analysis confirmed the spherical shape of the particles and DLS measured the hydrodynamic diameter of the particles as being 50.4 nm with a polydispersity index of 0.044 (Fig. 2B). Zeta potential measurements demonstrated that the nanospheres had a negative surface charge (−46.07 mV), which became closer to neutral (−8.41 mV) following PEG coating (Fig. 2B).

Nanoparticle labeling of cells

MSCs were incubated with PLL silica NRs and the uptake by the cells was qualitatively analyzed using microscopy and quantitatively analyzed using ICP-MS. Dark field microscopy demonstrated nanoparticle labeling of MSCs as evident by the yellow/green color of the cells as compared to control cells (Fig. 3A and C). In addition, dark nanoparticle aggregates within cells were visible under bright field (Fig. 3B and D). ICP-MS analysis...

Fig. 2 Characterization of (A) gold nanorods and (B) nanospheres using ultraviolet-light spectroscopy, transmission electron microscopy, and zeta potential analysis.
quantiﬁed the number of nanorods per MSC as $1.89 \pm 0.877 \times 10^5$ nanorods per cell (Fig. 4A). This large extent of cellular uptake of the nanorods can be attributed in part to both the silica coating, providing a more spherical shape to the particle which is more conducive for cellular uptake,\textsuperscript{16,17,41} and the PLL surface coating, which is commonly used as a transfection agent and has been used to promote cellular uptake of iron oxide nanoparticles.\textsuperscript{42–44}

Macrophages were incubated with 50 nm PEGylated gold nanospheres and dark ﬁeld and bright ﬁeld microscopy demonstrated a large extent of uptake by the cells (Fig. 3G and H) as compared to control cells (Fig. 3E and F). In addition, MSCs were incubated with the same nanoparticles and there was no evident uptake by the cells (Fig. 3I and J), which was veriﬁed by ICP-MS measurements (Fig. 4B). The reason MSCs did not uptake these particles can be attributed to the PEG surface coating, which passivates the particles and gives them a neutral charge, leading to minimal detection and uptake by cells.\textsuperscript{41,45} Macrophages, on the other hand, are characterized by their high rate of nonspeciﬁc uptake, and thus endocytose these particles to a large extent.\textsuperscript{46} Over time the mPEG–SH layer is slowly displaced by serum proteins, leading to preferential uptake by macrophages.\textsuperscript{47,48} Thus, 50 nm PEGylated gold nanospheres can be used as a contrast agent to preferentially label macrophages and identify the in vivo inﬁltration of macrophages.

Cell viability and function following nanoparticle labeling

LIVE/DEAD staining and an MTS proliferation assay were used to assess cell viability following nanoparticle labeling. Nanoparticle cytotoxicity is dependent upon the size, shape, and surface coating of the nanoparticles. In particular, CTAB, a surfactant used during nanorod synthesis and which forms a bilayer on the surface of the nanorod, is especially cytotoxic to cells, and thus removing or displacing CTAB is necessary to reduce the cytotoxic effects.\textsuperscript{31,49,50} Following labeling of MSCs with silica coated NRs, MSCs were viable up to 5 days and continued to proliferate, with no significant reduction in cell proliferation compared to the control cells (Fig. 6A and B). Labeling of macrophages with 50 nm PEGylated gold
nanospheres did not affect cell proliferation or viability (Fig. 6C and D) compared to controls.

The maintenance of MSC tubular network formation within 3D PEGylated fibrin gels was assessed following nanorod labeling and laser irradiation. This was performed to verify the process of photoacoustic imaging did not affect the cells’ ability to form tubular networks. MSCs without NR labeling or laser irradiation formed extensive tubular networks within the gels, and this network formation was maintained following NR labeling (Fig. 7A and B). The gels were also irradiated with a 800 nm laser to ensure photoacoustic imaging did not affect cell function. Both non-labeled cells and NR labeled cells continued to form networks following laser irradiation (Fig. 7C and D). These results demonstrate that the cells maintain their function of forming tubular networks following NR labeling and photoacoustic imaging.

In vitro photoacoustic imaging of nanoparticle labeled cells

In vitro photoacoustic imaging of nanosphere labeled macrophages and NR labeled MSCs was performed to evaluate the feasibility of distinguishing and quantifying the two cell types using photoacoustic imaging. The two cell types were mixed in various ratios and Fig. 8 shows the spectroscopic results in which the 100% macrophage inclusion was identified as having 50 nm PEGylated nanospheres (red) and the 100% MSC inclusion was identified as having NRs (green). A transition from nanospheres to NRs is evident as the concentration of macrophages decreases and of MSCs increases. These results demonstrate that the two cell types can be distinguished based on labeling with different nanoparticles, even when they are in close proximity. In addition, the ratios of the two cell types were calculated based on the photoacoustic imaging results, with the predicted cell ratios being a very good estimate of the actual cell ratios in the inclusions. Thus, photoacoustic imaging can be used in combination with the dual nanoparticle system presented here to quantify the infiltration of macrophages and the persistence of MSCs following delivery.

In vivo photoacoustic imaging of nanoparticle labeled cells

The dual nanoparticle system (consisting of NR labeled MSCs and nanospheres) was intramuscularly injected within PEGylated fibrin gel into the gastrocnemius of a rat following an ischemic injury. Combined ultrasound and photoacoustic imaging (US/PA) obtained at 800 nm shows a high photoacoustic signal located within the gastrocnemius muscle (Fig. 9A), which persisted until day 7. Spectroscopic analysis attributes the photoacoustic signal to NRs (shown in green) immediately after the injection on day 0. Over time, the contribution from 50 nm PEGylated nanospheres increases (shown in red) (Fig. 9B). In addition, there is a 20% relative increase in signal attributed to the nanospheres from day 0 to day 7 (Fig. 9C). This increase in signal from nanospheres can be attributed to infiltration of macrophages following the ischemic injury and subsequent endocytosis of the PEGylated nanospheres by the cells. The endocytosis of the nanospheres by macrophages leads to plasmon coupling of the particles, as demonstrated by the peak shift in Fig. 1, and thus an increase in photoacoustic signal within the range of wavelengths which were used for imaging (680–970 nm). A majority of the infiltrating macrophages are localized around the delivered gel.
system. This could be attributed in part to the fact that the nanospheres were delivered within the gel system containing the MSCs, and thus the nanoparticles which were available for phagocytosis by the macrophages were already localized in close proximity to the stem cells. In addition, others have shown that macrophages have important roles in wound healing and angiogenesis, with stem cells and macrophages potentially undergoing some sort of synergistic crosstalk to further promote the regeneration process. MSCs have been shown to secrete not only angiogenic factors, but also factors which influence macrophage function. Furthermore, others have found that macrophages are frequently associated with endothelial cells during the process of angiogenesis. In addition, chemotactic signals secreted from apoptotic stem cells could also promote macrophage migration to the area. Stem cell death following delivery for therapeutic purposes is a common issue, and thus it would not be surprising if macrophages are entering the area to scavenge apoptotic MSCs. Thus, one of the advantages of this dual nanoparticle system is the ability to distinguish the two cell types in vivo, providing information about the interaction between stem cells and macrophages. Such information could include if macrophages are phagocytosing MSCs leading to a transfer of contrast agent from MSCs to macrophages. Based on our imaging system, we would designate transfer of contrast agent from MSCs to macrophages if there is co-localization of the nanosphere and nanorod signal.

Histological analysis of the muscle sample following sacrifice at day 7 shows the presence of injected MSCs (fluorescently pre-labeled with CM-Dil) distributed throughout the muscle fibers (Fig. 10A). In addition, the presence of nanoparticles within the cells is evident under bright field. TUNEL staining was performed to assess cell apoptosis following injection. The TUNEL staining in Fig. 10B shows a large extent of cell death within the injection region corresponding to the delivered photodynamic agent.

Fig. 8 (A) Spectroscopic in vitro photoacoustic imaging of nanosphere labeled macrophages (red) and nanorod labeled MSCs (green) mixed in various ratios. (B) Quantification of the ratio of the two cell types based on the photoacoustic imaging results.

Fig. 9 (A) Longitudinal ultrasound/photoacoustic imaging (λ = 800 nm) and spectroscopic photoacoustic imaging of NR labeled MSCs and nanospheres injected within PEGylated fibrin gel into the gastrocnemius. (B) Ratio of macrophages/MSCs over time. (C) Quantification of the relative photoacoustic signal increase attributed to nanospheres compared to day 0.
MSCs. It has been largely reported in the literature that only a small proportion of stem cells survive following delivery.\textsuperscript{55,57,58} Immunostaining for macrophages (ED1) in Fig. 10C shows macrophage infiltration within the muscle and in particularly close proximity to the injected MSCs, which were fluorescently labeled with CM-Dil prior to delivery. The ED1 immunostaining data supports the \textit{in vivo} photoacoustic imaging results, which shows the presence of macrophages on day 7 surrounding the injected MSCs.

Fluorescence-activated cell sorting of macrophages

Macrophages were sorted from gastrocnemius muscle 7 days following injection. ICP-MS analysis confirmed that macrophages endocytosed gold nanoparticles within the muscle (Fig. 11). To confirm that macrophages were not just endocytosing MSCs labeled with gold NRs, a control condition consisting of unlabeled MSCs was used. In this condition gold was also detected within isolated macrophages, confirming that macrophages are endocytosing the gold nanospheres.

Conclusions

This study demonstrated the use of a dual nanoparticle system to monitor stem cells following delivery and to detect the presence of infiltrating macrophages to a wound area. Cellular uptake of the nanoparticles was qualitatively and quantitatively assessed \textit{in vitro} using microscopy and mass spectrometry, respectively. In addition, the viability and function of the cells was maintained following nanoparticle labeling. An \textit{in vitro} phantom imaging experiment verified that the two cell types could be distinguished and quantified using photoacoustic imaging. The stem cells were also successfully monitored following intramuscular injection into a hind limb ischemia model, and the increased infiltration of macrophages into the area was quantified over time. Histological analysis of macrophages confirmed the photoacoustic imaging results, and mass spectrometry analysis confirmed that the macrophages had endocytosed gold nanoparticles, and specifically gold nanospheres. These results represent important advancements in monitoring stem cells for therapeutic purposes and distinguishing the delivered cells from infiltrating immune cells, which also have important roles in the wound healing response.

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The TEM images were acquired at the Microscopy and Imaging Facility of the Institute for Cellular and Molecular Biology at the University of Texas at Austin. The ICP-MS data was acquired at the Mass Spectrometry Facility in the Department of Chemistry at the University of Texas at Austin and at the Quadrupole ICP-MS lab in the Jackson School of Geosciences at the University of Texas at Austin. The FACS samples were run at the Microscopy and Imaging Facility of the Institute for Cellular and Molecular Biology at the University of Texas at Austin. This work was supported in part by a National Science Foundation Graduate Research Fellowship awarded to Laura M. Ricles and by NIH under grant EB015007.

Notes and references

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