Function of mesenchymal stem cells following loading of gold nanotracers

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Background: Stem cells can differentiate into multiple cell types, and therefore can be used for cellular therapies, including tissue repair. However, the participation of stem cells in tissue repair and neovascularization is not well understood. Therefore, implementing a noninvasive, long-term imaging technique to track stem cells in vivo is needed to obtain a better understanding of the wound healing response. Generally, we are interested in developing an imaging approach to track mesenchymal stem cells (MSCs) in vivo after delivery via a polyethylene glycol modified fibrin matrix (PEGylated fibrin matrix) using MSCs loaded with gold nanoparticles as nanotracers. The objective of the current study was to assess the effects of loading MSCs with gold nanoparticles on cellular function.

Methods: In this study, we utilized various gold nanoparticle formulations by varying size and surface coatings and assessed the efficiency of cell labeling using darkfield microscopy. We hypothesized that loading cells with gold nanotracers would not significantly alter cell function due to the inert and biocompatible characteristics of gold. The effect of nanoparticle loading on cell viability and cytotoxicity was analyzed using a LIVE/DEAD stain and an MTT assay. The ability of MSCs to differentiate into adipocytes and osteocytes after nanoparticle loading was also examined. In addition, nanoparticle loading and retention over time was assessed using inductively coupled plasma mass spectrometry (ICP-MS).

Conclusion: Our results demonstrate that loading MSCs with gold nanotracers does not alter cell function and, based on the ICP-MS results, long-term imaging and tracking of MSCs is feasible. These findings strengthen the possibility of imaging MSCs in vivo, such as with optical or photoacoustic imaging, to understand better the participation and role of MSCs in neovascularization.

Keywords: neovascularization, gold nanoparticles, cell tracking, stem cells, fibrin gel, optical imaging

Introduction

Stem cells are advantageous in that they can potentially be used for cellular therapies due to the fact that they can differentiate into multiple cell types,1 including cardiovascular cells.2–5 Previous work by our group demonstrated that PEGylated fibrin gels promote mesenchymal stem cell (MSC) tubulogenesis and differentiation towards a vascular cell type.4 These findings suggest MSCs could potentially be delivered to an injury site after an ischemic event in order to assist in neovascularization and tissue repair.3–7 However, the exact mechanisms of vascular repair, as well as the role and extent of participation of MSCs in neovascularization, are not well understood.

To this end, there have been numerous proposed approaches for long-term tracking of stem cells in vivo. A majority of this research has focused on labeling MSCs in vitro,
delivering the MSCs in vivo, and sacrificing the animals at various time points and performing histology. However, a major issue with this methodology is the inability to track the cells within a single, live animal model over time. As a result, many investigators have turned to noninvasive imaging modalities which are capable of monitoring and tracking stem cells within a single model over time.

The imaging approaches previously investigated require use of contrast agents to label the cells. Efficient cell labeling with contrast agents is necessary in order to image and track MSCs adequately. For long-term imaging, it is desirable to have contrast agents which have long-term stability, are not toxic to the cells, and do not affect cell function. Radionuclides have short half-lives, and thus are not ideal for long-term imaging studies. Reporter genes indirectly label cells and are not diluted as a result of cell division. However, the cells need to be genetically manipulated, which could alter cell function, and a substrate must be administered intravenously for each imaging session. Nanoparticles provide an advantage over radionuclides, in that they can be used for long-term imaging and allow for repeated imaging over time. Although nanoparticle labeling can decrease over time due to cell division, certain nanoparticle materials, such as gold, are not toxic to the cells and do not alter cell function.

Taking into account the advantages and disadvantages of the various contrast agents which can be employed, we are interested in investigating gold nanoparticles to be used as contrast agents. We propose to use gold nanoparticles as contrast agents because of their optically tunable properties and inert characteristics. In addition, photoacoustic imaging can be used to image cells labeled with gold nanoparticles. Photoacoustic imaging can be used for in vivo imaging at reasonable depths, as well as for longitudinal studies. Specifically, the long-term goal of our study is to assess optically how MSCs delivered in vivo via a PEGylated fibrin gel participate in the process of neovascularization. In this study, we assessed the efficiency of MSC labeling with gold nanotracers and the effects of nanoparticle loading on cell function. We determined initial nanoparticle loading and retention over time in order to assess the feasibility of long-term imaging of MSCs. This study provides a first step in achieving the goal of tracking the migration of MSCs delivered in vivo using PEGylated fibrin gels in order to assess the role of MSCs in the process of neovascularization and tissue repair. A better understanding of the process of neovascularization, and the extent to which MSCs participate in this process, can potentially lead to the development of better therapies and therapeutics, and specifically therapies involving MSCs, for tissue repair.

**Methods and materials**

**Human MSC culture**

Human MSCs (Lonza, Walkersville, MD) were cultured in Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen, Carlsbad, CA) supplemented with 10% fetal bovine serum, 1% glutamax, and 1% penicillin-streptomycin. The cell culture medium was changed every 2–3 days. Cells were passaged using 0.25% trypsin/ethylenediamine tetra-acetic acid (Lonza), collected by centrifugation at 600 × g for seven minutes, and counted with Trypan Blue exclusion. For all experiments, cells were seeded at a density of 5000 cells/cm². Cells were grown under standard cell culture conditions (37°C, 5% CO₂). Passage 5–8 human MSCs were used in this study.

**Nanoparticle synthesis**

Gold nanoparticles were synthesized via citrate reduction of tetrachloroauric (III) acid (HAuCl₄) under reflux. Briefly, 100 mL of distilled water was heated to 100°C, and 1 mL of 10 mg/mL HAuCl₄ was added while stirring. Sodium citrate (11.4 mg/mL) dissolved in distilled water was then added. Gold nanoparticles of various sizes (20 nm, 40 nm, and 60 nm) were synthesized by varying the volume of sodium citrate, ie, 2 mL of sodium citrate solution was added for 20 nm nanoparticles, 1 mL of sodium citrate solution was added for 40 nm nanoparticles, and 0.6 mL of sodium citrate solution was added for 60 nm nanoparticles. Gold nanoparticles were coated with poly-L-lysine (1–5 kD) by adding 0.3 mg of poly-L-lysine (1–5 kD) solution and allowed to mix for 30 minutes. The nanoparticles were then centrifuged at 3200 × g for 30 minutes in order to remove the excess poly-L-lysine. Poly-L-lysine hydrobromide (Sigma-Aldrich, St. Louis, MO) was dissolved in distilled water prior to use.

**Nanoparticle characterization**

A DelsaNano (Beckman Coulter Inc, Brea, CA) was used to determine the size and zeta potential of citrate-stabilized and poly-L-lysine-coated gold nanoparticles in water and in media. Briefly, nanoparticle solutions were loaded into a glass cuvette and readings were taken at a temperature of 25°C. For size measurements, five repetitions were performed for each nanoparticle solution, with 60 readings per repetition.
For zeta potential measurements, three repetitions were performed for each nanoparticle solution, with five readings per repetition.

A FEI Tecnai transmission electron microscope fitted with a top mount AMT Advantage HR 1k × 1k digital camera was used to determine the size and shape of citrate-stabilized and poly-L-lysine-coated gold nanoparticles in water and in media. Images were taken at 135k× magnification. Prior to imaging, carbon-coated copper 300 mesh grids (Electron Microscopy Sciences, Hatfield, PA) were placed in the nanoparticle solutions and allowed to incubate for approximately two minutes. The grids were then removed from the solution and allowed to dry.

Nanoparticle incubation and assessment of cell uptake
Cells were cultured as described earlier. Nanoparticle medium was made by centrifuging the nanoparticle solutions (5000 × g for 15 minutes for 20 nm citrate-stabilized nanoparticles and 3220 × g for 30 minutes for all other nanoparticle solutions) and resuspending the pellet in phenol red free medium at a concentration of approximately 1012 nanoparticles/mL. MSC growth medium was aspirated from the flasks, and 200 µL/cm2 of nanoparticle medium was added to the cell culture and allowed to incubate for 24 hours. After 24 hours, the nanoparticle medium was removed and the cells were washed with phosphate-buffered saline. A Leica DMI2000B microscope equipped with a Leica DFC290 camera was used to obtain darkfield images (20× magnification) in order to assess nanoparticle uptake.

LIVE/DEAD staining
In order to assess cell viability after nanoparticle uptake, a LIVE/DEAD stain was performed. Briefly, stock solutions of calcine AM (4 mM) and ethidium homodimer-1 (2 mM) were diluted in phosphate-buffered saline to a final working concentration of 4 µM and 1 µM, respectively. MSC growth medium was removed from the wells, and the cells were washed twice in phosphate-buffered saline. The cells were then incubated in the LIVE/DEAD stain for 45 minutes at 37°C, after which they were washed twice in phosphate-buffered saline. The stained cells were then imaged using fluorescence microscopy (Leica DMI2000B microscope equipped with a Leica DFC290 camera) under 10× magnification. Control cells consisted of cells not incubated with nanoparticles. Cell viability was assessed at various time points over a two-week period.

MTT assay
A Sigma-Aldrich MTT cell proliferation assay kit was used to perform the MTT assay. Briefly, MTT reagent was added to cells at a ratio of 100 µL/mL of medium. The samples were incubated for three hours at 37°C. The detergent reagent was then used to lyse the cells. A universal microplate reader ELX800 (Bio-Tek Instruments Inc, Winooski, VT) was used to measure the absorbance values of the samples at 562 nm and a reference wavelength of 630 nm. Control cells consisted of cells not incubated with nanoparticles. A blank sample containing no cells was used and subtracted from all measurements. Cytotoxicity was assessed over a one-week period at various time points. Triplicate samples were used and the assay was performed twice.

The number of cells was calculated using a standard curve. A one-way analysis of variance, followed by t-tests for multiple comparisons, was used to compare the experimental conditions with the control condition at the same time point (P < 0.01).

MSC differentiation
The ability of the cells to exhibit bipotent proliferation (ie, adipogenic and osteogenic) after nanoparticle uptake was assessed for all nanoparticle conditions. Control cells consisted of cells not incubated with nanoparticles and which were induced to differentiate into the specified lineages. Negative control cells consisted of cells not incubated with nanoparticles and which were not induced to differentiate into the specified lineages.

Briefly, adipogenesis was induced by plateauing cells at a density of 2.1 × 104 cells/cm2 and allowing the cells to grow to confluence in MSC growth medium. After the cultures reached 100% confluence, the MSC growth medium was replaced with adipogenic induction medium (DMEM, 10% fetal bovine serum, 1% penicillin-streptomycin, 1% glutamax, 1 µM dexamethasone [Sigma-Aldrich], 10 µg/mL 3-isobutyl-1-methylxanthine [Sigma-Aldrich], 10 µg/mL insulin [Sigma-Aldrich], and 100 µM indomethacin [Sigma-Aldrich]). After three days, the adipogenic induction medium was replaced with adipogenic maintenance medium for one day (DMEM, 10% fetal bovine serum, 1% penicillin-streptomycin, 1% glutamax, and 10 µg/mL insulin). Three cycles of induction/maintenance were performed, after which the cells were incubated in adipogenic maintenance medium for seven days. Control cells were supplemented only with adipogenic maintenance medium. Oil red O
(Sigma-Aldrich) staining was used to assess adipogenesis. The cells were fixed in 10% formalin and then incubated in 60% isopropanol for four minutes. The cells were then incubated in oil red O staining solution for five minutes, rinsed in tap water, and counterstained with hematoxylin for one minute. After washing with tap water, the slides were mounted and viewed under phase contrast using a Leica DMIZ2000B microscope equipped with a Leica DFC290 camera (20× magnification).

Osteogenesis was induced by plating cells at a density of $3.1 \times 10^4$ cells/cm². The cells were allowed to adhere for 24 hours in MSC growth medium, after which the medium was replaced with osteogenic induction medium (DMEM, 10% fetal bovine serum, 1% penicillin-streptomycin, 1% glutamax, 50 µg/mL ascorbic acid, 100 nM dexamethasone [Sigma-Aldrich], and 10 mM beta-glycerophosphate disodium salt hydrate [Sigma-Aldrich]). Osteogenesis was induced over a period of two weeks, after which a Fisher von Kossa staining kit was used to assess osteogenesis. The cells were fixed in 10% formalin and then incubated in 5% silver nitrate for 40 minutes with exposure to ultraviolet light. The cells were washed in distilled water and then placed in 5% sodium thiosulfate for two minutes, after which they were rinsed in tap water and placed in nuclear fast red stain for five minutes. After washing with tap water, the slides were mounted and viewed under brightfield using a Leica DMI2000B microscope equipped with a Leica DFC290 camera (20× magnification).

Inductively coupled plasma mass spectrometry

Initial nanoparticle loading and retention over a two-week period was assessed for cells incubated with 20 nm citrate-stabilized nanoparticles using inductively coupled plasma mass spectrometry (ICP-MS). Cells were cultured and incubated with nanoparticles for 24 hours as described above. After 24 hours, the nanoparticle medium was removed and the cells were washed twice with phosphate-buffered saline. The cells were then incubated in MSC growth medium without nanoparticles and collected at the designated time points. Cell medium was changed every 2–3 days with MSC growth medium, and the cells were passaged after reaching about 80% confluence.

At the designated time points, the cells were collected, centrifuged, and resuspended in 300 µL of 70% nitric acid and incubated at 60°C for 14 hours, after which the cells were diluted 100× in distilled water. A standard curve was used to quantify the amount of gold in the cells. Briefly, standard solutions were made by diluting AAS gold standard (Sigma-Aldrich) in the same background solution as that of the cell solutions. All measurements were made using triplicate samples.

Results

Nanoparticle characterization

Nanoparticle size was analyzed using transmission electron microscopy (TEM) and dynamic light scattering (DLS). In addition, nanoparticle shape and zeta potential were assessed using TEM and DLS, respectively. Using DLS, the average size of the three nanoparticle formulations (ie, citrate-stabilized nanoparticles in water) were determined to be $23.1 \pm 4.14$ nm, $49.78 \pm 9.18$ nm, and $57.58 \pm 10.72$ nm (see Table 1), and were designated as 20 nm, 40 nm, and 60 nm, respectively. In addition, representative TEM images of 20 nm, 40 nm, and 60 nm citrate-stabilized nanoparticles in water are shown in Figures 1A, 1B, and 1C, respectively.

Coating the gold nanoparticles with poly-L-lysine caused the zeta potential of the nanoparticles in water to go from negatively charged (approximately $-30$ mV) to positively charged (approximately $+30$ mV for 40 nm and 60 nm poly-L-lysine and $+14$ mV for 20 nm poly-L-lysine), as determined using DelsaNano (results not shown). There was also a slight increase in the average size of the nanoparticles after coating with poly-L-lysine (see Table 1). It should be noted that 20 nm particles were most likely to undergo aggregation in the presence of poly-L-lysine, which may be attributed to cross-linking of nanoparticles by poly-L-lysine. This effect is observed for the smaller particles because, as the size of the nanoparticle decreases, the relative concentration of nanoparticles/mL upon synthesis increases as the third power of the ratio of nanoparticle diameters. As a result, a lower amount of poly-L-lysine per nanoparticle was added for the 20 nm particles compared with the 40 nm and 60 nm particles, which could result in insufficient coating of the smaller nanoparticles.

Table 1 Average nanoparticle diameter

| Average diameter |
|------------------|
| 20 nm CS (water) | $23.10 \pm 4.14$ nm |
| 20 nm PLL (water) | $1128.10 \pm 226.64$ nm |
| 40 nm CS (water) | $49.78 \pm 9.18$ nm |
| 40 nm PLL (water) | $52.02 \pm 9.73$ nm |
| 60 nm CS (water) | $57.58 \pm 10.72$ nm |
| 60 nm PLL (water) | $61.04 \pm 11.65$ nm |

Notes: Dynamic light scattering was used to determine the average size of the nanoparticle formulations. Results are shown as average nanoparticle size ± standard error.
sized nanoparticles and thus formation of aggregates. To
test this hypothesis, we varied the concentration of poly-L-lysine
used to coat 20 nm gold nanoparticles and found that
increased poly-L-lysine concentration resulted in decreased
nanoparticle aggregation (results not shown).

MSC nanoparticle loading
MSC nanoparticle loading was assessed using darkfield
microscopy. Control cells were not incubated with nano-
particles and appear as a faint blue (Figure 2A). Due to the
faint appearance of the control cells under darkfield, a phase
contrast image of the control cells is provided in Figure 2A1
to verify the presence of cells.

Nanoparticle-loaded cells appear a yellow-orange color
when observed under darkfield microscopy, as compared
with cells not loaded with nanoparticles, which appear
blue. The darkfield images show that cells were loaded
with 20 nm, 40 nm, and 60 nm citrate-stabilized nano-
particles after 24 hours of incubation (Figures 2B, 2D,
and 2F, respectively), as evidenced by the yellow-orange
appearance of the cells. In addition, cells were loaded
with 20 nm, 40 nm, and 60 nm poly-L-lysine-coated
nanoparticles after 24 hours of incubation (Figures 2C, 2E,
and 2G, respectively), as evidenced by the yellow-orange
appearance of the cells.

Cells incubated with 20 nm, 40 nm, and 60 nm gold
nanoparticles coated with polyethylene glycol did not exhibit
nanoparticle uptake, as evidenced by the lack of yellow-
orange color of the cells (results not shown). Instead, cells
incubated with polyethylene glycol coated nanoparticles
(PEGylated nanoparticles) more closely resembled the
control cells. However, MSC nanoparticle loading was not
expected for cells incubated with PEGylated nanoparticles
because of the well documented evidence for polyethylene
glycol as a stealthing agent.23,24 Due to the fact that the nano-
particle formulations used in this study did not incorporate
targeting agents (eg, targeting antibodies), nanoparticle
uptake by cells was passive, and thus MSCs were unable to
detect and passively uptake PEGylated nanoparticles.

MSC viability and cytotoxicity
MSC viability and cytotoxicity after nanoparticle loading was
assessed using a LIVE/DEAD stain and an MTT assay. The
effect of nanoparticle loading on MSC viability was assessed
over a two-week period at various time points. The time
points (day 1, day 7, and day 14) correspond to the number
of days after initial nanoparticle incubation. The results of the
LIVE/DEAD assay are shown in Figure 3A. Green cells were
considered live and red cells were considered dead. Control
cells were not incubated with nanoparticles. As evident from
the representative LIVE/DEAD images in Figure 3A, the
cells remained viable for all nanoparticle formulations over
a two-week period, and there was minimal cell death present
(indicated by white arrows).
Nanoparticle cytotoxicity was assessed at various time points over a one-week period using an MTT proliferation assay. The time points (day 1, day 4, and day 7) correspond to the number of days after initial nanoparticle incubation. The cell numbers were quantified using a spectrophotometer and a standard curve. Control cells were not incubated with nanoparticles. The results for the MTT assay are shown in Figure 3B. It should be noted that the nanoparticles loaded within the cells did not contribute to the ultraviolet-visible spectrophotometry readings (results not shown). Cell numbers for the various nanoparticle formulations were statistically compared with the control condition for the corresponding day. It was found for day 1 that cells loaded with 20 nm and 60 nm citrate-stabilized nanoparticles had a significantly lower cell number compared with control cells ($P < 0.01$). In addition, on day 7, cells loaded with 60 nm poly-L-lysine nanoparticles had a significantly lower cell number compared with control cells ($P < 0.01$). It should be noted that some conditions (day 4, 60 nm citrate-stabilized, 20 nm poly-L-lysine, 40 nm poly-L-lysine; day 7, 40 nm poly-L-lysine) had a significantly higher cell number compared with the control for the corresponding time point, the cause for which is not known at this time.

**MSC differentiation**

The bipotent ability of MSCs to differentiate into adipogenic and osteogenic lineages after nanoparticle loading was assessed in order to determine if MSC function was maintained after nanoparticle loading. Control (+) cells were not incubated with nanoparticles and were induced to differentiate, and control (−) cells were not incubated with nanoparticles and were not induced to differentiate.

Figure 4A shows the results of oil red O staining for assessing adipogenesis. Fatty lipid deposits were stained red (indicated by white arrows) and are indicative of adipogenesis. The ability of MSCs to differentiate into adipocytes was maintained for all nanoparticle formulations, as qualitatively assessed by comparing the control images with the nanoparticle-loaded MSC images. As expected, control (−) cells did not exhibit any lipid deposits, as indicated by the negative staining.

Figure 4B shows the results of von Kossa staining for assessing osteogenesis. Calcium deposits were stained black (indicated by red arrows) and are indicative of osteogenesis. The ability of MSCs to differentiate into osteocytes was maintained for all nanoparticle formulations, as qualitatively assessed by comparing the control images with the nanoparticle-loaded MSC images.
Figure 4 The bipotent ability of mesenchymal stem cells to differentiate into adipogenic and osteogenic lineages was assessed after 24 hours of nanoparticle incubation. Control cells were not incubated with nanoparticles. Control (+) cells were induced to differentiate and control (−) cells were not induced to differentiate. A) Oil red O staining was used to assess adipogenic differentiation by staining for lipid deposits (white arrows). B) von Kossa staining was used to assess osteogenic differentiation by staining for calcium deposits (red arrows). Scale bar 100 µm.

Abbreviations: CS, citrate-stabilized; PLL, poly-L-lysine.

nanoparticle-loaded MSC images. Control (−) cells did not exhibit any calcium deposits.

**MSC nanoparticle uptake and retention**

Initial nanoparticle loading and retention over a two-week period were determined for MSCs loaded with 20 nm citrate-stabilized gold nanoparticles using ICP-MS (see Figure 5A). The average number of nanoparticles/cell decreased by approximately an order of magnitude over a two-week period from $4.53 \times 10^5 \pm 3.98 \times 10^3$ nanoparticles/cell to $2.00 \times 10^4 \pm 2.18 \times 10^2$ nanoparticles/cell. Also, the number of nanoparticles/cell decreased exponentially over a two-week period. Figure 5B compares the average number of nanoparticles/cell and cell proliferation rate for cells loaded with 20 nm citrate-stabilized nanoparticles. The data were normalized to the maximum value for the corresponding data set. Day 7 was the only data point that showed a significant difference between the ICP-MS and cell proliferation data ($P < 0.01$), as determined using a Student’s $t$-test.

**Discussion**

Previous work has demonstrated that PEGylated fibrin gels promote MSC differentiation and tubulogenesis. Many investigators have suggested delivering stem cells to an injury site (eg, myocardial infarction) in order to promote wound healing and retain normal physiological function. However, it is not clear if the MSCs implanted within the gels are actively participating in neovascularization in vivo, and, if so, what the specific role of the delivered MSCs is in neovascularization. Thus, we are proposing to load MSCs...
with gold nanotracers in order to track and monitor the growth of MSCs over time in vivo after delivery. The specific aims of this study were to determine if nanoparticle loading of MSCs with gold nanotracers was possible, and to assess if MSC function was compromised as a result of nanoparticle loading. To this end, the results of this study demonstrate the feasibility of loading MSCs with gold nanotracers in order to track the MSCs in vivo. Specifically, MSCs were loaded with 20 nm, 40 nm, and 60 nm citrate-stabilized and poly-L-lysine-coated gold nanoparticles (see Figure 1). In addition, nanoparticle loading was not cytotoxic to the cells and did not substantially affect MSC viability. MSC differentiation to adipogenic and osteogenic lineages was also maintained.

MSC nanoparticle loading was accomplished via passive uptake of the nanoparticles by the cells because no targeting agents were used to promote nanoparticle uptake. The nanoparticles aggregated when resuspended in cell culture medium, as evident from the change in color of the nanoparticle solution from deep red to light blue, as well as the presence of peak broadening in the absorbance spectrum. Nanoparticle aggregation can most likely be attributed to the adsorption of serum proteins onto the nanoparticle surface. It is possible that protein adsorption could promote nanoparticle uptake by the cell.

It is interesting to note that the poly-L-lysine nanoparticles did not have a negative effect on MSC function. Other investigators have found that high concentrations of poly-L-lysine induce cell toxicity. However, poly-L-lysine is commonly used to coat substrates in low concentrations in order to promote cell adhesion, with no negative effects on cell viability. In this study, the cells were exposed to a low concentration of poly-L-lysine coating the nanoparticles, and, as a result, cell toxicity was not induced. In fact, previous investigators have also found that poly-L-lysine in low concentrations, such as that used for coating nanoparticles, is not toxic to cells.

The results of this study also demonstrate that nanoparticle loading decreased exponentially over time. Previous investigators have cited the dilution of nanoparticle concentration per cell as a result of cell proliferation and division. In addition, other investigators have suggested cell exocytosis of nanoparticles may occur. The exocytosis process of cells loaded with gold nanoparticles has been quantified, with the exocytosis process dependent on nanoparticle size and shape, cell type, and extracellular environment. Taken together, these studies demonstrate that decreased nanoparticle loading over time could be attributed to cell division, as well as other mechanisms, such as exocytosis of the nanoparticles by the cells. The normalized data for the average number of nanoparticles/cell and the inverse of the cell number over time (see Figure 5B) exhibit a similar trend, with what appears to be an exponential decrease over time. If the decrease in nanoparticle retention over time was due only to cell division, we would not expect a significant difference between the ICP-MS results and the cell proliferation data at the same time points. Of the three time points analyzed, only day 7 showed a statistically significant difference between the ICP-MS results and the cell proliferation data at the same time points. Of the three time points analyzed, only day 7 showed a statistically significant difference between the ICP-MS results and the cell proliferation data at the same time points. Of the three time points analyzed, only day 7 showed a statistically significant difference between the ICP-MS results and the cell proliferation data at the same time points.

Figure 5 A) Nanoparticle uptake and retention over a two-week period were assessed using inductively coupled plasma mass spectrometry (ICP-MS) for cells incubated with 20 nm citrate-stabilized (CS) nanoparticles for 24 hours. The results are shown as the average number of nanoparticles/cell ± standard error. B) Comparison of the normalized values for the average number of nanoparticles/cell and the inverse of cell proliferation over time for control and 20 nm citrate-stabilized cells. The results are shown as the mean ± standard error.
the results, a majority of the decrease in nanoparticle retention over time was due to cell division.

Regardless of the mechanisms responsible for the decrease in the number of nanoparticles/cell, the decrease in nanoparticle retention over time has implications for the ability to image cells over long time periods. That is, a significant decrease in nanoparticle loading over time corresponds to a substantial decrease in signal when imaging. However, previous investigators have demonstrated the ability to use photoacoustic imaging, which is a noninvasive imaging approach that is capable of imaging cells at sufficient depths in vivo, to image human epithelial carcinoma cells. The cells were also imaged in vitro and loaded with 50 nm epidermal growth factor receptor-targeted gold nanoparticles almost an order of magnitude lower than that exhibited by MSCs after two weeks. Thus, we are confident that MSCs loaded with gold nanotracers can be detected and imaged over a two-week time period using photoacoustic imaging.

As mentioned previously, the end goal of this study is to develop an imaging technique which can visualize and track MSCs at sufficient depths and spatial/temporal resolutions. Specifically, we are interested in using a noninvasive imaging approach which is capable of optically tracking MSCs in vivo over long time periods within a single animal model. Optical imaging modalities which could be employed to monitor and track MSCs include photoacoustic imaging, confocal microscopy, two-photon microscopy, and optical coherence tomography.

Conclusion

In summary, the work presented here demonstrates the ability to load MSCs with gold nanotracers of various sizes and surface coatings. In addition, this study also demonstrates that MSC function is not altered by nanoparticle loading. Furthermore, these findings strengthen the possibility of imaging MSCs in vivo in order to understand better the participation and role of MSCs in neovascularization. The concentration of nanoparticle loading and retention over a two-week period suggests that long-term imaging and tracking of MSCs is feasible. Specifically, we are interested in developing an imaging approach to track MSCs in vivo after delivery via a PEGylated fibrin matrix using MSCs loaded with gold nanotracers. Thus, our future research will include in vitro imaging studies to verify that nanoparticle-loaded MSCs can be imaged using optical imaging modalities (eg, photoacoustic imaging combined with ultrasound imaging) and that MSC viability is not affected as a result of imaging. In addition, in vivo studies are necessary to monitor the progression of MSC growth over time.

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Disclosure

The authors report no conflicts of interest in this work.

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